A Novel Imaging Spectrometer for
Energy-Distribution Measurements of
Photoelectrons from GaAs Cathodes

C. D. Schröter, A. Rudenko, A. Dorn, R. Moshammer and J. Ullrich

Max-Planck-Institut für Kernphysik, 69029 Heidelberg, Germany

Abstract. The investigation of the photoelectron-escape mechanism from GaAs cathodes with
negative electron affinity requires the detection of very low energy electrons. We have built a
novel, UHV-compatible, spectrometer where the photoelectrons are imaged by a homogeneous
electric field onto a position-sensitive detector. The time-of-flight of each single emitted electron
and its position on the detector is measured. From these informations energy-distribution curves
are extracted. The spectrometer has run successfully and preliminary energy-distribution curves
have been measured. The system is now under improvement. With the optimized spectrometer, an
excellent energy resolution (a few meV) can be achieved.

INTRODUCTION

To investigate the photoelectron-escape mechanism from a photocathode with negative
electron affinity, a spectrometer is required that allows the simultaneous measurement of
the energy and angular distributions of very low energy electrons. In the past, measure-
ments of the longitudinal energy distributions have been performed by several groups.
Quite recently, complete energy distributions have been studied as a function of longi-
tudinal and transverse energies. The employed method is based on selecting photoelec-
trons of a fixed longitudinal energy using a retarding field analyzer, and subsequently
measuring the associated differential transverse energy distribution by applying an adi-
abatic magnetic compression technique [1].

Recently we have built up a novel imaging spectrometer which allows the simultane-
ous measurement of the longitudinal and transverse momenta for each individual pho-
toelectron emitted from a GaAs surface [2]. Our spectrometer has run successfully and
preliminary energy-distribution curves (EDC's) have been extracted from this informa-
tion. However, the first tests have been performed under non-ideal conditions with high
extraction fields and, hence, the thus far measured EDC's have lower resolution than
what is ultimately achievable. Design changes have been made to be able to measure
EDC's of photoelectrons with an improved energy resolution in the near future.

EXPERIMENTAL SET-UP

A schematic diagram of the spectrometer set-up is shown in figure 1. The photocath-
ode is illuminated by a short-pulsed laser diode (pulse width ~ 100 ps). The laser focus
spot size and its position on the cathode is controlled by a CCD camera. For the mea-
measurements, highly doped p-type reflection mode GaAs/AlGaAs heterostructure crystals (6 \times 10^{18} \text{ Zn/cm}^2) are used with an active GaAs layer of 0.9 \mu m. For such a thin layer, the long tail of the electron pulse, generated by a \( \delta \)-pulse light excitation, extends out to less than 200 ps [3]. Hence, the photoelectron pulse width will be less than 300 ps and therefore short enough to not limit the final resolution of the electrons' time-of-flight (TOF) measurement. Details of the preparation technique of the photocathodes are described elsewhere [2, 4].

![Diagram of UHV chamber with spectrometer and MCP detector.](image)

Emitted photoelectrons are projected by a homogeneous electric field onto a position-sensitive micro-channel plate (MCP) detector. The homogeneous field is produced by means of 20 cylindrical electrodes on which equidistant potentials are applied. The TOF of each single emitted photoelectron and its position on the detector are measured. From the TOF, the longitudinal momentum of the electron can be determined and from the position on the MCP detector, its transverse momentum can be extracted.

Computer simulations show that with the spectrometer an excellent energy resolution can be achieved (see figure 2). But in order to reach the highest resolution a very homogeneous electric field is required to guide the electrons to the detector. In the transverse direction, high resolution can be obtained even at high extraction fields; however, in the longitudinal direction, high resolution is achieved only for the smallest electric-field strengths. Then, even small electric and magnetic stray fields will influence the parabolic trajectories of the photoelectrons. Thus, for the whole apparatus, only UHV-compatible materials with very low permeability have been used. External fields are compensated by three pairs of Helmholtz coils installed pairwise, perpendicular to each other. The magnetic stray fields originating from ion getter pumps are shielded by \( \mu \)-metal housings.
FIGURE 2. (a) Longitudinal and (b) transverse energy resolution of the spectrometer for electric field strengths of 150 V/m, 50 V/m and 10 V/m, respectively. The curves in (b) are plotted for $E_\perp = 0$ meV.

Great efforts have been undertaken to guarantee that the work functions of the different photocathode mounting components are homogeneous. Here, the path of a photoelectron is most sensitive to stray fields because the electron is not yet accelerated by the electric field. The potential of the photocathode mounts can be tuned relative to the potential of the first electrode of the spectrometer. This is important to minimize contact-potential differences in the photocathode mounting region. Moreover, the electrodes of the spectrometer are gold plated.

Using low-outgasing materials for all of the mounts, and pumping with ion getter pumps and volume getter strips should enable us to reach a vacuum base pressure in the $\sim 10^{-12}$ mbar range. This will ensure a long lifetime of the photocathodes and will lead to EDC measurements on undegraded cathodes in the near future.

ACKNOWLEDGMENTS

This work was partially supported by the Deutsche Forschungsgemeinschaft within the Leibniz-program. The heterostructure material was kindly put at our disposal by A. S. Terekhov. We are grateful to A. S. Terekhov as well as to D. A. Orlov and A. Wolf for their continuous support.

REFERENCES