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ABSTRACT

In photoelectron spectroscopy, the measured electron momentum range is intrinsically related to the excitation photon energy. Low photon energies (<10 eV) are commonly encountered in laser-based photoemission and lead to a momentum range that is smaller than the Brillouin zones of most materials. This can become a limiting factor when studying condensed matter with laser-based photoemission. An additional restriction is introduced by widely used hemispherical analyzers that record only electrons photoemitted in a solid angle set by the aperture size at the analyzer entrance. Here, we present an upgrade to increase the effective solid angle that is measured with a hemispherical analyzer. We achieve this by accelerating the photoelectrons toward the analyzer with an electric field that is generated by a bias voltage on the sample. Our experimental geometry is comparable to a parallel plate capacitor, and therefore, we approximate the electric field to be uniform along the photoelectron trajectory. With this assumption, we developed an analytic, parameter-free model that relates the measured angles to the electron momenta in the solid and verify its validity by comparing with experimental results on the charge density wave material TbTe3. By providing a larger field of view in momentum space, our approach using a bias potential considerably expands the flexibility of laser-based photoemission setups.

I. INTRODUCTION

Angle-resolved photoemission spectroscopy (ARPES) probes the electronic structure in momentum space and plays a fundamental role in our understanding of material properties.1 One important consideration when designing experiments is the accessible momentum range. In a conventional ARPES experiment, the in-plane momentum of the electrons in the solid, \( k_{\parallel} \), is determined by measuring the emission angle \( \theta_S \) at the sample surface and the kinetic energy \( E_K \) of the photoemitted electrons. Based on energy and momentum conservation, the in-plane momentum is then determined by:

\[
k_{\parallel} = \frac{1}{\hbar} \sqrt{\frac{2mE_K}{\cos \theta_S}},
\]

where \( m \) is the free electron mass. \( E_K \) varies with the excitation photon energies, which can vary from the ultraviolet to the hard x-ray regime. Higher photon energies result in higher kinetic energies and, therefore, in larger accessible momenta. In contrast, low photon energies access a narrower momentum range but typically have better momentum resolution.1

The measured momentum range also depends on the experimental geometry and the photoelectron detectors. Conventional hemispherical imaging analyzers are widely used in ARPES setups to measure the photoelectron’s emission angle and kinetic energy. The emission angle is measured along a single direction, as defined by the analyzer slit, and the angular range is geometrically limited...
by a fixed aperture at the entrance of the analyzer. The accepted range is typically about 30° while electrons are emitted over 180° from a flat sample surface. It is possible to acquire photoelectrons throughout the full angular range by rotating the sample relative to the analyzer and performing multiple measurements in different geometries. This approach is not only time-consuming, but can also detract from the data quality: the sample rotation can modify the photoemission matrix elements, the polarization geometry, the beam position on the sample, and the absorbed energy density of the pump pulse in ultrafast experiments, all of which prevent a direct comparison of the obtained spectra.

These considerations are especially relevant for setups operating at comparatively low photon energies, ∼10 eV, which limit the momentum field of view intrinsically. Prominent examples and motivation for this work are time-resolved photoemission setups that use 6 eV probe photons, which are generated by frequency upconversion of laser sources. When 6 eV photons excite a metallic sample with 4 eV work function, electrons of up to 2 eV kinetic energy are emitted over a momentum range of 1.45 Å⁻¹. This is only slightly smaller than the ∼1.6 Å⁻¹ Brillouin zone size of commonly studied materials with lattice constants of ∼4 Å. However, a 30° field of view of conventional analyzers covers only 0.37 Å⁻¹ momentum range. Collecting photoelectrons throughout the full angular range at once is, therefore, the most efficient way to observe the largest possible momentum range while keeping the benefits of low photon energies such as large photoemission cross-sections, least amounts of space charge and increased bulk sensitivity for photon energies below ∼10 eV.⁴,⁷,⁹

An electric field between the sample and the detector can be used to focus the photoelectron trajectories into the detector aperture and collect the complete angular range in a single, fixed geometry, see Fig. 1. Such an electric field can be generated by applying a bias voltage to the sample or by applying an extractor voltage to the entrance aperture of the analyzer, creating a potential difference between the sample and the detector. The latter approach is employed in photoelectron microscopy systems¹⁰ and more recently in momentum microscopes.

In contrast, the concept to bias the sample has rarely been used to measure a larger momentum range in systems with conventional hemispherical analyzers. However, sample bias has been applied for other purposes. For example, a sample bias was used in two-photon photoemission experiments in order to avoid

FIG. 1. (a) Illustration of the photoelectron trajectories emitted from the surface over a 2π solid angle. Only a small fraction of the electrons (indicated by thicker lines) pass through the circular aperture at the analyzer entrance (shown by a gray ring) as well as the vertical analyzer slit oriented along x and are detected. (b) and (c) The photoelectron trajectories, considering the same initial conditions as (a), are accelerated toward the analyzer entrance when a negative bias voltage is applied. The emission cone is effectively focused toward the analyzer entrance. (d) and (e) Two-dimensional view of the photoelectron trajectories with identification of the relevant variables. Without bias voltage as in (d), the electron trajectory is straight and the angle of emission at the sample is the same at the analyzer entrance. With a bias voltage as in (e), the electron trajectory is curved by the electric field between the sample and the analyzer entrance. The emission angle and the angle at the analyzer entrance are then different. Note that a difference between the sample and analyzer work functions creates an additional electric field, which is included in our model by considering an effective bias voltage $U_B$. For the simplicity of the illustration, the work functions of the sample and the analyzer are assumed to be identical in this figure ($U_B = U_B^*$).
complications at low kinetic energies,\textsuperscript{14-16} and it is useful to determine the work function of materials.\textsuperscript{17,18} The scarce use of this concept to expand the momentum field of view of conventional hemispherical analyzers might be explained by the challenge of establishing a well-defined and robust relation between the electron momenta in the solid and the measured angles that is applicable in the presence of an electric field. In particular, the electric field must be well-known to describe the photoelectron trajectories. Accordingly, recent implementations feature an electrically grounded mesh near the sample to minimize distortions of the electric field, however, putting constraints on the incidence of the light source.\textsuperscript{19-20}

In this work, we describe an implementation of sample bias for a setup with a conventional hemispherical analyzer as widely used in the photoemission community. We demonstrate access to the largest emitted momenta in a single experimental geometry using a 6 eV laser light source. Our design, presented in Sec. II, is an upgrade that minimizes modifications of existing systems. Furthermore, in Sec. II, we establish an analytical, parameter-free model to adapt the angle-to-momentum conversion [Eq. (1)] to the effect of a uniform electric field. Finally, in Sec. IV, we demonstrate experimentally the validity of both the technique and the model using a representative dataset of the charge density wave (CDW) material NbTe\textsubscript{3}. Our parameter-free model makes sample biasing generally applicable to increase the momentum field of view while keeping a quantitative angle-to-momentum relation.

II. MODEL

The angular range measured by a hemispherical analyzer is limited by the fixed aperture at its entrance. By applying a negative bias voltage between the sample and the electrically grounded analyzer, an electric field is generated in the photoelectron flight path. This accelerates the electrons and bends their trajectories toward the analyzer entrance, therefore, allowing for the detection of electrons with a larger photoemission angle, as schematized in Figs. I(a)–I(c). We are interested to establish a relation between the quantities measured by the detector and the ones at the sample surface in such a configuration. In a standard ARPES experiment without electric field, the angle $\theta_D$ and the kinetic energy $E_{K}^D$ recorded by the detector correspond directly to the emission angle $\theta_A$ and kinetic energy $E_{K}^A$ of the photoelectrons at the sample surface. In the presence of an electric field between the sample and the analyzer, this relation is modified. In the following text, we present a simple model that establishes this modified relation and the angle-to-momentum conversion equation, similar to Eq. (1), that applies in the presence of a uniform electric field.

Throughout this work, we distinguish the quantities at the sample surface, at the analyzer entrance (before the electron lenses), and recorded by the detector with indices S, A, and D, respectively. We assume that the kinetic energy at the analyzer entrance and the one recorded by the detector are identical, i.e., $E_{K}^S = E_{K}^D$. In contrast, the angles $\theta_S$ and $\theta_D$ are not necessarily the same, as discussed later.

First, we establish the relation between the kinetic energy at the sample surface $E_{K}^S$ and the one measured by the detector $E_{K}^D$. These kinetic energies are given by

$$E_{K}^S = h\nu - \Phi_S - E_B,$$

where $h\nu$ is the photon energy, $E_B$ is the electron binding energy in the solid, $U_B$ is the applied bias voltage, and $\Phi_S$ and $\Phi_A$ are the sample and analyzer work functions, respectively. This leads directly to the relation

$$E_{K}^D = E_{K}^S + eU_B,$$

where, for convenience, we have defined an effective bias voltage

$$U_B^* = U_B + (\Phi_A - \Phi_S)/e,$$

which includes the difference between the sample and analyzer work functions. It is interesting to note that this work function difference generates an electric field even in the absence of bias voltage and should be generally considered, particularly at low photon energies and large differences of sample and analyzer work functions.

With the energy relation established, we now turn to the angular relation. We approximate the electric field from the sample surface to the analyzer entrance by the uniform field generated in a parallel plate capacitor. This is applicable if the sample surface is normal to the electron lens axis, which is assumed throughout this work. With such a uniform electric field, the photoelectron momentum parallel to the sample surface is unchanged by the field, and consequently $k_{\parallel}^A = k_{\parallel}^D$. The parallel momentum at any position can always be expressed in terms of the kinetic energy and angle at that specific position. The angle-to-momentum conversion is then given in terms of quantities at the analyzer entrance by

$$k_{\parallel}^A = k_{\parallel}^D = 1/B \sqrt{2mE_{K}^D} \sin \theta_A. \tag{6}$$

While the value of $E_{K}^D$ is identical to $E_{K}^S$, the relation between $\theta_A$ and $\theta_D$ remains to be established for this equation to be applicable.

Figure 1(d) presents the configuration of a measurement in the absence of bias voltage, where $d$ is the distance between the sample surface and the analyzer entrance. This distance $d$ remains fixed to the analyzer focus distance throughout this work. In this configuration without bias voltage, the electron flight path is straight and the emission angle at the sample surface $\theta_A$ is equal to the angle $\theta_A$ measured at the analyzer entrance. By the design of the hemispherical analyzer, the angle $\theta_D$ measured by the detector is equal to $\theta_A$ in this standard configuration. We also define $x_A$ as the transverse position of the electron at the analyzer relative to normal emission. In Fig. 1(d), the values $\theta_A$ and $x_A$ provide redundant information as they are related by

$$x_A = d \tan \theta_A. \tag{7}$$

When the electrons are accelerated by an electric field, the photoelectron trajectory is bent as shown in Fig. 1(e). The values of $\theta_A$ and $x_A$ are both reduced in comparison to the field-free configuration. Equation (7) is not valid anymore and the relation of $\theta_A$, $\theta_D$, and $x_A$ is not directly obvious in this case. We can generally assume that $\theta_D = f(\theta_A,x_A)$. The unknown function $f$ represents the complex effect of the electrostatic lens before the hemispherical analyzer.\textsuperscript{21,22} Note that our model only evaluates analytically the electron trajectories before the electrostatic lens and does not attempt to model the trajectories within the lens. To describe the effect of the electrostatic lens, we, instead, limit ourselves to two simple limits to approximate the function $f$ and verify their validity by comparing
them to experimental results in Sec. IV. For both cases, we constrain \( f \) such that it always correctly describes the zero-bias limit, i.e., \( f(\theta_A, x_A) = \theta_A = \text{arctan}(x_A/d) \).

**A. Case I: Angular limit**

In the first case, we assume that only \( \theta_A \) is important for the imaging process of the lens system. In this angular limit, we simply have

\[
\theta_D = f(x_A) = \text{arctan}(x_A/d). \tag{8}
\]

It is then straightforward to obtain an angle-to-momentum conversion equation from Eq. (6),

\[
k_\parallel^A = \frac{1}{\hat{\mathbf{r}}} \sqrt{2meU^*_B} \sin \theta_D. \tag{9}
\]

This is simply the conventional conversion equation based on the quantities measured by the detector. Our results in Sec. IV indicate that this conversion is incorrect.

For later comparison with case II, we rewrite the angle-to-momentum conversion in terms of the kinetic energy at the sample surface,

\[
k_\perp^A = \frac{1}{\hat{\mathbf{r}}} \sqrt{2meE^*_{K}} F \sin \theta_D. \tag{10}
\]

Here, we defined the momentum-scaling factor \( F \) as

\[
F = \sqrt{1 + 2\alpha}, \tag{11}
\]

with the kinetic parameter \( \alpha \) defined in Eq. (18).

**B. Case II: Position limit**

In the second limit, we assume that only the value of \( x_A \) is important for the imaging process of the lens system. In this limit, the function \( f \) is given by

\[
\theta_D = f(x_A) = \text{arctan}(x_A/d). \tag{12}
\]

The value of \( x_A \) can be evaluated from basic electron kinematics,

\[
x_A = \frac{v_x}{a} \left( \sqrt{v_y^2 + 2ad} - v_y \right), \tag{13}
\]

where \( v = (v_x, v_y) \) is the initial velocity and \( a = (0, a) \) is the acceleration with the \( xy \) axes defined in Figs. 1(d) and 1(e) with the analyzer slit oriented along the \( x \)-axis.

Considering Eq. (12) and \( v_x^2 + v_y^2 = 2E^*_K/m \), we find

\[
\tan \theta_D = \frac{v_x}{a d} \sqrt{\frac{2E^*_K}{m} - v_x^2 + 2ad - \sqrt{\frac{2E^*_K}{m} - v_x^2}}. \tag{14}
\]

Solving this equation for \( v_x \), keeping only the solution that is non-zero in the limit \( a \to 0 \) and rewriting it in terms of the parallel momentum with \( mv_x = \hat{h}k_\parallel^A \), we obtain

\[
k_\parallel^B = \frac{1}{\hat{\mathbf{r}}} \sqrt{2meE^*_B} F \sin \theta_D, \tag{15}
\]

where the momentum-scaling factor \( F \) is defined as

\[
F = \sqrt{\frac{\alpha + 1 + \sqrt{2\alpha + 1 - \alpha^2 \tan^2 \theta_D}}{2}}, \tag{16}
\]

with the kinetic parameter \( \alpha \) being

\[
\alpha = \frac{mad}{2E^*_K}. \tag{17}
\]

The acceleration \( a \) is caused by the electric field generated by the effective bias voltage \( U^*_B \), combining the applied bias voltage and the difference between the sample and analyzer work functions [see Eq. (5)]. Considering the acceleration \( a = -eU^*_B/md \), the kinetic parameter \( \alpha \) is rewritten as

\[
\alpha = \frac{-eU^*_B}{2E^*_K}. \tag{18}
\]

The angle-to-momentum conversion for the position limit (case II) is, therefore, formed by the set of Eqs. (4), (5), (15), (16), and (18). We demonstrate in Sec. IV that this conversion can be successfully applied to our experimental results. Note that these equations are only physically relevant for positive values of the velocity \( v_y \). In the limit \( v_y \to 0 \), the kinetic energy at the sample is only given by the parallel momentum \( k_\parallel^B \) and defines a low-energy cutoff (LEC) in the photoemission spectrum at \( E^*_K,\text{LEC} = h^2(k_\parallel^B)^2/2m \). This LEC recorded by the detector can be expressed as

\[
E^*_K,\text{LEC} = -eU^*_B \left[ 1 + \left( \frac{\tan \theta_D^{\text{LEC}}}{2} \right)^2 \right]. \tag{19}
\]

by taking \( 2E^*_K/m = v_x^2 \) in Eq. (14).

In Fig. 2, we illustrate the momentum-scaling factor \( F \) [Eq. (16)] for different values of bias voltage. The momentum-scaling factor \( F \) is an indicator of the increased momentum range relative to a measurement without bias voltage. For example, the momentum range measured for \( E^*_K = 1 \text{ eV} \) is about four times larger at \( U^*_B = -50 \text{ V} \) (Fig. 2[d]) in comparison to \( 0 \text{ V} \). Overall, we see that, for a given bias, the factor \( F \) is mostly dependent on \( E^*_K \) and changes rapidly near the LEC. It has only a weak dependence on \( \theta_D \). The size of \( F \) becomes more important for larger bias voltages, representing a larger field of view in momentum space.

To compare the angle-to-momentum conversion from cases I and II, we trace their respective momentum-scaling factor [Eqs. (11) and (16)] in Fig. 2(e) at \( \theta_D = 0 \). It shows that the measured momentum range, in identical conditions, would be at least 50% larger for case I than for case II when \( a > 5 \). We show in Sec. IV that the position limit (case II) is consistent with our experimental results while the angular limit (case I) overestimates the momentum values.

**C. Scaling of photoemission intensities**

Experimentally, the measured quantity is the photoemission intensity as a function of angle \( N(E^*_K, \theta_D) \) while we are interested in the intensity in the range as a function of momentum \( N(E^*_K, k_\parallel^B) \). The coordinate transformation from \( \theta_D \) to \( k_\parallel^B \) derived above is accompanied by a transformation of the differential line element \( d\theta_D \to dk_\parallel^B \) that causes a scaling of the intensity,

\[
\frac{N(E^*_K, k_\parallel^B)}{N(E^*_K, \theta_D)} = \left( \frac{\partial k_\parallel^B}{\partial \theta_D} \right)^{-1} \frac{\partial k_\parallel^B}{\partial \theta_D}, \tag{20}
\]
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We implemented these requirements in our experimental setup in the following way. The bias voltage is generated from a DC Voltage Source DC205 from Stanford Research Systems and applied to the sample through a wire running from the top of the manipulator to the sample stage. The Kapton-insulated wire is electrically shielded by a grounded silver braid to avoid charging issues due to stray electrons. On the sample stage, this wire is connected to a hook made from a titanium sheet that can be attached to the copper post onto which the sample is fixed [Figs. 3(c) and 3(d)].

The oxygen-free high-conductivity (OFHC) copper post is screwed into a threaded 6061 aluminum insert, which is insulated from the titanium mounting piece by a sapphire cylinder and washer (Insaco Inc.) [Figs. 3(a) and 3(b)]. Sapphire insulators are chosen to optimize cryogenic performance. The combination of titanium and sapphire is favorable due to similar thermal expansion coefficients.\textsuperscript{25} The threaded aluminum insert and titanium piece are individually machined to match the sapphire inserts with tight tolerances, <10 μm. This assembly is fastened by application of a thin film of Loctite Stycast 1266 that has been outgassed in high vacuum before curing to achieve compatibility with ultra-high vacuum. The assembly is cured in a jig, which pulls on the threaded aluminum insert while pushing down on the sapphire washer and cylinder to ensure tight fits with optimal thermal contact. Aluminum washers are used to set the sample post orientation with respect to the titanium piece.

In order to form a parallel plate capacitor geometry, the top surface of the copper post is designed as a disk with a 1 cm diameter, the largest diameter possible to allow sample transfer in our experimental setup. The analyzer entrance is electrically grounded and acts as the second plate of the capacitor. The measurements are taken for a configuration near normal emission to maintain the parallel plate geometry and keep a uniform electric field.

In these experiments, a Spectra Physics Ti:sapphire oscillator operating at a repetition rate of 80 MHz generates 1.5 eV photons that are frequency quadrupled through two stages of second harmonic generation in β-BaB\textsubscript{2}O\textsubscript{4} non-linear crystals to provide a 6 eV photon source.\textsuperscript{24} The photoelectrons are detected by a Scienta R4000-WAL hemispherical analyzer, which has a work function of Φ\textsubscript{F} = 4.15 eV [determined using Eq. (3) with the measured value of ε\textsubscript{F} at the Fermi level]. The analyzer slit width was fixed to 0.1 mm throughout the measurements. Measurements were performed using a pass energy of 10 eV in the dither mode. The distance d between the sample and the analyzer entrance was fixed to the instrument-defined focus distance throughout the measurements.

To demonstrate the effects of sample bias and the validity of our model, we performed measurements on the compound TbTe\textsubscript{3}, a member of the well studied rare-earth tritelluride family featuring CDW order.\textsuperscript{25,26} TbTe\textsubscript{3} single crystals were grown using a Te self-flux technique,\textsuperscript{27} which ensures purity of the melt and produces large crystals with a high degree of structural order. Elements in the molar ratio Tb:Te = 0.03:0.97 were put into alumina crucibles and vacuum sealed in quartz tubes. The mixture was heated to 900 °C over the course of 12 h and kept at that temperature for a further 10 h. It was then slowly cooled to 650 °C over a period of 100 h. The remaining melt was decanted in a centrifuge. The resulting copper-colored crystals are malleable plates with dimensions of up to 5 × 5 × 0.4 mm\textsuperscript{3} and oriented with the long b axis perpendicular to the plane of the crystal plates. The nearly equal in-plane axes a and c are parallel to the crystal growth edges but must be

![Graph](image-url)

**Fig. 2.** (a)–(d) Momentum-scaling factor $F$ in the position limit [Eq. (16)] for relevant ranges of detector angle $\theta_D$ and kinetic energy $E^2_K$ at various bias voltage with $U_b = 0$. The angular dependence is weak while the energy dependence is pronounced near the low-energy cutoff (LEC). The factor $F$ is undefined below the LEC, as no photoelectrons can physically have those parameters. (e) Comparison of the momentum-scaling factor $F$ at $\theta_D = 0$ for the angular (case I) and the position (case II) limits with Eqs. (11) and (16), respectively.

where $\Delta k^I_D$ and $\Delta \theta_D$ are the bin sizes. The partial derivative from Eqs. (15) and (16) is given by

$$\frac{\partial k^I_D}{\partial \theta_D} = \frac{2mE_K^2}{\hbar} \left[ F \cos \theta_D + \sin \theta_D \frac{\partial F}{\partial \theta_D} \right]$$

with

$$\frac{\partial F}{\partial \theta_D} = -\alpha^2 \tan \theta_D \sec^2 \theta_D \left( \frac{1}{4} F \sqrt{2 \alpha^2 + 1} - \alpha^2 \tan \theta_D \right).$$

In general, the largest intensity correction occurs near the LEC. We note that this intensity scaling resulting from the coordinate transformation is a generic feature, present even without a bias voltage, and should generally be considered in any standard ARPES experiment, particularly when studying several eV wide spectra.

Without a bias voltage or difference of analyzer and sample work functions ($U_b = 0$), the partial derivative is simply given by

$$\frac{\partial k^I_D}{\partial \theta_D} = \frac{1}{\hbar} \sqrt{2mE_K^2} \cos \theta_D.$$
were performed for $U_B$ ranging from 0 to $-60$ V. The 6 eV flux was kept constant and its weak time-dependent drift was corrected for in the analysis. The measurement position was optimized at $U_B = -60$ V for normal emission, i.e., for the largest energy difference between the Fermi level ($E_F$) and the minimum of the LEC, as justified in the end of Sec. IV. The position remained fixed for measurements at all bias voltages. The sample work function was determined to be 5.14 eV.

**IV. EXPERIMENTAL RESULTS**

Measurements were performed in the $k_x$–$k_z$ plane with the analyzer slit along the diagonal of the Brillouin zone of TbTe$_3$, as illustrated in the inset of Fig. 4(f). Here, $k_x$ and $k_z$ indicate the reciprocal axes of the TbTe$_3$ crystal axes $a$ and $c$, respectively. The material is air sensitive, and crystals were stored in an oxygen and moisture-free environment.

TbTe$_3$ was cleaved in situ at a base pressure of $1 \times 10^{-10}$ Torr and measured at a constant temperature of 84 K. Measurements distinguished using, for instance, x-ray diffraction. The material is sensitive, and crystals were stored in an oxygen and moisture-free environment.

TbTe$_3$ was cleaved in situ at a base pressure of $1 \times 10^{-10}$ Torr and measured at a constant temperature of 84 K. Measurements...
FIG. 4. (a)–(e) Raw photoemission spectra for $U_B = 0$ to $-40$ V. The red line indicates the low energy cutoff (LEC) defined by Eq. (19). (f)–(j) Corresponding photoemission spectra after angle-to-momentum conversion [Eqs. (15) and (16)] and intensity scaling [Eq. (20)]. The red line indicates the LEC, below which the angle-to-momentum conversion is not defined. The inset in (f) is the schematic Fermi surface of $\text{TbTe}_3$. The red circles indicate the momentum at $E_F$ accessible with 6 eV photons. The solid black and dashed red lines indicate the measured cuts at 0 and $-40$ V, respectively.

The hole band centered at $k_\parallel = 0$ [Fig. 5(f)]. The separation $\Delta k$ between both peaks is nearly constant with $U_B$ [Fig. 5(g)]. There is, however, a clear $U_B$-dependent peak position shift that we again attribute to experimental limitations. To further illustrate the validity of the angle-to-momentum conversion model, we also present the mass of the hole band in Fig. 5(h). To determine the hole mass, the band dispersion was extracted by fitting energy distribution curves (EDCs) over the range of $\Delta k$. The effective hole mass does not deviate from the zero-bias value by more than 10% at all $U_B$. This result is remarkable considering that our simple angle-to-momentum conversion model has no adjustable parameter.

We note that the angular limit (case I) leads to a momentum-scaling factor at $U_B = -60$ V about twice as large as the one used here for the position limit (case II). It is, therefore, clear that the angular limit would lead to strongly $U_B$-dependent spectral features.
Furthermore, it would result in bands at momentum values larger than physically allowed from the sample work function and photon energy of the experiment. We can, therefore, conclude that the angular limit (case I) is not valid in the experimental conditions considered in this work.

We note that the increase in momentum range by a factor $F$ affects both the in-plane momentum parallel and transverse to the analyzer slit. This results in integrating a range of transverse momentum by a factor $F$. This increase in count rate at a constant 6 eV flux is directly evident from the color scale in Fig. 4 and the signal to noise ratio of the cuts in Fig. 5. Depending on the specific scientific question, this effective increase in analyzer transmission can be an additional important benefit.

We further characterize how the bias voltage affects spectral features on the energy axis. We first evaluate the intensity near $E_F$. Without any bands crossing $E_F$ for all $U_B$, in Fig. 6(a), we instead analyze the Fermi edge caused by electrons scattered in the final state. This region is marked in the center of the Brillouin zone as box 6 in Fig. 5(a). The Fermi edge was fitted to a Fermi–Dirac function convoluted with a Gaussian resolution function. The resulting position of $E_F$ and the full-width-at-half-maximum (FWHM) of the resolution function are presented in Figs. 6(b) and 6(c), respectively. In addition, the EDCs of the central hole-like band were fitted to a Lorentzian function with a linear background and the resulting FWHM is presented in Fig. 6(e). Overall, we observe no significant energy shift or broadening due to the bias voltage, therefore, demonstrating that the technique does not cause significant artifacts along the energy-axis.

Finally, we discuss experimental limitations of the bias voltage technique. The model is based on the assumption that a uniform electric field exists from the sample surface up to the analyzer. However, distortions of the electric field will occur when the configuration deviates from a parallel plate capacitor. In particular, the bias application apparatus can cause stray field and great care must be taken when designing the biasing mechanism to shield electric fields to the extent possible. Furthermore, the difference in work functions of the sample and of the holder can also create inhomogeneous fields near the sample surface.

Our model can only be applied for normal emission measurements as it relies on a parallel plate capacitor geometry. We, therefore, optimize for normal emission using the LEC. Specifically, we optimize for the lowest measurable kinetic energy defined by electrons without any in-plane momentum, as schematized by the red parabola in Figs. 7(a) and 7(b). In the presence of field distortions, electrons acquire a finite in-plane momentum transverse to the analyzer slit and one measures effectively off-normal emission while the sample and analyzer are oriented parallel. This is indicated by a higher energy minimum of the LEC, as illustrated with the blue parabola in Figs. 7(a) and 7(b). One can, therefore, optimize the experimental geometry to minimize the LEC energy relative to $E_F$ to compensate for field distortions. This ensures that the measured electrons at $k_F = 0$ are emitted normal to the sample surface.

In the measurements presented in this work, we optimized the LEC at a bias voltage of $U_B = -60$ V and confirmed that the spectrum near $E_F$ is symmetric around $k_F = 0$ after optimizing the geometry. This geometry was then retained for all bias values to provide a direct comparison. However, the field compensation changes with bias and the effective cut in momentum space can be modified. We, indeed, observe this effect, as illustrated in Fig. 7. In Fig. 7(c), we present the energy difference between $E_F$ and the minimum of the LEC. This quantity is maximal for $U_B = -60$ V and decreases with decreasing $|U_B|$. This is a signature that the measured electrons acquire a finite momentum transverse to the analyzer slit with decreasing $|U_B|$. The effective cut in momentum space, therefore, changes from a cut going through the center of the Brillouin zone at $U_B = -60$ V to a displaced cut for smaller $|U_B|$. This effect explains the observed asymmetry of the CDW gap for bias voltages different from $U_B = -60$ V [Figs. 7(e) and 7(f)]. Indeed, an
asymmetry is expected for cuts in momentum space displaced from the center of the Brillouin zone. In Fig. 7(d), the thickness of the black lines represents the size of the CDW gap in TbTe. While it is identical for positive $(+k)$ and negative $(-k)$ momentum for a cut going through the center of the Brillouin zone (red region), it will be larger at $-k$ than at $+k$ for a cut away from it (blue region). The combined observation of the CDW gap asymmetry and of the change in the LEC minimum strongly supports our interpretation that the effective cut in momentum space is modified with $U_B$. This change in the cut also explains some variations of the spectral features observed with different bias voltage in Fig. 5.

V. DISCUSSION AND CONCLUSIONS

Our experimental results show that the momentum field of view in ARPES experiments with hemispherical analyzers can be increased by applying a bias voltage to the sample. In addition, we derive an analytic, parameter-free expression for the conversion between measured angles and in-plane momenta that is applicable when the electric field is uniform. We confirm its validity with our experimental results on TbTe. As our model is parameter-free, it allows us to perform bias-measurements for a wide range of conditions without the need for any calibration.

Furthermore, the model has important implications even for measurements without bias voltage. Specifically, if the difference in sample and analyzer work functions is comparable to the photoelectron kinetic energies, the induced electric field will have observable effects on the photoelectron trajectories and our model should be considered to obtain accurate momentum and intensity values. Such a regime is commonly encountered in ARPES measurements using 6 eV photon energy.

We like to point out another approach to establish an angle-to-momentum conversion for a comparable experimental configuration. Jauernik developed a heuristic model that was calibrated to the well-known dispersion of the image potential state in front of the (001) surface of copper. The empirical bias-scaling factor of this model can be estimated using our parameter-free model. We find good agreement to the experimentally determined value, further supporting the general validity of our model.

While our model is limited to normal emission by definition, it could be generalized to off-normal emission by assuming that the electric field is generated by two infinite planes that intersect at a line.

The main experimental limitation of our implementation of sample bias is the distortion of the photoemission spectra caused by deviations from a uniform electric field, as defined by a parallel plate capacitor geometry. The general design of an ARPES system can limit the validity of this approximation, as well as the specific characteristics of different samples. As we demonstrate, a symmetric bias field between the sample, its holder, and the analyzer can be obtained by careful engineering of the experimental design. Other objects near the photoelectron trajectory, such as the capillary of a He lamp, can cause significant field distortions and a different design with its own angle-to-momentum conversion formalism might be necessary. We also note that samples with large flat surfaces are preferred to obtain a uniform electric field. The investigation of samples with rough surfaces is more challenging due to the irregular fields near the surface. A photoemission spot size smaller than the characteristic length scale of the sample inhomogeneities is beneficial. Field distortions can also be minimized by reducing the work function differences between the sample and its holder.

As evidenced in Fig. 7, the experimental configuration should be optimized for each specific bias voltage to compensate the field distortions that remain and obtain the most reliable results. Consequently, precise and stable positioning of the sample becomes more important as well.
A fundamental physical limitation of our approach remains the intrinsically limited momentum range at low photon energies. While it is possible to continue to compress the momentum range by increasing the bias voltage value, the maximum accessible momentum remains physically limited by the photon energy and the sample work function. For example, in our experiment, the complete physically allowed momentum range at $E_p$ is observed for $U_B = -40$ V and larger $|U_B|$ values do not provide additional information.

ARPES experiments at ~10 eV photon energies profit most from the increased momentum field of view a sample bias provides. A fixed sample orientation avoids issues caused by matrix elements, polarization effects, and beam walk on small samples, while retaining the advantages of high photoemission intensity, enhanced bulk sensitivity, and mitigation of space charge effects at low photon energies. The application of a bias voltage is not limited to photoemission studies but particularly benefit from measuring the electron dynamics over a large part of the Brillouin zone in a fixed configuration. In addition to all of the benefits applying in equilibrium, high photon energies and a fixed geometry avoids changes in absorbed excitation density in time-resolved ARPES experiments. The application of a bias voltage is not limited to photoemission studies but particularly benefit from measuring the electron dynamics over a large part of the Brillouin zone in a fixed configuration.

The recent development of an ultrafast 11 eV laser for time-resolved experiments relies on 6 eV probe photons and a quasi-continuous wave laser of 11 eV for high effective analyzer transmission due to a larger acceptance range of transverse momenta can be an important aspect when collecting statistics sufficient for high precision studies.

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AUTHOR DECLARATIONS

Conflict of Interest

The authors have no conflicts to disclose.

DATA AVAILABILITY

The data that support the findings of this study are openly available in Stanford Digital Repository at http://doi.org/10.25740/sk226xw9348.

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