Enhancing dipolar interactions between molecules using state-dependent optical tweezer traps

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We show how state-dependent optical potentials can be used to trap a pair of molecules in different internal states at a separation much smaller than the wavelength of the trapping light. This close spacing greatly enhances the dipole-dipole interaction and we show how it can be used to implement two-qubit gates between molecules that are 100 times faster than existing protocols and than rotational coherence times already demonstrated. We analyze complications due to hyperfine structure, tensor light shifts, photon scattering and collisional loss, and conclude that none is a barrier to implementing the scheme.

Electric dipole-dipole interactions can be used to entangle polar molecules. Ensembles of such molecules are a promising platform for quantum simulation [1–3], quantum computation [4] or the creation of many-body states for precision metrology [5]. Recent progress in the production and control of ultracold molecules [6–15] has brought these goals within reach of near-term experiments.

Molecules confined in arrays of optical tweezer traps are particularly attractive and have recently been realized [16–19]. The platform is scalable to several hundred sites, enables re-arrangement of the traps [20, 21] to reduce entropy or control which particles interact, and provides natural single-site addressability. Various authors have proposed protocols for two-qubit gates using rotational states of molecules [4, 22–27]. The number of possible gate operations is set by the ratio of rotational states. For conventional tweezer traps \(E_{\text{dd}}\), the coherence time of a trapped molecule in a superposition of rotational states. For conventional tweezer traps \(E_{\text{dd}}\) is limited by the minimum trap separation. This is roughly the wavelength of the trapping light, typically \(\sim 1 \mu m\), giving \(E_{\text{dd}}/h \sim 1 \text{kHz}\). Recent work has extended \(\tau_c\) to several milliseconds [28, 29] but, at these interaction strengths, only a few high-fidelity gates are possible. While prospects are good for further improvements—coherence times near 1 s have been demonstrated in hyperfine states of molecules [30] and electronic states of atoms in tweezers [31]—considerable advances are required to realize the full potential of this platform for quantum science.

Here we show how to increase the dipole-dipole interaction between two molecules by trapping them at reduced separations using the state-dependence of the molecule-light interaction. Our scheme has similarities to state-dependent optical lattices which have been used to control atoms on sub-wavelength scales [34–42], but benefits from the advantages of the tweezer platform noted above. For atoms, electric dipole-dipole interactions involve electronically excited states [34] whose short lifetimes severely limit \(\tau_c\). By using long-lived rotational states of molecules, we avoid this limitation entirely. Our method, shown in Fig. 1(a), uses two optical tweezers of different wavelengths, focused at the same position. The tight focussing of the light produces elliptical polarization components with opposite handedness on each side of the focus [43, 44]. A molecule with non-zero spin has an interaction with the light field that depends both on this handedness and on the orientation of the spin. Consequently, two molecules in different internal states are trapped at different positions in the trap and their separation can be controlled by varying the relative intensities of the two tweezers. This state-dependent potential allows \(E_{\text{dd}}\) to be enhanced by two orders of magnitude. We introduce these concepts and show how to apply them in practice to implement fast two-qubit gates.

**Method.**—The scheme can be illustrated using a simple \(^2\Sigma\) molecule with no hyperfine structure. We focus on the four states with total angular momentum \(J = \frac{1}{2}\),

\[
|0_\pm\rangle = |N = 0, J = \frac{1}{2}, m_J = \pm \frac{1}{2}\rangle, \\
|1_\pm\rangle = |N = 1, J = 1, m_J = \pm \frac{1}{2}\rangle,
\]

FIG. 1. (a) State-dependent trap formed from a two-color optical tweezer. Two molecules in different internal states are trapped in different locations due to opposite circular handedness (rotating arrows) on opposite sides of the focus. (b) Contour plot of \((I/I_{\text{max}})C\cdot z\) in \(y = 0\) focal plane for a single tweezer. Calculated using the vector Debye integral [32] for a lens with \(NA = 0.55\). The input beam is polarized along \(x\) and has \(1/e^2\) diameter equal to that of the lens. An approximate analytical approach to this calculation, based on Ref. [33], is given in the SM.
Calculations are for $\lambda \Delta$ the ground state, typically of our states by
where $d\delta x$ the excited state into two components separated by
is shown in Fig. 2(a). The spin-orbit interaction splits
cited electronic state. The relevant electronic structure
this case the vector shift is $z$ edness is along
gating along $x$ direction gives the handedness. For incident light propa-
isor parts cause state-dependent shifts. The vector shift
constants $\alpha$ has scalar, vector and tensor parts whose dependence on
E field of intensity $W$ the rotational energy
$\Omega = 1/2$, $\pi$. Dashed lines: energies of scalar (red) and vector (orange) tweezers.
(b) Scalar (blue) and vector (green dashed) polarizabilities calculated using Eqs. (2).
(c) Calculated potentials of scalar (red dotted), vector (orange) and combined (blue)
tweezers versus displacement along $x$ for $J = 1/2$ states of $N = 0$ and 1. Solid (dashed) line shows the potential for $m J = -1/2(1/2)$.
Calculations are for $\lambda vec = 0.8\lambda xc$, $\Delta sc = 50\delta fs$ and $I sc/I vec = 100$. (d) Blue line, left axis: $\delta x$ versus $I sc/I vec$. Green dashed line, right axis: enhancement of $E idl$ for point particles trapped at the two minima.

tum $N = 0$ are separated from the pair with $N = 1$ by
the rotational energy $E rot$.
Consider the interaction of this molecule with a light field of intensity $I$ and polarization $\epsilon$. The interaction has scalar, vector and tensor parts whose dependence on the frequency of the light can be factored out into three constants $\alpha(0)$, $\alpha(1)$ and $\alpha(2)$: the scalar, vector and tensor polarizabilities. The scalar interaction shifts all four of our states by $W_0 = -1/2e xc \cdot \alpha(0) I$. The vector and ten-
sor parts cause state-dependent shifts. The vector shift is non-zero when the field has ellipticity, described by $\hat{C}$ = Im($\epsilon$ $\times$ $\epsilon^*$). $|\hat{C}|$ gives the degree of ellipticity and its direction gives the handedness. For incident light propagating along $y$ and linearly polarized along $x$, this hand-
edness is along $z$ and is opposite either side of the focus (see Fig. 1(b) and Supplemental Material (SM) [45]). In this case the vector shift is $W_1 = \gamma sc 3/4g sc 1/2 \alpha(1) g jm (\hat{C} \cdot \hat{z}) I$, where $g jm = 1/[2J(J + 1)]$ and we have assumed $W_1$ is small compared to the spin-rotation interaction. $W_1$ is identical for $|0-/1-\rangle$ and opposite to that of $|0+/1+\rangle$. The tensor shift is zero for our $J = 1/2$ states; we return to it later.

The polarizabilities $\alpha(k)$ depend on the details of the electronic structure [46]. Here, for simplicity, we assume that they are dominated by interaction with the first excited electronic state. The relevant electronic structure is shown in Fig. 2(a). The spin-orbit interaction splits the excited state into two components separated by $\delta fs$, typically $\sim 2\pi \times 1$ THz. Their mid-point is $\omega AX$ above the ground state, typically $\sim 2\pi \times 500$ THz, and we define $\Delta$ as the detuning of the light field from this point. For $\Delta \ll \omega AX$, the polarizabilities can be written (see SM)

$$\alpha(0) \approx -\alpha(1) \approx -\alpha(2) \approx \frac{d^2 AX \Delta}{3\hbar(\Delta - \delta fs)(\Delta + \delta fs)},$$

where $d^2 AX$ is the dipole matrix element connecting the $X^2Sigma$ and $A^2Pi$ states.

We use tweezer traps at two different wavelengths $\lambda xc$ and $\lambda vec$, shown schematically in Fig. 2(a), which we call the scalar and vector traps. Their on-axis intensities are $I sc$ and $I vec$. The scalar trap light is red-detuned with $\Delta = \Delta sc \gg \delta fs$. In this regime, $\alpha(0) \gg \alpha(1)$ and the interaction is dominated by the scalar component. The vector trap light is tuned between the fine structure com-
ponents. Figure 2(b) shows $\alpha(0)$ and $\alpha(1)$ in this region. When $\Delta = \Delta vec = 0$, $\alpha(0) = 0$ while $\alpha(1) = -2d^2 AX / \hbar \delta fs$ which can be large.

Figure 2(c) shows the light shifts of the four $J = 1/2$ states of our model molecule as a function of position along the $x$ axis. The dotted red line shows the light shift of all four states in the scalar trap for $\Delta sc = 50\delta fs$. The tiny state dependence caused by the residual vector light shift at $\lambda xc$ is not visible on this scale. The solid (dashed) orange line shows the light shift of the $|0-/1-\rangle$ ($|0+/1+\rangle$) states in the vector trap where we have assumed $\lambda vec = 0.8\lambda xc$ and $I sc/I vec = 100$. Molecules in the $|0-/1-\rangle$ states and the $|0+/1+\rangle$ states are trapped on opposite sides of the focus. When both scalar and vector traps are present the potentials add, shown by the blue lines. Varying their relative intensity controls the separation of the minima, $\delta x$. The blue line in Fig. 2(d) shows how $\delta x$ depends on the intensity ratio while the green line shows the enhancement of $E idl$ for two point particles po-
tioned at the trap minima relative to those in separated scalar traps. The enhancement is ultimately limited by undesirable collisions that occur when the spacing is too small. As we will see, for realistic parameters, an en-
hancement of 2 orders of magnitude is achievable.

**Eigenstates.**—The dipole-dipole interaction Hamil-
tonian is

$$H idl = \frac{\vec{d}_A \cdot \vec{d}_B - 3(\vec{d}_A \cdot \hat{x}) \otimes (\vec{d}_B \cdot \hat{x})}{4\pi \varepsilon_0 |x_B - x_A|^3},$$

FIG. 2. (a) Schematic of relevant electronic structure. The excited state $A^2Pi$ is split by spin-orbit coupling into two states with $|\Omega| = 1/2, 3/2$. Dashed lines: energies of scalar (red) and vector (orange) tweezers. (b) Scalar (blue) and vector (green dashed) polarizabilities calculated using Eqs. (2). (c) Calculated potentials of scalar (red dotted), vector (orange) and combined (blue) tweezers versus displacement along $x$ for $J = 1/2$ states of $N = 0$ and 1. Solid (dashed) line shows the potential for $m J = -1/2(1/2)$. Calculations are for $\lambda vec = 0.8\lambda xc$, $\Delta sc = 50\delta fs$ and $I sc/I vec = 100$. (d) Blue line, left axis: $\delta x$ versus $I sc/I vec$. Green dashed line, right axis: enhancement of $E idl$ for point particles trapped at the two minima.
where \( \vec{d}_A \) and \( \vec{d}_B \) are the dipole moments of the two molecules, \( x_A \) and \( x_B \) their positions, and \( \hat{x} \) is a unit vector along \( x \). As we show in the SM, after restricting ourselves to states with one molecule trapped on either side of the focus, the eigenstates of the two-molecule Hamiltonian, including \( H_{dd} \), are

\[
|0\rangle = |0_\pm\rangle |0_+\rangle, \quad |1\rangle = |1_\pm\rangle |1_+\rangle, \\
|\Psi^\pm\rangle = \frac{1}{\sqrt{2}} (|0_\pm\rangle |1_+\rangle \pm |1_\pm\rangle |0_+\rangle),
\]

with energies 0, 2\( E_{\text{rot}} \) and \( E_{\text{rot}} \pm E_{dd} \) respectively. Here \( E_{dd} = \Lambda_{10}/4\pi\epsilon_0 |x_B - x_A|^3 \) and the quantity \( \Lambda_{ij} = \langle j_- | \hat{d}_A \cdot \hat{d}_B - 3(\hat{d}_A \cdot \hat{x}) \otimes (\hat{d}_B \cdot \hat{x}) | i_- \rangle \) can be positive or negative.

Since \( |E_{dd}| \) can approach or even exceed the motional energy spacing in the trap, \( \hbar \omega_x \), it is important to consider the motional degree of freedom of the two molecules. A 1D treatment is sufficient to elucidate the main points. When the upper and lower states in each pair have the same vector shift, so that the potential is the same for both, the eigenstates are (see SM) \( |\psi\rangle |n_{cm}\rangle |\phi(x_{rel})\rangle \). Here \( |\psi\rangle \) is one of the internal eigenstates of Eqs. (4) and \( |n_{cm}\rangle \) is a harmonic oscillator eigenstate for the center of mass coordinate \( x_B + x_A \). The relative motional state \( |\phi(x_{rel})\rangle \) is an eigenstate of the state-dependent dimensionless Hamiltonian

\[
H_{rel} = \frac{p_{rel}^2}{2} + \frac{1}{2}(x_{rel} - \delta x)^2 + q \frac{r^3}{|x_{rel}|^3},
\]

Here \( x_{rel} = \sqrt{\frac{M \omega_x}{2\hbar}} (x_B - x_A) \) is the relative motional coordinate, \( p_{rel} \) the conjugate momentum, \( \delta x = \sqrt{\frac{M \omega_x}{2\hbar}} \), \( r = \sqrt{\frac{M \omega_x}{2\hbar}} (\Lambda_{10}/4\pi\epsilon_0 \hbar \omega_x)^{1/3} \) is the separation, in reduced units, at which \( E_{dd} = \hbar \omega_x \), and \( M \) is the mass of the molecule. The factor \( q \) reflects the state-dependence of the dipole-dipole interaction, and is equal to \( \{0, -1, 1, 0\} \) for \( |\psi\rangle = \{00, |\Psi^-\rangle, |\Psi^+\rangle, |11\rangle\} \) respectively. The relative motional states for \( q r^3 > 0 \) are examined in the SM and show two important effects. First, the finite extent of the wavefunction means that \( \langle 1/x_{rel}^3 \rangle > 1/3 \). Second, the molecules are pushed apart by their interaction so their mean separation is larger than \( \delta x \). In the motional ground state, the first effect dominates at larger \( \delta x \) increasing \( E_{dd} \), while the second effect dominates at small \( \delta x \), reducing \( E_{dd} \) below the value for fixed point dipoles.

**Complications in real molecules.**—The addition of nuclear spin introduces a hyperfine interaction which can mix states of different \( J \). For these mixed states, the vector Stark shift depends on the relative size of the hyperfine and spin-rotation interactions, which differs from one rotational state to the next. Consequently, the position of the potential minimum for \( |0_\pm\rangle \) is shifted relative to \( |1_\pm\rangle \). As shown in the SM, for a shift \( \xi \) in reduced units, the resulting imperfect overlap of the spatial wavefunctions reduces the dipole-dipole energy by \( e^{-\xi^2} \), the square of the overlap integral. As we will see, this reduction is typically small.

A second complication is that states outside \( N = 0 \) can have a tensor Stark shift due to the light at \( \lambda_{\text{sc}} \). The most relevant effect of this is to couple states with \( \Delta m \leq 2 \) near the center of the trap, allowing tunneling between the left and right potentials. This coupling is eliminated when the incident polarization of the scalar and vector traps are orthogonal. At other angles the tunneling is proportional to the wavefunction overlap so becomes negligible when the molecules are well separated.

**Realistic example.**—To illustrate the power and practicality of our method, we show how to implement a simple two-qubit gate using CaF molecules. CaF has been confined in optical tweezer traps [18] and has a structure similar to the model molecule but with a fluorine nuclear spin of \( 1/2 \). We map the states of the model molecule to our specific case as follows: \( |0_\pm\rangle = |N = 0, F = 1, m_F = \mp 1\rangle, |1_\pm\rangle = |1, 1, \pm 1\rangle \). We also introduce \( |2_\pm\rangle = |2, 2, \pm 2\rangle \) [47]. The states \( |0_\pm\rangle \) and \( |2_\pm\rangle \) form our computational basis, while \( |1_\pm\rangle \) are used to implement the gate. Many other choices of states are possible and may have different advantages. Figure 3(a) shows the potentials for the three pairs of states and for the parameters given in the caption. The closely spaced traps can be loaded adiabatically and without collisional loss from two separated tweezers using simple intensity ramps as shown in the SM. We suppose the molecules have been cooled to the motional ground state [46], resulting in the wavefunctions shown for each potential. The trap frequency is within 2% of 160 kHz for all states, and the corresponding rms wavepacket size is 23 nm.

Figure 3(b) shows the two-molecule states relevant for the gate. The matrix elements of \( H_{dd} \) are zero between states of our computational basis \( |0_\pm\rangle \) and \( |2_\pm\rangle \). The states \( |2_-, 1_+\rangle \) and \( |1_-, 2_+\rangle \) are mixed by \( H_{dd} \) giving the pair of entangled states \( |\Psi_1\rangle = (|2_-, 1_+\rangle \pm |1_-, 2_+\rangle)/\sqrt{2} \) split by \( 2E_{dd} \). A microwave pulse resonant with the \( |2_-, 2_+\rangle \leftrightarrow |\Psi_1^+\rangle \) transition and of sufficient duration to resolve it from \( |2_-, 2_+\rangle \leftrightarrow |\Psi_2^+\rangle \) entangles the two molecules. As we will see, \( E_{dd} \) and \( \hbar \omega_x \) are similar in size for our choice of parameters so a gate which resolves the dipole-dipole splitting will also resolve the motional sidebands so that motional heating can be avoided. Note that the transitions \( |2_-, 0_+\rangle \leftrightarrow |\Psi_1^0\rangle \) and \( |0_-, 2_+\rangle \leftrightarrow |\Psi_1^1\rangle \) are detuned because \( \Lambda_{21}/\Lambda_{10} \approx 2.9 \). A 2π pulse implements the two-qubit gate \( a |0_-, 0_+\rangle + b |0_-, 2_+\rangle + c |2_-, 0_+\rangle + d |2_-, 2_+\rangle \rightarrow a |0_-, 0_+\rangle + b |0_-, 2_+\rangle + c |2_-, 0_+\rangle - d |2_-, 2_+\rangle \). This gate is universal in combination with single-qubit operations which can be carried out rapidly using two-photon microwave pulses [13, 29, 48, 49]. In an array of such qubits, single-qubit addressability is obtained through a combination of microwave polarization and tweezer intensity. The polarization determines which molecule in a pair is addressed, and a small change in intensity of the selected tweezer relative to all others ensures that only the molecule in that tweezer is addressed.
The blue lines in Fig. 3(c) show the energy shift of $|\Psi_{2\pm}\rangle$ as a function of the separation of the potential minima for the $|2\pm\rangle$ states. The dashed and solid lines show results for fixed point dipoles and the full 1D calculation respectively (see SM for details). For a separation of 106 nm, as shown in Fig. 3(a), the combined effect of the dipole-dipole interaction pushing the molecules apart and the imperfect overlap of the motional wavefunctions, reduces $E_{dd}$ by $\sim 45\%$. Also shown are the expected dominant loss mechanisms in the trap. We calculate the collisional loss rate $R_{col}$ using the coefficient measured in Ref. [49] for CaF in a 780 nm tweezer trap. $R_{col}$ decreases with increasing separation and is largest when both molecules are in $|2\pm\rangle$ where their overlap is largest. The photon scattering rate $R_{ph}$ is dominated by scattering from the vector trapping light. We have assumed a fixed $I_{sc}$ so $R_{ph}$ increases with separation since larger separations require larger $I_{sc}$. Over the range shown, the ratio of $E_{dd}/h$ to the sum of the loss rates is large. Choosing the separation of the $|2\pm\rangle$ states to be 106 nm, $R_{col} \approx 280\,s^{-1}$ and $R_{ph} \approx 200\,s^{-1}$ while $E_{dd}/h = 130\,kHz$, almost 500 times larger. This is also 100 times larger than the maximum interaction energy achievable with separate tweezers. For a fixed vector Stark shift, $R_{ph}$ scales inversely with the fine-structure interval, so will be smaller for heavier molecules. For example, it is reduced by factors of $\sim 4, 6$ and 19 in SrF, YO and YbF respectively. $R_{col}$ may be very different in other systems or for the same system at different wavelengths [50, 51]; this is an important topic for investigation.

To scale our scheme to many molecules, traps can be rearranged to implement gates between different pairs. A useful metric is the time required to move a pair from two separated potentials into a single, combined trap ready for the fast gate. In the SM we describe a simple adiabatic protocol which takes 50 µs. More sophisticated non-adiabatic transport protocols [52, 53] can be completed more rapidly and without heating, as demonstrated for ions [54].

Summary.—We have proposed a new scheme which uses state-dependent optical tweezer traps to confine pairs of polar molecules at distances much smaller than the wavelength of the trapping light, and shown how to engineer a greatly enhanced dipole-dipole interaction between them. We have analyzed an example in detail, including the effects of hyperfine structure and tensor light shifts. We find that two-qubit gates can be implemented at least 100 times faster than existing protocols with characteristic figure of merit $E_{dd}\tau_c/h \sim 10^3$ for rotational coherence times already demonstrated for molecules [28, 29]. Thus, our work enables useful quantum information processing without further improvements to coherence times. Because the gate is so much faster, the effects of fluctuating magnetic fields or tweezer intensity matter less. We have designed a specific two-qubit gate, but our scheme provides a similar speedup for any gate that uses the dipole-dipole interaction e.g. [25]. Shaped microwave pulses that produce remarkable robustness to various experimental imperfections [27] can also be utilized in our scheme. Our method will work for all molecules laser cooled so far, and the heavier ones have a reduced scattering rate which may be an important advantage. The method should also work for heteronuclear alkali molecules prepared in the $^3\Sigma$ state [55].

As well as quantum information processing, the enhanced dipole-dipole interactions will be useful in quantum simulation. For example, a linear chain of tweezers with a pair of molecules in each can implement an SSH model [56] in a natural way. Furthermore, the ability to control the wavefunction overlap between two molecules with such precision is unique and offers a new tool for studying collisions and quantum chemistry with unprecedented precision and control.

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