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<td>Author(s)</td>
<td>Laussy, Fabrice P.; Taylor, Thomas.; Shelykh, Ivan A.; Kavokin, Alexey V.</td>
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Superconductivity with excitons and polaritons: review and extension

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Superconductivity with excitons and polaritons: review and extension

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Abstract. A system where a Bose-Einstein condensate of exciton-polaritons coexists with a Fermi gas of electrons has been recently proposed as promising for realization of room-temperature superconductivity. In order to find the optimum conditions for exciton and exciton-polariton mediated superconductivity, we studied the attractive mechanism between electrons of a Cooper pair mediated by the exciton and exciton-polariton condensate. We also analyzed the gap equation that follows. We specifically examined microcavities with embedded n-doped quantum wells as well as coupled quantum wells hosting a condensate of spatially indirect excitons, put in contact with a two-dimensional electron gas. An effective potential of interaction between electrons was derived as a function of their exchanged energy \( \hbar \omega \), taking into account the retardation effect that allows two negatively charged carriers to feel an attraction. In the polariton case, the interaction is weakly attractive at long times, followed by a succession of strongly attractive and strongly repulsive windows. Strikingly, this allows high critical temperature solutions of the gap equation. An approximate three-steps potential is used to explain this result that is also obtained numerically. The case of polaritons can be compared with that of excitons, which realize the conventional scenario of high-\( T_c \) superconductivity where a large coupling strength accounts straightforwardly for the high critical temperatures. Excitons are less advantageous than polaritons but may be simpler systems to realize experimentally. It is concluded that engineering of the interaction in these peculiar Bose–Fermi mixtures is complex and sometimes counter-intuitive, but leaves much freedom for optimization, thereby promising the realization of high-temperature superconductivity in multilayered semiconductor structures.


Keywords: excitons; micro-optics; superconductivity.

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1 Introduction

The electron gas undergoes, in some conditions, a phase transition to bound pairs of electrons (the so-called Cooper pairs), which replace electrons as the fundamental agent of the electronic properties. Cooper pairs are, from the point of view of their electric charge, objects qualitatively identical to the underlying electrons. From the point of view of their spin, on the other hand, they become integer-spin particles, that is, from the spin-statistics theorem, bosons rather than fermions. This shift of statistical paradigm of the carriers—from Fermi to Bose-statistics—results in the...
outstanding behavior of superconductivity, that is, conduction of electric charge by a macroscopic coherent wavefunction (akin to a Bose–Einstein condensate). It has taken some time to capture the fundamental and universal features of this phenomenon and set them apart from particularities of certain cases only. The gap of excitations, responsible for zero resistivity, for instance, results from the long-range nature of the Coulomb interaction, but gapless superconductivity is also possible. One of the central, fundamental concepts of superconductivity is that of a coherent quantum state of charged bosons. Although superconductivity was discovered empirically, and its theoretical construction consisted in assembling a puzzle, it is now possible to envisage engineering superconducting phases in other systems, based on this understanding of condensation of charged bosons. If superconducting phases can be identified in other systems, progresses will be quick for the understanding of cuprate superconductivity, which still eludes compelling theoretical explanation of its intrinsic mechanism.

A system that is making rapid and impressive progress in terms of creating and controlling macroscopic quantum states is that of microcavity exciton-polaritons (see Ref. 2 for a review). These quasi-particles which combine properties of light (cavity photons) and matter (quantum well excitons) have been noted for their predisposition to accumulate in macroscopic number in a single or few quantum states. They have many advantages from a practical point of view, such as their 2D geometry, which allows straightforward manipulation by lasers impinging at an angle, and their short lifetime, which allows continuous monitoring of the system, reconstructing its internal dynamics also by angle-resolved spectroscopy. The pumping can be either coherent (driving states in parametric scattering configurations) or incoherent (with a constant flow of unrelated particles relaxing into the ground state). In nitride systems, the formidable claim has been made of room-temperature Bose–Einstein condensation. Recently, there has been great interest in propagation of polariton fluids and their superfluid properties, with reports of quantized vorticity and persistent currents. These rising stars of macroscopic coherence have also been proposed to realize another much sought after quantum phase at high-temperature: superconductivity. Polariton condensates cannot conduct electric current themselves, being neutral particles. One of the proposed implementations involves “quatrons” (or quadrions) rather than polaritons. Quatrons are bound states of two electrons with a polariton. They remain bosons from spin-addition rules but carry an electric charge. A Bose condensate of quatrons would, through its superfluid propagation, exhibit superconductivity. To date, however, the existence of the quatron, predicted theoretically, has not been found experimentally. Recently, we have approached the problem from another, and more conventional, angle, that of the Bardeen, Cooper, and Schrieffer (BCS) mechanism with an important new feature: replacement of phonons by Bose-condensed exciton-polaritons in the role of a binding agent between electrons. In the present paper, we generalize the model proposed in Ref. to describe a wide range of hybrid Bose–Fermi semiconductor systems where a Bose–Einstein condensate of neutral quasi-particles (excitons or exciton-polaritons) coexists with a Fermi sea of electrons. We show that indeed such systems are promising for observation of superconductivity. Moreover, as the superconducting gap and critical temperature are very sensitive to the concentration of bosons in the system and the latter may be controlled by direct optical excitation, light-induced superconductivity in semiconductor heterostructures appears to be possible. In this work we closely follow the BCS approach, generalized and adapted to the case of superconductivity mediated by a Bose–Einstein condensate. Solving the gap equation in this case turns out to be a nontrivial problem, requiring careful analysis.

1.1 BCS with a Bose–Einstein Condensate as a Binding Agent

BCS is a pillar of superconductivity theory, which relies on three main tenets:

1. Instability of the Fermi sea,
2. Existence of an attractive interaction,
3. Condensation of charged bosons.

These are the three insights that were mainly contributed by Cooper, Bardeen, and Schrieffer, respectively, and that they could assemble into the BCS theoretical edifice and exploit to reproduce strikingly or predict successfully most of the superconductivity phenomenology.
The first point follows from Cooper’s observation\textsuperscript{27} that an arbitrarily small attractive interaction between two electrons on top of the Fermi sea leads to a bound state (the Cooper pair), thanks to the truncation of the momentum space for states with wavevector \( k > k_F \) (Fermi wavevector). This is a general result, which follows from the Bethe-Goldstone equation for the two-electron problem.

The second point is the identification of an effective attraction between electrons that normally experience bare Coulomb repulsion. This attraction is attributed for conventional superconductors to an interaction through phonons, the Bardeen-Pines potential,\textsuperscript{28} that consists, vividly, of one electron wobbling the lattice at a first time, which affects another electron at a second time (and at a larger timescale, since the lattice dynamics is much slower than that of electrons).\textsuperscript{29} If the frequency of the lattice vibration is smaller than that of the propagating electron, the net effect results in an effective attraction [Leggett offers an insightful toy model of coupled oscillators to capture the essence of the interaction character (attractive or repulsive)].\textsuperscript{30}

The last point is the so-called BCS state, which is a coherent superposition of paired bound states and which brings the two-particle Cooper effect to a collective behavior of all electrons in the system.\textsuperscript{31}

With these three ingredients put together, the BCS theory is complete.\textsuperscript{32} For our purposes, points 1 and 3 will be regarded as fundamental and well-established features of (BCS type of) superconductivity. Point 2, which might appear a mere desiderata for point 1 to apply, leaves us room for identifying and designing new types of attractive potentials, optimizing the range of applicability and strength so as to obtain robust superconductivity (e.g., holding at high temperatures or high magnetic fields) or its manifestation in a new class of systems (in microcavities). Bardeen himself, with coworkers,\textsuperscript{33} investigated possibilities to engineer a more robust BCS state in a bilayer structure where excitons replace phonons as mediators of the interaction. The idea of substituting phonons by excitons was pioneered by Little\textsuperscript{34} and developed by Ginzburg,\textsuperscript{35} who coined the term and theorized the possibility of high-temperature superconductivity, much before it came to fruition with cuprates.\textsuperscript{36} Cuprates, however, exploit another (still unknown) mechanism different to BCS.\textsuperscript{37}

In the following, we revisit the Ginzburg mechanism, based on BCS, with emphasis on maximizing the strength of interaction between electrons, so as to maintain their binding, and therefore superconductivity, to higher temperatures. The text is organized as follows: in Sec. 2, we give a short overview of the BCS mechanism and the exciton (Ginzburg) mechanism, outlining the points of special interest in our case. In particular, we introduce the gap equation. In Sec. 3, we introduce our hybrid Bose–Fermi system configuration, its Hamiltonian and microscopic interactions, and the effective electron-electron Hamiltonian that results from a mean-field approximation for the condensate and the usual Frölich transformation. We obtain the shape of the effective electron-electron interaction \( U \), that we find to be quite different in character to the Cooper (square well) potential. We also consider possible variations of our scheme, namely, a microcavity with a condensate of exciton-polaritons and a condensate of indirect excitons in coupled quantum wells.\textsuperscript{38} The system of coupled quantum wells explored by several groups\textsuperscript{39,40} might be easier to realize and study (it does not need a cavity) and presents some interesting differences as compared to the polariton system. On the other hand, polaritons condense at much higher temperatures.\textsuperscript{18} In Sec. 3.2, we study the gap equation for a Bose condensate-mediated effective interaction. Because the potential is not positive-definite, the problem is not well-posed numerically. We propose a simplified potential and an approximate solution of the gap equation, which we motivate by studying its validity on well-established approximations. We obtain the critical temperature in this case. In Sec. 4, we give our conclusions and perspective on this new application of excitons and polaritons and discuss how to measure the effect experimentally.

## 2 BCS and Ginzburg Mechanisms

Superconductivity is a fundamental property of solids. At low enough temperatures, most metals superconduct. As the mechanism is rooted in quantum mechanics, temperatures are expected to be very low and indeed this is the case for all metals. After considerable theoretical efforts
from various groups, a compelling theoretical model was assembled by Bardeen, Cooper, and Schrieffer, the so-called BCS theory. The model provides the critical temperature:

$$k_B T_C = \Theta e^{-\frac{1}{2}}$$

(1)

where, in conventional superconductivity, $\Theta$ is the Debye energy $\hbar \omega_D$, and $g = N(0) V$ with $N(0)$ the density of electrons at the Fermi energy and $V$ the electron-phonon coupling strength. The Debye energy is, in good approximation, the maximum energy that can be carried by a phonon. The mechanism therefore relies heavily on phonons, as was realized empirically before the advent of BCS (good conductors, for instance, are bad superconductors, since $V$ is small; also through the isotope effect, which correlates critical temperature with mass of the crystal atoms, and thus with resonance frequency). The exponential form and the presence of $N(0)$ show that the effect is a collective one involving all electrons, which have formed a new phase of matter that cannot be approached perturbatively by the independent electron pictures (since $f(z) = e^{-1/2}$ has no Taylor expansion around zero). Based on the theory, and accumulated experience, it was widely accepted that critical temperatures would not exceed a few tens of Kelvins, since the Debye energy, which can be quite large in some systems (hundreds of Kelvins), is exponentially reduced. In all conventional superconductors $N(0) V \ll 1$ (the so-called weak-coupling regime).

Ginzburg made the obvious but daring assumption that to achieve higher critical temperatures—crucial for technical applications which can easily be understood to be momentous—it is enough to find a system where $\Theta$ and/or $g$ are increased. Replacing phonons by excitons, for instance, Ginzburg found that values $\Theta/k_B \approx 10^3$, $10^4$ K as well as $g \approx \frac{1}{3}$, $\frac{1}{2}$ are obtained, yielding temperatures of several hundred Kelvin. High critical temperatures have been later reported in cuprates and nowadays, temperatures as high as 125 K are obtained in systems such as Ti-Ba-Cu-oxide. Cuprate superconductivity does not appear to follow the BCS pattern. On the other hand, all attempts to date to realize the exciton mechanism or a variation of it have remained fruitless. This is this mechanism, rooted in BCS, which we consider in this text. Since it consists in substituting the phonons of conventional BCS—or the excitons of Ginzburg scheme—by a Bose–Einstein condensate (BEC) of excitons or exciton-polaritons, the starting point for its microscopic description starts with the same zero temperature gap equation:

$$\Delta_k = -\sum_{k'} U_{kk'} \frac{\Delta_{k'}}{2E_{k'}}$$

(2)

where $U_{kk'}$ is the effective interaction between electrons with wavevectors $k$ and $k'$ and energy $E$. The gap $\Delta$ can be identified as the macroscopic wavefunction of a Cooper pair, which is also the order parameter for the superconducting phase. If it is nonzero, the system is in the superconducting state. A realistic microscopic treatment of $U$ is very complicated. A simplified version is provided by the Jellium model, which is a toy model of a metal that gives predominance to electron-electron interactions, that is, in particular, the underlying crystal is approximated as a uniform (structureless) background (like a “jelly”) in which the interacting electron gas evolves under its own self-interactions and the overall charge cancellation of the background. A popular effective interaction is given by the Bardeen–Pines potential which is derived from the microscopic form of the electron-lattice interaction:

$$U_{BP}(\omega, q) = \frac{\kappa_0}{1 + \omega^2/\omega_T^2} \left[ 1 + \frac{\omega_{ph}(q)^2}{\omega^2 - \omega_{ph}(q)^2} \right]$$

(3a)

$$= \frac{\kappa_0}{\left( 1 + \frac{\omega^2}{\omega_{Te}^2} \right) \left( 1 - \frac{\omega^2}{\omega_{ph}^2} \right)}$$

(3b)

with $q$ the phonon wavevector. In Eq. (3a), $\omega_{Te}$ is the Thomas-Fermi screening parameter and $\omega_{ph}$ is the phonon dispersion. The first term between the brackets is bare Coulomb repulsion, and the
second term, which is frequency dependent, follows from the perturbative coupling to the lattice, tracing out the phonons.

The physical meaning of Eq. (3a) is at the heart of the phonon-mediated mechanism. The interaction is \( \omega \) dependent, which means, in Fourier transform, time dependent. This reflects the famous retardation effect in superconductivity. This effect is based on the strong difference between the electron Fermi velocity in metals and the sound velocity. Roughly speaking, a fast electron from the Fermi surface creates a slow phonon and goes away. After some time, another fast electron arrives and absorbs the slow phonon. The average distance between these two electrons remains of the order of 100 nm, the distance at which the screened Coulomb repulsion can be safely neglected. Due to the retardation effect, a weak phonon-mediated attraction of electrons wins over their Coulomb repulsion and provides formation of Cooper pairs at low enough temperatures.

3 Interaction Hamiltonian

A sketch of the structure we propose appears in Fig. 1(c). A QW is doped negatively (with density of electrons \( n \)) and is put in contact with another QW where excitons are formed and are stable. The first (lower) QW hosts the 2DEG which is to undergo superconductivity while the second (upper) QW hosts the condensate of excitons that is to mediate it. This structure can also be placed at the maximum of the optical field confined in a microcavity \(^{26}\) (formed by two Bragg mirrors facing each other). The latter configuration also has the advantage that two excitonic-QWs can sandwich the 2DEG-QW since in this case the condensate is delocalised in the entire structure thanks to the photon fraction of the polariton. This allows an increase by a factor of 2, or more if the multilayer structure is further repeated, of the densities achievable in this system. The drawback of microcavities is a short radiative lifetime of polaritons. In order to maintain the polariton condensate one needs to pump it resonantly by a high-intensity laser, which may lead to undesirable heating of the system. The alternative system which we consider is a condensate of long-living spatially indirect excitons in coupled QWs, separated by a thin but high barrier from a QW containing a two-dimensional electron gas (2DEG). This system is simpler to realize as it does not contain the optical cavity. The condensate of indirect excitons may be maintained by a low-intensity optical excitation. Moreover, indirect excitons in coupled QWs possess very significant dipole moments which strengthens their interaction with electrons from a 2DEG. However, Bose condensates of indirect excitons have been found only at temperatures below 1 K.\(^{38}\) while condensates of exciton-polaritons are now routinely produced at room-temperature.\(^{43}\) In the following, we compare the advantages of the two respective schemes. Most of the underlying model applies to both equally. We focus more particularly on the microcavity system since it is more general. The exciton system is recovered by decoupling the photons.

![Fig. 1](image_url)

Fig. 1 One of the possible designs to evidence exciton-polariton mediated superconductivity. (a) Polariton (thick blue solid line) and exciton (magenta dashed line) dispersions with schematic representation of scattering of the boson that mediates the interaction between electrons. Also shown in thin blue solid line is the renormalised dispersion \( E_{\text{bog}} \), essentially identical to \( E_{\text{pol}} \). (b) Corresponding scattering of two electrons on the surface of the Fermi sea, exchanging momentum \( q \) through scattering of a boson in panel (a). (c) detail of the sandwich structure, showing the doped well (below) containing free electrons and the well hosting the condensate (above) of excitons or of exciton-polaritons. This layer can be sandwiched in different ways.
In Fig. 1(a), the exciton (resp. polariton) dispersion is shown in dashed magenta (resp. solid blue). The difference between the bare polariton dispersion (thick blue) and the bogolon dispersion (thin blue) is very small over the range of exchanged momenta of interest (of the order of the Fermi wavevector). The condensate in both cases is at \( k = 0 \). Scattered particles at any wavevector between \(-2k_F\) and \(2k_F\) mediate electron-electron interactions on the Fermi sea, as sketched on Fig. 1(b). The model microscopic Hamiltonian is taken as:

\[
H = \sum_k [E_{\text{pol}}(k) \alpha_k^\dagger \alpha_k + E_{\text{el}}(k) \sigma_k^\dagger \sigma_k] + \\
+ \sum_{k_1,k_2,q} [V_C(q) \sigma_{k_1,q}^\dagger \sigma_{k_2,q} + X^2 V_X(q) \sigma_{k_1,q}^\dagger \sigma_{k_2,q}^\dagger \sigma_{k_1,q}^\dagger] + U a_{k_1}^\dagger a_{k_2}^\dagger a_{k_1} a_{k_2},
\]

(4)

with \( E_{\text{pol}}(k) \) and \( E_{\text{el}}(k) \) the polariton and 2DEG dispersions for the in-plane wavevector \( k \), respectively. In the exciton case (without the microcavity), it suffices to replace \( E_{\text{pol}} \) with \( E_{\text{ex}} \) in the above. \( V_X \) is the electron-polariton interaction, \( U \) the polariton-polariton interaction and \( V_C \) the electron-electron repulsion. We now consider these terms in turn. Parameters assumed are listed in Table 1.

### 3.1 Electron-Electron Interaction

In the original BCS mechanism, electron-electron repulsion is either neglected altogether or overcome by the attractive mechanism and not manifested outside of the attractive window. We take it into account here since it is a detrimental factor for binding and most of our concern for experimental realization is to optimize this value. The full form of the potential is given by the Yukawa potential:

\[
V_C(q) = \frac{e^2}{2\epsilon A \vert q \vert + \kappa},
\]

(5)

with screening constant \( \kappa \). We get rid of the momentum dependence by averaging the potential \( V_{\text{eff}}(\omega, q) \) over the Fermi surface (FS), where:

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<tr>
<th>Parameter</th>
<th>Meaning</th>
<th>Value</th>
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<tr>
<td>( \epsilon )</td>
<td>Permittivity</td>
<td>( 7\epsilon_0 \approx 6.2 \text{ As/(mV)} )</td>
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<tr>
<td>( \beta_e )</td>
<td>Electron reduced mass</td>
<td>( 0.22 \approx 1.15 )</td>
</tr>
<tr>
<td>( \beta_h )</td>
<td>Hole reduced mass</td>
<td>( 0.23 \approx 0.85 )</td>
</tr>
<tr>
<td>( L )</td>
<td>Distance between wells</td>
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</tr>
<tr>
<td>( \kappa )</td>
<td>Coulomb screening length</td>
<td>( \approx 1.2 \times 10^9 \text{ m}^{-1} )</td>
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<tr>
<td>( m_x )</td>
<td>Exciton mass</td>
<td>( (0.22 + 1.25)m_e \approx 1.3 \times 10^{-30} \text{ kg} )</td>
</tr>
<tr>
<td>( m_p )</td>
<td>Photon mass</td>
<td>( 10^{-5}m_e \approx 9.1 \times 10^{-36} \text{ kg} )</td>
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<tr>
<td>( 2g )</td>
<td>Rabi splitting</td>
<td>45 meV [0]</td>
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<tr>
<td>( X )</td>
<td>Hopfield coefficient (exciton weight)</td>
<td>( 1/\sqrt{2} ) [1]</td>
</tr>
<tr>
<td>( k_F )</td>
<td>Fermi wavevector</td>
<td>( 5 \times 10^{10} \text{ m}^{-1} )</td>
</tr>
<tr>
<td>( a_B )</td>
<td>Exciton Bohr radius</td>
<td>( 1.98 \times 10^{-9} \text{ m} )</td>
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<tr>
<td>( R_y )</td>
<td>Exciton Rydberg</td>
<td>32 meV</td>
</tr>
<tr>
<td>( d )</td>
<td>Dipole moment</td>
<td>4 nm [12 nm]</td>
</tr>
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(4)

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\]

(5)

with screening constant \( \kappa \). We get rid of the momentum dependence by averaging the potential \( V_{\text{eff}}(\omega, q) \) over the Fermi surface (FS), where:
The second electron scatters between states in 3D:

$$k_1, k_2$$

and $$k_{1}', k_{2}'$$ are the initial and final states of the electrons scattered with the exciton (polariton) from the condensate. The second electron scatters between states $$k_{1}', k_{2}'$$ such that $$k_1' - k_2' = q$$. The characteristic energy $$E(q)$$ is that of a bogolon (an elementary excitation of the condensate).

In 3D, the FS where these electrons that scatter lie, is the surface of a sphere, while in 2D, it is a circle (we will speak of surface in both cases). The vector difference is therefore joining the two end-points on the surface. If the potential has spherical symmetry ($$V(q) = V(|q|)$$), the average of all two vectors on the FS reduces to that where $$q$$ is pinned at one point of the surface (the south pole in Fig. 2) and runs over the FS. This average, in the particular choice of Fig. 2, is the usual polar integration with $$k_2$$ describing the surface as $$\theta$$ (and $$\phi$$ in 3D) are varied, with:

$$q^2 = 2k_F^2(1 + \cos \theta),$$  \hspace{1cm} (7)

from Al Kashi’s theorem, so that the average potential $$\bar{V}_{\text{eff}}$$ reads, in 3D:

$$\bar{V}_{\text{eff}}(\omega) = \int_0^{2\pi} \int_0^\pi V_{\text{eff}}(q, \omega)k_F^2 \sin \theta d\theta dq / N,$$  \hspace{1cm} (8)

where $$N$$ is the normalization, i.e., the same integral where $$V_{\text{eff}}$$ is replaced by unity. This gives, in 3D:

$$\bar{V}_{\text{eff}}(\omega) = \frac{1}{2} \int_{1}^{1} V_{\text{eff}} \left[ \sqrt{2k_F^2(1 + \cos \theta)}, \omega \right] d \cos \theta,$$  \hspace{1cm} (9)

$$= \frac{1}{4} \int_{0}^{4} V_{\text{eff}}(k_F\sqrt{\theta}, \omega) d\theta,$$  \hspace{1cm} (10)

where we integrate over $$\theta = 2(1 + \cos \theta)$$ since this is a natural variable in Eq. (7), and, in 2D:

$$\bar{V}_{\text{eff}}(\omega) = \frac{1}{2\pi} \int_{0}^{2\pi} V_{\text{eff}} \left[ \sqrt{2k_F^2(1 + \cos \theta)}, \omega \right] d\theta.$$  \hspace{1cm} (11)

Our system is 2D, in which case, from Eqs. (5) and (11):

$$\bar{V}_C = \frac{e^2}{4\pi \epsilon A} \int_{0}^{2\pi} \frac{d\theta}{\sqrt{2k_F^2(1 + \cos \theta) + \kappa}} = \frac{e^2}{2\pi \epsilon A} \ln \frac{2k_F - \sqrt{4k_F^2 - \kappa}}{2k_F - \sqrt{4k_F^2 - \kappa}}.$$  \hspace{1cm} (12)

When $$\kappa > 2k_F$$, both the numerator and the denominator become pure imaginary so their quotient remains real. When $$\kappa = 2k_F$$, $$\bar{V}_C = e^2 / \Phi^2 \backslash \epsilon A \backslash \kappa$$. This is plotted in Fig. 3. Since we are trying to maximize attraction, that is, minimize repulsion, systems with small screening length and large wavevectors should be favored (but these parameters play critically on other aspects of the mechanism and the optimum is not compulsorily $$k_F / \kappa > 1$$).
3.2 Electron-Exciton Interaction

The electron-exciton or exciton-polariton interaction is one of the most important ingredients of the mechanism, as it ultimately determines the shape of the effective potential. In the microcavity, an electron (from the 2DEG) interacts with a polariton (from the condensate) through its excitonic component, so this is really the electron-exciton interaction that is to be computed, weighted by the Hopfield coefficient (the excitonic fraction) $X$. Let us consider, therefore, the scattering of an electron in one of the parallel QW, separated by a distance $L$ from the QW with excitons. The matrix element of the direct interaction between excitons and electrons reads:

$$V_X(q) = \int \Psi_X^*(Q, r_e, r_h) \Psi^*(\mathbf{k}, \mathbf{r}_1) V(\mathbf{r}_1, \mathbf{r}_e, \mathbf{r}_h) \Psi_X(Q, r_e, r_h) \Psi(k, \mathbf{r}_1) d\mathbf{r}_1 d\mathbf{r}_e d\mathbf{r}_h,$$

where $\mathbf{r}_1$, $\mathbf{r}_e$, $\mathbf{r}_h$ correspond to the 2D coordinates of the 2DEG electron, the exciton electron and the exciton hole respectively. The 2DEG electron is described by a plane wave while the electron/hole in the condensate are assumed to be in the 1s bound state with plane wave center-of-mass motion:

$$\Psi(q, \mathbf{r}_1) = \frac{1}{\sqrt{A}} e^{i\mathbf{k}\mathbf{r}_1},$$

$$\Psi_X^*(Q, \mathbf{r}_e, \mathbf{r}_h) = \sqrt{\frac{2}{\pi A}} \frac{U_e(z_e) U_h(z_h)}{a_B} e^{iQ\mathbf{R}_e} e^{-e\mathbf{r}_e/2a_B},$$

where $\mathbf{R}_e = \beta_e \mathbf{r}_e + \beta_h \mathbf{r}_h$, $\mathbf{R}_h = \mathbf{r}_e - \mathbf{r}_h$ are in-plane coordinates of the center-of-mass of the exciton and relative coordinate of electron and hole in the exciton, $U_e(z_e)$ and $U_h(z_h)$ are normal to the QW plane electron and hole envelope functions, respectively. We also consider the existence of a dipole moment $d$ for the exciton, which can be intrinsic to the structure, because of spatial separation of electrons and holes in coupled QWs, or (in the case of microcavities) be induced by an internal piezo-electric field, or result from an externally applied electric field. To account for all these possibilities, one can consider the layers of electrons and holes in the exciton shifted in the $z$-direction with respect to the position of the center-of-mass by a distance $l = d/e \ll L$. The matrix element of the interaction is then computed to be:

$$V_{de}(q) = \frac{e^2}{2eA} \frac{e^{-ql}}{q} \left\{ \frac{1}{1 + (\beta_e q a_B/2)^2} - \frac{1}{1 + (\beta_h q a_B/2)^2} \right\},$$

$$+ \frac{e d}{2eA} e^{-ql} \left\{ \frac{\beta_e}{1 + (\beta_e q a_B/2)^2} + \frac{\beta_h}{1 + (\beta_h q a_B/2)^2} \right\}.$$

---

**Fig. 3** Average electron-electron repulsion (in natural units) in the QW. This should be made as small as possible, which is obtained for higher Fermi wavevectors (for a given screening length).
where Eq. (15a) is the direct electron-exciton interaction that exists even in the absence of a dipole moment of the exciton, and Eq. (15b) is the dipolar interaction. The direct interaction vanishes at small exchanged momenta, while the dipolar-induced one assumes its maximum value here of $2d/(2\varepsilon_0\varepsilon_A)$. Overall, the dipolar interaction is naturally much larger than the direct one, since the exciton is electrically neutral.* These facts are summarized in Fig. 4.

3.3 Polariton-Polariton Interaction

We treat the polariton-polariton interaction within the $s$-wave scattering approximation with strength $U = \alpha_B^2 R_{\text{ex}} X^4/\lambda$ (where $\alpha_B$ is the exciton Bohr radius, $R_{\text{ex}}$ the exciton binding energy and $\lambda$ the normalization area. $X$ is the exciton Hopfield coefficient, the square of which quantifies the exciton fraction in the exciton-polariton condensate). For interaction between bare excitons, $X = 1$. Exciton-exciton (polariton-polariton) interactions are repulsive, in general. They result in linearization of the elementary excitation spectra, the Bogoliubov dispersion, but its role is not crucial to the mechanism of superconductivity we discuss, since at the wavevectors of interest, the changes brought by this term are very small compared to the kinetic energy of noninteracting excitons (exciton-polaritons).

3.4 Effective Interaction

We now proceed to bring our microscopic model toward a form suited to study Cooper-pairing and superconductivity, that is, we apply the canonical Fröhlich transformation that will result in an effective BCS Hamiltonian. Just as in the case of phonons, we start by getting rid of polaritons. We assume a condensate is formed with mean population $N_0$. We do not consider which mechanism, coherent or incoherent, is responsible for creating and maintaining this state. We do assume, however, it is coherent and with a definite phase, so that we can apply the mean-field approximation $\langle a_k^\dagger + q a_k \rangle \approx \langle a_k^\dagger \rangle \langle a_k \rangle + \langle a_k^\dagger \rangle \langle a_k \rangle \approx N_0 \delta_{k,0}$ with $N_0$ the density of polaritons in the condensate. This allows us to obtain the following expression for the Hamiltonian, after diagonalizing the polariton part by means of a Bogoliubov transformation (that leaves the free propagation of electrons and their direct interaction, $H_C$, invariant):

$$H = \sum_k E_{\text{el}}(k) a_k^\dagger a_k + \sum_k E_{\text{bog}}(k) b_k^\dagger b_k + H_C + \sum_{k,q} M(q) a_k^\dagger a_{k+q} (b_{-q}^\dagger + b_q), \quad (16)$$

where $E_{\text{bog}}(k)$ describes the dispersion of the elementary excitations (bogolons) of the interacting Bose gas, which is very close to a parabolic exciton dispersion at large $k$:

*Note at this point that there is an error in Ref. 26 where only the direct exciton interaction has been taken into account with an incorrect power in the parenthesis, which led to an expression similar to the dipolar interaction. The direct interaction by itself turns out to be too small to evidence superconductivity with the parameters chosen in Ref. 26, therefore a dipole moment should be induced in this case, say by applying an external electric field, to restore the effect.
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\[
E_{\text{bog}}(\mathbf{k}) = \sqrt{\tilde{E}_{\text{pol}}(\mathbf{k})(\tilde{E}_{\text{pol}}(\mathbf{k}) + 2UN_0A)},
\]

where \(\tilde{E}_{\text{pol}}(\mathbf{k}) \equiv E_{\text{pol}}(\mathbf{k}) - E_{\text{pol}}(0)\) and with the renormalized bogolon-electron interaction strength:

\[
M(\mathbf{q}) = \sqrt{N_0A}\sqrt{\frac{E_{\text{bog}}(\mathbf{q}) - \tilde{E}_{\text{pol}}(\mathbf{q})}{2UN_0A - E_{\text{bog}}(\mathbf{q}) + \tilde{E}_{\text{pol}}(\mathbf{q})}}.
\]

The last term of Eq. (16) coincides with the Fröhlich electron-phonon interaction Hamiltonian, which allows us to write an effective Hamiltonian for the bogolon-mediated electron-electron interaction. This results in an effective interaction between electrons, of the type \(\sum_{\mathbf{k}, \mathbf{k}_1} V_{\text{eff}}(\mathbf{q}, \omega)\sigma_{\mathbf{k}}^c \sigma_{\mathbf{k} + \mathbf{q}} \sigma_{\mathbf{k}_1}^c + \mathbf{q}\sigma_{\mathbf{k}}^c \sigma_{\mathbf{k}_1} \sigma_{\mathbf{k}}\). The effective interaction strength reads \(V_{\text{eff}}(\mathbf{q}, \omega) = V_C(\mathbf{q}) + V_\Theta(\mathbf{q}, \omega)\), with:

\[
V_\Theta(\mathbf{q}, \omega) = \frac{2M(\mathbf{q})^2E_{\text{bog}}(\mathbf{q})}{(\hbar\omega)^2 - E_{\text{bog}}(\mathbf{q})^2}.
\]

Equation (19) recovers the boson-mediated interaction potential obtained for a Bose–Fermi mixture of cold atomic gases,\(^{47}\) in the limit of vanishing exchanged wavevectors. It describes the BEC induced attraction between electrons. Remarkably, it increases linearly with the condensate density \(N_0\). This represents an important advantage of this mechanism of superconductivity with respect to the earlier proposals of exciton-mediated superconductivity,\(^{33,34,48}\) as the strength of Cooper coupling can be directly controlled by optical pumping of the exciton-polariton condensate. The attractive potential is displayed for various exchanged energies in Fig. 5, as a function of \(\theta\), defining the exchanged momentum expressed directly through the angle \(\theta\) defined in the Fermi circle (cf. Fig. 2). As commented earlier, the negative part corresponds to attraction, and the potential alternates between repulsive and attractive character, obtained at different exchanged momenta. We do not want to keep track of such complicated wavevector dependence, and therefore will average the interaction over the Fermi sea. A notable feature is, for most values of \(\omega\), the presence of a pole \(\theta_0\), where \(E_{\text{bog}}[\sqrt{2k_F^2(1 + \cos \theta)}] = \hbar\omega\). As is seen in the figure, \(\theta_0\) separates the attractive part from the repulsive part. The average will bring the additional convenience of canceling such divergencies. We note as well that the spectrum of excitations of the exciton-BEC may be changed in the presence of the electron gas, so that their eventual dispersion may be different to \(E_{\text{bog}}\).\(^{49}\) This has no effect on the Cooper-pairing of electrons which we discuss here.

We therefore wish to perform the average

\[
U_0(\omega) = \int_0^{2\pi} V_\Theta(\mathbf{q}, \omega) d\theta.
\]

![Fig. 5 Effective electron-electron interaction as a function of \(\theta\), defining the exchanged momentum \(q = \sqrt{2k_F^2\sqrt{1 + \cos \theta}}\) on the Fermi sea at energies \(\hbar\omega = 40, 50, \text{ and } 60 \text{ meV}\), respectively. The potential is symmetric around \(\pi\). In the first case, the whole dependency (over \([0, 2\pi]\)) is shown, with a first zoom in the inset showing two poles and another the attractive region that is of small amplitude but extends over a large range. With increasing energies, the poles recede towards smaller values of \(\theta\) with a dominating effect of the repulsive (positive) energy. In the central panel, in dashed magenta, the regularized potential is superimposed.](image)
where $q = \sqrt{2k_E^2(1 + \cos \theta)}$, as seen previously. Since $V_A$ is symmetric around $\pi$ we perform the integral $\int_0^{\pi}$ only. The integral would be easily computed numerically if there were no pole. There are dedicated numerical methods to compute principal values numerically, but in our case, since the pole is first order, it is enough to isolate it analytically by defining

$$f(\theta) = (\theta - \theta_0) V_A[q(\theta), \omega],$$

so that

$$U_0 = \int_0^{2\pi} f(\theta) - f(\theta_0) \frac{d\theta}{\theta - \theta_0} + f(\theta_0) \ln \frac{2\pi - \theta_0}{\theta_0},$$

where $f(\theta_0) = \lim_{\theta \to \theta_0} f(\theta)$ and the first integral is regular (it is shown in Fig. 5 in dashed magenta). The integration (average) is then straightforward and produces the results shown in Fig. 6.

If we take instead of $E_{pol}$ a quadratic dispersion for the excitation

$$E_x(k) = \frac{\hbar^2k^2}{2m_x}$$

and assume all-excitonic interactions, $X = 1$, we can consider the same effect in the absence of a microcavity, relying on a purely excitonic (rather than polaritonic) BEC. In this case, the same procedure as detailed above leads to an effective potential as shown in Fig. 7. We kept all parameters the same for comparison except for the dipole moment, which we have taken three times as large ($d = 12$ nm). This corresponds to the spatial separation of electrons and holes in the

**Fig. 6** Exchanged-momentum averaged interaction $U_0(\omega)$ between electrons of the 2DEG, as $N_0$ is increased. It is dimensionless and is attractive (resp. repulsive) when negative (resp. positive). Densities $N_0$ are shown in units of $10^{12}/\text{cm}^2$. In insets (a) and (b), a zoom of the regions delineated on the left panel, firstly in the small energies (long time) range, where the attraction is seen to be repulsive for smaller densities, because of the Coulomb repulsion (shown in red), and secondly for the densities $\approx 10^{12}/\text{cm}^2$ in the area where the character of the interaction changes abruptly from attractive to repulsive (higher densities are shown in dotted lines).

**Fig. 7** Same as Fig. 6 (also in units of $10^{12}/\text{cm}^2$) but for the case of an exciton condensate. A larger dipole moment $d$ has also been assumed. The potential is different in character, much closer to Cooper's potential with larger attraction at longer times.
system of indirect excitons studied by Butov et al.\textsuperscript{38} The potential we obtain in the exciton case is very different in character from the polariton case, and is more closely related to the Cooper (conventional) shape of a square well, or the Bogoliubov potential including repulsive elbows.

From these potentials, one can proceed to solve the gap equation.

4 Gap Equation

4.1 Cooper Potential

The BCS gap equation (2) is easier to tackle as a continuous equation:

\[
\Delta(\xi, T) = -\int_{-\infty}^{\infty} U_0(\xi - \xi')\Delta(\xi', T) \tanh(\frac{E}{2k_B T}) \frac{d\xi'}{2E},
\]  

(24)

\[E = \sqrt{\Delta(\xi', T)^2 + \xi'^2}\]where we have also introduced a finite temperature \(T\) from the Fermi-Dirac distribution of elementary excitations.\textsuperscript{32} With the BCS approximation of a step potential, the gap equation at zero temperature simplifies to \(\Delta(\xi, T) = -U_0 \int_{-\infty}^{\xi - \hbar \omega_D} \Delta(\xi', T)/2E d\xi'\). If \(\Delta \gg \hbar \omega_D\), the \(\xi\) dependence in the integral boundaries can be neglected (or, coming back to Eq. (2), one sees that in the initial gap equation, \(\Delta_0\) is exactly constant if \(U_{kk'} = U_0\) and is zero otherwise). Thus, we can assume in this case the gap to be of the form:

\[
\Delta(\xi) = \begin{cases} 
\Delta(0) & \text{if } |\xi| \leq \hbar \omega_D, \\
0 & \text{otherwise}.
\end{cases}
\]  

(25)

In this case, simplifying \(\Delta(0)\) on both side of Eq. (24), we obtain:

\[
\Delta(0) = \frac{\hbar \omega_D}{\sinh(1/U_0)}.
\]  

(26)

Equation (26) is better known as its approximation when \(U_0 \ll 1\), in which case it takes the form of the famous BCS gap expression,

\[
\Delta(0) \approx 2\hbar \omega_D \exp(-1/U_0)
\]  

(27)

Solving exactly the gap equation calls for some numerical method. Equation (24) is a non-linear integral equation, of the type studied by Hammerstein, i.e.,

\[
\Delta(\xi, T) = \int K(\xi, \xi')f(\Delta(\xi'))d\xi'
\]  

(28)

where in our case \(K(x, y) = -U_0(x - y)/2\) and \(f[y, z] = z \tanh(\sqrt{y^2 + z^2})/\sqrt{y^2 + z^2}\). There are strong conditions of existence of non-trivial (nonzero) solutions when \(U_0 \leq 0;\textsuperscript{53,54}\) however, the case when the kernel \(K\) is not positive-definite, that is, in presence of repulsion,\textsuperscript{1} has been much less studied mathematically.\textsuperscript{55} Cooper’s potential, being always negative, falls in the category of potentials which admit a unique non-trivial solution, with a stable numerical technique to obtain it, namely, since the mapping is contractive, by iterations of the gap equation: an initial (nonzero) function \(\Delta_0\) is used to compute the rhs of Eq. (24), providing \(\Delta_1\) which is injected back until the function converges. The gap computed in this way for a given Cooper potential is shown in Fig. 8(a). As one can see, the BCS approximation [Eq. (26)] remains a rather coarse approximation, since the gap turns out in this case to be bell-shaped rather than being a step function. Surprisingly, \(\Delta(0)\) is however in much closer agreement with its approximation [Eq. (28)], as shown on Fig. 8(b) and 8(c) in the weak and strong coupling regime, respectively. There are small quantitative deviations in (b) between the numerical points and the formula when using the exact parameters of the potential. By fitting the numerical results, a perfect agreement can be found for slight variations of \(\hbar \omega_D\) and \(U_0\). The BCS approximation therefore turns out to be an exceedingly good one as compared to an exact solution of the gap equation. For the

\[\text{as well as attraction, since the case of only repulsion admits only } \Delta = 0 \text{ as a solution.}\]
procedure to make sense, similar results should be obtained for a smoothed well that approximates the BCS square well.\textsuperscript{41} We will not address this point here, but go directly to the case where the potential is not always attractive.

4.2 Bogoliubov Potential

The potential is not always attractive when, for instance, some overall repulsion, such as direct Coulomb interaction, is superimposed on the attractive Cooper potential, as shown on Fig. 9. The Coulomb interaction is time independent and should therefore extend to all $\omega$ but here also a cutoff $\omega_C$ is introduced to avoid divergencies. This results in an attractive, Cooper-like potential, flanked by two repulsive windows. Such a potential is known as the Bogoliubov potential.\textsuperscript{56,57}

This approximation has been used to show extremely counterintuitive behavior of the gap equation and justify a-posteriori another heavily criticized approximation of BCS, neglecting Coulomb repulsion: the BCS mechanism indeed assumes only attraction between electrons, which can be dimmed by Coulomb repulsion, but which never explicitly appears as such (like in the Bogoliubov scenario). The great result of Cooper was that binding occurs at arbitrarily small attraction. An important result of the Bogoliubov potential is to show that the detrimental effects of Coulomb repulsion are greatly reduced in the gap.\textsuperscript{57}

We now present a linearization of the gap equation that allows one to obtain an approximate solution for the critical temperature.\textsuperscript{56} By assuming the gap equation to be a two-step valued function $\Delta = (\Delta_1, \Delta_2)^T$, the gap equation becomes $(I - 1)\Delta = 0$ with

$$I = -\begin{pmatrix} -U_0 I_1 & V_C I_2 \\ V_C I_1 & V_C I_2 \end{pmatrix},$$

where

$$I_1 = \int_{-\hbar \omega_0}^{\hbar \omega_0} \frac{\tanh \left[ \frac{\xi}{(2 k_B T_C)} \right]}{\sqrt{\xi^2 + \Delta_1^2}} d\xi \approx \ln \left( \frac{1.13 \hbar \omega_0}{k_B T_C} \right).$$

Fig. 8 Numerical solution to the gap equation with the BCS square well potential (a). The gap $\Delta(\xi)$ is not step-wise as approximated in the BCS model, but its value at zero exchanged energy, $\Delta(0)$, is in close agreement with the analytical expressions, as seen in (b) and (c): $\Delta(0)$ as obtained from numerically solving the gap equation (dashed blue) and from Eq. (26) (solid magenta), (b) in the weak-coupling limit when $U_0 \ll 1$, with small deviations as coupling is increased, and (c) in the strong-coupling where the agreement becomes perfect again. Note that $\Delta$ increases linearly with the coupling strength out of the weak-coupling limit.

Fig. 9 Adding an overall repulsive Coulomb repulsion $V_C$ (red, left) until a cutoff $\omega_C$ to the BCS potential $V_0$ (green, left) results in the Bogoliubov step-wise potential (right). We take the convention $V_0$, $V_C$, $U_0$ positive in the above representation and $U_0 = V_0 - V_C$, so that, e.g., $V_C < 0$ means attractive contribution of the “Coulomb repulsion.”
In Eq. (30a), the BCS approximation has been applied while in Eq. (30b), the fact that \( |\xi| \gg 0 \) has been used to neglect \( \Delta \) in the denominator and the temperature in the numerator. The parametrization of the matrix in terms of the potential depends on the particular configuration (for example the relative widths of the various layers of the structure). Here we have adopted the original parametrization of Bogoliubov, which assumes narrow repulsive elbows surrounding a large attractive central region. Solving the linear equation for \( I_1 \) and then for the critical temperature, we find:

\[
k_B T_C \approx 1.13 \hbar \omega D \exp \left( \frac{1}{V_0 - V_C} \right). \tag{31}\]

In this form, one can see how Coulomb repulsion indeed introduces a small correction to the original BCS formula. The expression also seems to indicate that \( U_0 \) could be repulsive \((V_0 - V_C < 0)\) and still lead to a gap as long as the denominator in Eq. (31) remains positive.

In the following, we compare these predictions with numerical solutions of the gap equation, keeping in mind that the iterative procedure is not assured, mathematically, to converge. We have observed that indeed, it sometimes encounters problems and exhibits strong instabilities, with bifurcations of solutions, for example.

In Fig. 10, we show the evolution of the gap function as \( V_C \) is increased, from zero (BCS) to a point where the overall potential is essentially repulsive. With onset of the repulsion, the gap acquires two negative sides and becomes a highly distorted function for large \( V_C \). Note that the approximation of constant gap over the various regions is at least as good as for the case of BCS.

In Fig. 11, we now show the case where \( U_0 = V_0 - V_C \) is held constant as the strength of the repulsion is varied independently. In foresight of what is to come later, we also allow the elbows to be negative, that is, to contribute an additional attraction to the conventional mechanism (for now we do not consider physical justification of this). Another unexpected result is obtained: the repulsive potential is in this case favoring a larger gap, as can be seen by comparing (a), where the gap function is highly oscillatory, to (d) where it recovers the BCS bell-shape. In (h), the distorted but overall attractive potential still results in a BCS type gap, but wider and larger. Note how the repulsion, by “squeezing” the gap, allows it to achieve much higher values than for the case of smaller or no repulsion.

These unexpected results are confirmed phenomenologically by the Bogoliubov approximation Eq. (31), which reproduces qualitatively the trend of the gap \( \Delta(0) \) as shown in Fig. 12. Here we should emphasize that the two quantities are not meant to be compared quantitatively, since one, \( \Delta(0) \) (computed numerically) is the gap at zero temperature while the other, \( k_B T_C \) is the temperature at which the gap vanishes. There is a monotonous relationship between the two, that is, increasing \( \Delta(0) \) implies increasing \( T_C \), so one can appreciate the consistency of the results by observing similar trends. The obstacle to conducting an extensive numerical comparison is that it is an intensive task numerically to compute \( T_C \), since this requires solution of the gap equation for various temperatures until the curve \( \Delta(T) \) is obtained and its intersect with zero is found.

\[\text{Fig. 10 Gap (thickness magenta and, magnified, thickness khaki) of the Bogoliubov potential (filled blue) solved numerically for the parameters: } h\omega_D = 1, \ h\omega_C = 2, \ V_0 = 1 \text{ and } V_C \text{ taking values from (a) to (d) of: } 0 \text{ (BCS), 0.1, 0.3 and 0.495. Coulomb repulsion results in a dip in the gap function, that, for increasing values, results in oscillations in the gap function. Paradoxically, even when it is large and dominating attraction, repulsion does not prevent a gap, as seen in (d).} \]
A critical slowing down phenomenon makes the iterative process slower as the critical temperature is approached. In addition, numerical instabilities are stronger at nonzero $T$. Therefore, although it is relatively straightforward to compute $\Delta(0)$ numerically, it is not convenient to use this method to obtain $T_C$. On the other hand, the Bogoliubov approximation gives a fair estimate of $T_C$, but is not able to provide the gap at zero temperature, since at the core of its method, there is an assumption of vanishing $\Delta$. Therefore, we have two complementary methods, each suited to provide a relevant aspect of the problem. We note that the gap at zero temperature is an important quantity which can be measured independently from $T_C$ by Andreev reflection in conductivity experiments.

### 4.3 Polariton Potential

In the case of the polariton problem, we have seen that, even when neglecting Coulomb repulsion (as in the original BCS formulation), the potential $U(\omega)$ departs strongly from the Cooper potential and features two large attractive regions far from small energies, immediately followed by two strong repulsive windows. We extend the Bogoliubov method to a three-step approximation of this potential, such as displayed in Fig. 13, with, in reference to previous potentials, notations $\omega_D$, $\omega_C$ and $\omega_R$ for the boundaries of the central, shallow attractive region, narrow, deep attractive region and repulsive region, respectively. This is a notation only and is not mean to be understood as referring to Debye, Coulomb, or Bogoliubov in any strict sense. Following the same premises, we approximate the gap equation by a three-step valued function $\Delta = (\Delta_1, \Delta_2, \Delta_3)^T$. This approximation turns out to be an exceedingly good one in certain cases, such as the one displayed in Fig. 13, where the gap itself is also a three-steps function in good approximation. Here there is even more room to choose a parametrization of the $I$ matrix. We now give general guidelines on how to build this matrix. The simplest method is to fix $\xi$ on the lhs of Eq. (25) at the
center of each region and, in the corresponding row, take for each column the potential that is sampled more by the difference $j\xi - \xi_0^j$. Refinements are possible, such as weighting elements of $I$ by coefficients which reflect how much time the variables $\xi$ and $\xi_0^j$ spend in the regions that determine the matrix equation. This problem has the following mathematical expression: how is the random variable $X - Y$ distributed when $X$ (resp. $Y$) is uniformly distributed in an interval $[g_l, g_{l+1}]$ (resp. $[h_l, h_{l+1}]$). The solution is easily obtained as proportional (normalize to unity) to:

$$P(X - Y = \theta) \propto \sqrt{[\max(g_l, h_l - \theta)] - \min(g_{l+1}, h_{l+1} - \theta)]^2 + [\max(\theta + g_l, h_l) - \min(\theta + g_{l+1}, h_{l+1})]^2.}$$

(32)

This is easily obtained geometrically (the square root comes from Pythagoras’ theorem) and the problem results in finding the intersect of a line with the grid. There are two configurations. Working out the cases shows that Eq. (32) reduces to a triangular or a top-head truncated triangular distribution. The coefficients entering $I$ can then be taken as the potentials weighted by the area intersecting their corresponding region. The soundness of such an approach can be checked numerically to compare quantitatively various parametrizations. For simplicity, we shall here consider cases where only the dominant potential is considered. An example of such a gap equation $(I - 1)\Delta = 0$ is defined with:

$$I = \left( \begin{array}{ccc} V_0 I_1 & V_1 I_2 & -V_C I_3 \\ V_1 I_1 & V_0 I_2 & V_0 I_3 \\ -V_C I_1 & V_0 I_2 & V_0 I_3 \end{array} \right),$$

(33)

where

$$I_i = \int \frac{\hbar \omega_i^c \tanh [\hbar \omega_i^c / (2k_B T_C)]}{\sqrt{\omega_i^c + \Delta_i^2}} d\xi_i,$$

(34)

and $I_i$ is integrated on the respective steps, as defined by the integral boundary conditions, i.e., $(\omega_1^c, \omega_2^c, \omega_3^c, \omega_4^c, \omega_5^c) = (0, \omega_D, \omega_D, \omega_C, \omega_C, \omega_D)$. As before, we estimate the critical temperature by the condition that gives vanishing values of the gap on the whole interval. The first integral $I_1$ is evaluated in the same approximation as for the usual BCS (or Bogoliubov) potential:

$$I_1 = \ln \left( 1.13 \frac{\hbar \omega_D}{k_B T_C} \right).$$

(35)

Fig. 13 (solid blue) Three-step potential that models the form of effective electron-electron interaction when polariton-mediated with the three parameters $\omega_B, C, D$ and, (thick magenta) the gap $\Delta(\xi)$ as computed numerically for this potential. In good approximation, the gap is also itself a three-steps function.
\(\mathcal{I}_2\) and \(\mathcal{I}_3\) are also essentially logarithmic in their energy range and we take:

\[
\mathcal{I}_2 = \ln(\omega_C/\omega_D), \quad \text{and} \quad \mathcal{I}_3 = \ln(\omega_B/\omega_C).
\]

(36)

We solve the gap equation by setting its determinant to zero and solving for \(T_C\), giving

\[
k_B T_C = 1.13\hbar \omega_D \exp \left[ -\frac{1}{V_0} \ln \left( \frac{\omega_C}{\omega_D} \right) \ln \left( \frac{\omega_C}{\omega_B} \right) \frac{V_1 + V_C}{V_0} \left( \frac{\omega_B}{\omega_C} \right)^2 \right].
\]

(37)

We see that the impact of the polariton potential shape on the gap is rather intricate, here as well, with some unexpected behavior, produced both numerically and from this formula. In Fig. 14, we show the effect of widening the central, attractive region of the potential. Naively one would expect this to increase the gap (or critical temperature), since \(\Delta(0)\) increases linearly with \(\hbar \omega_D\). However, Eq. (37) predicts a decrease of the critical temperature, as shown in Fig. 15. The reason why is understood by computing the gap at zero temperature, as seen in Fig. 14, where the repulsive barrier is shown to behave as a trap for the gap function, which results, as the width is increased, in a loosening of the gap strength, akin to the quantum mechanical situation of a bound state in a square potential. The numerical and linearized models display a good qualitative agreement, as seen in Fig. 15. Let us repeat that we are not considering here quantities that can be directly compared, since \(\Delta(T_C)\) should be used for this purpose.

Another example is shown in Fig. 16, where both the attractive and repulsive parts of the potential are increased together. This results in a rapid increase of the gap. The result is also confirmed by numerical simulations, as shown in Fig. 17, and a good agreement is obtained. Unexpectedly, if the repulsive part grows twice as fast as the attractive one, not only does this also result in an increase of the gap, but even a faster one. This, too, is confirmed by the numerics. Note that in this case, we reach a region where our numerical procedure jumps to

![Fig. 14](http://nanophotonics.spiedigitallibrary.org/)

Fig. 14 Potential (filled blue) and its corresponding gap equation, solved numerically, as the size of the central attractive plateau is increased. Somewhat unexpectedly, the gap decreases as a result. The repulsive barriers have the effect of squeezing the gap up.

![Fig. 15](http://nanophotonics.spiedigitallibrary.org/)

Fig. 15 \(\Delta(0)\) (joined points, blue) and \(k_B T_C\) (magenta) from the three-step Bogoliubov approximation, in the case where the width of the central attraction is increased. Unexpectedly, this results in a decrease of the critical temperature and of the gap.
other solutions (usually of a highly oscillatory character, such as those reported in \cite{57}), but behaves as expected until then. In this regard, the value of having an approximate analytical solution is obvious.

This analysis is finally applied to the case of the gap equation with the numerical results of the critical temperature obtained in both the polariton and exciton cases for the cases of Figs. 6 and 7. The parameters are gathered in Table 1 and the results plotted in Fig. 18. Both cases show a strong variation of the critical temperature with moderate variations of the condensate density (one order of magnitude). The polariton case is steeper and roughly linear while the exciton case increases less quickly but begins earlier. In both cases, temperatures are very high, and other effects will surely break the mechanism, for instance the loss of the condensate. This shows, however, the robustness of the mechanism in the conditions where it should apply.

5 Conclusions

We have studied possible mechanisms of superconductivity in semiconductor heterostructure systems where a Bose–Einstein condensate mediating the effective electron-electron interaction...
leads to enhancement of the coupling, yielding very high critical temperatures. We have considered the case of an exciton-BEC, consisting of a sandwich of an $n$-doped QW containing the superconducting electrons, in contact with coupled QWs where the BEC of indirect excitons is formed, for instance by optical excitation. We have also considered the case of a polariton BEC, where $n$-doped QWs and undoped QWs hosting excitons are embedded in a microcavity. We have computed in these two cases the effective electron-electron interaction $U(\omega)$ as a function of the exchanged energy $\hbar \omega$, showing how the retardation effect acquires a peculiar character in the polariton case, namely, yielding a weakly attractive potential at long times, followed by a succession of strongly attractive and strongly repulsive windows. The gap equation in this case exhibits strong differences as opposed to the case of the Cooper potential (of conventional BCS and exciton-BEC mechanism). To understand the physical mechanism leading to large gaps and bypass numerical instabilities, we studied in detail the gap equation in the cases of simplified step-wise potentials, and offered an analytical method to obtain the critical temperature, which is in qualitative agreement with the numerical results. Our results suggest record-breaking critical temperatures in these systems. We stress, however, that the path toward achievement of exciton-mediated superconductivity at high temperatures may be long and full of obstacles. Of the two experimentally relevant systems that we have considered, one (coupled QWs) only shows the BEC of excitons at very low temperatures (less than 1 K). Consequently, one cannot expect high-temperature superconductivity in this system, although at low temperatures the superconducting gap may be very large. In microcavities, polariton BEC or polariton lasing have indeed been demonstrated at room-temperature. However, embedding a high-quality $n$-doped QW inside the cavity, the proper choice of the experimental geometry in order to minimize optical absorption in the doped QW, and especially fabrication of quantum contacts for selective injection of carriers in the QW of interest, may pose technological difficulties. It is possible that hybrid metal-semiconductor or semi-metal-semiconductor systems may appear more suitable for the observation of the predicted effects. A conventional superconductor put in contact with a semiconductor also seems promising. Collective quantum phenomena in Bose–Fermi mixtures are extremely complicated and we foresee breakthroughs in their study in multilayer structures combining a Fermi gas of electrons and an exciton-BEC.

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