

Some recent advances in *ab initio* calculations of nonradiative decay rates of point defects in semiconductors

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Abstract: In this short review, we discuss a few recent advances in calculating the nonradiative decay rates for point defects in semiconductors. We briefly review the debates and connections of using different formalisms to calculate the multi-phonon processes. We connect Dr. Huang's formula with Marcus theory formula in the high temperature limit, and point out that Huang's formula provide an analytical expression for the phonon induced electron coupling constant in the Marcus theory formula. We also discussed the validity of 1D formula in dealing with the electron transition processes, and practical ways to correct the anharmonic effects.

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1. Introduction

It is a great honor to write this short review in memorial of Dr. Kun Huang, one of the founding persons of Chinese semiconductor science, and a great pioneer in studying electron-phonon coupling in semiconductors. I didn't have the opportunity to meet with Huang in person, but have the fortune to learn from and collaborate with many of his students and student's students. The favorite topic of Huang is electron-phonon coupling and its consequences in semiconductor physics. In this short review, I will discuss a few recent developments in this area, mostly based on my own works, some of them in collaborations with colleagues in the Institute of Semiconductors in Chinese Academic of Science for which Huang has served as its first director. This by no mean is a complete review, but rather some personal views about this topic.

One topic is the nonradiative decay of free carriers caused by defects in semiconductor, most notably the Shockley-Read-Hall (SRH) process. This is an important process in the operation of semiconductor devices, especially for deep defect states. The electrons (or holes) hop from the band edge to the defect state through multiple phonon processes, and it can subsequently lead to carrier annihilation. The basic formalisms of such multiphonon processes have been worked out in 50's by S. K. Pekar^[1] and K. Huang, A. Rhys^[2]. These derivations are based on Franck-Condon approximation of the electron-phonon wave functions, where the electron-nuclear wave function is approximated as $\Psi_{i,n}(r, R) = \psi_i(r, R)\phi_{i,n}(R)$, here i and n are electron and phonon eigen state index, and r and R are electron and nuclear degree of freedom. While the electron wave function $\psi_i(r, R)$ satisfies the electron Schrodinger's equation at a given nuclear coordinate R :

$$H(r, R)\psi_i(r, R) = \varepsilon_i(R)\psi_i(r, R). \quad (1)$$

The nuclear wave function $\phi_{i,n}(R)$ satisfies its own Schrodinger's equation using $\varepsilon_i(R)$ as its potential energy:

$$\left[\sum_R -\frac{1}{2M_R} \nabla_R^2 + \varepsilon_i(R) \right] \phi_{i,n}(R) = E_{i,n} \phi_{i,n}(R). \quad (2)$$

Here $E_{i,n}$ is the total eigen energy of the electron-nuclear wave function $\Psi_{i,n}(r, R)$, M_R is the nuclear mass. However, the $\Psi_{i,n}(r, R) = \psi_i(r, R)\phi_{i,n}(R)$ is not a true eigen state of the electron-nuclear total Hamiltonian due to a cross derivative term: $\frac{\partial \psi_i(r, R)}{\partial R} \frac{\partial \phi_{i,n}(R)}{\partial R}$ as well as a second derivative term: $\frac{\partial^2 \psi_i(r, R)}{\partial R^2}$. Usually the cross derivative term is much larger than the second derivative term, and it causes coupling between the initial electron state $\psi_i(r, R)$ and the final electron state $\psi_j(r, R)$. This naturally leads to the adiabatic state coupling formalism^[1, 2].

Although elegant in its derivation, very soon it was realized that the adiabatic state coupling formalism gave too small transition rates compared with experiments. To solve this problem, Kovaskiy, Sipdsvskiy^[3, 4] and Passler^[5, 6] proposed to use a static formalism, where both $\psi_{i,n}(r, R)$ and $\psi_{j,n}(r, R)$ are replaced by their counter parts $\psi_{i,n}(r, R_0)$ and $\psi_{j,n}(r, R_0)$ where R_0 is chosen as the equilibrium position at one electronic state (e.g., R_0 will be either $R_0(i)$ or $R_0(j)$). Thus although the state $\psi_{i,n}(r, R_0)$ and $\psi_{j,n}(r, R_0)$ (the states before and after the transition) are the electron eigen states when $R = R_0$, when R deviates from this R_0 , they are no longer eigen states due to the dependence of the electron Hamiltonian on R . Thus, the phonon displacement away from the equilibrium position R_0 causes the electronic state coupling.

For a long time, there were debates for the validity and the meaning of using static coupling formalism^[3-8]. Huang gave a very interesting derivation in early 1980^[8, 9]. He shown that, in a perturbation representation of the electronic state $\psi_{i,n}(r, R)$ using the static state $\psi_{j,n}(r, R_0)$ as the basis set, if one replaces the static eigen energies $\varepsilon_i(R_0)$, $\varepsilon_j(R_0)$ in the perturbation theory denominator $1/(\varepsilon_i(R_0) - \varepsilon_j(R_0))$ by the R dependent eigen energies: $1/(\varepsilon_i(R) - \varepsilon_j(R))$, then one can arrive

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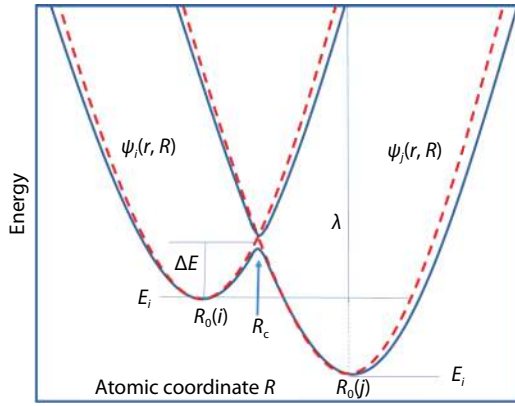


Fig. 1. (Color online) A schematic energy diagram of the phonon degree of freedom at the initial electronic state $\psi_i(r, R)$ and final electronic state $\psi_j(r, R)$.

at the static coupling formalism starting from the adiabatic coupling formalism. This in a sense unifies these two formalisms within the framework of perturbation theory^[8]. At the end, the static coupling formalism will be the preferred formalism.

One can view this problem from other angles. The Franck-Condon approximation is good only at R positions where there is no near degenerated i and j electron eigen states. Unfortunately, the degeneracy (or say the energy crossing of these two states) is exactly what happens at the coordinate R_c where the transition happens (see Fig. 1). So, the Franck-Condon approximation is invalid at the transition nuclear coordinate R_c . This is the same situation as for the conical point in quantum chemistry treatment of molecular systems. This can also be seen from Huang's perturbation treatment. If $1/(\epsilon_i(R) - \epsilon_j(R))$ is used, then at the crossing point R_c , the perturbation theory is diverging. Actually, if one follows exactly the adiabatic state definition, at the crossing point R_c , the physical identity of the i and j state will change, one must be very carefully in identifying which state is which when describing the transition. For example, there will be a large and sharp spike in $\frac{\partial \psi_i(r, R)}{\partial R}$ at R_c due to the identity change (if i is used to indicate one continuous adiabatic state, as indicated by one solid line in Fig. 1). This makes the Franck-Condon electron and phonon wave function separation treatment a bad approximation at this R_c , and perhaps more importantly, it makes the commonly used approximation in the adiabatic coupling treatment: replacing $\frac{\partial \psi_i(r, R)}{\partial R}$ by $\frac{\partial \psi_i(r, R)}{\partial R}|_{R_0}$, totally invalid at R_c . All these justify the use of static formalism.

Fig. 1 also connects the nonradiative state transition to the Landau-Zener transition^[10, 11] and Marcus theory^[12]. The Landau-Zener transition is mostly a 1D model. It has been used by Lang *et al.*^[13] to treat the nonradiative transition. Similar approach has also been used by Alkauskas *et al.*^[14] more recently in an *ab initio* treatment. However, the multiphonon process is not necessarily a 1D process, in the sense that the transition cross point R_c (the energy conservation modes) and the coupling (the promoting modes) are not caused by the same phonon degree of freedoms^[15]. More specifically, the energy conservation (accepting) modes (or say the phonon modes which lead to the transition point R_c) come from the displacement $\Delta R_{ij} = R_0(i) - R_0(j)$. This displacement ΔR_{ij} comes from the diagonal elements of the electron-phonon coupling:

$\langle \psi_i | \frac{\partial H}{\partial R} | \psi_i \rangle - \langle \psi_j | \frac{\partial H}{\partial R} | \psi_j \rangle$ (together with the phonon frequencies). On the other hand, the promoting modes comes from the cross terms of the electron-phonon coupling: $\langle \psi_i | \frac{\partial H}{\partial R} | \psi_j \rangle$. The ΔR_{ij} and the cross coupling can both be viewed as multidimension vectors, and the directions of these two vectors can be rather different. In the high dimensional space of R , their dot product could be close to zero^[16]. For example, in the $\text{Zn}_{\text{Ga}} + \text{V}_{\text{N}}$ defect complex in GaN, the energy accepting modes are consisted with acoustic phonon modes, while the promoting modes come from optical phonons^[15].

Marcus theory is also often used to describe electron transitions from one state to another, especially for electronic states located at different positions^[17]. While the energy barrier of the Marcus theory also comes from atomic displacements (hence the diagonal elements of electron-phonon coupling), the electronic coupling can come from different sources, e.g., an applied electric field, or more intrinsic coupling. In the case the coupling is also caused by phonon modes (e.g., the pure multiphonon process as described above), Huang has derived a very interesting high temperature classical approximation formula^[9], which essentially gives the coupling constant of Marcus theory based on the electron-phonon coupling constants and phonon mode frequencies. Even when the main coupling is not caused by electron-phonon coupling, it might be possible that the coupling constant provided by Huang's formula is related to the environmental effects during the Landau-Zener transition^[18, 19]. This will be an interesting research topic for the future.

2. *Ab initio* calculations of nonradiative carrier decay rates

Modern theoretical defect studies are mostly depending on *ab initio* calculations, especially using density functional theory (DFT)^[20–22]. In recent years, the use the hybrid functional^[23] has made the defect level calculations more accurate, and can yield energy level results consistent with experiments^[24]. One current frontier is to study the dynamic properties of such defects, including the nonradiative decay rates. As discussed above, the basic formalism (e.g., the static coupling formalism) already existed. Thus a contemporary challenge is to use *ab initio* method to calculate the electron-phonon coupling, and compared the results with experiments. The straight forward calculation of $\langle \psi_i | \frac{\partial H}{\partial R} | \psi_j \rangle$ requires the calculation of the change of $V_{\text{tot}}(r)$ (the total selfconsistent potential in DFT method) due to the displacement of one atomic coordinates. Thus, if there are N atoms in a supercell (as in a defect calculation), there need to be $3N$ self-consistent field (SCF) calculations. As N can be in the order of 1–2 hundreds, this makes the calculation rather expensive. We have developed an approach which can yield all the electron-phonon coupling constants for a given ψ_i, ψ_j pair in one SCF calculation^[15]. This makes the multiphonon process calculation more practical.

More specifically, to calculate $C(i, j, R) = \langle \psi_i | \frac{\partial V_{\text{tot}}(R)}{\partial R} | \psi_j \rangle$, during the SCF calculation, one can add one extra term in the totalelectronicchargedensity: $\rho(r) = \sum_i |\psi_i(r)|^2 o_i + a \text{Re}[\psi_i^*(r) \times \psi_j(r)]$, here o_i is the occupation number of the Kohn-Sham eigen state ψ_i . The first term is the usual formula to calculate the charge density, while the second term is the additional charge

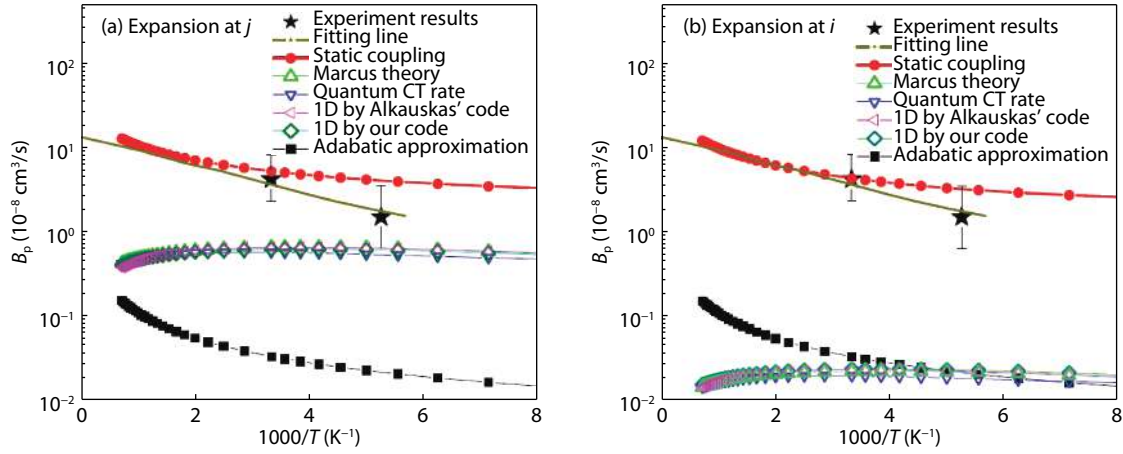


Fig. 2. (Color online) The comparison of transition rates calculated using different formulas for the nonradiative transition of electron from the conduction band of bulk GaP to the $\text{Zn}_{\text{Ga}} + \text{O}_{\text{p}}$ point defect. Marcus theory, quantum CT rate, 1D by Alkauskas's code, and 1D by our code, are all one dimensional models. They all give very similar results. Compared with experiment, the multiphonon static coupling formula gives the best results, while the adiabatic coupling results are almost two order of magnitudes smaller. In (a), the calculations are done using the $R_0(j)$ as the perturbation starting point, while in (b), $R_0(i)$ is used as the starting point. As one can see, the results of these two treatments are similar for the multiphonon formula of static coupling and adiabatic coupling. On the other hand, for all the 1D formula, the results are very different. The details of the calculations are described in Ref. [31]. The images are taken from Ref. [16] with permission.

density with α being a small parameter. Besides the charge density, all other formulas are kept the same as in a normal Kohn-Sham calculation. Furthermore, the atomic force is calculated using Hellmann-Feynman formula as: $F_{R,\alpha} = \int \rho(r) dV_{\text{ion}}(r, R) / dR d^3r$ (V_{ion} is the nuclear ionic potential). Then we can show that [15]:

$$C(i, j, R) = \frac{dF_{R,\alpha}}{d\alpha}. \quad (3)$$

The derivative regarding to α can be done numerically, by using $\alpha = 0$ (the original SCF calculation results) and a small α like 0.1. In doing so, we can get all the coupling constant $C(i, j, R)$ in one extra SCF calculation. The same variational approach can be applied to hybrid functional calculations. In that case, during the SCF calculation, besides the above additional term in the charge density, one also needs to add one extra term in the Fock exchange integral as:

$$\hat{P}\psi_k = \alpha \int [\psi_i(r)\psi_j^*(r') + \psi_j(r)\psi_i^*(r')] f(|r-r'|) \psi_k(r') d^3r', \quad (4)$$

here $f(|r-r'|)$ is the truncated Coulomb interaction kernel used in the hybrid functional. Besides the electron-phonon coupling constant, one also needs to calculate the phonon spectrum of the defect system in order to use the analytical formulas derived by Huang *et al.* One way to calculate the phonon spectrum of the supercell is to calculate the dynamic matrix $M(R_1, R_2) = \frac{\partial^2 E_{\text{tot}}}{\partial R_1 \partial R_2} = \frac{\partial F_{R_2}}{\partial R_1}$. Once again, this requires the numerical displacement of all N atoms within the supercell. Thus the benefit of the above variational calculation of the electron-phonon coupling constant will be lost if such dynamic matrix needs to be calculated directly. Fortunately, we found that [15] if both R_1 and R_2 are away from the point defect, beyond a cutoff radius R_0 , then $M(R_1, R_2)$ can be approximated by the counter part from the perfect crystal. For crystal, due to the translational symmetry, one only needs to displace the atoms within one primary cell, instead of all the atoms within a supercell. As a result, we only need to calculate $\frac{\partial F_{R_2}}{\partial R_1}$

for R_1 within R_0 . Typically 10–50 numerical displacements are needed to carry out the SCF calculations to yield all needed $\frac{\partial F_{R_2}}{\partial R_1}$. It has been shown this procedure yield very accurate phonon spectrum for the supercell system containing one point defect [15].

After we obtain both the electron-phonon coupling constant and phonon spectrum, the multi-phonon nonradiative process can be calculated directly. There are different ways to derive the analytical formula, all based on the use of Fermi Golden rule between the initial and final states with Franck-Condon separation of the electron wave function and phonon wave function. The coupling Hamiltonian in the static coupling approximation is a perturbation term proportional to $\sum_R C(i, j, R)(R - R_0)$ (note, the R in the summation \sum_R and in $C(i, j, R)$ is used as an index, while in $(R - R_0(i))$ it is a vector). R_0 is the starting point for the static coupling calculation. There is an ambiguity for what to choose for R_0 . Usually one either choose it as $R_0(i)$ or $R_0(j)$. However, for most cases, since the promoting mode direction of $C(i, j, R)$ is almost perpendicular to the accepting mode direction of $\Delta R_{ij} = R_0(j) - R_0(i)$, the choice of $R_0(i)$ or $R_0(j)$ does not really matter (as well be shown later in Fig. 2). What left is to evaluate $\langle \varphi_{i,n}(R) | (R - R_0(i)) | \varphi_{j,m}(R) \rangle$ under a thermodynamic assembly. The harmonic approximation is taken to describe phonon modes $\varphi_{i,n}(R)$ and $\varphi_{j,m}(R)$. Very often it is assume they are the same phonon modes (and frequencies) but with a zero point displacement of ΔR_{ij} , although analytical equations can also be obtained if different harmonic phonon modes at i and j are used [25]. To evaluate the $\langle \varphi_{i,n}(R) | (R - R_0(i)) | \varphi_{j,m}(R) \rangle$ under a thermodynamic assembly and a delta function for energy conservation, one first converts the delta function for energy conservation into an integral using Dirac distribution function:

$$\delta(\omega) = \frac{1}{2\pi} \int_{-\infty}^{\infty} e^{i\omega t} dt. \quad (5)$$

This always leads to an dt integration for the final formula. After subsequent derivations based on matrix manipulations^[26–28], one can obtain a formula for the i to j non-radiative transition rate as:

$$W_{ij} = 2\pi \sum_{k_1, k_2} C_{ij}^{k_1} C_{ij}^{k_2} A_{ij}^{k_1, k_2}, \quad (6)$$

here C_{ij}^k is just the electron-phonon coupling constant $C(i, j, R)$, however, converted into the phonon mode coordinate k . The matrix $A_{ij}^{k_1, k_2}$ can be evaluated as:

$$A_{ij}^{k_1, k_2} = \frac{1}{2\pi Z} \int_{-\infty}^{\infty} X_{ij}^{k_1, k_2}(t, T) e^{-i(E_i - E_j)t} dt, \quad (7)$$

here $Z = \sum_n \exp(-\beta E_{i,n})$ is the phonon partition function and $\beta = (k_B T)^{-1}$, the E_i and E_j are the defect state energies of the electronic state i and j at their equilibrium atomic positions $R_0(i)$ and $R_0(j)$ respectively (Fig. 1). The expression for the matrix $X_{ij}^{k_1, k_2}(t, T)$ is a bit complicated^[16]. It is expressed by several other matrices with phonon frequencies and the atomic displacement ΔR_{ij} as their variables.

In our previous work^[16], we have adopted the above formalism by Borrelli *et al.*^[28]. However, we later found that Huang gave a different derivation earlier in 1981^[9]. He explicitly integrated out the harmonic phonon wave functions using their Gaussian representations. He arrived at a more concise formalism:

$$W_{ij} = 2\pi \int_{-\infty}^{\infty} \left\{ \left[\sum_k C_{ij}^k \Delta Q_{ij}^k (\cos(\omega_k t) + i \coth(\beta \omega_k / 2) \sin(\omega_k t)) \right]^2 + \frac{1}{2} \sum_k |C_{ij}^k|^2 \frac{1}{\omega_k} (\coth(\beta \omega_k / 2) \cos(\omega_k t) + i \sin(\omega_k t)) \right\} \times \frac{1}{2\pi} \exp \left[-it(E_j - E_i) - \sum_s \frac{\omega_s}{2} |\Delta Q_{ij}^s|^2 (\coth(\beta \omega_s / 2) \times (1 - \cos(\omega_s t) - i \sin(\omega_s t))) \right] dt. \quad (8)$$

Here ΔQ_{ij}^k and ΔQ_{ij}^s are the atomic displacements ΔR_{ij} between state i and j converted into phonon mode coordinates k and s respectively. We have numerically tested this concise equation versus the more complicated equation derived by Borrelli *et al.*^[28], they give the exactly same results.

The concise formalism allows Huang to apply the steepest decent approximation to get an closed analytical formula (without the dt integration) for high temperature approximation. To do that, one finds the maximum of the exponent as a function of t , and expands it with a second order approximation. The resulting Gaussian exponential can be integrated over t , yielding in a closed analytical formula. Such closed analytical formulas exist for adiabatic approximation of multiphonon transition^[7], as well as for quantum mechanical treatment of the Marcus theory^[29] where the transition coupling between the two electron states ψ_i and ψ_j is assumed to be a constant, independent of the phonon degree of freedom. For the static coupling approximation, we only found such closed formula in Huang's work^[9]. We can thus call it the Huang's formula, which is:

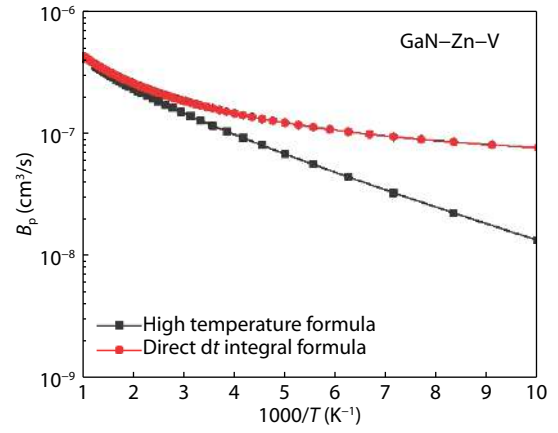


Fig. 3. (Color online) The nonradiative electron transition rates for a hole from the valence band edge to a $\text{Zn}_{\text{Ga}} + \text{V}_{\text{N}}$ complex defect state in GaN. The high temperature formula result is compared with direct t integration formula. When the temperature is smaller than 300 K, the high temperature formula under estimates the transition rate.

$$W_{ij} = \left(\frac{\pi k T}{S \bar{\omega}} \right)^{1/2} \left(\sum_k \frac{1}{\omega_k^2} |C_{ij}^k|^2 \right) \exp \left(- \frac{(E_i - E_j - S \bar{\omega})^2}{4 k T S \bar{\omega}} \right). \quad (9)$$

Here S is the Huang-Rhy's factor^[2] $S = \sum_k |\Delta Q_{ij}^k|^2 \frac{\omega_k}{2}$, and $S \bar{\omega} = \sum_k |\Delta Q_{ij}^k|^2 \frac{\omega_k^2}{2}$. The $S \bar{\omega}$ is nothing but the reorganization energy λ used in classical formula like the Marcus theory. This reorganization energy equals $\epsilon_i(R_0(j)) - \epsilon_i(R_0(i))$ as shown in Fig. 1. The above formula can be directly compared with the Marcus theory which is:

$$W_{ij} = \left(\frac{\pi k T}{\lambda} \right)^{1/2} \frac{1}{k T} |V_c|^2 \exp \left(- \frac{(E_i - E_j - \lambda)^2}{4 \lambda k T} \right). \quad (10)$$

Thus, Huang's formula provides an expression for the coupling constant $|V_c|^2$ in the Marcus theory as: $k T \sum_k \frac{1}{\omega_k^2} |C_{ij}^k|^2$ this is valid for the phonon induced coupling between the electron states i and j . As discussed before, one can think about other causes for the coupling in a more general case. Nevertheless, the Huang's formula can be used to estimate the phonon contribution to such coupling.

We have tested Huang's formula of Eq. (9) versus direct dt integration of Eq. (8). As shown in Fig. 3, the high temperature formalism is valid when the temperature T is higher than 300 K for the case of $\text{Zn}_{\text{Ga}} - \text{V}_{\text{N}}$ defect in GaN and a hole transition from the valence band edge to the defect state.

We can now numerically calculate the multiphonon transition rate using *ab initio* methods, either using the direct integration as in Eq. (8), or using the high temperature Huang's formula of Eq. (9). Such calculation procedure has been implemented within the PWmat code package^[30], in an automatic way to calculate the electron-phonon coupling, defect phonon spectrum, and the static coupling formula for the nonradiative transition rate. The calculated results are usually in good agreement with the experiments. Fig. 2 shows a comparison of different calculation methods, both with the explicit multiphonon mode calculation, and 1D models along the ΔR_{ij} direction. As we can see that, in this case, the 1D model all give very different results compared with the explicit multiphonon static coupling formula. The static coupling results are in good agree-

ment with the experiment. Furthermore, the 1D model sensitively depend on which equilibrium point is chosen for R_0 : $R_0(i)$ or $R_0(j)$. On the other hand, the static coupling result is insensitive to such choices.

3. Remaining challenges and direct dynamic process simulations of electron–phonon coupling problems

There are still challenges for accurate prediction of the non-radiative decay rate and the related studies. The first is an experimental one, as there is a scarcity of the experimentally measured nonradiative decay rate for different defects. The commonly used method: deep level transient spectroscopy (DLTS) can miss some deep defect with low concentrations, and results can also be influenced by factors like Coulomb repulsion between the charged defect and the band edge free carriers. Although there are other alternative techniques, both optical measurements and non optical measurements^[32], due to possible multichannel competition, the interpretation of the results can still be challenging. In terms of calculation, the anharmonicity of the phonon oscillation is one major uncertainty. In *ab initio* calculation, it is often found that the directly calculated the atomic relaxation energy λ after the electron transfers from i to j state (e.g. from band edge state to the defect state), can be larger than the relaxation energy as calculated by $S\bar{\omega}$. If the relaxation and the transition coupling happens in the same direction, then an 1D approximation can be used, and direct numerical calculation can be used to study the transition process as a Landau-Zener transition^[33]. However, if that is not the case, such 1D approximation cannot be used. In such cases, one approximation is to rescale all the phonon frequency so that $S\bar{\omega} = \sum_k |\Delta Q_{ij}^k|^2 \frac{\omega_k^2}{2}$ will yield the same results as the numerically calculated reorganization energy λ . This also allows us to evaluate the Eq. (8) in the low temperature situation. Another challenge is that, the phonon modes at i and j can be different. For Huang's formula, while one can use the average formula for the effective coupling constant, what more critical is the exponential term, which is $\exp(-\Delta E/kT)$, here ΔE is the barrier between i and j , as shown in Fig. 1. One approximation is to re-evaluate this ΔE , the lowest valley crossing point for the two multi-dimension parabolas between i and j , and using that to replace the exponential term in Huang's high temperature formula. As a matter of fact such correction can be even applied to the low temperature integration formula. We found that, after such correction, the calculated transition rate can increase by almost an order of amplitude in some cases, bringing the result further closer to the experimental values.

We also like to mention that sometime the SRH electron-hole recombination can happen through a multi-step process, with several intermediate transitions corresponding to different occupations and charge states for the defect level^[34]. Although analytical multi-phonon formula can still be used in such cases, one alternative approach could be to do direct real-time time dependent density functional theory (rt-TDDFT) simulations^[35]. In the rt-TDDFT calculation, the nuclear movement follow the Ehrenfest dynamics. It is a classical description for the nuclear movement, thus it is likely adequate in high temperature limit. Since the electron movement follows

the time dependent Schrodinger' equation, it can be used to describe the Landou-Zener transition. Compared to the analytical formalism, one advantage of direct rt-TDDFT is its ability to describe anharmonic nuclear movement and strong electron-phonon coupling. On the other hand, due to the classical description of the nuclear movement, it lacks the detail balance between the i to j transition and j to i transition. Recently, we have added such detailed balance within rt-TDDFT, as a result, it can be used to describe multiphonon nonradiative decay. For example, we have used such direct simulation to study the molecule dissociation caused by electron ionization. This is particularly useful to describe very strong electron-phonon coupling, e.g., inside a small molecule, or for a carrier on a localized defect state and going through multiple stages in such defect state. Future investigation of such problems will be interesting. For larger systems, one can ignore the back reaction from the electron movement to the nuclear movement, thus can use the ground state *ab initio* molecular dynamics trajectory to describe a time dependent Hamiltonian, then use such Hamiltonian to study the carrier dynamics. This approach is called nonadiabatic molecular dynamics (NAMD), which allows the simulations of much larger systems (e.g., a few hundred atoms) for much longer times^[36] (e.g., 10 ps). All these provide alternative approaches to study the electron-phonon coupling and its related carrier dynamics behavior. Compared with the analytical formula, one limitation is the simulation time. If the decay lifetime is much longer than 10 ps, direct simulation will become rather difficult. So they are only good for strongly electron-phonon coupled systems (for rt-TDDFT simulations) or for problems involve many electronic state transitions (for NAMD simulations).

4. Conclusions

In summary, in this short review, some current developments in calculating nonradiative decay rates of defects in semiconductors are presented, as well as a brief review for the derivation of Huang's formula. In particular, one procedure was developed to calculate all the electron-phonon coupling constant within one self-consistent field calculation. Another approximated way is introduced to calculate all the phonon modes within a supercell containing a point defect. Such developments, together with Huang's formalism, allow us to calculate the nonradiative decay rate at *ab initio* level, and the results agree well with the experiment. We have also discussed some of the remaining challenges and possible approaches to overcome them. These include the anharmonic phonon effects and different phonon modes at electronic state i and j . Finally, we discussed modern direct simulation methods, either rt-TDDFT, or NAMD, which can be used to study problems with strong electron-phonon coupling and strong anharmonicities of the phonon modes, or to study carrier dynamics involving many electronic states.

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