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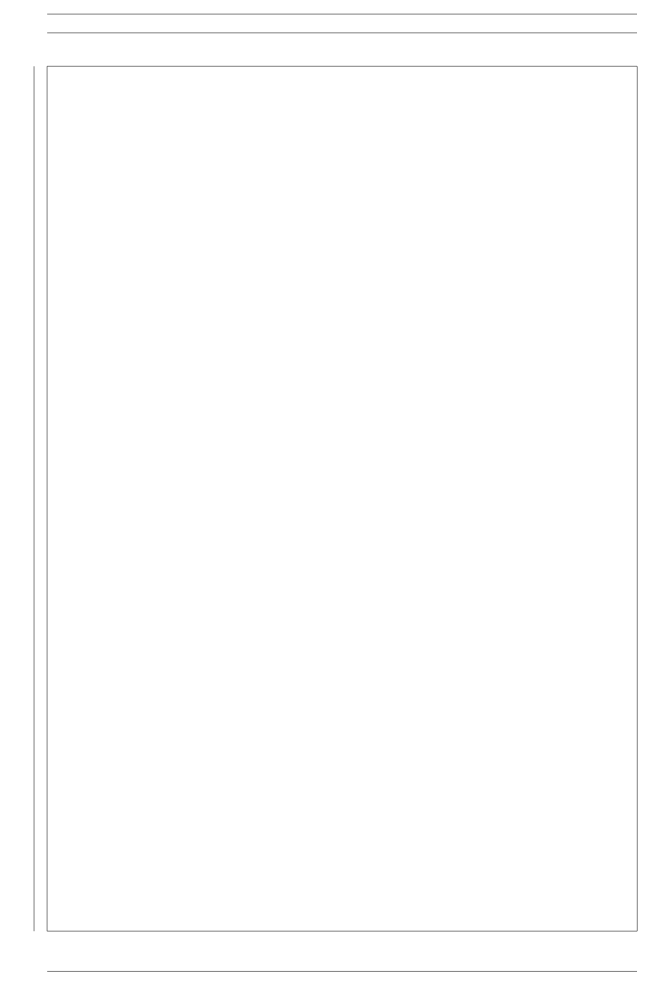
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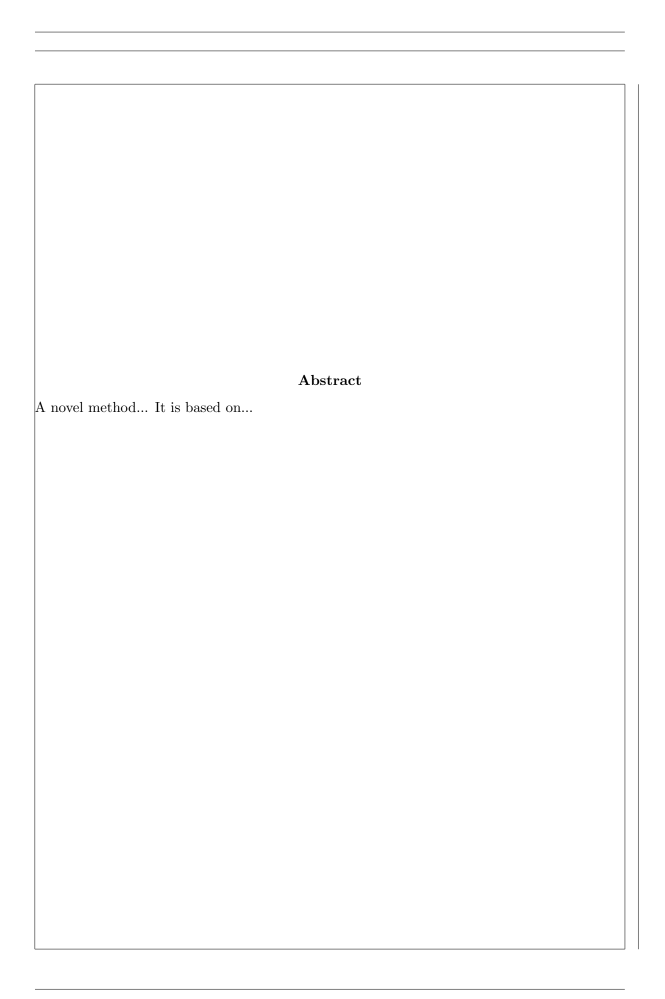
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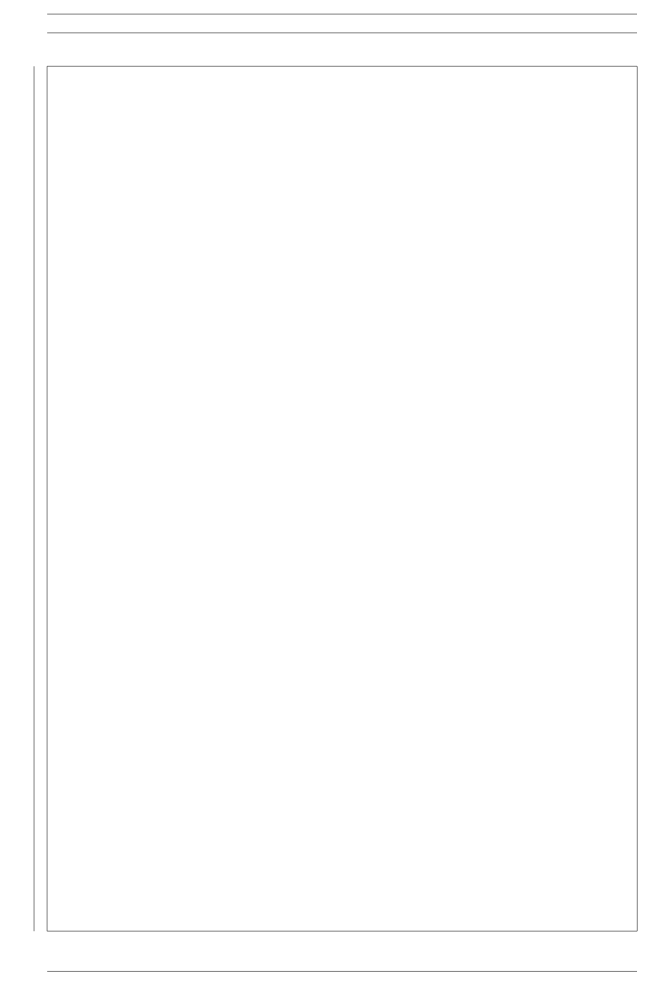
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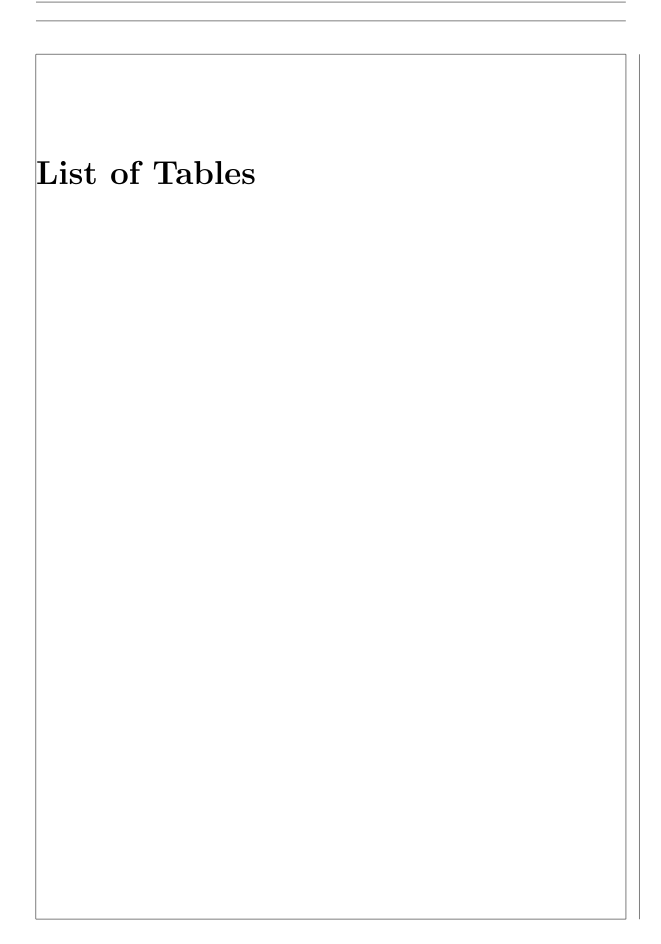


Contents

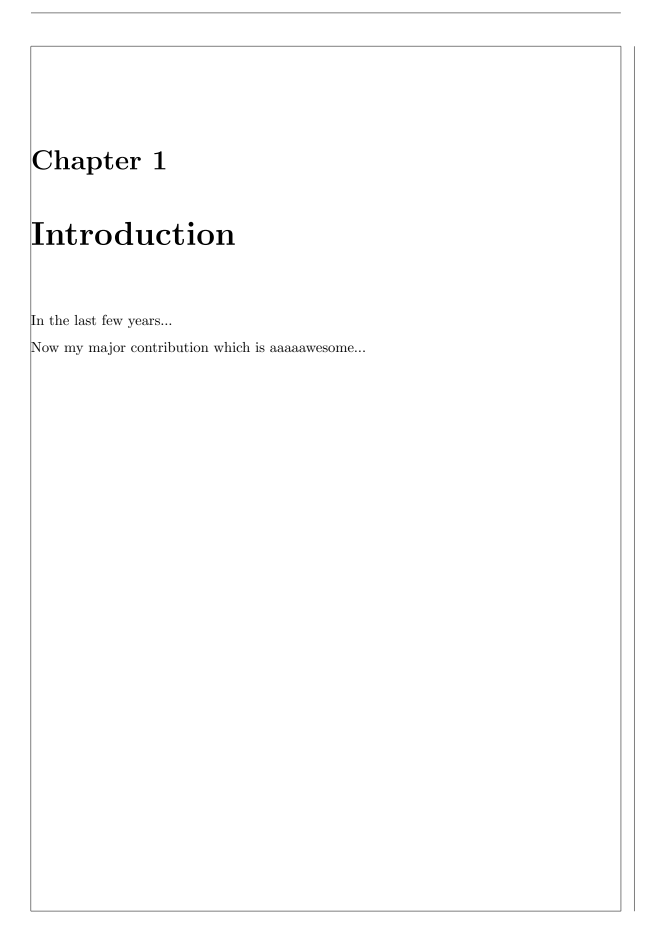
Ta	able of Contents	i
Li	st of Figures	i
Li	st of Tables	ii
\mathbf{A}_{0}	${f cknowledgements}$	iii
1	Introduction	1
${f 2}$	The Awesome Theory	2
3	Confocal Setup	3
4	Experimental Results 4.1 Diamond Characteristics 4.1.1 Raman Measurements 4.1.2 TEM 4.2 Photoluminescence spectra 4.2.1 Sideband 4.3 g2 Measurements 4.4 Photostability discussion	6 6 6 6 6 6 6 7
6	Conclusion	8
Bi	Text of Minor Interest ibliography	9 11 12
		- -

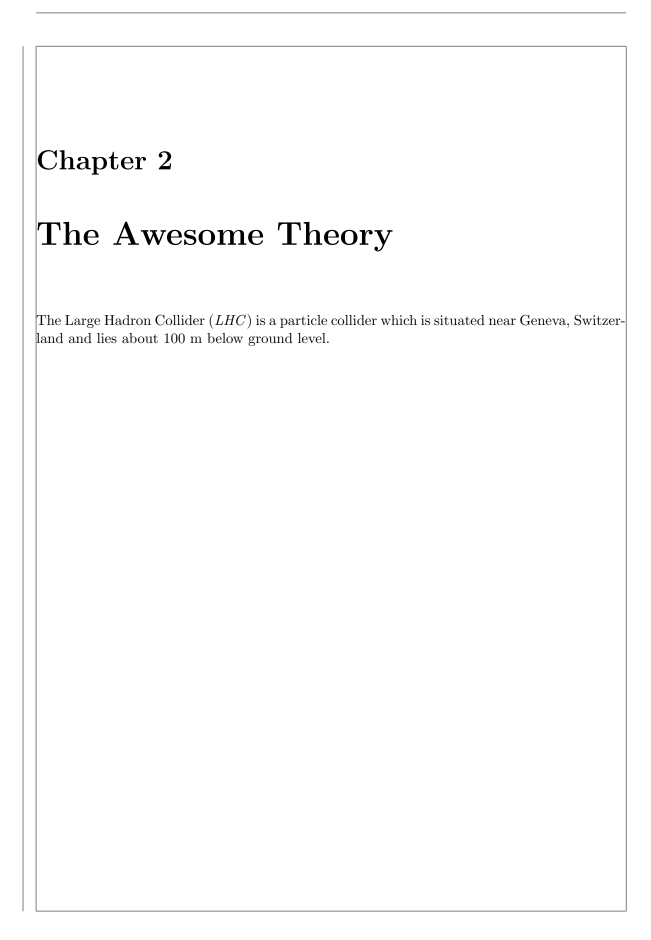
List of Figures

3.1	Confocal setup	 4
3.2	HBT, spectrometer	 Ę,



Acknowledgements First of all, I want to thank my supervisor... I am very grateful for the guiding help of... I am grateful to...





Chapter 3

Confocal Setup

The key measurements of this thesis are fluorescence measurements of SiV centers in nanodiamonds. For this aim, a home-built confocal setup is used, which is described in this chapter.

The confocal setup serves to perform a series of measurements on fluorescence light: scanning the sample to find SiV centers, recording luminescence spectra of the aforementioned, determine the saturation count rate , and determine whether the emitter in question is a single emitter by performing photon autocorrelation measurements. The two key components for these measurements are

- A Hanbury-Brown and Twiss setup to investigate the single photon character. It is built up of two avalanche photo diodes (APDs) which also serve to scan the sample in order
- A grating spectrometer to investigate the spectral properties.

Figure 3.1 depicts a sketch of the confocal setup. Except for the laser and the sample stage, the whole setup is fixed to a vertical breadboard. This design allows for easy exchanging of the samples, without the need of gluing them to a vertical stage.

to find emitters on the sample surface; and to perform saturation measurements.

The friction between the sample and the aluminum surface of the stage is sufficient that the sample does not move during scanning. If it is important that the sample has a defined orientation, it is put inside of an aluminum angle. The stage is powered with two stepper motors () in the horizontal x and y directions. The objective is fixed to another stage which in turn is fixed to the vertical breadboard. In this way, the vertical z direction is implemented for focusing the laser light on the sample.

The bright red color at the left-hand side of the sketch represents the excitation beam path. The sample is excited with a continuous wave diode laser (Schäfter-Kirchhoff, 58FCM) which emits at a wavelength of 660 nm. The outlet of the light is through a pigtail fiber, the light is outcoupled and collimated exploiting an aspheric lens. To suppress sideband emission from the laser, a bandpass filter with a window of 10 nm around a center of 660 nm is used. The excitation beam then hits a glass plate (fabricator Halle Germany) to be guided through a microscope objective and focused on the sample. The microscope objective is of the type Olympus, LMPlanFLN 100x and has a numerical aperture of 0.8. As the luminescence light

saturation

introduced

hickness

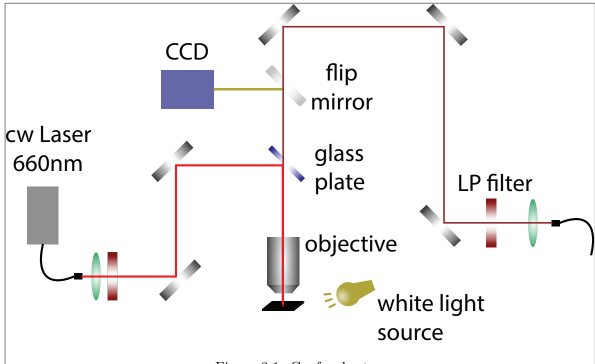


Figure 3.1: Confocal setup

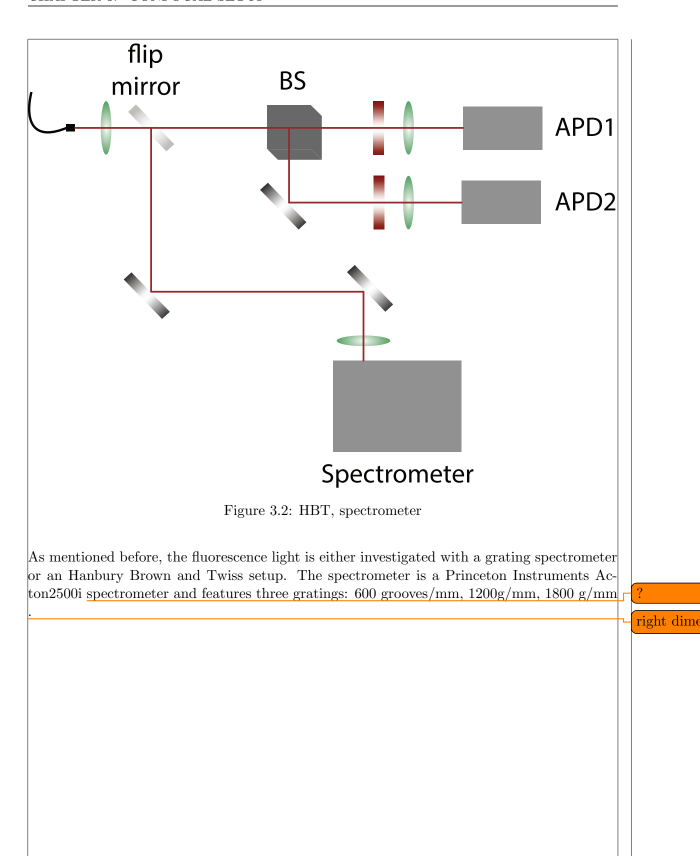
from the emitter is in the same focus as the excitation laser light, it is effectively collected by the objective. Hence also the name confocal setup.

The collected light then follows the detection beam path depicted in a dark red color in Figure 3.1. Both the excitation light reflected from the sample surface and the fluorescence light pass through the glass plate. In the usual useage, the flip mirror just after the beamsplitter is lowered, allowing the light coming from the sample to move on towards a single mode fiber. In front of the single mode fiber there is a longpass filter to filter out the residual excitation light and also ambient light. The fluorescence light is fed into a single-mode fiber (Thorlabs SM600) with an aspheric lens. The single-mode fiber serves two purpuses: First, to connect the confocal microscope with the Hanbury Brown and Twiss setup and the spectrometer. Second, its about 4.3 µm diameter serves as a pinhole to ensure to collect only fluorescence light in the focal spot besser erklren).

According to the experimental necessities, instead of the mentioned glass plate a dichroic mirror () can be employed. The glass plate features a high transmission of 90% and therefore a high collection efficiency of fluorescence light, the dichroic mirror allows for a higher excitation intensity using the same excitation laser. However, a high excitation intensity may cause permanent fluorescence intermittence of the SiV centers (for further detail, refer to). In general, if a high excitation is necessary, for instance for saturation measurements, the dichroic mirror is used; otherwise, the glass plate is used to collect as much fluorescence light as possible.

Another feature of the setup is that it is possible to have a look at the sample surface before starting the fluorescence measurements. For this purpose, the sample is illuminated with white light from a halogen lamp and the flip mirror after the glass plate is brought into an upright position to guide the light onto a CCD camera ().

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Chapter 4

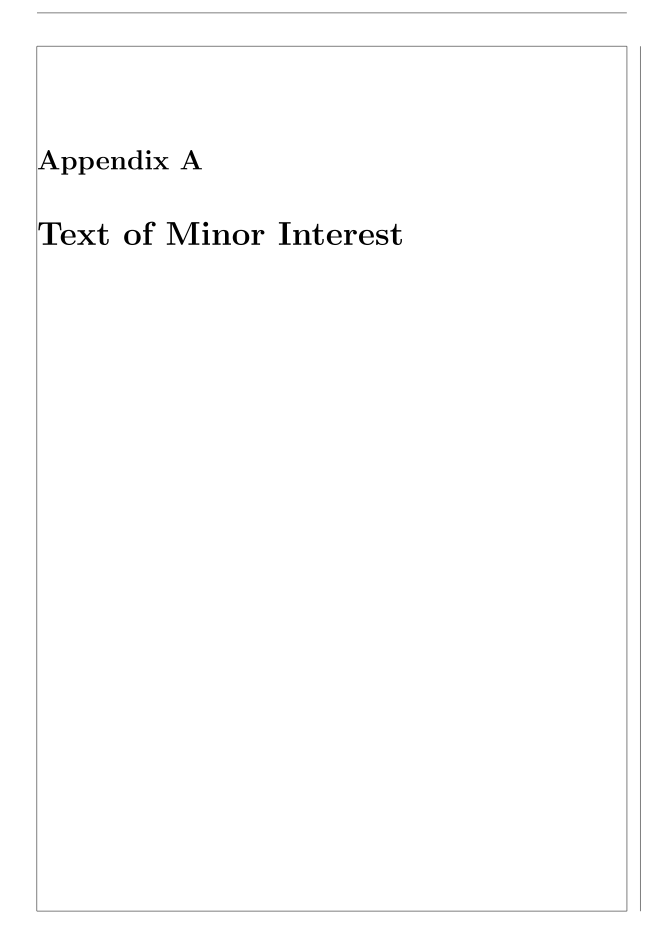
Experimental Results

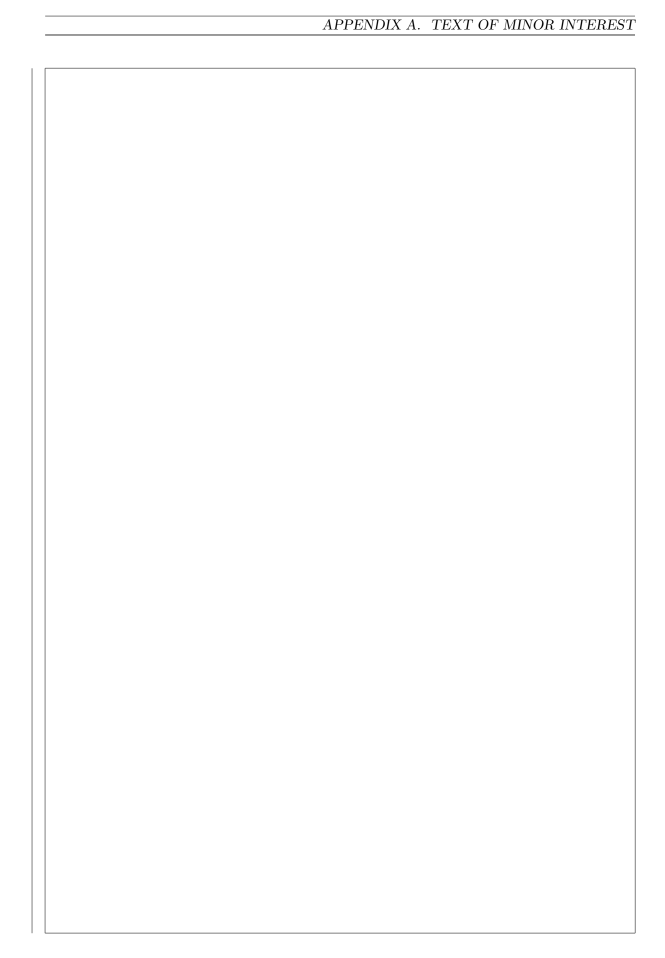
In the following paragraphs, both phenomenological of the nanodiamonds and spectroscopic measurement of the SiV centers are described. Unless explicitly otherwise stated, the results in this paper report measurements of the milled nanodiamonds containing *in-situ* incorporated SiV centers.

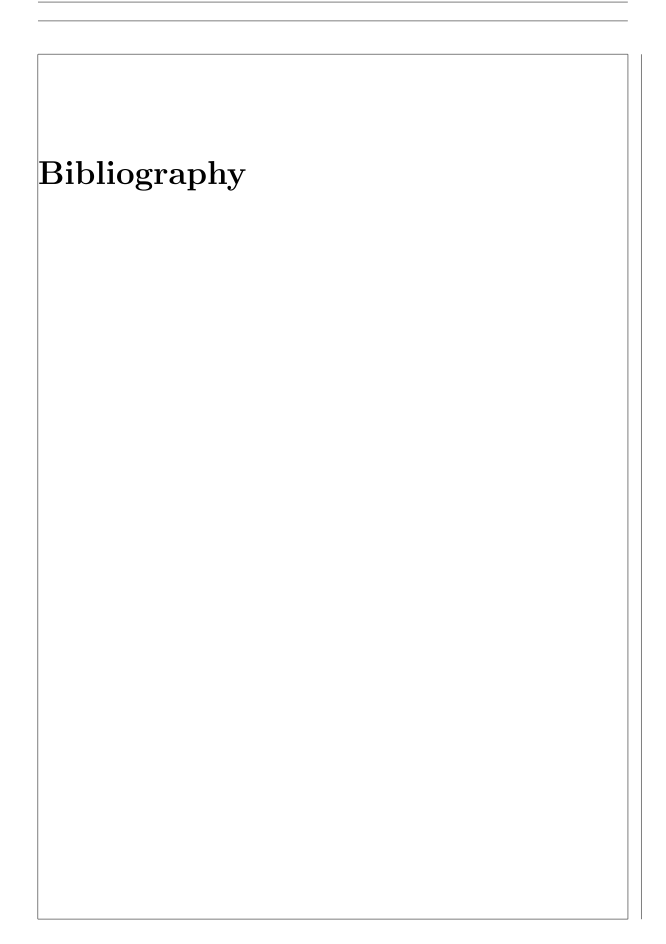
- 4.1 Diamond Characteristics
- 4.1.1 Raman Measurements
- 4.1.2 Transmission Electron Microscopy
- 4.2 Photoluminescence spectra
- 4.2.1 Sideband
- 4.3 Photon correlation measurements
- 4.4 Photostability

Chapter 5		
discussion		

Chapter 6
Conclusion
In conclusion







Index		
LHC, 2		