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A dark-field microscope for background-free detection of resonance fluorescence from single semiconductor quantum dots operating in a set-and-forget mode

Andreas V. Kuhlmann,1, a) Julien Houel,1 Daniel Brunner,2 Arne Ludwig,1,3 Dirk Reuter,3,4 Andreas D. Wieck,3 and Richard J. Warburton1
1Department of Physics, University of Basel, Klingelbergstrasse 82, CH-4056 Basel, Switzerland
2Instituto de Física Interdisciplinar y Sistemas Complejos, IFISC (CSIC-UIB), Campus Universitat Illes Balears, E-07122 Palma de Mallorca, Spain
3Lehrstuhl für Angewandte Festkörperphysik, Ruhr-Universität Bochum, D-44780 Bochum, Germany
4Department Physik, Universität Paderborn, Warburger Strasse 100, D-33098 Paderborn, Germany

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Optically active quantum dots, for instance self-assembled InGaAs quantum dots, are potentially excellent single photon sources. The fidelity of the single photons is much improved using resonant rather than non-resonant excitation. With resonant excitation, the challenge is to distinguish between resonance fluorescence and scattered laser light. We have met this challenge by creating a polarization-based dark-field microscope to measure the resonance fluorescence from a single quantum dot at low temperature. We achieve a suppression of the scattered laser exceeding a factor of $10^3$ and background-free detection of resonance fluorescence. The same optical setup operates over the entire quantum dot emission range (920–980 nm) and also in high magnetic fields. The major development is the outstanding long-term stability: once the dark-field point has been established, the microscope operates for days without alignment. The mechanical and optical designs of the microscope are presented, as well as exemplary resonance fluorescence spectroscopy results on individual quantum dots to underline the microscope’s excellent performance. © 2013 AIP Publishing LLC.

[I. INTRODUCTION]

Semiconductor quantum dots, in particular self-assembled InGaAs quantum dots, are very attractive as the building blocks for quantum light sources and spin qubits.2 Self-assembled InGaAs quantum dots (operating at wavelengths around 950 nm at low temperature) exploit technologically advanced GaAs heterostructures and have become the workhorse system in the field. It is hugely advantageous to explore the physics using resonant rather than non-resonant laser excitation. On the one hand, non-resonant excitation introduces sources of noise resulting in excitation and spin dephasing.3 On the other hand, resonant (but not non-resonant) excitation allows a spin to be initialized,4, 5 manipulated,6 and read-out7 optically. Resonant excitation, i.e., coherent laser spectroscopy, on single InGaAs/GaAs quantum dots was first developed with differential transmission detection,8 using Stark-shift modulation of the transitions energy along with lock-in detection for noise rejection.9 The detection scheme exploits an interference between the laser field and the field associated with coherently scattered photons:10 it provides a sensitive detection scheme but does not provide direct access to the resonance fluorescence, the single photons scattered or emitted by the quantum dot. These photons are crucial to develop a high-fidelity single photon source and, further afield, in developing a quantum dot-based quantum network with applications in quantum communication.11

Recently, the resonance fluorescence of a semiconductor quantum dot3, 12–21 has been observed. The challenge experimentally is to distinguish quantum dot-scattered light from scattered laser light. With non-resonant excitation, this separation is trivial to achieve on account of the widely different wavelengths. With resonant excitation, this scheme fails. One scheme for the detection of resonance fluorescence exploits the different wave vectors of the laser light and the resonance fluorescence,12–15 This is very much in the spirit of the original ensemble experiments in atomic physics in which resonance fluorescence was detected in a direction orthogonal to the carefully defined propagation direction of the laser.22, 23 In a semiconductor context, one implementation of this scheme involves coupling laser light to a waveguide containing quantum dots with edge illumination, detecting the resonance fluorescence in the orthogonal vertical direction.12, 13, 18 Another scheme exploits a further property of light: its polarization. The idea is to operate in the dark-field as defined by the polarization: the laser and the detection are defined to have orthogonal polarization states. Provided laser scattering preserves the polarization, the crossed polarizer configuration ensures that scattered laser light is prevented from entering the detection mode. Success has been achieved using crossed linear polarizations.3, 16–21

In our experiments, we have pursued the polarization-based dark-field technique as, first, it does not require a specially fabricated waveguide, and second, space limitations in the bore of a superconducting magnet limit the possibilities for efficient edge illumination. It is clear that achieving sufficient laser rejection based on polarization requires both high

a)andreas.kuhlmann@unibas.ch
II. DESCRIPTION OF THE DARK-FIELD MICROSCOPE

The design of the dark-field microscope makes no particular demands on the sample although a flat, smooth surface is best. Once the wavelength range of the optics is adapted to the emission range, the dark-field concept operates equally well with the sample at room temperature or at low temperature. Here, as an example of a two-level system in the solid state, we study self-assembled InGaAs quantum dots emitting at wavelengths around 950 nm at low temperature. The microscope combines both high spatial resolution, implemented by means of orthogonal excitation/collection polarization states: the linear polarizer sets the laser polarization to s-polarized laser light; the linear polarizer and quarter-wave plate define and control the state of light polarization. In this scheme, the PBSs define linear s-polarization and linear p-polarization for excitation and detection, respectively. The linear polarizer sets the polarization of the laser light to s-polarization before striking the PBS, the quarter-wave plate controls the polarization thereafter. In particular, the quarter-wave plate allows for a compensation should an ellipticity be inadvertently induced. The back-scattered s-polarized laser light is reflected by both the first and second PBSs by 90° such that the s-polarization is highly suppressed in transmission. The p-polarized component of the quantum dot emission, however, is transmitted and can be detected.

The confocal configuration improves the microscope’s dark-field performance. Both light scattering at surface imperfections in the detection beam path and the p-polarized field component in the focal spot are highly suppressed. The more field confinement at the focus the larger is the p-polarized field component of the focal spot of an incident s-polarized laser beam. However, the intensity distribution of the p-polarized component has a clover-leaf pattern with
an antinode at the centre, i.e., it is highly suppressed by the confocal detection.

B. Dark-field microscope design

Experiments on single semiconductor quantum dots typically require low temperatures. The dark-field microscope is therefore integrated into a free-beam microscope system developed for low temperature experiments. The microscope optics apart from the objective lens remain under ambient conditions, as shown in Fig. 1. The construction frame for the microscope “head” is a 30 mm cage system that allows a modular design: the lower horizontal microscope “arm” provides the excitation laser, the vertical arm is used for light detection, and the upper horizontal arm to image the sample surface. Each module is attached to a central cage, hosting the PBSs.

The lower horizontal microscope arm provides a link between remote excitation sources and the microscope. Its output is a well collimated beam of coherent laser light, precisely controlled in linear polarization, and used to excite a single quantum dot resonantly. A single mode (SM) fibre (FONT Canada SM fibre NA = 0.12, mode field diameter (MDF) 5.2 μm) interconnects the microscope and the excitation laser (Toptica DL pro 940). By adjusting an x/y-translation stage (Thorlabs CP1XY), the fibre core can be centered on the optical axis defined by the collimator (Thorlabs C280TME-B NA = 0.15, f = 18.4 mm), which is mounted in a z-translation stage (Thorlabs SM1Z). Aspheric lenses are used to collimate/focus the laser beam, as they provide diffraction limited performance for monochromatic applications. A metallic nanoparticle linear fiber polarizer (Thorlabs LPVIS050-MP) mounted on a rotary stepper positioner (attocube ANR240) polarizes the excitation laser linearly and additionally allows the axis of linear polarization to be precisely controlled. The polarizer’s transmission is 82% and its extinction ratio exceeds 8 orders of magnitude at a wavelength of 950 nm. The piezo-driven rotary stepper positioner provides both 360° endless rotation and a step size as small as 1 mdeg. Furthermore, after aligning the polarizer position by means of the control electronics (attocube ANC300), the piezos are grounded and their position is locked, providing outstanding long-term stability. The four cage rods of the excitation arm are connected to a tilt stage (Thorlabs KC1-T/M) which is attached to the central beam splitter cage and allows for a compensation of any angular displacement of the beam.

The vertical microscope arm is designed to collect light efficiently with a confocal rejection of any stray light. This relies on coupling into a SM fibre (FONT Canada SM fibre NA = 0.12, MDF 5.2 μm) which interconnects the microscope and the detectors. The same optical and opto-mechanical components as for the light collimation unit of the horizontal arm are used. The vertical arm is assembled directly on to the PBS cage. The lower tilt stage allows to correct for a misalignment with respect to the optical axis of the objective lens (Thorlabs 352330-B NA = 0.68, f = 3.1 mm).

The upper horizontal microscope arm provides the possibility of monitoring the objective lens focal plane, i.e., the sample surface. An achromatic lens (Thorlabs AC254-150-B-ML, f = 150 mm) focuses light onto the chip of a complementary metal oxide semiconductor (CMOS) camera (Allied Vision Technologies Guppy F-503B), resulting in a magnified image (magnification of 48) of the sample surface. Again a tilt stage allows angular control of the optical axis.

All modules of the microscope are attached to a central cage made from a solid piece of aluminum. It provides stability to the microscope and at the same time hosts two PBSs (B. Halle & Nachfolger PTW 2.10), crucial to implement the polarization filtering. The PBSs allow beam splitting sensitive to the polarization of the incident beam. Two right angle prisms made of flint glass are cemented together to form a cube. A dielectric beam-splitter coating which is deposited on one of the prisms provides a close to unity transmission for p-polarized and close to zero transmission for s-polarized light. The polarization suppression exceeds 4 orders of magnitude.

A quarter-wave plate (B. Halle & Nachfolger RZQ 4.10) is mounted beneath the PBSs on a second piezo rotary stage. (Note that the quarter-wave plate behaves as a half-wave plate for the reflected laser light as the laser beam passes it twice.) On the one hand, it is useful during the setup procedure to misalign the quarter-wave plate deliberately and allow some reflected laser light into the detection arm. On the other hand, the quarter-wave plate represents an extra degree of freedom and it turns out that this is crucial: it compensates for any distortion from linear to elliptical polarization in the two polarization states. It is not exactly clear where these small distortions arise, but they are probably related to a birefringence of the sample (GaAs with thin metal layer), solid immersion lens, objective lens, or the cryostat window. The quarter-wave plate used here is a zero order wave plate designed for 946 nm and was chosen because, first, it is less temperature sensitive than the multi-order counterparts; and, second, its performance at these particular wavelengths 950 ± 20 nm surpasses the performance of achromatic wave plates. The accuracy on the path difference of the quarter-wave plate is ±2 nm. Again a crucial point for the long-term behavior of the dark-field microscope is that the quarter-wave plate is mounted on a piezo positioner, as for the linear polarizer.

The microscope is inserted into a 2 in. bore stainless steel tube, evacuated, and filled with ~25 mbar He gas (exchange gas) at room temperature. The tube is then slowly inserted into a He bath cryostat equipped with a 9 T superconducting solenoid. The optics at 300 K are possibly subject to thermal drift but these are minimized by working in a ±1 °C temperature stabilized laboratory.

C. Dark-field microscope alignment

The microscope operates in both confocal and dark-field modes. For confocal performance, the excitation and collection beams must be concentric and parallel to the optical axis of the objective lens. While monitoring the focal spots on the sample surface, the tilt stages are aligned in order to superimpose the focal spots. The z-position of the sample relative to the objective focal plane is adjusted by moving the sample with nanometer precision. During this alignment step,
laser light is also coupled into the fibre of the vertical microscope arm. Subsequently, once the confocal condition has been achieved, the linear polarizer and quarter-wave plate are aligned to suppress the back-reflected laser light. The linear polarizer is aligned to define the polarization of the laser to $s$. A rough alignment is done by monitoring the transmitted signal at the PBS as the polarizer is moved. A minimum in transmission is required. The dark-field point is set by monitoring the back-reflected laser intensity. First, the linear polarizer is fine aligned by minimizing the back-reflected light, and second, the quarter-wave plate. We find that iterative fine tuning of the polarizer and quarter-wave plate angles enhances the rejection further, typically by a factor of 10 after $\sim$3 iterations. Piezo-electronics allow remote control of both the angles of the linear polarizer and the quarter-wave plate. Once the angles are set, the piezos are grounded.

**III. DARK-FIELD MICROSCOPE PERFORMANCE**

The performance of the dark-field microscope is characterized under real, experimental conditions: the laser is focused on a quantum dot sample in a low temperature experiment.

**A. Quantum dot sample**

The InGaAs quantum dots are grown by molecular beam epitaxy utilizing a strain-driven self-assembly process and are embedded in a Schottky diode. They are separated from an $n^+$ back contact by a 25 nm thick GaAs tunnel barrier. On top of the quantum dots is a capping layer of thickness 150 nm, followed by a blocking barrier, an AlAs/GaAs superlattice of thickness 272 nm. The samples are processed with Ohmic contacts to the back contact, grounded in the experiment.

**B. Laser suppression and long-term stability**

In order to observe resonance fluorescence with a high signal-to-background ratio, the microscope’s laser suppression has to be high. The laser rejection can be determined by rotating the quarter-wave plate, switching between laser rejection maximally on and maximally off. The back-reflected laser light intensity depends periodically on the quarter-wave plate angle with a period of $\pi/4$. A laser suppression exceeding $10^8$, corresponding to an optical density (OD) of 8 is achieved. (The OD is defined as $OD = -\log (1/T)$ with transmission $T$.) Fig. 2 shows a time trace of the detected laser light with and without laser rejection. An initial count rate of 580 MHz is reduced to 4 Hz by switching on the suppression. A single PBS achieves an extinction ratio of OD 5, a second PBS enhances the laser suppression. However, it is not increased by a further 5 orders of magnitude. We believe this difference is due to a stress induced birefringence of the PBSs, defining an upper limit for the laser suppression.

The effort to align the dark-field microscope is low. However, how stable is the alignment? Fig. 3 shows how the optical density depends on the quarter-wave plate angle: it is an extremely sensitive dependence. A change as small as a few mdeg can worsen the rejection by one order of magnitude. On the one hand, it emphasizes the need for a mdeg positioning resolution, and on the other hand, the need for an extreme
mechanical and thermal stability to achieve good long-term dark-field performance.

Despite the high sensitivity to the quarter-wave plate angle (Fig. 3), the long-term stability of the microscope is outstanding. It can be operated in a set-and-forget mode: an optical density close to 7, see Fig. 4, is achieved over an arbitrary period of time, exceeding typical measurement times by orders of magnitude.

IV. RESONANCE FLUORESCENCE ON A SINGLE QUANTUM DOT

Once the required high laser suppression is realized, the resonance fluorescence signal-to-background ratio on a single quantum dot is measured. Resonance fluorescence spectra of the single negatively charged exciton X\textsuperscript{1−} recorded at different laser powers and zero magnetic field are shown in Fig. 5. The lineshape of the optical resonance is Lorentzian, the full-width at half-maximum (FWHM) is 1.6 μeV at “low” power and 7.1 μeV at “high” power. The increase in linewidth with power reflects power broadening. Whereas the background, the residual laser signal, increases linearly with laser power, the quantum dot emission saturates and, thus, the signal-to-background ratio is power dependent. At an excitation power below quantum dot saturation the signal-to-background ratio is as high as 39 000:1 (Fig. 5(a)). Above saturation, a ratio >10\textsuperscript{3}:1 (Fig. 5(b)) is achieved.

One experiment which requires a high signal-to-background ratio and long integration times (and hence a stable setup) is a g\textsuperscript{(2)} measurement, i.e., an intensity correlation experiment. Laser light and a stream of single photons exhibit quite different g\textsuperscript{(2)}(\tau = 0) values, 1 and 0, respectively, such that a leakage of laser light into the single photon stream is very detrimental. The time-dependence of g\textsuperscript{(2)} was measured with a Hanbury Brown-Twiss interferometer (Fig. 6). There is a very clear dip at time delay zero, demonstrating anti-bunching in the photon statistics of the neutral exciton X\textsuperscript{0}. Note that even with a single channel count rate of 250 kHz, an

![FIG. 4. Long-term behaviour. The microscope is aligned to reject the laser reflected at the quantum dot sample (GaAs plus thin metal layer), and the residual counts are recorded by a single photon detector. The optical density (OD), defined as OD = \(-\log (1/T)\) with transmission \(T\), is plotted as a function of time. The microscope is stable over many hours with an OD > 6.8.](image1)

![FIG. 5. Resonance fluorescence on a single InGaAs quantum dot with different optical Rabi couplings. Resonance fluorescence spectra are recorded with a single photon detector at constant laser frequency. Detuning is achieved by sweeping the gate voltage with respect to the laser frequency. (a) Below quantum dot saturation, at an excitation power corresponding to a Rabi energy \(\Omega\) of 0.7 μeV, a signal-to-background ratio of 39 000:1 is achieved. (b) At high pump power, where power broadening dominates the optical linewidth, a signal-to-background ratio >10\textsuperscript{3}:1 is realized. Solid red lines show Lorentzian fits to the data (black points), blue dashed lines indicate the background.](image2)

![FIG. 6. g\textsuperscript{(2)} measurement of the resonance fluorescence from the neutral exciton X\textsuperscript{0} in a single InGaAs quantum dot. A clear dip at zero time delay demonstrates photon anti-bunching. The red curve shows the convolution of the two-level atom result, \(g_2(t) = 1 - [\cos(\lambda t) + 3(2\tau_\lambda \sin(\lambda t))] \exp(-3t/(2\tau))\) with \(\lambda = \Omega^2 - 1/(4\tau^2)\). Rabi frequency \(\Omega\), and radiative lifetime \(\tau\), with the response of the detectors (Gaussian with FWHM 0.67 ns) and provides a very good description of the data (black points). The blue curve shows the two-level atom response only. A lifetime of \(\tau = (1.0 \pm 0.1)\) ns and a Rabi frequency \(\Omega = (0.9 \pm 0.1)\) μeV were determined by fitting the data to the convolution. The measurement time was 9 h with a single channel count rate of 250 kHz.](image3)
integration time of \( \sim 9 \) h was required to achieve a high signal-to-noise ratio in the \( g^{(2)} \) measurement: the stability of the dark-field microscope was clearly important. It turns out that the residual value \( g^{(2)}(\tau = 0) \) is determined entirely (within the signal:noise) by the jitter in the detectors (\( \sim 0.6 \) ns), which is comparable to the radiative decay time (\( \sim 1 \) ns). Within error (\( \sim 1\% \)), the true quantum dot \( g^{(2)}(\tau = 0) \) is 0.00.

The resonance fluorescence, presented in Figs. 5 and 6, was measured on different excitons, the single negatively charged exciton \( X_{1}^{-} \) and the neutral \( X_{0} \), respectively. The resonance fluorescence of the \( X_{0} \) is linearly polarized (\( \pi \) or \( \pi_{0} \)); the resonance fluorescence of the \( X_{1}^{-} \) is unpolarized in the absence of a magnetic field. \( B = 0 \), circularly polarized (\( \sigma ^{+} \) or \( \sigma^{-} \)) for \( B \neq 0 \). The optics of the dark-field microscope define linear \( s \) for the excitation and linear \( p \) for the detection polarization. Nevertheless, resonance fluorescence of both optically active excitons can be measured independent of the selection rules, provided that the sample and microscope axes are not aligned. Ideally, the \( s/p \) basis is rotated by 45\(^{\circ} \) with respect to the \( \pi_{1}/\pi_{0} \) basis.

The dark-field microscope works well across the entire ensemble of quantum dots spanning a bandwidth of about 60 nm in wavelength. The dark-field point is so sensitive to the polarization axes that small achromatistics in the polarizers play a role: a change in wavelength \( \Delta \lambda \) requires a re-adjustment of the quarter-wave plate and linear polarizer alignments for optimum dark-field performance, typically a few tens of mdeg for \( \Delta \lambda = 1 \) nm. Furthermore, resonance fluorescence on a single quantum dot can be recorded not just at \( B = 0 \) but also at high \( B \). At high \( B \), a high suppression of scattered laser light can be achieved. As for a change in wavelength, for optimum dark-field performance the quarter-wave plate and polarizer alignment have to be re-adjusted as the magnetic field increases. Crucial for the performance at high magnetic field is the linear polarizer angle, differing significantly (\( \sim 10\% \)) from the zero field angle probably due to a Faraday effect\(^{29} \) of the objective lens, solid immersion lens or sample. Rotating the polarizer introduces a small \( p \)-polarization to the dominantly \( s \)-polarized beam propagating to the sample. A 10\(^{\circ} \) rotation results in a rotation of the polarization axis by \( \sim 1\% \) on account of the properties of the PBS. A resonance fluorescence spectrum of an \( X_{1}^{-} \) recorded at a magnetic field \( B \) of 4 T is shown in Fig. 7. The lineshape of the optical resonance is clearly non-Lorentzian, and there is a hysteresis between forward (red) and backward (blue) detuning. A dynamic nuclear spin polarization locks the quantum resonance to the laser energy as the gate voltage is tuned, an effect referred to as dragging.\(^{30,31} \)

As an outlook, we comment that the microscope can be developed further in some simple ways. For instance, given that the quantum dot basis (\( \pi_{1}/\pi_{0} \)) is dot-dependent,\(^{32} \) it may be valuable in the future to include also a way of rotating the microscope basis (\( s/p \)) relative to the \( \pi_{1}/\pi_{0} \) basis, either by inserting an additional wave plate or by rotating the sample. It may also be interesting to develop the capability of operating the microscope not with \( s/p \) polarizations but with \( \sigma^{+}/\sigma^{-} \) polarizations. Finally, we note that the quantum efficiency of the resonance fluorescence collection is limited by the high refractive index of the sample: light is refracted to such large angles at the GaAs/vacuum interface that it is collected inefficiently. In this experiment, this situation was rectified to some degree (factor of \( \sim 5 \) in signal strength) by the solid immersion lens. Despite this low quantum efficiency, the rejection of the scattered laser light in our dark-field microscope is more than sufficient to observe background-free resonance fluorescence from single quantum dots. The next step is therefore to increase the collection efficiency: the dark-field performance is already more than good enough. Candidate structures are resonant micro-cavities, photonic nanowires, or following the spirit of these experiments, ultra-high index solid immersion lenses.

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FIG. 7. Resonance fluorescence spectra of a single InGaAs quantum dot in a magnetic field. The laser suppression at high magnetic field is as good as that achieved at zero magnetic field. At \( B = 4 \) T the lineshape is a top hat and there is a hysteresis between forward and backward scanning directions. This effect is referred to as dragging.\(^{30,31} \)