The effect of impurities on spin-polarized Zeeman bound states in dilute magnetic semiconductor-superconductor hybrids

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We investigate the effect of single and multiple impurities on the Zeeman-localized, spin-polarized bound states in dilute magnetic semiconductor hybrid system. Such bound states appear whenever a dilute magnetic semiconductor showing giant Zeeman effect is exposed to an external magnetic field showing nanoscale inhomogeneity. We consider the specific example of a superconductor-dilute magnetic semiconductor hybrid, calculate the energy spectrum and the wave functions of the bound states in the presence of a single impurity, and monitor the evolution of the bound state as a function of the impurity strength and impurity location with respect to the center of the Zeeman trapping potential. Our results have important experimental implications as they predict robust spin textures even for ideal samples. We find that for all realistic impurity strengths the Zeeman bound state survives the presence of the impurity. We also investigate the effect of a large number of impurities and perform ensemble averages with respect to the impurity locations. We find that the spin-polarized Zeeman bound states are very robust, and they remain bound to the external field inhomogeneity throughout the experimentally relevant region of impurity concentration and scattering strength. © 2010 American Institute of Physics.

I. INTRODUCTION

Due to its potential use in spintronics and quantum information applications, the controlled manipulation of individual spins has been a subject of great interest in recent years. In the search for appropriate materials for device fabrication, one possible route is the use of III-V or II-VI diluted magnetic semiconductors (DMS).

In II-VI DMSs, the exchange interaction between band charge carriers and the localized magnetic spins of the magnetic dopant (such as Mn) gives rise to a giant splitting between band states with different spin components. This giant Zeeman spin splitting has been extensively investigated and characterized.1,2 Given the linear dependence of this splitting on the external applied magnetic field for fields lower than roughly 1 T, it can be described in terms of a very large effective g-factor for the band carriers. For example, Dietl et al.3 reported an electron effective g-factor of about 500 at sub-Kelvin temperatures in a CdMnSe sample, which implies a value of about 2000 for the effective g-factor of a hole in this material. In previous works,4–8 we proposed to take advantage of this large effective g-factor by combining it with a spatially inhomogeneous applied magnetic field, to create a spatially varying Zeeman potential that acts as a confining potential for only one spin orientation. In suitable conditions, this leads to a single charge carrier with a well defined spin being trapped in the region of large local magnetic field. This spin-polarized charged carrier can then be manipulated through external control on the applied inhomogeneous magnetic field.

The needed nanoscale inhomogeneous magnetic field can be generated in various ways. One possibility is to use nanoscale magnets placed above a DMS quantum well (QW).4–6,9,10 Depending on the shape and orientation of the nanomagnet, different nonuniform fields are generated, giving rise to various types of confined states.1,2,11

Another possibility is the use of Abrikosov vortices that appear in type-II superconductors (SCs). Above the lower critical field, $B_{c1}$, vortices populate the SC, forming a vortex lattice. Nanoengineering can be used to ensure that the vortices nucleate in well defined positions.3,11 The field of a single vortex is nonuniformly distributed around a core of radius $r \sim \xi$ ($\xi$ is the SC coherence length) and decays away from its maximum value at the vortex center over a length scale $\lambda$ ($\lambda$ is the penetration depth). If a SC layer that hosts such vortices is placed above a DMS layer (QW), the inhomogeneous magnetic field of the SC vortices creates an inhomogeneous magnetic field in the DMS layer. According to our previous calculations,7,8 these fields are sufficiently large to result in the confinement of band carriers with a given spin orientation in the small region of the DMS QW that is located directly under a SC vortex core. Thus, spin textures are formed and can be manipulated by moving the source of the magnetic field, i.e., the SC vortex. In other words, the SC vortices act as spin and charge tweezers and can be used for a wide array of applications, from investigation of the Hof-
stader butterfly and making spin shuttles\textsuperscript{7} to generating anions of interest for topological quantum computing.\textsuperscript{15} The inhomogeneous field can also be generated by non-Abrikosov vortex structures of SC, for example, a SC layer with holes, blind holes, or magnetic nanodots on top.\textsuperscript{4,3,7,13}

In our previous work, we assumed that the DMS QW is perfectly clean and with Mn impurities distributed in a perfectly homogeneous fashion (for instance, by digital doping\textsuperscript{14}), so that the Zeeman potential is a smooth function mirroring the applied magnetic field. For such systems, we obtain typical binding energies of the spin-polarized charge carrier on the order of a few meV.\textsuperscript{7,8} As we show in this paper, such perfect homogeneity is not necessary. This is a very important result from experimental detection and application point of view, since it opens up the possibility of investigating Zeeman localization effects in samples grown during standard molecular beam epitaxy (MBE) conditions.

Since the effective g-factor is proportional to the local concentration of Mn, inhomogeneities in their distribution will result in a spatially varying $g(\mathbf{r})$ which will modify the trapping potential. Even more detrimental are charged impurities which exist in any sample. Charge dopants are needed to introduce charge carriers in the II-VI DMS because the Mn is isovalent with the valence-II element it replaces. The disorder due to these charged dopants can be minimized in the usual way used for two-dimensional (2D) electron systems, by doping at some distance away from the DMS QW and using gates to control the concentration of charge carriers in the DMS QW. However, small concentrations of undesired charged dopants in the DMS QW cannot be avoided.

In this article, we investigate the role played by charged scattering centers on the confinement of spin-polarized carriers in the DMS QW. We focus on repulsive centers, which are expected to be most detrimental to the binding of the spin-polarized carrier in the Zeeman trap, and then briefly discuss the other types of scattering centers mentioned above. We show that due to the difference between the length scale of the confining potential and the repulsive scattering potentials, the bound-state energy is not significantly changed even in the presence of a large number of impurities. This result indicates that the binding of spin-polarized charge carriers in Zeeman traps created in a DMS QW is very robust against such defects, with immediate implications for spintronics applications based on this scenario.

It is important to also emphasize that our conclusions apply to a much wider range of systems than the one of interest to us because these conclusions are based on quite simple physics, as we show below. We argue that in any system (such as quantum dots and QWs) where localization occurs on a lengthscale much larger than that of typical scattering potentials, the effect of such scattering centers is very small even if their potential is very large.

This paper is organized as follows: In Sec. II, we describe in more detail the system we study, our theoretical approach, and the approximations we have made. In order to build intuition and to help interpret later calculations, in Sec. III, we analyze the results for a simplified one-dimensional (1D) model in the presence of a single impurity. This allows us both to gauge our computational scheme against exact solutions and to gain some intuition about the effect of the scattering centers. In Sec. IV, we repeat the analysis for a single scattering center for the 2D of interest to us and discuss the influence of the symmetries in our results. Finally, in Sec. V, we present the results for a random distribution of impurities and various impurity concentrations. We conclude in Sec. VI with a summary and discussion.

II. THEORETICAL MODEL

The hybrid structure of interest to us is sketched in Fig. 1(a) and consists of a SC layer in the vortex phase placed above a DMS QW. The two materials are separated by a thin insulating layer. The SC can be Pb or Ni, for example, both these metals can be grown on top of semiconductors using MBE techniques.\textsuperscript{15} The QW consists of a weakly p-doped DMS. An example of such a DMS QW is a p-doped (Cd,Mn)Te well, doped with N by a modulation doping technique that avoids inhomogeneity effects and increases the mobility of the carriers.\textsuperscript{16} The carriers of interest to us are confined in the 2D DMS QW. The giant Zeeman effect in the DMS is described by a very large g-factor. This combines with the inhomogeneous magnetic field generated by each vortex to give rise to an effective potential that binds a spin-polarized hole under each vortex, in a clean sample. As stated, we are now interested in the effect of repulsive scattering centers on these spin-polarized bound states. A typical potential well in the presence of a single and of several charged scatterers is illustrated in Figs. 1(b) and 1(c), respectively.

In a parabolic approximation, the Hamiltonian for a charge carrier in the 2D DMS QW which interacts with the magnetic field of the vortex and with scattering centers is

$$H = \frac{[p - q\vec{A}(\mathbf{r})]^2}{2m} - \frac{1}{2}g_{\text{eff}}\mu_B\vec{B}(\mathbf{r}) \cdot \vec{B}(\mathbf{r}) + \sum_i V_i(\mathbf{r}),$$

where $m$ and $q$ are the effective mass and charge of the carrier, $\vec{B}(\mathbf{r})$ is the magnetic field generated by a single vortex, $\vec{A}(\mathbf{r})$ is its vector potential, and $V_i(\mathbf{r})$ are the repulsive potentials from the $i$ scattering centers. For simplicity, we assume that the motion of the carrier in the QW is 2D. In the calculations shown here, we use $m = 0.5m_e$ and $g_{\text{eff}} = 500$. 

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig1}
\caption{(Color online) (a) Illustration of the magnetic field generated by a single vortex in a hybrid system made of a type II SC on top of a DMS QW which contains impurities. (b) Potential profile of the Zeeman trap plus a single repulsive impurity and (c) the same, but for a random distribution of impurities (figures not to scale).}
\end{figure}
these being typical values for holes in II-VI semiconductors. Because of the large $g_{\text{eff}}$, for a single vortex, the effect of the $A(x,y)$ term is negligible compared to that of the Zeeman interaction and we ignore it from now on.\(^7\)

The experimental results show that the Zeeman splitting for holes in a DMS is anisotropic, depending on the direction of the magnetic field with respect to film plane.\(^17\) Using a Luttinger Hamiltonian,\(^18\) one can also show that in a 2D QW, the $g_{\text{eff}}$ of heavy holes is highly anisotropic, with an in-plane component much smaller than that perpendicular to the film plane.\(^6\) With these results in mind, we can further simplify our problem by considering only the effect of the $z$ component of the magnetic field, which is most strongly coupled to the charge carrier. This approximation allows us to decouple the Hamiltonian in the spin-up and spin-down sectors, and consider them separately. We will only focus on the spin component for which this Zeeman potential acts as a trap.

As discussed extensively in Ref. 7, the size of the confining region defined by the inhomogeneous magnetic field depends on the SC’s parameters and the distance between the SC and DMS layers. It is typically several tens of nanometers in size. On the other hand, the impurity potential of the scattering centers is considerable only in a much smaller range of a few angstroms in the immediate vicinity of the impurity. It is the effect of this strong repulsive potential on the bound state that is of interest to us, since the long-range Coulomb repulsion is too weak to be relevant (although its effects can also be studied with our method, if so desired).

Given the large difference between the characteristic length scales, we model the scattering potential as a delta function. After all these approximations, the Schrödinger equation for the trapped spin component is

$$\left[\frac{p^2}{2m} - g_{\text{eff}}\mu_B \frac{\sigma}{2} B_z(\vec{r}) + a \sum_i \delta(\vec{r} - \vec{R}_i)\right] \psi_\alpha(\vec{r}) = e \psi_\alpha(\vec{r}),$$

where $a$ characterizes the strength of the impurity potential and $\vec{R}_i$ is the location of the $i^{th}$ impurity.

We solve for the eigenvalues in the usual way, by expanding the wave function in a complete basis set and finding the appropriate coefficients from a matrix equation. Our basis functions are B-spline polynomials on a nonuniform knot sequence adjusted optimally for each specific arrangement of charged impurities. This method has been widely used in atomic physics\(^19\) and is ideally suited for problems such as this. Details of the method and its advantages are discussed in the Appendix.

### III. 1D MODEL

We first study a 1D model with a single delta function impurity inside a square well potential. This provides an intuitive understanding of the physics we want to explore, and also a numerical test of our B-spline scheme since it permits the comparison between the numerical results and the exact solution.

We consider a 1D square well potential

$$V_0(x) = \begin{cases} -a_0 & \text{if } -L/2 \leq x \leq L/2 \\ 0 & \text{otherwise}, \end{cases}$$

where $L$ is the lateral size of the well. To it, we add a single impurity potential

$$V_1(x) = a \delta(x - x_0),$$

where $x_0$ is the impurity position. In this case, the Schrödinger equation is given by

$$-\frac{\hbar^2}{2m} \frac{d^2}{dx^2} \psi(x) + V_0(x) + V_1(x) = e \psi(x).$$

To understand the effects of the impurity on the bound states for different strengths of the impurity potential, we calculated the eigenvalues of this Hamiltonian. The energy of the lowest two bound eigenstates is shown in Fig. 2 versus the strength of the impurity potential expressed as a dimensionless quantity $\alpha=a/(a_0 L)$. The parameters are $L=100$ nm and $a_0=3.5\hbar^2/(mL^2)$, and the impurity is located in the center of the well, $x_0=0$. Figure 2 reveals that the ground-state energy first increases with $\alpha$ and then saturates to the value of the first excited state. On the other hand, the energy of the first excited state is independent of $\alpha$. The reason for this behavior is straightforward to understand. Since the first excited state has a node at the location of the impurity, its energy and wave function are naturally unaffected by its presence. On the other hand, the original ground-state wave function has a maximum at the impurity location, so its energy increases with $\alpha$. However, for large enough $\alpha$, the ground-state wave function also develops a dip precisely at the location of the impurity and becomes insensitive to its strength. This is demonstrated by the insets of Fig. 3, which show the ground-state wave function at $\alpha=0$ and $\alpha=100$. In essence, when $\alpha \gg 1$, the impurity potential splits the original well into two isolated wells which are degenerate, given the symmetry of the problem.

When the impurity is placed off-center, as in Fig. 3, the system loses this symmetry and the $\delta$ function separates the original well into two distinct wells of different sizes. Since the impurity is now placed in a location where both the origi-
The ground-state and the first excited-state wave function were finite, both energies now increase with $\alpha$ when $\alpha < 1$. At higher values of $\alpha$ both the ground-state and excited-state energies saturate. As shown in the insets, the wave functions develop dips at the location of the impurity and become progressively more localized on one or the other smaller wells. The new ground-state wave function becomes confined to the larger new well. These results agree with those of previous work on a mathematically similar problem arising in a very different context.

While these results are very easy to understand, they do provide key insight into the effects of the impurity on the bound states. First, it is clear that symmetry plays a very important role. While this influence is maximized by the 1D character of the model, we expect analogous features in the 2D case.

More importantly, we already see that even if the impurity potential is large, its overall effect on the bound-state energy and wave function is limited. Because it is extended over a much larger area, the bound wave function simply avoids the impurity by developing a dip (or a zero) in its vicinity, at relatively low energetic cost. In 1D the impurity effectively “cuts” the wave function. This will not happen in $d=2$ or higher dimensions. However, as we demonstrate below, the limited effect on the binding energy is a general feature observed in higher dimensions.

The results shown so far also illustrate why B-splines are an ideal basis for such problems. If one tried to use a more traditional basis of orthonormal polynomials such as harmonic oscillator eigenstates, one would have to mix in very many basis states in order to be able to accurately describe smooth wave-function variations on a short length scale, near the impurity, as shown in the right inset of Fig. 2. Describing regions where the wave function essentially vanishes, as shown in the right inset of Fig. 3, would be even more difficult. On the other hand, use of B-splines allows one to just slightly increase the number of basis states by sampling the vicinity of the impurity on a smaller mesh. As a result, the calculation with one or more impurities is very comparable, in terms of numerical computational costs, with the one in the absence of impurities. A more detailed discussion of these issues is given in the Appendix.

IV. 2D ZEEMAN POTENTIAL WITH A SINGLE IMPURITY

We now can consider the 2D system in the presence of a single charged impurity. As we are not interested in the details of the Zeeman potential, we take the magnetic field generated by the vortex to have a Gaussian profile. This approximation is reasonably close to the vortex core, where the magnetic field decays exponentially on a length scale set by $\xi$. We therefore solve Eq. (2) where the magnetic field profile is taken to be

$$B_z(\vec{r}) = B_0 \exp\left(-\left(\vec{r} - \vec{r}_0\right)^2\xi^2\right),$$

where $\vec{r}_0$ is the location of the vortex core, and $B_0 = 0.206$ T and $\xi = 50$ nm are the strength and the range of the magnetic field (these are typical values for a dirty Pb film in a type II regime). We solve the equation numerically for a system of finite size $200 \times 200$ nm$^2$, with periodic boundary condition and a grid of $51 \times 51$ knots. This large lateral size is chosen so as to eliminate the finite size effects in our bound-state energies. For more details about the implementation of this numerical method and the periodic boundary conditions, see the Appendix.

We follow a procedure similar to the one in Sec. III and begin with a single impurity located at the center of the Gaussian and vary its potential strength. Again, we define a dimensionless quantity $\alpha = 2a/(g_{eff}\mu_B B_0/\pi \xi^2)$ related to the strength of the impurity potential. When the impurity is located at $\vec{r}_0$, as shown in Fig. 4, the ground-state energy increases with $\alpha$ and saturates at a finite value for $\alpha > 0.01$. In the insets of Fig. 4, we can see that the ground-state wave function maintains its s-type symmetry as we increase $\alpha$ but develops a dip at its center. Similarly to the 1D case, the first excited state, which is twofold degenerate, is not affected by the impurity potential, as can be seen in Figs. 5(a) and 5(c). This is because its eigenfunctions have p-type symmetry, with a node at the origin where the impurity is located. These results mirror those of the 1D case but are less...
dramatic: the energy separation $\Delta E$ between the two lowest eigenstates changes by less than 30% for arbitrarily strong repulsive impurity potentials even when the charged impurity is placed where it should do the most damage.

Next, we investigate the dependence of the bound energies and wave functions on the location of the impurity, for a fixed strength $\alpha$. The results are shown in Fig. 6. The ground-state energy decreases monotonically as the impurity moves away from the vortex center because it costs less energy to create a dip at its location (the ground-state wave function is shown in the inset). Once the impurity is farther away than the characteristic length scale of the bound state, the ground-state energy saturates to its unperturbed value. Due to the symmetry of the problem, the degeneracy of the first excited state is lifted. The state which has its nodal line where the impurity is located is unaffected by it. The other eigenstate has a finite wave function at the impurity location, and its energy is increased. However, if the impurity is placed far enough, its effect vanishes and the two excited eigenstates become degenerate again. The excited wave functions are shown in Figs. 5(b) and 5(d).

Finally, we analyze how the energies of the lowest eigenstates change when the impurity is off-center at a fixed distance $r_I=5$ nm and we vary the impurity strength. The results are shown in Fig. 7. As expected, the ground-state energy increases but very little, since for large $\alpha$ a dip appears at the location of the impurity and its effect saturates, irrespective of its strength. The excited state with a nodal line at the impurity location continues to be unaffected by it, while the other excited state behaves similarly to the ground state: its energy increases with $\alpha$, but only by a finite amount until its wave function acquires a zero where the impurity is.

These results clearly demonstrate that a single impurity is unable to unbind the Zeeman trapped charge carrier, irrespective of how large its repulsive potential is. All it can do is to simply “poke a hole” in the wave function, and this costs a relatively small energy. Even if the impurity is placed at the center of the Zeeman potential well, the energetic cost is little. This suggests that one could have many impurities in the vortex area without affecting the bound state significantly. Our expectations are indeed verified in Sec. V.

Finally, we note that similar results are expected if the applied magnetic field is weaker. This results in a smaller binding energy and therefore a more extended wave function which is even less sensitive to the effect of local impurities. We have verified explicitly that, indeed, no qualitative changes are observed even for a $B_0$ as low as 0.08 T.

V. RANDOM DISTRIBUTION OF IMPURITIES

We now address a more realistic situation of a random distribution of multiple impurities inside the QW. In this case, as detailed in the Appendix, the use of B-splines is particularly useful to minimize computational costs.

For a large number of impurities, the energy and wave function will depend on the distribution of impurities. We follow the standard procedure and average our results over various impurity configurations. We performed averages over 1000 configurations at different impurity concentrations. The digitally doped Cd$_{1-x}$Mn$_x$Te has a zinc-blende structure with a lattice constant of about 6.5 Å. There are four Cd in a unit cell, and we take $x \sim 1.6$. Assuming a single layer of Cd$_{1-x}$Mn$_x$Te gives a Mn areal concentration of 0.15 nm$^{-2}$. As already discussed, the number of charged dopants can be minimized by the use of codoping techniques. While it is impossible to eliminate all these scattering centers, we expect that their concentration is much smaller than the Mn concentration.
at a very small energetic cost. As a result, even for large concentrations of very strong scatterers, the overall effect on the bound state is very limited.

This result has obvious positive implications for any applications based on such trapped states, since they show that it is not necessary to worry about the effects of impurities or to use expensive methods to eliminate them. Samples grown in reasonably clean conditions should suffice for devices based on these spin-polarized, Zeeman-trapped states. Also we notice that for systems with arrays of holes on the SC layer, the critical current can be enhanced, which opens more possibilities toward application.

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**APPENDIX: B-SPLINES**

B-splines have long been employed successfully in atomic and molecular physics but are not as frequently used in condensed matter physics. The aim of this appendix is to give a brief intuitive picture of the B-splines and their usefulness for solving Schrödinger equations with complicated potentials. Many more details on B-splines and their uses are available in the literature.

B-splines are piecewise polynomials and therefore are well suited for interpolation, having been extensively used in fitting tools, including many commercial software applications. In the context of interest to us, they are used as a basis in which to expand the eigenfunctions. Of course, there are many possible finite basis sets and finite elements methods that can be used to solve Schrödinger equations. The main advantage of using B-splines is the flexibility to choose the grid points on which the B-splines are defined. If we need to describe slowly varying functions, a large mesh suffices, resulting in a small basis set. In regions where there are fast variations, one can use a finer local mesh, optimized to give the desired accuracy for the minimum increase in the number of basis functions. Furthermore, because the B-splines are piecewise polynomials, matrix elements can be efficiently evaluated to machine accuracy with Gaussian integration. Finally, the banded nature and sparsity of the resulting matrices allow for the use of very large basis sets, if need be.

Suppose that we need to approximate a 1D function in a given interval \( x \in [a, b] \). We first define a knot sequence of points in this interval \( \{ x_i \mid a = x_0 \leq x_1 \leq x_2 \leq \ldots \leq x_N = b \} \). The location of the points as well as their number will be chosen so as to optimize the process. For instance, the knot sequence

![Graph showing the ground-state energy vs the impurity concentration.](image)

**FIG. 8.** The ground-state energy vs the impurity concentration. The energy was averaged over 1000 different impurity distributions and we chose \( \alpha = 0.005 \).

**VI. CONCLUSIONS**

In this article, we studied the effect of single and multiple impurities on the spin-polarized state bound by a trapping potential generated by an inhomogeneous magnetic field in a DMS QW. We demonstrated that this effect is very limited because of the large difference in length scales between the size of the wave function and the typical size of a strong scattering potential. For very large scattering potentials, each impurity pokes a hole in the bound wave function, especially in digitally doped layers.
can consist of regions of equally spaced knots combined with regions where the knots are space in an exponential fashion or any other suitable scheme. On this knot-sequence, we define the normalized B-splines of rank \( k \) by the following recursion relation:

\[
B_{i,k}(x) = \begin{cases} 
1 & \text{if } x_i \leq x < x_{i+1} \\
0 & \text{otherwise}
\end{cases} \tag{A1}
\]

and

\[
B_{i,k}(x) = \frac{x-x_i}{x_{i+k-1}-x_i} B_{i,k-1}(x) + \frac{x_{i+k}-x}{x_{i+k}-x_{i+1}} B_{i+1,k-1}(x), \tag{A2}
\]

where \( i \) is the index of a knot point and \( k \) is the order of the spline. Thus, the \( B_{i,k}(x) \) is a polynomial of order \( k-1 \) defined piecewise in the interval \([x_i, x_{i+k}]\) and vanishing outside it. Moreover, all derivatives up to the order of \( k-2 \) are also continuous. Since we use these functions to expand eigenstates, which are continuous and have first and second continuous derivatives, it follows that we should use a cubic (\( k=4 \)) or higher order B-splines. Here we use cubic splines, as they are the simplest splines with the desired properties. From now on, we simplify our notation and use \( B_i(x) \) to mean the cubic B-spline. The function of interest is then expanded in terms of these splines:

\[
f(x) = \sum_i \alpha_i B_i(x) \tag{A3}
\]

where \( \alpha \) are the \( N+3 \) coefficients of our expansion (for the \( N+1 \)-point knot sequence).

To solve an eigenvalue problem, we expand the wave function in terms of B-splines and reduce the problem to a general matrix system:

\[
\mathbf{H} - \varepsilon \mathbf{S} \mathbf{v} = 0, \tag{A4}
\]

where \( \mathbf{H} \) is the Hamiltonian matrix, \( \mathbf{S} \) is the overlap matrix, and \( \mathbf{v} \) is the eigenvector containing the unknown coefficients for the various B-splines. For a 1D Schrödinger equation with a potential \( V(x) \), the matrix elements of \( \mathbf{H} \) and \( \mathbf{S} \) are equal to

\[
H_{i,j} = \frac{1}{2m} \int \frac{dB_i(x)}{dx} \frac{dB_j(x)}{dx} dx + \int B_i(x)V(x)B_j(x) dx,
\]

\[
S_{i,j} = \int B_i(x)B_j(x) dx, \tag{A5}
\]

where we used an integration by parts in the kinetic energy. The integrals for the kinetic energy and the overlap can be evaluated analytically, while those of the potential may need numerical integration, if \( V(x) \) is a complicated function. However, given the finite support of each B-spline, such matrix elements are nonvanishing only if \( |i-j| < k \), so the number of needed integrals scales like the number of basis functions, not like its square, as is the case for most other basis.

The choice of the knot sequence plays an important role in these solutions. A good choice of knot distribution can assure a fast convergence to the true eigenenergy with a small basis set. To illustrate this, we consider a Gaussian potential and we calculate the ground-state eigenvalue and eigenfunction using two different knot sequences. As illustrated in Fig. 9, one is a uniform distribution of knots, while the other has an exponential distribution of knots from the center of the potential. In the inset of Fig. 9, we compare the convergence of the energy as a function of the number of knots. It is clear that the nonuniform distribution is much more efficient in this case, giving a very accurate wave function and eigenenergy for a sequence with very few knots, i.e., a very small basis set.

In the atomic physics, rigid boundaries are normally used when working with B-splines. Since in solid state physics we sometimes want to investigate bulk and transport properties, we opted to generalize the B-spline method to periodic boundary conditions. The periodic conditions for B-spline are constructed in the following way (for cubic splines, although the generalization to higher orders is straightforward), suppose we want B-splines on the support of the knot sequence \( \{x_i|a=x_0 \leq x_1 \leq x_2 \leq \ldots \leq x_N=b\} \).

(i) Expand the knot sequence \( \{x_i\} \) to \( \{x'_i\} \) by adding the knots

\[
x'_2 = x_{N-2} - (b-a), \tag{A6}
\]

\[
x'_1 = x_{N-1} - (b-a). \tag{A7}
\]

Note that since \( x'_1 \) and \( x'_2 < a \), they are outside the interval of interest.

(ii) Construct the B-splines \( \{B'_i\} \) with the usual procedure on the new support \( \{x'_i\} \). Note that the first two B-splines, which we call \( B'_0, B'_1 \), have finite support outside our desired interval.

(iii) Now construct the B-splines \( \{B_i\} \) by moving the pieces of \( B'_i \) defined outside the interval of interest to inside \([a,b]\) by a translation of period \((b-a)\).

Following the above procedure, all \( \{B_i(x)\} \) are periodic functions with period \((b-a)\). Hence the functions in the Hilbert space expanded by \( \{B_i(x)\} \) with coefficients \( c'_i \)'s:
are all periodic, thus ensuring the periodic boundary condition. Of course, for our problem of interest, we choose $b - a$ to be large enough that the localized wave function is all fully contained inside it. In other words, increasing $b - a$ has no effect on the eigenenergies we calculate.

The generalization to 2D or higher-dimensional systems is straightforward. The wave function needs to be expanded in products of B-splines

$$f(x) = \sum_i c_i B_i(x) \quad (A8)$$

and all the procedures above can be repeated. For our problem, we simply choose a high-density of knots in the immediate neighborhood of each impurity, where the wave functions change rapidly, and a wide exponential mesh everywhere else, where the function changes slowly. The mesh near each scatterer has been optimized until convergence is reached.

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