



Leibniz
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Hannover



Master's Thesis

**Camera based electroluminescence measurements of
perovskite solar cells.**

A thesis presented for the degree Master of Science.

Jonas Oberröhrmann
Institute for solar energy research Hamelin (ISFH)

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1 Introduction

lorem ipsum.

2 Theory of Electroluminescence

This chapter explains the theoretical background of Electroluminescence (EL). Starting with the rate of radiating recombination, depending on the material's band structure (see section XY). The rate depends on the carrier concentration, so section XY explains the carrier density profile. Section XY ends with the light path of the radiation out of the sample.

2.1 Rau's relationship

The basis for our discussion of EL emission is the relationship given by Rau (2007):

$$\frac{d\Phi}{d\vec{r}}(\vec{r}) = \alpha(\vec{r}) n(\vec{r})^2(\vec{r}) \Phi_{BB} u(\vec{r}). \quad (2.1)$$

Because all the volume elements $dV(\vec{r})$ contribute to the emitted photon flux we integrate the photon flux:

$$\Phi_{em}(\vec{r}_S, \Omega_S, E_\gamma) = \int T(\vec{r}, \vec{r}_S) \alpha(\vec{r}) \Phi_{BB}(E_\gamma) u(\vec{r}) d\vec{r}, \quad (2.2)$$

for the electromagnetic flux at \vec{r}_S , into spacial angle Ω_S at photon energy E_γ . Inserting Donolato's reciprocity theorem (?) for $u(\vec{r})$ yields (CITE):

$$\Phi_{em}(\vec{r}_S, \Omega_S, E_\gamma) = \int T(\vec{r}_s, \vec{r}) \alpha(\vec{r}) f_c(\vec{r}) d\vec{r} \Phi_{BB}(E_\gamma) \left[\exp\left(\frac{qV}{kT}\right) - 1 \right]. \quad (2.3)$$

2.2 The Principle of detailed balance

W. van Roosbroeck and W. Shockley presented 1954 the principle of detailed balance, showing the relation between the rate of radiative recombination and optical properties, such as the absorbtion coefficient α (CITE).

2.3 Quantum theoretical theory for the absorption and emission of light

¹ BEGIN WITH VECTOR POTENTIAL

The quantum theoretical treatment of light emission or absorption starts with the interaction of electrons with electromagnetic radiation, described by the Schrodinger Equation

$$H\Psi = E\Psi \quad (2.4)$$

with the Wavefunction Ψ , Eigenenergy E and Hamiltonian H (SOURCE):

$$H = \frac{(p + eA)^2}{2m} + V. \quad (2.5)$$

Neglecting terms of order $(eA)^2$, the Hamiltonian can be separated into the unperturbed part H_0 and the electromagnetic interaction part:

$$H = H_0 + H_i = H_0 + \frac{(eA)^2}{2m} \quad (2.6)$$

and H_0 :

$$H_0 = \frac{p^2}{2m} + V. \quad (2.7)$$

In crystals the potential is periodic in space and energy bands are formed, consisting of allowed states and energy gaps. The solution to the unperturbed Hamiltonian H_0 can be calculated approximately by ... and confirmed by experimental techniques (UPS, ...)(PICTURE). That way the Energies E_0 are known and the solution to the perturbed hamiltonian Equation 2.6 can be approximated by perturbation theory (SOURCE). This leads to interaction terms, mixing different states and allowing transistions obeing transistion rules. First order perturbation theory gives a transition propability from the initial state $|i\rangle$ to final state $|f\rangle$:

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} |\langle f | H_i | i \rangle|^2 \delta(E_f - E_i \mp \hbar\omega). \quad (2.8)$$

Inserting Equation 2.6 yields

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} \left(\frac{eA}{mc} \right)^2 |\vec{e} M_{fi}(\vec{k})|^2 \delta(E_f - E_i \mp \hbar\omega), \quad (2.9)$$

with $M_{fi}(\vec{k})$ being the matrix dipolemoment for states f and i :

$$\vec{e} M_{fi}(\vec{k}) = \langle f | \vec{e} \vec{p} | i \rangle = e \int \bar{\Psi}_f (-i\hbar \nabla) \Psi_i d\vec{r}. \quad (2.10)$$

To account for all possible transistions one has to integrate over all possible values \vec{k} and summing over all initial and final states, denoted by valence band v and

¹BassaniPastoriParravicini-ElectronicStatesandOpticalTransitionsinSolidsPergamonPress1975

conduction band c:

$$W(\hbar\omega) = \frac{2\pi}{\hbar} \left(\frac{eA}{mc} \right)^2 \sum_{v,c} \int \frac{2d\vec{k}}{(2\pi)^3} |\vec{e} M_{fi}(\vec{k})|^2 \delta(E_f - E_i \mp \hbar\omega). \quad (2.11)$$

Returning to the connection between transition propability and absorption coefficient α , which is defined as the ratio of absorbed energy in unit time and unit volume to incoming energy flux:

$$\alpha(\hbar\omega) = \frac{\text{energy absorbed in unit time and unit volume}}{\text{Energy flux}}. \quad (2.12)$$

The absorbed energy is the photon energy times the total transition rate $W(\hbar\omega)$. The incoming energy flux is calculated by the Poynting vector from vector field \vec{A} . This yields for α :

$$\alpha(\hbar\omega) = \frac{\hbar c 2\pi}{A_0^2 \omega n} W(\hbar\omega). \quad (2.13)$$

3 Perovskites and perovskite solar cells

This chapter gives an overview over the material perovskite, it's properties and current research, and perovskite solar cells. PSCs consist of several materials, each providing specific functionalities and need to be chosen accordingly to the perovskite.

3.1 Perovskite materials

3.2 Perovskite Solar Cells

4 Setup

This chapter presents the experimental setup for electroluminescence measurements. Successful EL measurements consist of the injection of charges into the recombination layer, charge recombination and luminescence detection. Therefore section 4.1 explains the general setup. The following section 4.2 explains the Perovskite Cell layout and electrical contacting of the cells, which is used to inject charges into the perovskite layer. The emitted radiation is detected by a camera with the use of additional optics, explained in section 4.3. The chapter concludes with the consideration of noise and measurement errors.

4.1 General Setup

The setup is enclosed in a black housing, shielding the inside from outside light and noise (see Figure 4.1). All parts inside the chamber are painted black to minimize internal reflections and thereby the detection of stray light. A vacuum pump, voltage source, multimeter and operating table with a computer are connected to the setup from the outside, providing further utility. Inside the enclosure are the probe holder and the camera. The camera is mounted on rails and can be moved in three dimensions, enabling position according to the used optics and samples.

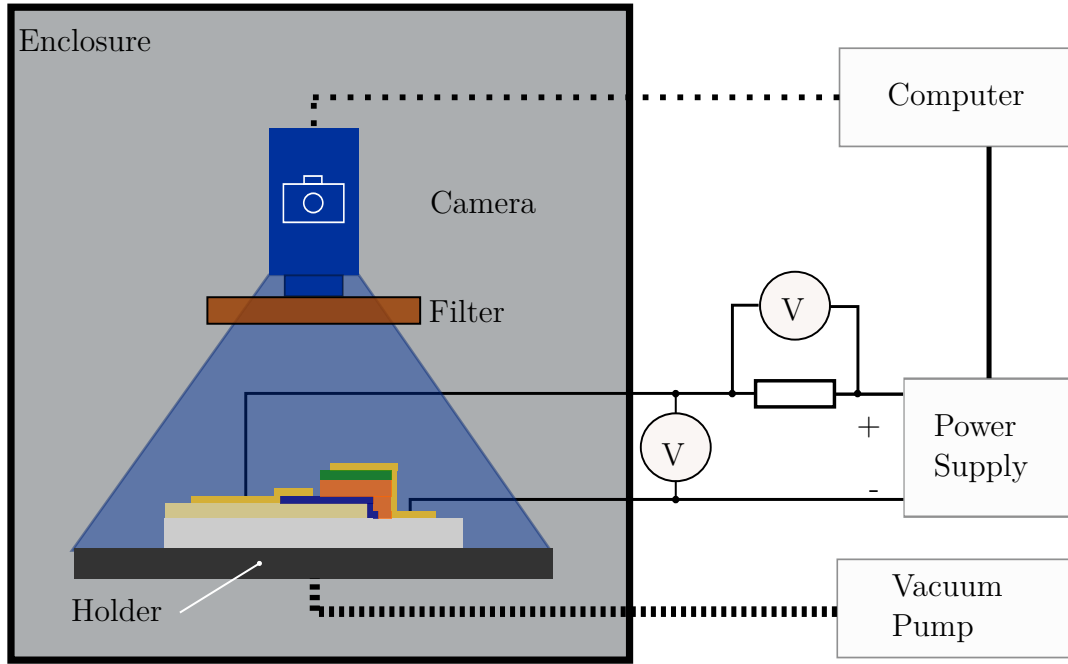


Figure 4.1: Schematic of the EL measurement setup. Camera and Probe are positioned inside the setup, and the computer, power supplies and measurement devices are provided from the outside. Two multimeters are used to measure the voltage drop and current sourced.

4.2 Electrical Connection

Perovskite Solar Cells (PSCs) are manufactured on 25 mm x 25 mm glass substrates (see Figure 4.2). Four cells are deposited on one substrate, each single cell having an active area of 4 mm x 3.5 mm. The cell stack consists of a 0.7 mm thick glass substrate, with a 100 nm thick layer of Indium Tin Oxide (short ITO) deposited on top. The ITO is structured by laser etching, isolating the individual cells from each other. Onto the ITO, 10 nm of Spiro-TTB are used as a Hole Transport Layer (HTL), followed by 500 nm of methylammonium lead triiodide (short MAPI) as the absorbing material. The top side of the MAPI is contacted with 8 nm of fullerene (short: C_{60}) and 23 nm bathocuproine (short: BCP), acting as the Electron Transport Layer (short ETL). In the end of the fabrication process, 100 nm Gold are deposited by thermal vapor deposition on top of the ITO and the C_{60} and BCP. To successfully contact the PSC a holder with four pins is used. The PSC is face down, with the glass substrate above, onto the holder, positioning the gold areas on all contacts. The holder uses two pins to check proper contacting of the PSC and the other two to source voltage.

The holder is electrically connected to a power supply¹, in series with a 100 m Ω resistance, R , and two multimeters² (see Figure 4.1). One multimeter measures the voltage drop across the PSC. The other multimeter measures the voltage drop across

¹Kokusuki

²Keithley 2000 Multimeter

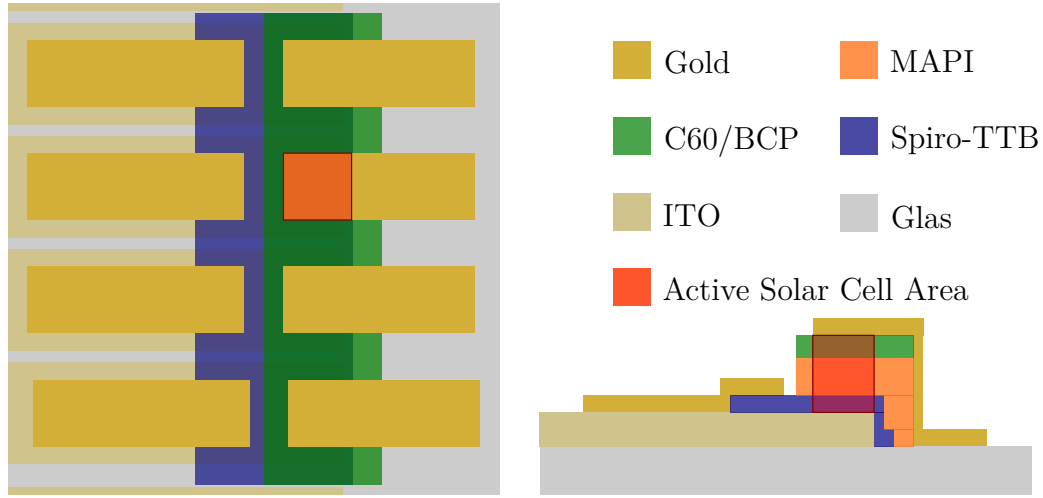


Figure 4.2: Schematic layout of the perovskite solar cell. Not to scale.

the resistor and the flowing current I is then calculated by Ohm's law:

$$I = \frac{U}{R}. \quad (4.1)$$

The surface of the holder is structured with rills and connected to a vacuum pump, fixing the PSC on the holder by underpressure. This ensures mechanical- and electrical contact stability throughout the measurement.

This setup enables safe contacting of the probes inside the enclosure and thus enables reliable EL measurements.

4.3 Luminescence detection

In the setup a CCD camera is used to detect the emitted electroluminescence. The camera detects radiation over a wide range of wavelengths, while the perovskite emits luminescence only at a relatively short range of wavelengths. Therefore filters are used to detect only the EL radiation. Optics focus the emitted radiation at a specific distance onto the imaging sensor. This chapter explains the used components and physical processes.

4.3.1 Filters

Between probe and sensor two lowpass filters³ are used to filter the wavelengths reaching the detector. In filters absorption or interference are used to either transmit or reflect specific wavelengths. The filter in the setup is a bandpass filter and chosen accordingly to the luminescence emission spectrum of the PSC (see Figure 4.3). This reduces the detected wavelengths to 780 nm to 800 nm, the interval with maximal

³LP714, LP718

intensity of the PSC, which enables maximum sensitivity to the emitted radiation and limits the detection of stray light. The filters are part of the photoluminescence capability of the setup, where light emitting diodes (LEDs) optically generated charges in the perovskite. Because of the high intensity and shorter wavelength the lowpass filters need to be included, for a higher SNR.

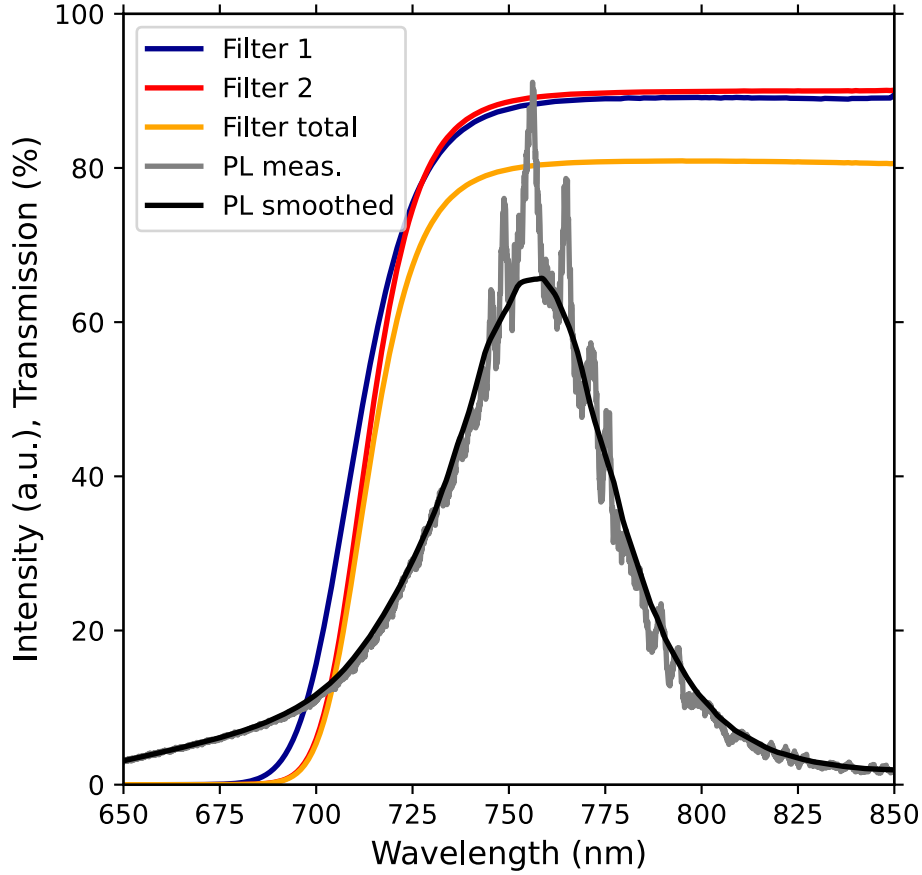


Figure 4.3: Emitted luminescence spectrum of the perovskite and transmission function of both filters.

4.3.2 Optics

Optics focus the radiation onto a charge coupled sensor (CCD). This setup uses a lens⁴ with a focus length of 25 mm and an aperture of 1.4, as listed by the manufacturer. The lens is positioned at the minimal object distance of 25 cm away from the PSC. An extender is used in between lens and camera, mounted directly to the camera with a C-mount, doubling the resolution of the setup. This decreases the intensity at the CCD camera, because of absorption.

⁴Pentax, C2514-M (KP)

SKETCH FOR LENS SYSTEM

4.3.3 Charge Coupled Detectors

The radiation is focused onto a charge coupled detector (CCD) sensor⁵ in a camera⁶. CCD sensors are silicon chips structured into small squares, called wells [SchnellCCD1993](see Figure 4.4). The number of wells correspond to the number of pixel in the taken picture. Radiation generates charges, electrons and holes, and externally applied voltages separate and trap the charges in the wells. For a specific time, called exposure time, radiation hits the sensor and charges accumulate in the wells. The amount of generated electrons per incoming photon is called the quantum efficiency (QE), and depends on the sensor material and energy of the photon. The QE for the chosen sensor peaks at 500 nm and decays for larger wavelengths (compare ??). For the wavelengths transmitted by the filter (see section before), the QE drops to about 10 %, which reduces the sensitivity. After the exposure time a series of voltages is applied to shift the charges from the light sensitive wells to opaque covered wells. Then the charges are extracted row by row and the voltage of each well is measured. An A/D converter converts the voltage into a digital signal, which is saved and later processed. The camera is connected to a computer by optical fibers and the acquired pictures are processed by the software LabView. The software enables an automatic control of the setup and the acquisition of multiple images in a row.

This process measures the EL distribution over the PSC surface. However several

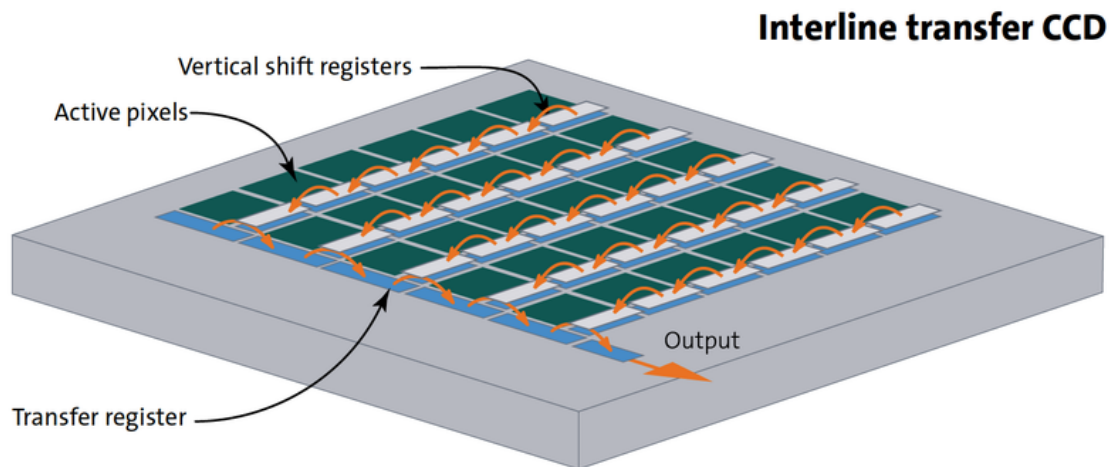


Figure 4.4: Schematic representation of a CCD sensor. Radiation generates charges in the active pixels and is transferred through vertical shift registers to the transfer register. Taken from [StemmerCCD].

⁵Sony ICX285-AL

⁶PCO, sensicam qe

sources of errors occur, which are discussed in the next section.

4.4 Noise and measurement errors

Several sources of errors deviate the measured signal from the physical value. Common errors in the camera image acquisition are dark noise, readout noise and hot or cold pixel. Dark noise refers to the thermal generation of electrons, depending on the temperature and the material's properties. To reduce thermal generation the CCD sensor is cooled to -12 C. To further reduce dark noise, Images without illumination or applied voltage are taken, and subtracted from the EL image.

Readout noise occurs, when the electrons are shifted from well to well. The manufacturer specifies the read out noise of 5 electrons. This relates to about one count, with an analog to digital conversion efficiency of 4 electrons per count.[**ManualSensicam**] EL images are taken at around 3000 counts, 75 % of the possible 4096 counts, showing that readout noise can in most cases be neglected.

ERROR BUDGET FOR MEASUREMENT

5 Experimental part

This chapter presents the experimental results of EL measurements on PSCs. Observed where the qualitative interpretation of perovskites, transient behaviour and the localization of defects in different materials of the PSC.