

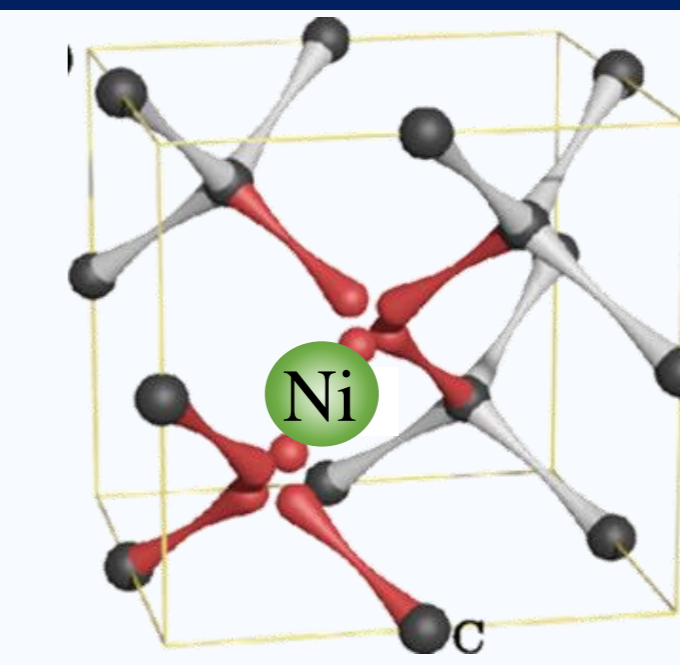


## Abstract

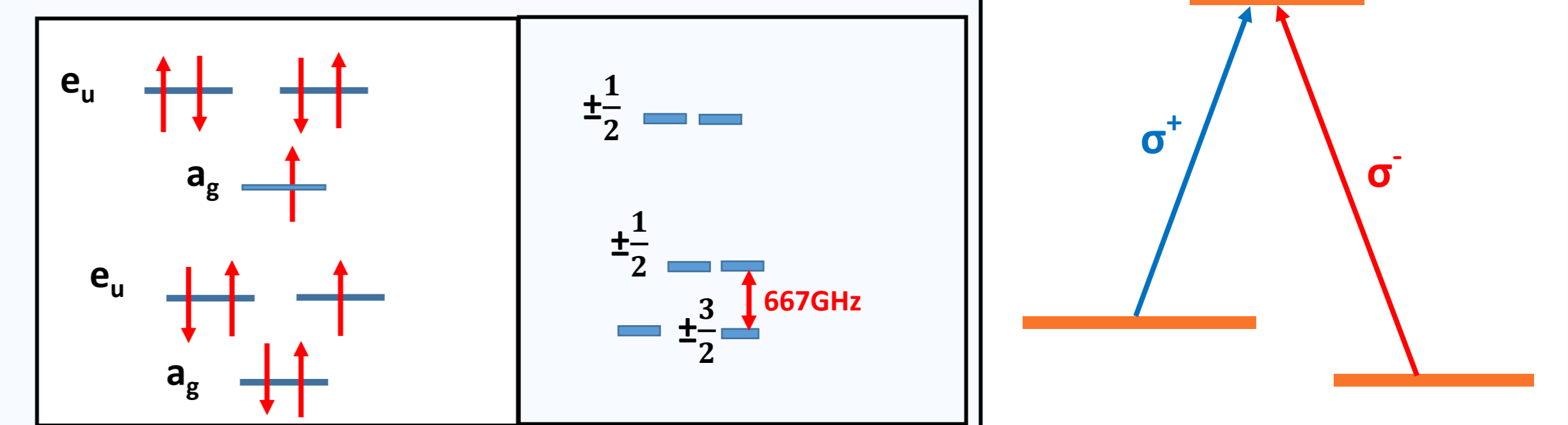
We study **nickel-related color centers** in diamond towards their application in a **solid-state light-spin interface**. These centers have strong spin-orbit interaction, thus are potentially suitable for high-temperature operation. Using polarized resonant excitation we observe signs of both spin and charge optical pumping at 10K.

## Background

- Diamond is rich in point defects that are able to change the optical and electrical properties of the host material, often called color centers.
- Nickel is a typical contaminant in high pressure high temperature (HPHT) diamonds.
- The 883/885 nm (1.4 eV) color center, is attributed to NiV<sup>-</sup> (S=1/2)
- +1/2 and -1/2 spin ground states couple through a single **polarized lambda system**.
- Light-spin interaction is mediated by spin-orbit coupling. For coherent operation it has to surpass the dephasing rate.
- Due to the Ni atom, the spin-orbit coupling rate is ~700 GHz – equivalent to a temperature of 35K



Polarized Lambda System



Atomic and electronic structure of the nickel-vacancy defect in diamond

## Rational

### Goals:

- Spin pumping by polarized resonant excitation
- Stabilizing -1 charge state by off-resonant excitation

### Research question:

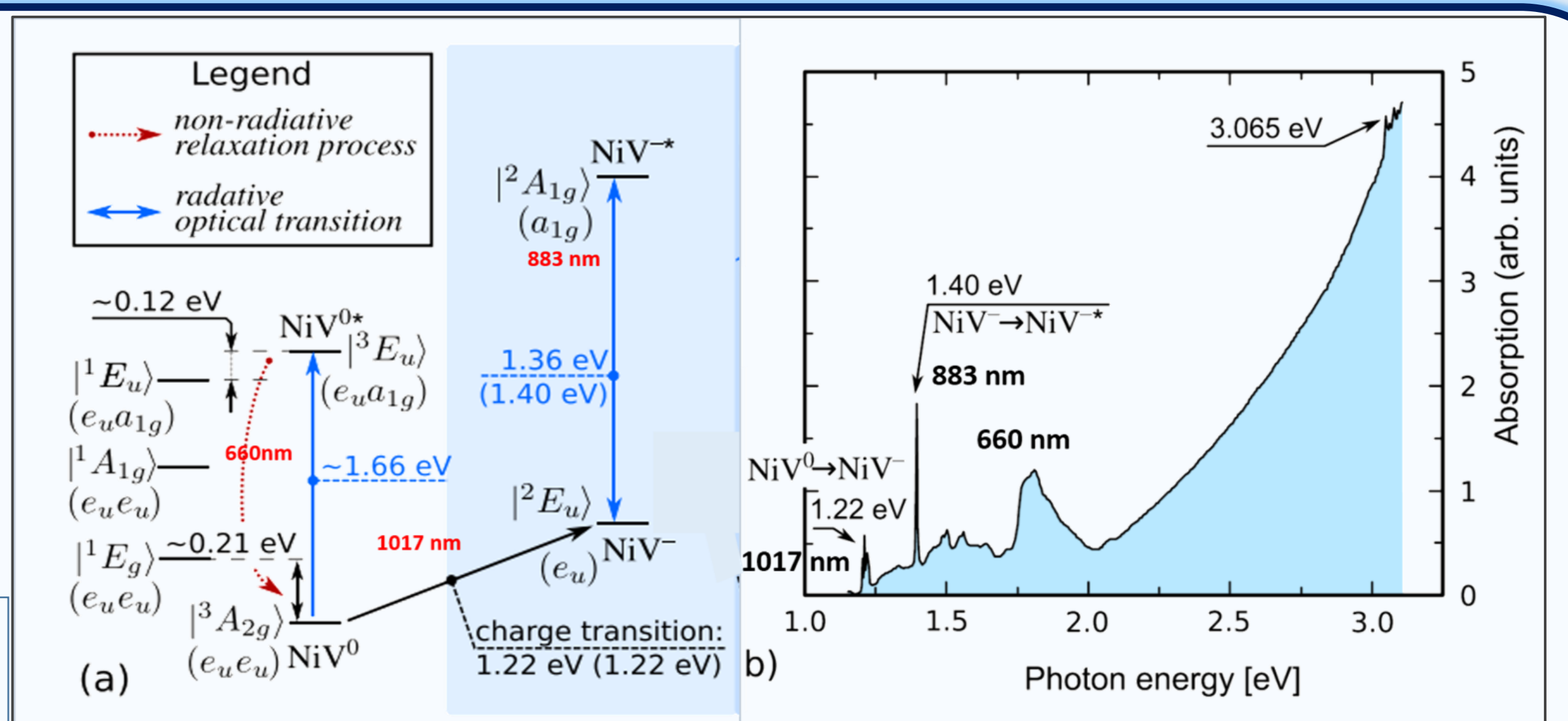
What wavelength and power are required for optimal charge stabilization?

- Candidates: 660 nm and 1017 nm (two sharp absorption resonances)

### Experimental methods:

- Simultaneous and sequential two-color excitation
- Polarization pulsed excitation

Proposed model: Gergő Thiering, and Adam Gali. "Magneto-optical spectra of the split nickel-vacancy defect in diamond." *Physical Review Research* 3, 043052 (2021).



## Sample and Experimental Setup

### Sample

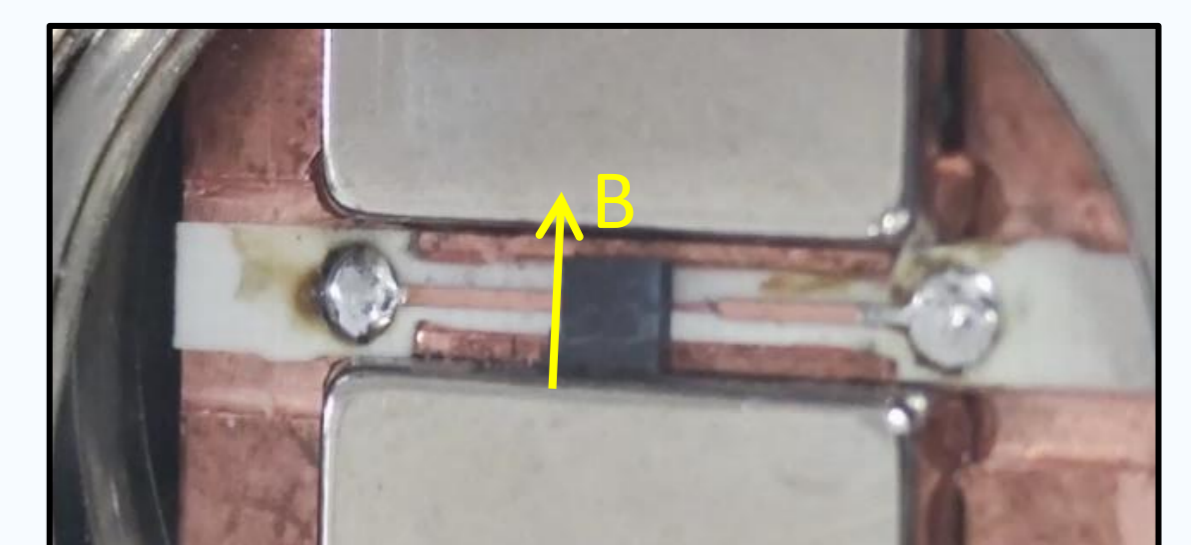
- The substrate is a <111> single-sector type IIb HPHT-grown artificial diamond (New Diamond Technology)
- Nickel was ion-implanted at 2 MeV at a dose of  $5 \times 10^{12} \text{ cm}^{-3}$  (Uni. Surry).
- The sample was then HPHT treated at 2000C and 8 Gpa for 2 hours (Element Six).

### Experimental setup

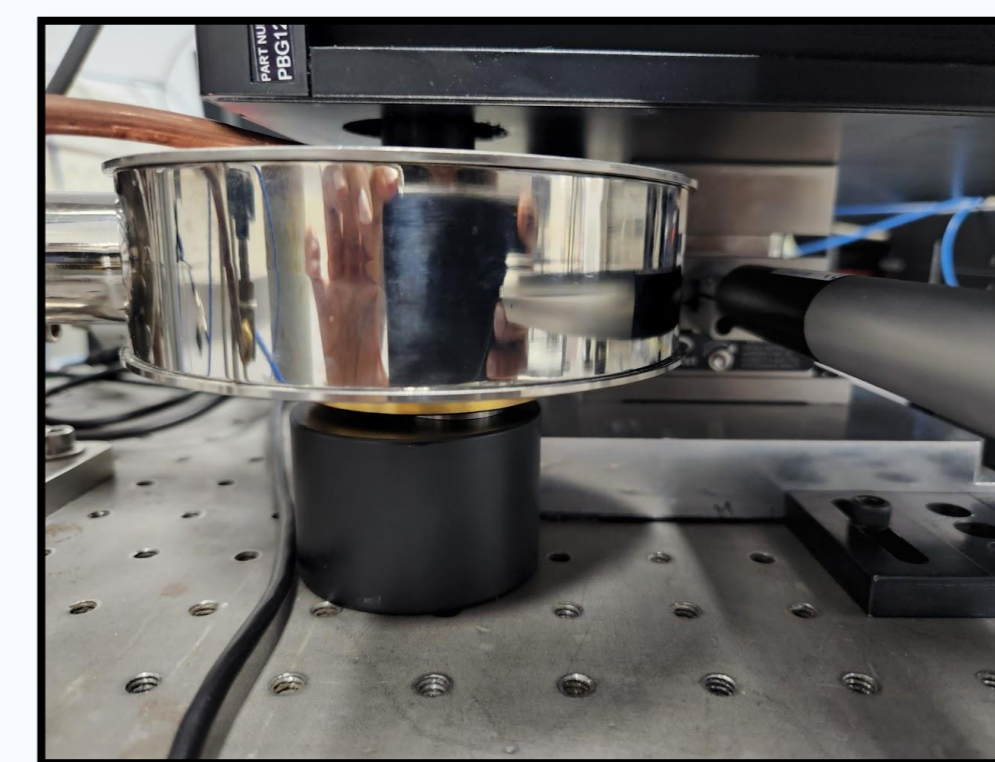
- 10K closed-cycle cryostat (ARS). Permanent magnets inside.
- Confocal microscope with three laser sources at 660 nm, 883 nm, and 1017 nm.
- 0.75 m spectrometer with a cooled ccd camera (Teledyne-PI)
- Avalanche photo detector and time-tagger electronics (Swabian Instruments)



Ni-implanted <111> diamond samples



Blue IIb sample between the magnets inside the cryostat



Microscope objective above sample chamber and magnet below it

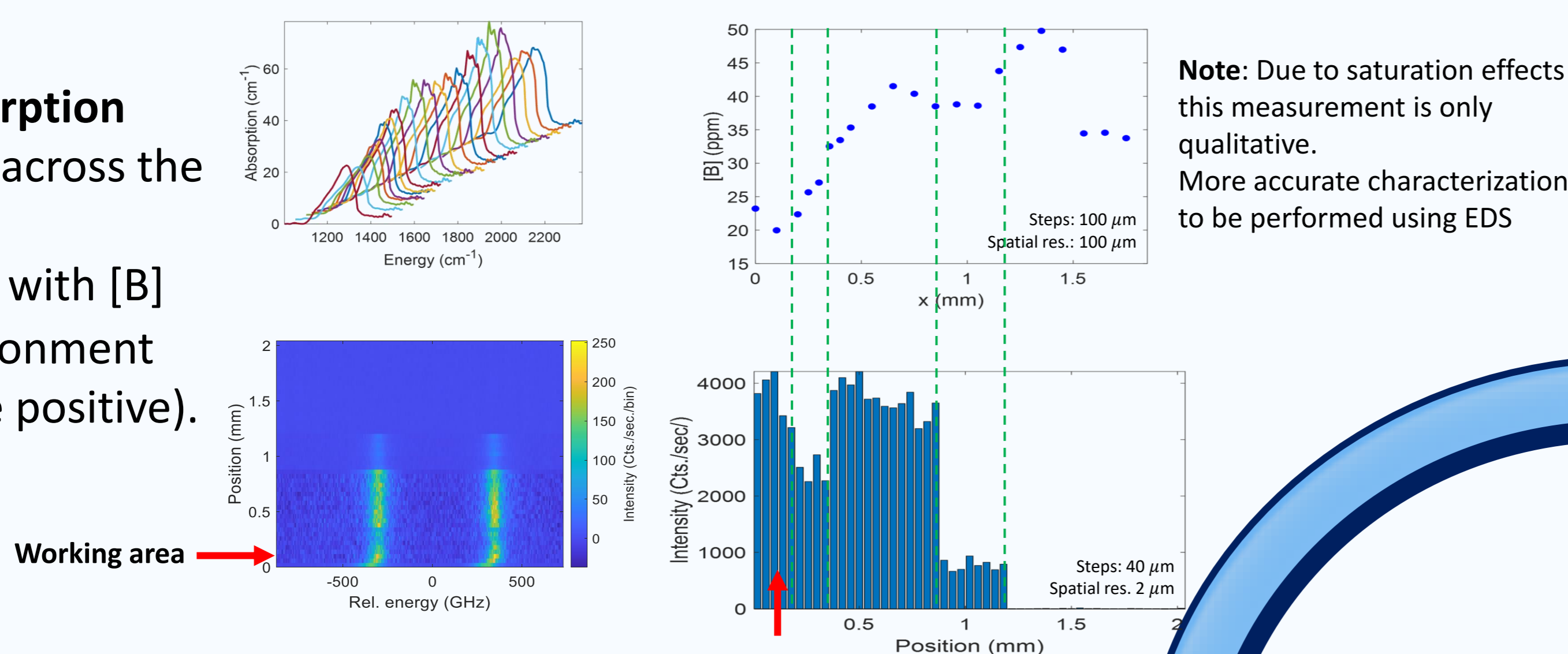
## Results

### I. Controlling the charge environment by adding boron (an acceptor)

#### FTIR absorption

[B] varies across the sample.

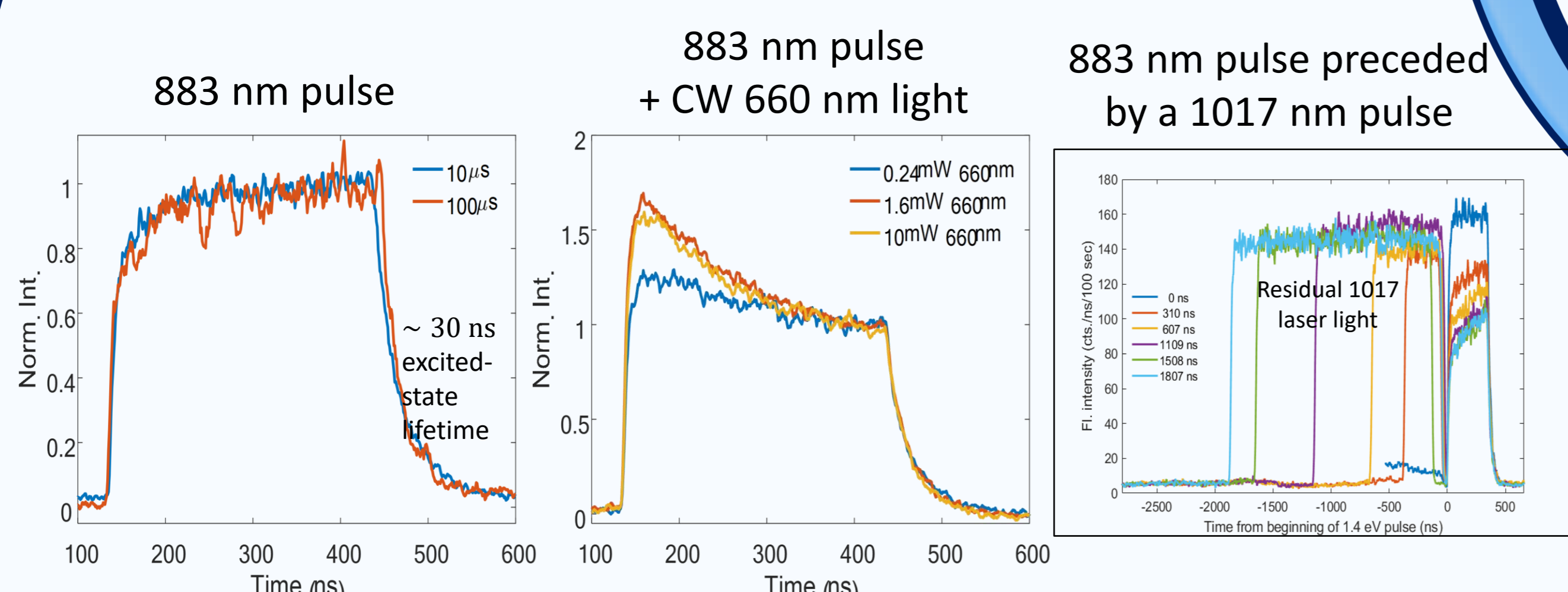
$E_f$  lowers with [B] (the environment gets more positive).



Note: Due to saturation effects this measurement is only qualitative. More accurate characterization to be performed using EDS

1.4 eV emission decreases across the sample: it comes from a charge state that is unstable in a too-positive environment

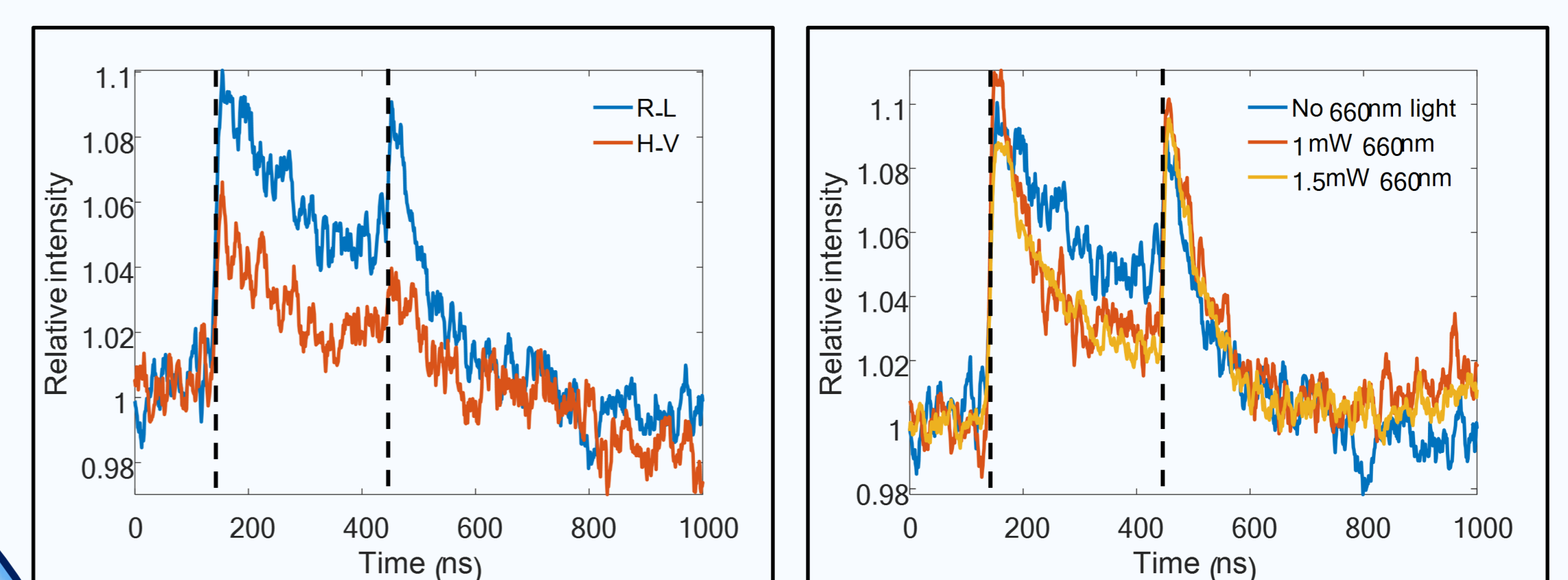
### II. Controlling fluorescence dynamics using an additional color



883 nm partially pumps out of the emitting state into a long-lived (>100 us) dark state. 660 nm light shortens the dark state's lifetime by pumping back into bright state. 1017 nm light pumps more population out of the bright state.

### III. Polarization pulse

Change only the polarization of the 883 nm light



Polarization alone cannot affect charge state

→ The 'spikes' are an evidence for spin pumping induced by circularly-polarized resonant excitation

## Combined charge and spin dynamics

### Conclusions and future work

#### Three simplest options :

- Dark state is an opposite **spin state**, with very long lifetime. 660 scrambles the spin state.
- Dark state is a different **charge state**. 1017 charges to that state, and 660 charges back.
- There are **two dark states**, one of a different spin and the other of a different charge. 660 resets both.

#### How to discriminate?

- Introduce a 660 nm 'reset' pulse, just before the 883 pulse.
- Look for polarization dependence of the fluorescence dynamics.
  - If there's none, it's only charge pumping.
  - Otherwise, spin pumping is present as well, and their ratio can be extracted