Wide-range electrical tunability of single-photon emission from chromium-based colour centres in diamond

T Müller¹,⁷, I Aharonovich²,³, L Lombez¹, Y Alaverdyan¹, A N Vamivakas¹, S Castelletto²,⁴, F Jelezko⁵, J Wrachtrup⁶, S Prawer² and M Atatüre¹

¹ Cavendish Laboratory, University of Cambridge, Cambridge CB3 0HE, UK
² School of Physics, University of Melbourne, VIC 3010, Australia
³ School of Engineering and Applied Science, Harvard University, Cambridge, MA 02138, USA
⁴ Centre for Micro-Photonics, Faculty of Engineering and Industrial Science, Swinburne University of Technology, Mail H 34 Hawthorn, VIC 3122, Australia
⁵ Institut für Quantenoptik, Universität Ulm, Albert Einstein Allee 11, 89069 Ulm, Germany
⁶ 3. Physikalisches Institut, Universität Stuttgart, 70550 Stuttgart, Germany
E-mail: tm373@cam.ac.uk

New Journal of Physics 13 (2011) 075001 (9pp)
Received 25 January 2011
Published 5 July 2011
Online at http://www.njp.org/
doi:10.1088/1367-2630/13/7/075001

Abstract. We demonstrate electrical control of the single-photon emission spectrum from chromium-based colour centres implanted in monolithic diamond. Under an external electric field, the tunability range is typically three orders of magnitude larger than the radiative linewidth and at least one order of magnitude larger than the observed linewidth. The electric and magnetic field dependence of luminescence gives indications of the inherent symmetry, and we propose Cr–X or X–Cr–Y-type non-centrosymmetric atomic configurations as the most probable candidates for these centres.

Room-temperature operation of diamond-based colour centres offers a unique platform for the field of quantum photonics [1]. The nitrogen-vacancy (NV) colour centre in diamond has attracted interest in diverse research fields, ranging from spin-based quantum information [2–5]
to nanoscale magnetometry [6–9], within the last decade. The impressive level of spin coherence [5], the fast microwave spin manipulation [10] and the recently demonstrated spin–photon interface [11] are essential to the proposed goals. However, the NV centre’s broad spectrum is unfavourable for applications involving spectrally indistinguishable photons and efficient spin–photon entanglement. Although emission from the zero-phonon line (ZPL) has been shown to be radiative lifetime broadened at 1.8 K [12], it only accounts for about 4% of the total spectrum, owing to the dominance of phonon-assisted decay. This motivated the search for alternative centres, where potentially a similar degree of spin control could coexist with superior photonic properties. Consequently, silicon-vacancy (Si-V) [13, 14] and nickel-based centres, such as NE8 [15], were studied in the last few years for this purpose. Recently, chromium-based colour centres were reported as being among the brightest single-photon emitters in the near-infrared spectrum [16, 17]. Here, we show that these centres have an impressive level of spectral tunability of the order of a few meVs, while sustaining their oscillator strength. Furthermore, this is the first report of electric field tunability from impurities in diamond other than NV. This electrical tunability is an important asset for generating spectrally indistinguishable single photons from multiple colour centres. In addition, while the NV centre’s atomic structure has been known for decades [18], the chromium-related defect has been fabricated only recently and even its basic components need to be identified. The particular dependence of the luminescence from chromium-related emitters on an external electric field allows us to infer a non-centrosymmetric atomic configuration.

The colour centres under study were engineered by the ion implantation of chromium and oxygen into pure type IIA ([N] < 1 ppm, [B] < 0.05 ppm, Element Six) diamond crystals based on the protocol discussed in [19]. The implantation energies of the chromium and the oxygen atoms were 50 and 19.5 keV, respectively. Based on SRIM simulations, such energies will maximize the overlap of these two ions in the diamond lattice. All implantations were performed at room temperature in high vacuum (∼10−7 Torr). After implantation, the samples were annealed at 1000° C in a forming gas ambient (95% Ar–5% H2) for 2 h to repair damage caused by the implantation. The centre is only observed in samples with the right concentration of nitrogen, and co-implanting oxygen (or alternatively sulphur) increased the yield of optically active centres by at least one order of magnitude. Implantation of chromium into ultrapure diamond crystals ([N] < 5 ppb) did not lead to formation of the reported centres. The measurements reported here were performed on a sample with co-implantation of chromium and oxygen only. All of our measurements were carried out at liquid helium temperature (4 K) using a home-built confocal microscope, where the optical axis was aligned with the [001] crystallographic axis of the sample. The defects were optically excited non-resonantly using a continuous-wave titanium:sapphire laser tuned to 710 nm. The laser light was focused on the sample with a high numerical aperture lens (NA = 0.65), which was also used to collect the emission from the defects. Residual pump laser light was filtered out using a 740 nm long-pass filter and photoluminescence (PL) was then focused into a single-mode fibre that acted as a pinhole. A spectrometer with a liquid-nitrogen-cooled CCD was used to analyse the PL, and time-correlated photon counting was used for lifetime and intensity-correlation measurements. Isolated bright spots on the sample were found by scanning the piezoelectric stage on which the sample was mounted. Magnetic fields were applied with fixed direction along the optical axis using a superconducting magnet. Electrical gates were fabricated using electron-beam lithography and consist of 10 nm of chromium and 40 nm of gold.
Figure 1. (a) Resolution-limited ZPL spectrum of a chromium-related centre at 4 K under non-resonant excitation at 710 nm. The linewidth $\Gamma$ is well below 10 GHz, and no phonon sidebands can be resolved. (b) An example of a broadened ZPL spectrum of a chromium-related centre, with a linewidth of about 60 GHz. Again, phonon sidebands cannot be resolved. (c) Magnetic field dependence of the resolution-limited ZPL at 793.1 nm shown in panel (a). The magnetic field was applied parallel to the optical axis, and spectra were taken every 500 mT. (d) Polarization dependence of excitation (blue) and emission (red) of the ZPL transition at 765.1 nm shown in panel (b). The solid circles show experimental values and the solid curves are fits to the data with sine square law. The visibility $(I_{\text{max}} - I_{\text{min}})/(I_{\text{max}} + I_{\text{min}})$ is 96.6% for emission and 97.8% for excitation. Emission and excitation are polarized along the same axis.

Figure 1(a) displays typical PLs of a single chromium-related centre when excited non-resonantly with a 710 nm laser. The spectrum shows spectrometer-resolution-limited (well below 10 GHz) photon emission strongly concentrated in the ZPL. A very weak phonon-assisted luminescence can be observed for long timescale integration, which is consistent with the previously reported Debye–Waller factor well exceeding 0.9 [17]. Not all centres had resolution-limited ZPL emission, and figure 1(b) displays an example of broadened emission from a single centre. The excited state lifetime for such centres is around 1 ns [16, 19], implying the presence of broadening mechanisms, such as spectral diffusion, fast dephasing or a finite lifetime of the final state. While previous work on chromium centres suggests that the predominant contribution to broadening may be spectral diffusion at 4 K [17], field correlation measurements would be able to reveal the relevance of each mechanism. Luminescence from
(a) Histogram of ZPL centre wavelengths of 65 emitters under non-resonant excitation. (b) Temporal behaviour of the ZPL spectrum in 1 s time intervals. Spectral jumps of the order of a nanometre are observed, as well as a temporary loss of fluorescence. (c) Intensity line cuts at 785.5 nm (red) and 786.4 nm (green) of the spectra shown in panel (b), displaying the anticorrelated emission intensity of the two lines. The two emission wavelengths do not coexist, but occasionally spectral jumps within the integration time of 1 s lead to reduced contrast.

all centres studied was linearly polarized to better than 90%, as shown in figure 1(c). We find that absorption is also linearly polarized and that the emission and the absorption dipoles point along the same direction. The direction of the excitation and emission dipole with respect to the crystallographic axes varies from centre to centre. About 50% of the investigated emitters are polarized along the [100] crystal axis; others display angles of 60° and 120° with respect to the [100] axis. However, we also observe emitters that do not fall into any of these categories. These polarization properties are maintained under the application of both magnetic and electric fields. Figure 1(d) displays typical magnetic field responses of these centres up to 7 Tesla external field along the optical axis of the microscope (corresponding to the [001] axis of the sample). Within the resolution of our experiments (≈5 GHz), the ZPL does not show any shift, splitting, broadening or loss of emission intensity. These measurements could indicate a transition between states with commensurate Landé factors, as in the case of NV centres. Alternatively, the non-resonant excitation could give rise to spin polarization of the centre in a state unaffected by magnetic field. A weak spin–orbit coupling can furthermore make potential transitions to different spin states undetectable within our signal-to-noise limit, such that no final information can be extracted concerning the spin part of the electronic wavefunction.

The two centres presented in figure 1 differ in the actual wavelength of their ZPL emission. Figure 2(a) shows a histogram of single ZPL emission wavelengths from the chromium-based defects grouped in 2 nm bins. The wavelength distribution of observed ZPLs ranges from 740 up to 800 nm, particularly concentrated around 748 and 775 nm. It is interesting to note that while the NV centre exhibits a very well-defined spectral signature, other defects, such as the Si-V, show a similar inhomogeneous broadening of ZPL centre wavelength [14]. The lack of discrete accumulation of the histogram to a few centre wavelengths suggests that we are not looking at a family of microscopic structures, each having its own ZPL wavelength, but rather the same microscopic structure under a variation of local external effects. For the implanted
chromium-related centres, the lattice surrounding each colour centre is quite rich, containing a significant number of impurities. The implantation of high doses ($\sim 10^{10}$ ions cm$^{-2}$) of chromium results in a high chromium concentration in the diamond lattice. In addition, the fabrication process generates a high concentration of substitutional nitrogen, oxygen, vacancies and NV centres while forming a lower density of chromium-based centres [20]. A PL under a 532 nm laser excitation indeed shows that the NV density is too high to optically resolve individual NV centres. The density of impurity atoms and ions is therefore quite high. We expect that the histogram of figure 2(a) occurs due to the individual local electric field and strain experienced by each centre due to alien atoms. In addition, the mere presence of a high density of non-carbon atoms can modify the diamond band structure in the vicinity of each centre, contributing to the dispersion we observe in the transition wavelength. One could further speculate that the two suggestive lobes of the histogram may correspond to two charging states of the centre similar to the NV$^0$ and NV$^-$ states, where the charging energy here would be around one third that for NV centres, suggesting slightly weaker confinement. In addition to ZPL wavelength inhomogeneity, most centres show switching between multiple discrete lines and/or dark periods of the order of seconds. Figure 2(b) displays temporal dynamics of the ZPL spectrum for a single centre. This switching behaviour does not change the polarization direction of the emission, and all lines originating from the same emitter are polarized along the same direction. The excitation laser power strongly influences these dynamics such that the centres become more stable under lower excitation laser power. This is again consistent with optically induced changes in the charge configuration in the vicinity of each centre, possibly related to photoionization of the defects in the diamond lattice. Intensity plots along 785.5 nm (red curve) and 786.4 nm (green curve) of figure 2(c) display the anticorrelated behaviour of emission originating from the same ZPL. We note that some centres do not display switching and remain stable over the whole measurement time. Also, on rare occasions, we observe loss of fluorescence, possibly due to ionization of the colour centre.

The temporal signature of each centre is unique, indicating the high level of sensitivity of these artificially fabricated centres to their local environment. As strain and electric field influence the local surroundings of defects significantly, we proceed to investigate the luminescence of the centres under an externally applied electric field. Emitters are exposed to electric fields using interdigitated gates on the diamond surface, as shown in figure 3(a). The spacing between the fingers and their width is 3 $\mu$m. We fabricated structures both parallel to the crystal axes [010] and [100], and at 45$^\circ$ with respect to the axes. Applying 100 V across the fingers results in local electric field $F$ of up to $8.5 \times 10^6$ V m$^{-1}$ and the polarity can be reversed by applying a reverse bias. To determine the local field just beneath the diamond surface, we use the Lorentz local electric field approximation $F = V(\epsilon + 2)/2d$, with $\epsilon$ being the dielectric constant for diamond, $V$ the voltage applied and $d$ the gate separation. The Stark shift observed in the optical transition should then be given by $\hbar \Delta \nu = -\Delta \mu \cdot F - \frac{1}{2} F \cdot \Delta \alpha \cdot F$ in second-order perturbation theory, where $\hbar$ and $\Delta \nu$ are Planck’s constant and the change in transition frequency. $\Delta \mu$ and $\Delta \alpha$ represent the change in dipole moment and the change in polarizability between the excited and the ground states, respectively.

Figures 3(b)–(e) present the response of the ZPL spectrum to the applied electric field for four different centres. A significant part of the observed centres exhibits a linear dependence, as seen in figures 3(b) and (c). A predominantly quadratic response was observed for a few centres, as seen in figures 3(d) and (e), and about half of the centres remained unchanged by the electric field. The Stark shift (both linear and quadratic) can cause either an increase or a decrease in
Figure 3. (a) Interdigitated gold gates on single-crystal diamond. (b) Resolution-limited ZPL emission spectrum as a function of the applied electric field. Negative field values result in a linear dipole coefficient $\Delta\mu = 2.1 \text{ GHz MV}^{-1} \text{ m}^{-1}$; positive field values give $\Delta\mu = 0.2 \text{ GHz MV}^{-1} \text{ m}^{-1}$. The linewidth of the centre remains unchanged under an electric field, whereas the intensity increases for large negative field values (inset). (c) A centre with a strong linear Stark-shift coefficient $\Delta\mu = 8.36 \text{ GHz MV}^{-1} \text{ m}^{-1}$ for positive applied field values. The inset shows two spectra taken at 0 MV/m (blue) and 72.8 MV/m (red), with a peak separation of 1.49 nm. At high-field values, this centre exhibits line broadening, whereas the overall intensity remains constant. (d) A centre with ZPL at 778.1 nm displaying a predominantly quadratic dependence on the applied electric field with a positive polarizability difference $\alpha = 0.36 \text{ GHz MV}^{-1} \text{ m}^{-2}$ and linear coefficient $\Delta\mu = 0.19 \text{ GHz MV}^{-1} \text{ m}^{-1}$. (e) This centre shows a negative polarizability difference $\Delta\alpha = -0.075 \text{ GHz MV}^{-1} \text{ m}^{-2}$, with a linear coefficient $\Delta\mu = 0.054 \text{ GHz MV}^{-1} \text{ m}^{-1}$, much weaker than those reported in panel (d).
transition energy when the magnitude of the electric field is increased. Note that the strength and the sign of the observed Stark shift per centre both depend on the respective orientation of the electric dipole and the gate structure. We have investigated 12 emitters with the gate parallel to the [100] axis of the crystal, and 26 emitters with the gate at an angle of 45° with respect to the crystal axis. We did not find a significant difference in the electric field response between the two directions or any obvious correlation between the electric field response and the angle between absorption dipole and gate structure. For most of the observed centres, the linewidths remain largely unaffected under the application of an electric field, as displayed in the inset of figure 3(b).

The responses of the chromium-based emitters to electric fields are striking for two reasons. Firstly, the centres exhibit a large tunability of the sharp emission lines, typically two to three orders of magnitude larger than the extracted radiative linewidth and still sustaining the oscillator strength. The range over which the ZPL emission is tunable is at present limited by the maximum applied voltage. This is remarkable and of extreme importance for application of these emitters in the generation of indistinguishable photons from different sources [21–23]. Using the model presented above, we extract typical linear dipole moments of the order of $\Delta \mu = 8.36 \text{GHz m MV}^{-1}$ (1.66 Debye). This value is comparable to that for NV centres in diamond. With such performance, the investigated chromium-related defect becomes comparable with other solid state systems, such as self-assembled quantum dots [24] or conjugated polymer molecules [25] for electrical tunability.

Secondly, we observe different linear dipole moments for positive or negative field directions. The linear dipole moments can even be effectively zero for one field direction but finite for the other, as seen in figure 3(b). Since we do not observe a change of slope for finite electric field values for either polarity, it is unlikely that this surprising feature is environment mediated, but rather inherent to the centre. As the exact geometry of the present defect is unknown, it is not possible to compare this result with a simple theoretical prediction for a two-level system. Nevertheless, we can establish an intuitive picture for our findings using an atomic physics model. Considering the level structure of a simple atom in free space, a linear Stark shift only occurs for hybridized energy levels due to the well-defined parity of the pure electronic states. We can therefore expect at least one of the probed electronic energy levels of the investigated chromium-related defect to have orbital degeneracy. Furthermore, the heavy dependence of the observed Stark shift coefficient on the electric field polarity for linearly responding centres is only possible for a centre lacking inversion symmetry. This argument can be illustrated by looking at the well-studied NV centre whose symmetry group ($C_{3v}$) does not contain inversion symmetry either. Recent theoretical work on the excited state level structure of the NV centre [26] indicates that an electric field can indeed cause mixing within the excited state manifold leading to eigenstates with different linear dipoles. Assuming a similar structure for the chromium-related centre in the excited (ground) state of the optical transition, and the presence of a relaxation mechanism within the excited (ground) state manifold to whichever state has lower (higher) energy for a given electric field, our findings become more plausible. Irrespective of whether this behaviour originates from the ground or from the excited state of the optical transition, the lack of inversion symmetry of the defect compound has consequences for the atomic structure of the centre. We can infer that the centre cannot consist of a single chromium atom only, since inversion symmetry would not be broken for either a substitutional or an interstitial single impurity atom in diamond. Given that the centre is controllably fabricated with a finite yield, we speculate further that the centre is formed by no more than two atomic
species and vacancies. We therefore propose a Cr–X or X–Cr–Y-type atomic structure for these Cr-based centres, where X and Y are likely to be oxygen, nitrogen or sulphur. We note that one of the atomic species may indeed act purely as a donor or acceptor for a centre nearby. A vacancy for X in Cr–X is unlikely due to the stringent dependence of centre formation on oxygen (sulphur) and nitrogen densities simultaneously. However, either X or Y in X–Cr–Y formation may be occupied by a vacancy still preserving the lack of inversion symmetry. These restrictions on the atomic structure of the centre open the door for simulations of the centre using DFT models to gain further insight into the electronic level structure.

In short, we have observed a large range tunability of chromium-related centres in diamond via the Stark effect. The asymmetric nature of the linear dipole response allows us to infer an atomic configuration lacking inversion symmetry. This most likely leads to an electronic level structure with orbital degeneracy that is lifted under electric fields. Given the strong linear dipole present in the system, charge fluctuations in the environment of the defect will have an impact on the emission properties. Due to the fabrication procedure, the chromium centres are embedded in an environment that hosts a number of non-carbon atoms and vacancies. This will lead to not only inherent electric field per centre due to charge distribution, but also a slight bandgap modification for the diamond in the vicinity of the centres. The optical performance of the defects is likely to be significantly enhanced if fabricating defects with a cleaner matrix at the end of the protocol can be achieved, thus making them available to applications in various quantum photonics technologies. Resonant excitation, optically detected magnetic resonance measurements on a single centre, as well as electron spin resonance measurements on an ensemble of centres will shed more light on the level structure under magnetic field.

Acknowledgments

We gratefully acknowledge financial support from the University of Cambridge, the European Research Council (FP7/2007–2013)/ERC Grant agreement no. 209636, the Australian Research Council, The International Science Linkages Program of the Australian Department of Innovation, Industry, Science and Research (project no. CG110039), the European Union Sixth Framework Program under program no. EQUIND IST-034368, the DFG (SFB/TR21, FOR1482) and the BMBF (EPHQUAM and KEPHOSI). IA acknowledges the Grant-In-Aid of Research from Sigma Xi, The Scientific Research Society. We thank C Matthiesen and C-Y Lu for technical assistance, and H Tureci and D D O’Regan for helpful discussions.

References


New Journal of Physics 13 (2011) 075001 (http://www.njp.org/)
Balasubramanian G et al 2010 Ultralong spin coherence time in isotopically engineered diamond Nat. Mater. 8 383


Degen C L 2008 Scanning magnetic field microscope with a diamond single-spin sensor Appl. Phys. Lett. 92 243111


Balasubramanian G et al 2008 Nanoscale imaging magnetometry with diamond spins under ambient conditions Nature 455 648–51


Togan E et al 2010 Quantum entanglement between an optical photon and a solid-state qubit Nature 466 730

Tamarat Ph et al 2006 Stark shift control of single optical centres in diamond Phys. Rev. Lett. 97 083002


Aharonovich I, Castelletto S, Simpson D A, Stacey A, McCallum J, Greentree A D and Prawer S 2009 Two-level ultrabright single photon emission from diamond nanocrystals Nano Lett. 9 3191–5


