Electric and magnetic form factors of the proton

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**Electric and magnetic form factors of the proton**


(A1 Collaboration)

1Institut für Kernphysik, Johannes Gutenberg-Universität Mainz, 55099 Mainz, Germany
2Department of Physics, University of Zagreb, 10002 Zagreb, Croatia
3Jožef Stefan Institute, Ljubljana, Slovenia
4LPC-Clermont, Université Blaise Pascal, CNRS/IN2P3, F-63177 Aubière Cedex, France
5CERN, CH-1211 Geneva 23, Switzerland, on leave of absence from Physik-Department, Technische Universität München, 85748 Garching, Germany
6Department of Physics, University of Ljubljana, Slovenia

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This paper describes a precise measurement of electron scattering off the proton at momentum transfers of $0.003 \lesssim Q^2 \lesssim 1$ GeV$^2$. The average point-to-point error of the cross sections in this experiment is $\sim 0.37\%$. These data are used for a coherent new analysis together with all world data of unpolarized and polarized electron scattering from the very smallest to the highest momentum transfers so far measured. The extracted electric and magnetic form factors provide new insight into their exact shape, deviating from the classical dipole form, and of structure on top of this gross shape. The data reaching very low $Q^2$ values are used for a new determination of the electric and magnetic radii. An empirical determination of the two-photon-exchange correction is presented. The implications of this correction on the radii and the question of a directly visible signal of the pion cloud are addressed.

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**I. INTRODUCTION**

Form factors of extended quantum systems are traditionally considered as a means to access the distribution of charge, magnetism, and weak charge through their Fourier transforms. However, this traditional interpretation is only approximately valid for a light system as the proton. More recently it was realized through the generalization of quark-gluon structure functions that an interpretation on the light-cone frame is not only mandatory but also fruitful since it offers better insight into the quark-gluon structure of the nucleon (for a summary see Ref. [1]). However, for such an application of form factors, good knowledge from the smallest to the highest momentum transfers is needed. Though common fits of the form-factor models to the measured cross sections: the classic Rosenbluth separation and a direct fit of form-factor models to the experimental results, it produces form factors without any model assumption. The second method is often used in fits for at least a decade. It has many advantages, especially when coupled with an experiment optimized for this style of analysis.

(i) The traditional Rosenbluth separation of the electric and magnetic form factors uses measurements of the cross section at constant $Q^2$ and varied values of the polarization parameter $\varepsilon$. This method, however, limits unnecessarily the kinematical range since the range of beam energy and scattering angle are larger than the constant-$Q^2$ domain. Aiming the second approach, the experiment is not bound by this constraint. All data at any kinematical point can be used for the fit, even if the covered range of $\varepsilon$ at a given $Q^2$ is not wide enough to separate the form factors.

(ii) It is a notorious problem to measure an absolute cross section to better than about $\pm 2\%$. This was already
the case in the old fits of Hofstadter [10] and is highly relevant for all older measurements since the uncertainty of the absolute normalization may have been as large as \( \pm 5\% \) mostly due to the uncertainty in the determination of the electron beam current, target thickness, and solid angle of the spectrometers. A common fit to the world data has to account for the uncertainty in the cross-section normalization of the different experiments. To this end one may introduce the normalization as parameters in the fits, as has been done in previous extractions (e.g., Refs. [3–5,7]). We can employ the same technique to determine the normalizations. Having groups of measurements with good relative normalization internally given by the experiment, the fit can determine the relative normalization of the groups with regard to each other and the global normalization via the extrapolation to the known form-factor values at \( Q^2 = 0 \).

(iii) While the classic Rosenbluth approach gives a correct error estimate for the form factor themselves, traditionally numbers for the anticorrelation between \( G_E \) and \( G_M \) are not given. Without this information, uncertainties of fits to the form factors, for example, for the extraction of the radius, cannot be calculated correctly. A direct, global fit to cross sections does not have this problem.

(iv) We find that the robustness of the fits is increased (see Sec. VI B). This is, on the one hand, caused by the smoothing effect of the fit and, on the other hand, the fit allows us to separate the form factors even if the measurements do not coincide in \( Q^2 \). Therefore, the effective density of measurements in \( Q^2 \) can be increased.

The experiment presented here, to our knowledge, is the first specifically optimized for this method of analysis. This approach and the precise measurements extending down to very low momentum transfer made some distinctive improvements over previous extractions possible as follows:

(a) The form-factor normalization at \( Q^2 = 0 \): The momentum region covered by the new measurement at MAMI, the details of which are described in this paper, is \( 0.003 \text{ GeV}^2 \lesssim Q^2 \lesssim 1 \text{ GeV}^2 \). The small statistical error of \( 0.2\% \) at low \( Q^2 \) reduces dramatically the uncertainty of the normalization at \( Q^2 = 0 \).

(b) Normalization of different data sets: The new data presented here have excellent relative normalization in a large \( Q^2 \) range, tying together the normalizations of the different data sets in the overlap region. Together with the first point, the absolute normalization is fixed with small uncertainty for a broad range in \( Q^2 \). Of course, the normalization factors extracted by the fit have to be independent of the specific fit model. The analysis presented here has this feature.

(c) Two-photon-exchange (TPE) correction: The cross-section data and the asymmetry measurements with polarized electrons give inconsistent results for the form factors. This inconsistency is believed to be caused by the unconsidered TPE contribution which is deemed to be more important for the Rosenbluth formula, containing the electric and magnetic form factors \( G_E^2 \) and \( G_M^2 \) as a sum weighted by kinematical factors, than for the asymmetry formulas, which give the ratio \( G_E/G_M \). Both methods are based on the one-photon-exchange approximation only [11]. We fit the cross-section data together with the polarization data using a simple empirical model to parametrize the inconsistency. This empirical ansatz can reconcile the measurements and can easily be compared to theoretical calculations of TPE.

Some of the results of this paper are topical theoretically and at the center of recent controversies as follows:

(1) Electric and magnetic radii: A very precise determination of the electric radius of the proton through the Lamb shift of muonic hydrogen [12] has given a 4\% smaller value than both the CODATA value [13] and the result of the present experiment [8]. The smaller muonic Lamb shift value has been confirmed recently with an updated result [14]. This discrepancy is a so-far unresolved puzzle. A similar discrepancy existed for the magnetic radius between the determination from the hyperfine splitting of electric hydrogen [15] and some of the fits of the electron scattering world data [16]. However, as will be discussed in Sec. VII C, this discrepancy disappears with the result of this experiment [8].

(2) TPE correction: This correction has been calculated by several groups but with diverging results (see Ref. [17] and references therein). After a controversy [16,18,19] we present an experimental method in this paper showing the size of the effect possibly assigned to TPE and make it directly accessible to a comparison with theoretical calculations.

(3) Possible signal of a pion cloud: The common idea is that the proton form factors are smooth and show no narrow structure of small \( Q^2 \) scale. In an outdated fit of the pre-2003 data Friedrich and Walcher hypothesized the existence of a bump-dip structure which they attributed to the signal of a pion cloud [2]. This bump-dip could not be confirmed by the results of this experiment [8], which, however, shows some other similar and more significant structure. Though the idea of such structures is not very welcome [20] as it is considered to be a “popular fantasy” on theoretical grounds, the fits presented here will show further evidence for it.

The resolution of these controversies is partially possible through the high-precision results for the form factors presented here and as discussed later.

The paper is organized as follows. In the second section we present the relevant details of the experiment at MAMI followed by the description of the theoretical basis in the next section. The fourth section describes the extraction of cross sections from the measured count rates. The analysis of the cross section and some peculiar aspects of the statistics needed for the determination of errors and for the construction of
confident bands is described in the fifth section. It follows a discussion of the results in the frame work sketched above, concluding with the new determination of the proton form factors in Sec. VI and of the radii in Sec. VII.

Because of the size of the data set, cross sections and tabulated fits are not included as an appendix in the paper but are available electronically as part of the Supplemental Material [21], on the arXiv, and on request from the authors.

II. EXPERIMENT

In this section, we give an overview of the accelerator and detector facilities used in the experiment and describe the hydrogen target and the program of the measurement.

A. Accelerator

MAMI, the Mainz Microtron [22–24], is a normal conducting continuous-wave electron accelerator. It consists of a cascade of three racetrack microtrons (RTMs) and a fourth stage, a harmonic double-sided microtron (HDSM). The accelerator is equipped with two electron sources: a thermionic source, which can provide currents in excess of 100 μA, and a polarized source that makes use of the photoelectric effect on a GaAs crystal using polarized light which can provide more than 30 μA.

A linear accelerator injects the electrons with 3.97 MeV into the first RTM. Each of the three RTMs contains a normal conducting accelerator segment and two large high-resolution conventional magnets which recirculate the beam back into the accelerator segment. In the first RTM, the beam is recirculated 18 times, raising the electron energy to 14.86 MeV. The second RTM boosts this to 180 MeV in 51 turns.

The beam may now bypass the rest of the accelerator and may be directed to the different experimental sites. Alternatively, it can enter RTM 3, which can boost the energy up to 855 MeV in 90 turns. Every other recirculation path can be instrumented with a kicker magnet which deflects the beam to the exit beam line system. Thus, the energy can be selected in 15-MeV steps.

The beam may then be injected into the fourth stage. The HDSM stage comprises two antiparallel accelerator segments, one of which is operated at the doubled frequency to suppress instabilities. The beam is recirculated by four magnets. The HDSM stage raises the energy up to 1.6 GeV in 43 recirculations.

The absolute beam energy uncertainty is 150 keV and the root-mean-square (rms) energy spread is 30 keV at 855 MeV and 110 keV at 1.5 GeV. For the measurements described in this work, an unpolarized beam with beam energies of 180, 315, 450, 585, 720, and 855 MeV was used.

B. Detector setup

The detector setup of the A1 collaboration at MAMI is called the three-spectrometer facility. The three high-resolution magnetic spectrometers, labeled A, B, and C, can be operated in single, double, or triple coincidence mode. A detailed description can be found in Ref. [9]. The spectrometers can be rotated around a central pivot to measure at different scattering angles. The scattering angle can be read out with an absolute accuracy of 0.01°, leading to an uncertainty in the cross section of around 0.02%.

The magnetic system of spectrometer A and C is composed of a quadrupole, a sextupole, and two dipoles. This complex system facilitates a high-precision measurement of particle momentum and angle inside a relatively large acceptance of up to 28 msr. Spectrometer B consists of only a single dipole in a clamshell configuration, leading to a slim design with higher spatial resolution but smaller acceptance (5.6 msr); for out-of-plane measurements, spectrometer B can be tilted by up to 10°.

Each of the three spectrometers is equipped with similar detector systems consisting of two scintillator planes, two packets of two vertical drift chamber layers (VDC), and a gas-Čerenkov detector. The scintillators are used for triggering, particle identification, and for a time reference. The drift chambers are used for the reconstruction of the particle trajectory. The Čerenkov detector distinguishes between muons (and heavier particles) and electrons.

C. Target system

The target system is enclosed in a vacuum scattering chamber located on the rotation axis of the spectrometers and directly connected to the beam vacuum tube. A target ladder holds several interchangeable solid-state materials like graphite, polyethylene, HAVAR foil, copper, etc., of varying thicknesses. Additionally, a luminescent screen (an Al2O3 plate with a crosshair printed on it) is mounted; it is used for beam position calibration. The target ladder has a vertical translation degree of freedom that is actuated by an electric motor to select the target material.

The normal lid of the barrel-shaped scattering chamber can be exchanged for two different target constructions: A high-pressure gas target and a cryogenic target. The present experiment used the latter filled with liquid hydrogen as a proton target.

The cryogenic target system is composed of two loops. An inner loop (the “Basel loop”) is filled with the target gas, which is liquefied before the beginning of the beam time. The completely liquefied material is continuously recirculated by a fan. The loop contains an interchangeable target cell; two types were used in this experiment: a 5-cm-long, cigar-shaped cell with its axis in the beam direction and a cylindrical cell with a diameter of 2 cm and the axis perpendicular to the scattering plane. A heat exchanger couples the inner loop to the outer loop, which is coupled to a Philips compressor. The outer loop is also filled with hydrogen and works like a heat pipe: Hydrogen is liquefied at the Philips compressor. It flows down to the target, cooling down the target heat exchanger. The warmed up hydrogen then evaporates and returns to the Philips compressor.

The hydrogen inside the inner loop is subcooled to ensure that the beam load does not substantially change the density of the hydrogen by local heating above the boiling point. Nevertheless, for higher currents the beam is rastered in the transverse directions to reduce the effective power density (in
D. Additions to the standard experimental setup

1. pA meter

In a typical experiment at the three-spectrometer facility, the beam current is measured with a Förster probe located in a part of RTM 3 where all recirculations of the beam pass through. Accordingly, the accuracy of the measurement is best with the highest number of recirculations, i.e., for a beam energy of 855 MeV. For 180 MeV, the beam does, however, not pass the probe.

Therefore, for measurements at small energies, a pA meter was installed at a collimator right before the linear accelerator segment. When the beam is deflected on the collimator, a current proportional to the beam current can be measured precisely. The pA meter has been calibrated versus the Förster probe with a range of beam currents using a beam energy of 855 MeV, where the Förster probe is most sensitive. Since the pA meter is situated in a section of the accelerator where the beam energy is constant, the calibration is valid for all beam energies. The pA-meter system also has a better precision for low beam currents, which are needed for the measurements at small scattering angles.

2. Beam position stabilization

A shift of the beam position on the target results in a drift of the measured cross section [25]. The beam is normally stabilized by the circulation in RTM 3, which dampens beam position changes introduced in the earlier stages of the accelerator. This self-stabilization is less effective with lower recirculation number, i.e., lower energies, and is absent in the case of an incident beam energy of 180 MeV when the beam bypasses RTM 3. To eliminate beam position drifts, a beam-position control system has been installed by the MAMI group [26]: The beam position is measured with two cavities in front of the target. Their signal is digitized and a correction current for the beam steering dipoles in the beam line is generated. The cavities need high beam currents for adequate sensitivity. Therefore, the beam has to be switched to a diagnostic mode where the beam is modulated as a train of high current pulses with a low duty cycle. These periods have to be excluded from the cross-section measurements. During the data taking, the A1 computer system periodically disable the data acquisition and generates a signal to the MAMI control system to start the adjustment process. When the correction has been performed, MAMI signals back to the A1 system and data acquisition is resumed. The analysis codes have been modified to account for these pauses in the data acquisition.

The system was installed in the beginning of the second measurement period (see Table I) and was used for all later measurements. After the installation of the system, there was no beam position drift detectable. For the first period, we observed beam position shifts of less than 0.3 mm, leading to a change in the cross section of less than 0.1%. Because this change is matched by the luminosity measurement, the error in the ratio of the two is negligible.

### Table I. Overview of the beam times. Setup changes are changes of momentum and/or angle of at least one spectrometer.

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<td>2 days</td>
<td>6 days</td>
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<td>180, 720</td>
<td>315, 450, 720</td>
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<td>Short</td>
<td>Long</td>
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<tr>
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<td>173</td>
<td>217</td>
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<tr>
<td>No. of measurements</td>
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<td>490</td>
<td>574</td>
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<tr>
<td>Beam currents</td>
<td>585: 11 nA to 5.5 μA to 10 μA</td>
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<tr>
<td>to 855: 0.8 μA to 720: 90 nA to 450: 30 nA</td>
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<tr>
<td>to 315: 28 nA to 1.6 μA</td>
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<td>720: 8–10 μA</td>
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E. Kinematic coverage

At a given (negative) four-momentum transfer squared,

\[ Q^2 = 4E E' \sin^2 \frac{\theta}{2}, \quad (1) \]

where \( E (E') \) is the energy of the incoming (outgoing) electron and \( \theta \) is the electron scattering angle, the relative contributions of \( G_E \) and \( G_M \) to the cross sections depend on the polarization of the virtual photon,

\[ \varepsilon = \left[ 1 + 2 \left( 1 + \frac{Q^2}{4m_p^2} \right) \tan^2 \frac{\theta}{2} \right]^{-1}. \quad (2) \]

In order to extract the form factors from the measured cross sections using the traditional Rosenbluth technique, it is mandatory to measure at different \( \varepsilon \) for a given, constant \( Q^2 \) value. While this constraint does not have to be fulfilled when performing the mentioned global fit, which is in focus here, the range and number of different \( \varepsilon \) values in a given \( Q^2 \) range determine the accuracy of the separation.

Figure 1 displays the part of the \( \varepsilon-Q^2 \) plane accessible by the accelerator and detector setup in this experiment. To help the readability, only the centers of the overlapping acceptances are marked. The shaded areas are excluded because of the various experimental limitations (see the figure caption).

In order to vary \( \varepsilon \) at constant \( Q^2 \), both the scattering angle and the incident beam energy have to be changed. A beam energy change takes about 6 h at MAMI, which is quick in comparison to other accelerator facilities but still too costly to be done frequently. Thus, the measuring program was organized to minimize beam energy changes. Since the energy gain in RTM 1 and RTM 2 is fixed, the minimum beam energy of MAMI is achieved when the beam passes RTM 3 without further acceleration, resulting in a 180-MeV beam. The beam time allocation permitted to measure at six energies, spread out evenly between 180 and 855 MeV in 135-MeV steps. In this experiment, no use was made yet of the 1.6-GeV stage of MAMI. While the HDSM stage (MAMI C) was already commissioned for productive use, there was no experience with the quality of the beam, and it was not yet possible to extract the beam at energies between 0.855 and 1.6 GeV.
FIG. 1. The accessible kinematical region in the $e$-$Q^2$ plane. Shaded regions are excluded because of minimum or maximum beam energy (darkest gray), maximum detectable momentum of spectrometer A and B (medium grays), and minimum angle of B and maximum angle of C (lightest gray). The centers of the acceptances of the different kinematical settings are denoted with symbols (squares, circles, and triangles correspond to spectrometer A, B, and C). The symbols are slightly shifted vertically to distinguish overlapping measurements. The upper black lines represent a possible future extension using MAMI C energies. To stretch the low-$Q$ correspond to spectrometer A, B, and C). The symbols are slightly shifted vertically to distinguish overlapping measurements. The upper black lines represent a possible future extension using MAMI C energies. To stretch the low-$Q$ range.

At each beam energy, the measured angle range was maximized. The geometric designs of target and spectrometers allow each spectrometer to cover different but overlapping angular ranges. To maximize the angular range covered by the data set and the internal redundancy in the data, all three spectrometers were used in parallel. Spectrometers A and C were used alternately as the “production spectrometer” (changing angle from run to run) and “luminosity monitor” (at a fixed angle). The spectrometer angle is changed each time only by one-fourth of the acceptance, i.e., by 2° for A and C and by 0.5° for B, so at each angle the cross section is measured four times with the same spectrometer but with different parts of the spectrometer acceptance. While the quality of the reconstruction is not good enough to split up the acceptance of a single measurement in smaller bins with the aim-for precision, this fourfold oversampling of the scattering angle allows us to recover the $Q^2$ dependence inside the acceptance of a single measurement. Spectrometer A is situated on the opposite side of the beam line from spectrometers B and C. Hence, a comparison in the overlap region between spectrometer A and the others is testing the beam, target, and rotation-axis alignment. Spectrometer A can be used in the range between 25° and 110°. Due to the construction of the target, the angle of A had to be limited to 90° for the long target cell and to 110° for the short target cell. Spectrometer C extends the angle range to over 130°. The slim construction of spectrometer B allows it to reach scattering angles down to 15.5°.

When the field of a spectrometer is changed, the magnetic fields change due to eddy currents and have to stabilize. Since this takes some time, the momentum was adjusted only every second angle change in order to keep the elastically scattered electrons at roughly a constant position on the focal plane of the spectrometers.

For most of the individual points, the required time to achieve the envisaged statistics of below 0.2% was around 30 min. Most settings were divided into two 5- to 20-min-long submeasurements. This reduces the statistical accuracy per submeasurement (which is compensated by the higher number of measurements) but increases the accuracy of the luminosity determined by the pA meter, which measures the beam current before and after each submeasurement and facilitates the search and elimination of time dependent effects.

The measurements were performed during three beam time periods, summarized in Table I.

### III. THEORY FOR ELASTIC ELECTRON PROTON SCATTERING

#### A. Cross section in the first Born approximation

The kinematical parameters of the elastic scattering of an electron on a target at rest is depicted in Fig. 2.

The incident electron has a four-momentum $k_1 = (E_1 = E, \vec{p}_1)$. It is scattered in the direction $\Omega = (\theta, \phi)$ with four-momentum $k_2 = (E_2 = E', \vec{p}_2)$. In the scattering process, the four-momentum $q = k_1 - k_2$ is transferred to the target via the exchange of a virtual photon. The target of mass $M$ is initially at rest, $P_t = (M, 0)$.

The unpolarized cross section is independent of the azimuthal angle $\phi$. Therefore, there are only two degrees of freedom and the cross section can be expressed in terms of the energy $E$ of the incoming electron and the scattering angle $\theta$ or, equivalently, by the (negative) four-momentum transfer squared $Q^2$ and the photon polarization $\varepsilon$.

In the one-photon-exchange approximation, the unpolarized cross section for the elastic scattering of an electron on a
proton with internal structure is given as follows:

\[
\left( \frac{d\sigma}{d\Omega} \right)_0 = \left( \frac{d\sigma}{d\Omega} \right)_{\text{Mott}} \left[ F_1^2 + \tau (\kappa F_2)^2 \right] + 2\tau (F_1 + \kappa F_2)^2 \tan^2 \frac{\theta}{2},
\]

with the dimensionless quantity \( \tau \equiv Q^2/(4m_p^2) \) and where

\[
\left( \frac{d\sigma}{d\Omega} \right)_{\text{Mott}} = \frac{4\alpha^2 E^2}{Q^2} E' \left[ 1 - \beta^2 \sin^2 \left( \frac{\theta}{2} \right) \right]
\]

is the recoil-corrected Mott cross section, which is the cross section for the scattering of a pointlike spin-\( \frac{1}{2} \) particle on a scalar pointlike target. The internal structure is expressed here in terms of the Dirac and Pauli form factors, \( F_1 \) and \( F_2 \), respectively. The relations

\[
G_E = F_1 - \tau \kappa F_2, \\
G_M = F_1 + \kappa F_2,
\]

translate the Dirac \( F_1 \) and Pauli \( F_2 \) form factors into the Sachs form factors \( G_E \) and \( G_M \). They were first proposed by Yennie et al. [27]. Sachs et al. [28,29] proposed that this choice provides a more physical insight than \( F_1 \) and \( F_2 \) since in the Breit frame, defined as \( P_1^B + P_2^B = (2E_B,0) \), the transition current reduces to

\[
J_B = e \chi_p^T (2MG_E,i\vec{\sigma} \times \vec{q} \cdot G_M) \chi_p.
\]

In this frame, \( G_E \) and \( G_M \) are the Fourier transforms of the spatial charge and magnetization distributions. A frequent criticism of this idea is that the Breit frame is equivalent to an infinitely heavy proton or, alternatively, a proton affixed to the coordinate origin and, therefore, the charge and magnetization distributions are not "real," i.e., frame dependent. A more detailed discussion of this point can be found in Ref. [1].

With the Sachs form factors, the cross section is given by

\[
\left( \frac{d\sigma}{d\Omega} \right)_0 = \left( \frac{d\sigma}{d\Omega} \right)_{\text{Mott}} \times \left[ G_E^2(Q^2) + \tau G_M^2(Q^2) + 2\tau G_M^0(Q^2) \tan^2 \frac{\theta}{2} \right] \]

\[
= \left( \frac{d\sigma}{d\Omega} \right)_{\text{Mott}} \varepsilon G_E^2 + \tau G_M^2. \tag{7}
\]

The choice of the Sachs form factors eliminates the mixed term in the cross section, which now depends on the squares of \( G_E \) and \( G_M \) only.

In the static limit \( Q^2 = 0 \), the form factors normalize to the charge and magnetic moment of the proton in units of the electron charge and of the nuclear magneton \( \mu_K \), \( G_E(0) = 1 \) and \( G_M(0) = \mu_p \).

The standard method to extract the form factors from measured cross sections is the Rosenbluth separation [30]. It exploits the linear structure in \( \varepsilon \) of Eq. (7) and separates the form factors by measurements at constant \( Q^2 \) but different \( \varepsilon \) values.

The somewhat unfamiliar method used in this paper consists in inserting many distinctly different form-factor models into Eq. (7) and fitting their parameters directly to the measured cross sections. This will be discussed in detail in Sec. V B and following subsections.

As mentioned above, in the Breit frame the Sachs form factors are the Fourier transforms of the charge and magnetization distributions. Expanding the kernel of the Fourier integrals in terms of \( Q^2 \) yields

\[
G(Q^2)/G(0) = 1 - \frac{1}{12}(r^2) Q^2 + \frac{1}{135}(r^4) Q^4 \ldots , \tag{8}
\]

where \( \langle r^n \rangle \) is the \( n \)-th moment of the electric or magnetic distribution. Therefore, the second moments can be determined by

\[
\langle r^2 \rangle = -6 \frac{dG(Q^2)}{dQ^2} \bigg|_{Q^2=0}, \tag{9}
\]

i.e., from the slope of the form factors at \( Q^2 = 0 \).

### B. Radiative corrections

It is not possible to measure the lowest-order cross section directly since higher-order diagrams, as depicted in Fig. 3, always contribute to the elastic scattering process. It is common practice to divide these contributions into groups with an additional virtual (v1–v5 in Fig. 3) or real photon (r1–r4). However, this grouping is problematic: Divergences in one group cancel against divergences in the other group, hence all graphs have to be evaluated at once. This leads to a correction \( \delta \) to the one-photon-exchange calculation,

\[
\frac{d\sigma}{d\Omega} = \left( \frac{d\sigma}{d\Omega} \right)_0 (1 + \delta). \tag{10}
\]

Here \( \left( \frac{d\sigma}{d\Omega} \right)_0 \) is the cross section for one-photon exchange alone [Fig. 3(b)] as given by Eq. (7), while \( \left( \frac{d\sigma}{d\Omega} \right)_1 \) is the cross section when next-to-leading-order contributions are taken...
into account (graphs v1–v5 and r1–r4 in Fig. 3). Conversely, the nonradiative cross section \( \frac{d\sigma}{d\Omega} \) can be determined in a first-order approximation by identifying the experimental cross section with \( \frac{d\sigma}{d\Omega} \) and dividing it by \( (1 + \delta) \).

The integrals over the internal four-momenta of the graphs v1–v3 are logarithmically divergent for large momenta. This can be treated theoretically by charge and mass renormalization. Details can be found in Refs. [31,32]. Graph v2 leads to an infrared divergence, but it can be shown [33,34] that this cancels with corresponding divergences of the graphs r1 and r2.

In the following, the formulas for the contributions from different groups of diagrams used in this work will be presented. For details of the calculation see Refs. [31,32].

The vacuum polarization (v1) gives rise to the term

\[
\delta_{\text{vac}} = \frac{\alpha}{\pi} \left\{ \left( v^2 - \frac{8}{3} \right) + \frac{3}{2} v^2 \ln \left( \frac{v + 1}{v - 1} \right) \right\}, \quad (11)
\]

\[
\frac{Q^2 \gg m_t^2 \to \frac{\alpha}{\pi} \left\{ -\frac{5}{3} + \ln \left( \frac{Q^2}{m_t^2} \right) \right\}, \quad (12)
\]

with \( v^2 = 1 + \frac{4m_t^2}{Q^2} \), where \( m_t \) is the mass of the particle in the loop. The approximation (12) is valid for loop electrons. However, at the energy scales of this experiment and within the envisaged accuracy, the vacuum polarization via muon and tau loops has to be accounted for and must be evaluated with Eq. (11).

The finite part of the electron vertex correction (v2, the infinite part cancels later on) is given in the ultrarelativistic limit by

\[
\delta_{\text{vertex}} = \frac{\alpha}{\pi} \left\{ \frac{3}{2} \ln \left( \frac{Q^2}{m^2} \right) - 2 - \frac{1}{2} \ln^2 \left( \frac{Q^2}{m^2} \right) + \frac{\pi^2}{6} \right\}. \quad (13)
\]

In the same limit, the contribution from real photon emission by the electron (r1, r2) yields the following:

\[
\delta_R = \frac{\alpha}{\pi} \left\{ \ln \left( \frac{(\Delta E_x)^2}{E E'} \right) \left[ \left( \frac{Q^2}{m^2} \right) - 1 \right] - \frac{1}{2} \ln^2 \eta \right\}
\]

\[
+ \frac{1}{2} \ln^2 \left( \frac{Q^2}{m^2} \right) - \frac{\pi^2}{3} + \text{Sp} \left( \cos^2 \frac{\theta_e}{2} \right) \right\}, \quad (14)
\]

where \( \eta = E/E', \Delta E_x = \eta \Delta E'. E' \) is the energy of an electron scattered elastically through an angle \( \theta \) when no photon is emitted. An electron which radiates a photon has a lower energy than \( E' \). \( \Delta E' \) is the maximum difference to \( E' \) allowed by the radiative tail cutoff; it is called the cut-off energy. Details about the Spence function \( \text{Sp}(x) \) can be found in Appendix B of Ref. [35].

The terms where the proton contributes (v3–v5, r3, and r4) are complicated and an exact calculation requires the knowledge of the internal structure of the proton. Maximon and Tjon [31] divide the correction in three parts, one proportional to the charge \( Z (\delta_1) \), one to \( Z^2 (\delta_2) \), and a third part in which they include all of the structure dependence \( (\delta_3^{(1)}) \). The last part is believed to be small for the kinematics of this work and is therefore neglected. The other two correction terms (the \( Z \) dependence is divided out) are given by

\[
\delta_1 = \frac{2\alpha}{\pi} \left\{ \ln \left( \frac{4(\Delta E_x)^2}{Q^2 x} \right) \ln \eta + \text{Sp} \left( 1 - \frac{\eta}{x} \right) - \text{Sp} \left( 1 - \frac{1}{\eta x} \right) \right\}, \quad (15)
\]

\[
\delta_2 = \frac{\alpha}{\pi} \left\{ \ln \left( \frac{4(\Delta E_x)^2}{m_p^2} \right) \left( \frac{E_p'}{|p'|} \ln x - 1 \right) + 1 \right\}
\]

\[
+ \frac{E_p'}{|p'|} \left[ -\frac{1}{2} \ln^2 x - \ln x \ln \left( \frac{\rho^2}{m_p^2} \right) + \ln x \right.
\]

\[
- \text{Sp} \left( 1 - \frac{1}{x} \right) + 2\text{Sp} \left( \frac{1}{x} + \frac{\pi^2}{6} \right) \right\}, \quad (16)
\]
with

\[ x = \left( \frac{Q + \rho}{4m_p^2} \right)^2, \quad \rho^2 = Q^2 + 4m_p^2. \]  

(17)

For vanishing cut-off energies and, hence, \( \Delta E_1 \), the corrections \( \delta_R, \delta_1, \text{and } \delta_2 \) get infinitely large. In this case, however, more photons than just one are emitted in each scattering event. It has been shown in Refs. [33,36] that this can be approximately taken into account by exponentiation of the corresponding correction terms as well as for the vertex correction. For the vacuum polarization contribution, Vanderhaeghen et al. [32] iterate the first-order contribution to all orders, which does not lead to an exponentiation. In total, they find

\[ \left( \frac{d\sigma}{d\Omega} \right)_{\text{exp}} = \left( \frac{d\sigma}{d\Omega} \right)_{0} e^{\delta_{\text{vac}} + \delta_R + Z\delta_1 + Z^2\delta_2}, \]  

(18)

which, for the kinematics used in the present work, differs marginally (below 0.05%) from the fully exponentiated form, which will therefore be used in the analysis of the measured cross sections as follows (\( Z \) has been set to 1):

\[ \left( \frac{d\sigma}{d\Omega} \right)_{\text{exp}}(\Delta E') = \left( \frac{d\sigma}{d\Omega} \right)_{0} e^{\delta_{\text{vac}} + \delta_{\text{mass}} + \delta_R + \delta_1 + \delta_2}(\Delta E'). \]  

(19)

It will be described in Sec. IV B 1 how these higher-order contributions are accounted for in the determination of the first-order cross section from the measured data.

C. Coulomb distortion and two-photon exchange

The Coulomb distortion, i.e., the scattering process via the exchange of many soft photons, and the related two-photon exchange, where both photons have a sizable momentum, is not fully included in the radiative corrections. There is yet some theoretical uncertainty in the modeling of these two effects. For the two photon effect, the off-shell nucleon and its excited states have to be modeled. For a discussion of the Coulomb distortion, see Ref. [35]. A complete treatment of these effects is not the topic of this paper.

Nevertheless, the Coulomb distortion cannot be ignored completely, especially for the determination of the radius, as has been shown by Rosenfelder [37]. He finds that the extracted radius is enlarged by about 0.018 fm when Coulomb distortion is accounted for. For \( Q^2 = 0 \), the correction is in agreement with the simple additional correction factor \( (1 + \delta_F) \), the so-called Feshbach correction [38,39], by McKinley and Feshbach as follows:

\[ \delta_F = Z\alpha\pi \frac{\sin^2 \frac{\theta}{2} - \sin^2 \frac{\theta}{2}}{\cos^2 \frac{\theta}{2}}. \]  

(20)

This correction has been applied to the measured cross sections and is maximal for 180° scattering yielding a downward correction of 1.2%.

The TPE becomes more important at larger \( Q^2 \) and may explain the difference between polarized and unpolarized measurements at large \( Q^2 \) [11]. Therefore, a lot of theoretical work focuses on the energy scales above 1 GeV. In 2007, Arrington et al. [4] have reanalyzed the world data set with a model for two-photon-exchange corrections and made two fits, one with the corrections applied and one without. The ratio of these fits represents an estimate of the two-photon effect on the form-factor ratio in the higher \( Q^2 \) region of the present experiment, which can be used to compare the form-factor ratio from fits to Rosenbluth data with previous polarized measurements. However, one has to keep in mind that these fits rely on just one model [18,19] and a generalization is uncertain. Therefore, we have chosen a different phenomenological approach (Secs. VI C and V C 4).

IV. DETERMINATION OF THE CROSS SECTIONS

A. Overview

In order to calculate the cross section from counting rates one has to know the luminosity and the acceptance of the detector. However, the acceptance is not just a fixed number given purely by the collimator geometry; it also depends on the target length and position and on the spectrometer angle and the momentum of the particles. The only feasible way to determine the cross section from the measured number of scattering events is by comparing this number to the result of a full simulation of the experiment, \( \sigma_{\text{sim}} \), including all aspects of the detector response, external energy loss of the electrons in the target material, and all radiative corrections. The measured cross section is then found as

\[ \sigma_{\text{rel,exp}} = A - B \sigma_{\text{sim}} \mathcal{L}. \]  

(21)

Here \( A \) is the number of counts in the peak region integrated to the cut-off energy \( \Delta E' \), \( B \) is the estimated background in this region, \( \sigma_{\text{sim}} \) is the simulated cross section including radiative corrections integrated over the acceptance of the spectrometers, and \( \mathcal{L} = \int \mathcal{L}_{\text{eff}} dt \) is the time-integrated effective, i.e., prescaling and dead-time corrected, luminosity. For the calculation of \( \sigma_{\text{sim}} \), one has to make use of an assumed cross section which should be sufficiently close to the true one. As a result, Eq. (21) yields the measured cross section relative to the assumed one.

We define the data taken at one energy and angle over some time as a “run.” The runs are grouped into a “set of runs,” where the relative normalizations of the runs to each other in the same set are determined by the luminosity-monitor measurements (see Sec. IV C 4). With the setup of this experiment, as well as with any other, the absolute normalization of each run, i.e., the luminosity, which comprises the absolute knowledge of the target length, the absolute current calibration, and the absolute detector efficiencies, can be determined only to the level of a few percent. Therefore, the relative normalization between different sets of runs will be left floating in the fits of the final analysis. The absolute normalization, finally, is fixed by the known values of the form factors at \( Q = 0 \), as already mentioned.

B. Cross-section simulation

Instead of correcting the measured number of events for efficiencies, acceptance problems, and the radiative processes, as was the case in the classical electron-scattering experiments,
these ingredients are better incorporated into the simulated cross section as discussed in the following.

1. Internal radiative corrections

As described above, the extraction of the first-order Born cross section (\( \frac{d\sigma}{d\Omega} \)) of the process \( ep \rightarrow e' p' \) also requires the calculation of a (radiation) correction factor \( f_{\text{corr}} \). This factor depends on the kinematics and cut-off energy \( \Delta E' \) in the spectrum of the scattered electrons,

\[
\left( \frac{d\sigma}{d\Omega} \right)_{\text{exp}} = \left( \frac{d\sigma}{d\Omega} \right)_{0} f_{\text{corr}}(\Delta E', E, \theta),
\]

or, differential in the cutoff energy,

\[
\left( \frac{d\sigma}{d\Omega d\Delta E'} \right)_{\text{exp}} = \left( \frac{d\sigma}{d\Omega} \right)_{0} \frac{df_{\text{corr}}}{d\Delta E'},
\]

where \( \frac{d\sigma}{d\Omega} \) is the radiative tail. A generator algorithm for a Monte Carlo calculation has to produce events replicating this cross section in an efficient manner, i.e., without too great computational costs and with a choice of the kinematic quantities so the variance of the weights of the individual events is minimized. The generator is based on a generator for events for a virtual Compton-scattering experiment described in Ref. [40]. In its original form, it was limited to a description of the shape of the tail without correct global normalization. In the course of the present work, it was extended to also describe accurately the peak region and to have the correct normalization. The generator first generates a vertex position, a scattering angle, and an azimuthal angle from pseudorandom normalization. The generator first generates a vertex position, a scattering angle, and an azimuthal angle from pseudorandom number sequences. Then, it follows the principles laid out in Vanderhaeghen et al. [32] in calculating the energy of a radiated photon. In the next step, a direction of this photon has to be generated. Since the cross section depends strongly on the photon angle, peaking close to the directions of the electrons, but vanishing at the exact directions, it is important to generate the photon direction with importance sampling, that is, generating more events where the cross section is higher so the weights of the events are nearly constant. To this end, one generates the directions using a suitable approximation of the Bethe-Heitler part of the cross section. Such an approximation is given by the sum of individual cross sections for radiation off the incoming or outgoing electron, neglecting the interference,

\[
\left( \frac{d\sigma}{d\Omega} \right)_{\text{approx.}} = \frac{1}{2} \left( \frac{d\sigma}{d\Omega} \right)_{e} + \frac{1}{2} \left( \frac{d\sigma}{d\Omega} \right)_{e'},
\]

\[
\left( \frac{d\sigma}{d\Omega} \right)_{e} = \frac{1}{N(E, \bar{p})} \frac{1 - \cos^{2} \theta_{e,\gamma}}{(|\bar{p}| - \cos \theta_{e,\gamma})^{2}},
\]

\[
N(E, \bar{p}) = -4 - 2 \frac{E}{|\bar{p}|} \ln \left( \frac{E}{|\bar{p}|} - 1 \right) \left( \frac{E}{|\bar{p}|} + 1 \right),
\]

and the same expressions for \( E', p', \) and \( \theta_{e',\gamma} \). \( E (E') \) is the incoming (outgoing) electron energy, \( \bar{p} (p') \) the corresponding momentum, and \( \theta_{e,\gamma} (\theta_{e',\gamma}) \) the angle between the incoming (outgoing) electron and the photon. The generator selects with equal probability whether the photon is radiated from the incoming or outgoing electron.

Then the transformation method is used to generate random values with a distribution according to Eq. (24). The cumulative distribution is given by

\[
F(\theta_{e,\gamma}) = \frac{\cos(\theta_{e,\gamma})}{\frac{d\sigma_{1}}{d\Omega_{\text{rad}}}} \frac{d\sigma}{d\Omega} \frac{d \cos \theta_{e,\gamma}}{d \Omega_{\text{rad}}},
\]

\[
= \frac{1}{N(E, \bar{p})} \left( \frac{E}{|\bar{p}|} - \cos \theta_{e,\gamma} \right)^{2} - 2 + \frac{E}{|\bar{p}|}.
\]

A uniformly distributed number \( r \) between 0 and 1 is now transformed by solving \( r = F(\theta_{e,\gamma}) \) to the new random variable \( \theta_{e,\gamma} \) with the correct distribution (see Ref. [41], Sec. 7.2). The required inversion of \( F \) is realized numerically via a bisecution method.

The innermost part of the generator calculates the Feynman graphs of the lowest order describing the Bethe-Heitler (radiation from the electron) + Born (radiation from the proton) processes for the now-fixed kinematics. Here a Jacobian for the transformation \( d\Omega_{1}^{\text{ab}} \) to \( d\Omega_{1}^{\text{cm}} \) has to be taken into account. It is calculated numerically using finite differences.

The cross section calculated with these graphs is infrared divergent. This is accounted for by a modification of the propagators. Their denominators are

\[
\text{Bethe-Heitler} : 2k_{1}q^{\text{rad}} - 2k_{2}q^{\text{rad}},
\]

\[
\text{Born} : -2pq^{\text{rad}}, p'q^{\text{rad}},
\]

where \( k_{1}(k_{2}) \) is again the four-vector of the incoming (outgoing) electron and \( q^{\text{rad}} \) the four-vector of the radiated photon evaluated in the center-of-mass (c.m.) system. Here, \( q^{\text{rad}} \) is replaced with \( q_{\text{mod}}^{\text{rad}} = q^{\text{rad}}/|q^{\text{rad}}| \). Hence, the calculation yields the correct cross section multiplied with a factor \( K^{2} = |\vec{q}^{\text{rad}}|^{2} \) since the matrix element enters quadratically into the cross section. One order of \( K \) is then divided out at the cross-section level, and the remaining order has to be accounted for later when the different parts of the generator are combined.

2. External radiation

In an extended target, the material in the path of the particles before and after the scattering inflict an energy loss in addition to the internal processes. When the cryogenic target is used, the incoming beam has to pass through different layers of matter until the scattering process occurs. This includes the walls of the target and the liquid hydrogen inside the target. Additionally, the cold target acts as a cold trap. “Snow,” i.e., frozen water and nitrogen from the residual gas inside the vacuum chamber, can build up on the target surface and is easily identified as additional peaks in the scattering spectrum. Switching to high current melts the snow where the beam enters and exits the target. However, snow remains on the sides of the cell. The outgoing electron has to pass part of the hydrogen, the wall of the target, possibly snow, and then
the windows between the spectrometer and vacuum chamber and a short distance of air between them. In all these layers, the electron loses energy by external bremsstrahlung and ionization of the atoms. These processes have to be folded with the internal bremsstrahlung spectrum; the simulation does this numerically.

3. Resolution

The resolution of the drift chambers, the characteristics of the electronics, and the knowledge of the transport inside the magnetic system give rise to specific error distributions for the extracted kinematical variables of the detected particles at the target. In the simulation, we employ a simple Gaussian to model the error distribution in the reaction vertex and the sum of two Gaussians with different weights and width to model the error distributions in the in-plane and out-of-plane angles and in the momentum. With two Gaussians, the longer tails in the distributions of the errors in these variables are better reproduced than by a single Gaussian.

4. Test of the description of the radiative tail

In the classical electron scattering experiments the radiative correction was calculated from a cutoff energy by Eq. (18). By contrast, we calculate in the simulation the full radiative tail, accounting for the convolution over the complicated acceptance of the spectrometers, for the convolution of internal and external energy loss, and for the dependence of the cross section on the energy. With the classical radiative correction one meets a dilemma: A large cutoff is favored by the then-small correction; however, there comes in the unaccounted dependence of the cross section on the energy. The latter disadvantage is minimized by a small cutoff, but then one has fewer events, the correction is large, and there enters an uncertainty from the resolution of the spectrometer. Compared to the evaluation of the old measurements, the specific choice of the cutoff energy is considerably less important in our method, where the measured cross section is given by a comparison of the simulated and measured spectrum, giving full account to the acceptance and to the energy dependence of the cross section. We demonstrate the quality of the description of the radiation tail in Fig. 4, which shows a comparison of the measured \( \Delta E'_{\text{exp}} \) spectrum and the simulated tail. For this comparison, the background in the measured spectrum is suppressed by a vertex cut. The upper panel compares the two spectra directly in a logarithmic scale. The downward bend at large \( \Delta E'_{\text{exp}} \) is due to the finite momentum acceptance of the spectrometer. The lower panel shows the ratio of the integrals over the two distributions which, finally, gives the experimental cross section: Over the wide range of cut-off energies the result depends on \( \Delta E'_{\text{exp}} \) by less than 10\(^{-3}\).

C. Determination of the experimental cross sections

In this section we discuss the determination of the number of events \( A \), the background \( B \), and of the luminosity \( \mathcal{L} \) for Eq. (21) and of some further corrections and anomalies.
deviation of the particle momentum from the spectrometer’s central momentum, to have a well-defined momentum acceptance. Figure 5 shows a typical measured spectrum of $\Delta E_{\text{exp}}'$ and the effect of these cuts, which are summarized in Table II. To select the elastic reaction for the extraction of the cross section, we apply a cut in $\Delta E_{\text{exp}}'$ accepting events only in the elastic peak region up to a certain cut of $\Delta E_{\max}$.

For spectrometer B, an additional cut has to be applied. With this spectrometer particles can be detected whose initial trajectory between target and spectrometer lies outside the acceptance defined by the collimator. These particles hit the long snout in front of the collimator and may be scattered back into the acceptance and arrive at the focal plane. They are identified by the horizontal coordinate at the entrance of the snout, $y_{\text{snout}}$. Figure 6 shows a two-dimensional histogram of the horizontal coordinate at the collimator, $y_{\text{coll}}$, versus $y_{\text{snout}}$. The events around $y_{\text{snout}} = 0$ correspond to good events. On the other hand, one identifies a shadow on the right and a dimmer shadow on the left side belonging to events from snout scattering. In the final analysis, a cut with $|y_{\text{snout}}| < 30$ mm was applied. As one can see in Fig. 7 showing the distribution of the events that are suppressed by this cut, the electrons scattered by the entrance snout are located in the region of the radiative tail.

### 2. Determination of resolution and central momentum

In the simulation, the accuracy of the determination of the particle coordinates is parameterized in the target reference frame, i.e., the simulation contains parametrizations for the resolution in the vertex position, in the momentum and in the in-plane and out-of-plane angles. These parametrizations include parameters for the width of the distributions, which

![Figure 5](image_url)  
**FIG. 5.** Typical $\Delta E_{\text{exp}}'$ spectrum for a measurement with spectrometer A at $90.5^\circ$ at an incident energy of $E = 585$ MeV. (a) Spectrum without any cut. (b) After all cuts except a cut in $\Delta E_{\text{exp}}'$. (c) Events that are rejected by the cuts. Random events give rise to the nearly constant level between $-30$ and $15$ MeV. The bump around $30$ MeV originates from events detected near the edges of the detector plane.

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<td>$\Delta p_c$</td>
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![Figure 6](image_url)  
**FIG. 6.** Histogram of $y_{\text{coll}}$ versus $y_{\text{snout}}$ of the measurement with spectrometer B at $32.5^\circ$ at an incident energy $E = 180$ MeV. The events around $y_{\text{snout}} = 0$ correspond to good events, and the sidebands result from back scattering from the snout walls. The black vertical lines indicate the cut used in the analysis. The gray scale is logarithmic to emphasize the events in the left and right sideband. The diagonal ridges seen in the good events are caused by scattering off the target walls.
depend on the kinematics and which are determined for each run individually.

It is particularly important that the peak position in the $\Delta E_{\mathrm{exp}}$ histogram matches between experiment and simulation, otherwise the cut in $\Delta E_{\mathrm{exp}}$ would fail to select the equivalent part of the peak and tail region in both the experiment and the simulation. This is governed by the knowledge of the central momentum of the spectrometer, determined by the strength of the magnetic field.

The determination of the central momenta and resolutions is done in a two-step process. The first step is to find the vertex resolution and, together with this, a possible target offset. For this, a standard nonlinear least-squares optimization is performed: For each variation of the parameters, the simulation is run, and the vertex histograms of experiment and simulation are compared. Since the reconstruction of the measured data is not dependent on these values, only the simulation has to be updated at each fit iteration.

In the second step, the central momentum and the remaining resolutions are optimized. To this end, the spectra of $\Delta E_{\mathrm{exp}}$, of the angles, and of $\Delta p_t$ (the momentum relative to the central momentum) are compared. Since the central momentum value affects the reconstruction of the electron kinematics from the measured data, both data analysis and simulation have to be updated at each step.

3. Background subtraction

As mentioned in Sec. II, the liquid hydrogen is contained in a cryo cell. The electron beam has to pass through the walls of this cell, a 10-μm-thin foil made of HA V AR, an alloy of several metals, and also possibly pass through a layer of snow. Scattering off the wall or snow nuclei produces background in the energy region of the elastic peak of the hydrogen, both from the radiative tail from elastic scattering off the proton, and from quasielastic scattering.

The shapes of the background from both elastic and quasielastic scattering are taken from simulations based on a physical model, while their amplitude is fitted to the data. As verified by empty target measurements, the inelastic peaks are either so small that they can be ignored (<0.035% of the elastic hydrogen peak in the worst case) or they are outside the region around the elastic hydrogen peak accepted for the cross-section determination. In the latter case, the region of the inelastic peak is excluded from the fit of the background amplitudes.

Figure 8 displays a measured spectrum and the difference spectrum, i.e., the data histogram minus the three simulated and scaled spectra, elastic off hydrogen, elastic off wall or snow atoms, and quasielastic off wall or snow atoms. One sees the excellent agreement in the region of the radiative tail from elastic scattering off the proton, while there are slight
imperfections at $\Delta E'_{\text{exp}} = 0$ around the steep fall offs of the hydrogen peak, which, however, level out to zero in the integral (cf. Fig. 8 bottom). The first measurement at this kinematic, shown in the upper plots and in black in the bottom plot, has about half of the statistics of the second measurement, which was taken shortly after. The depression close to the peak is caused by a minuscule offset in the peak position—the second measurement is better in this regard. It has to be noted that for both measurements, the integral ratio varies less than $\pm 0.15\%$ between 7 and 30 MeV. (N.B.: In Ref. [8], the same data were erroneously labeled differently.)

4. Luminosity

In order to determine the integrated luminosity, the beam current is continuously measured during data taking with a fluxgate magnetometer. At the start and end of each run, the beam current measurement is compounded with the pA-meter measurement on the collimator as described above. This reduces the uncertainty for small currents and short runs.

The target density is calculated from continuous pressure and temperature measurements. In order to avoid local overheating of the liquid hydrogen due to the heat load of the passing electron beam, at high currents the beam is rastered over the (curved) frontal face of the target. The small change in the effective target thickness due to this rasterization is accounted for by the simulation. Still, the absolute length of the cooled-down cryo cell is hard to determine to better than 1%. Uncertainties in its determination, however, enter as a constant factor in the normalization which will be taken as a fit parameter, as already mentioned.

For all measurements, one of the three spectrometers was used as a luminosity monitor, i.e., this spectrometer stayed with the same field at the same angle, thus measuring the count rate for a fixed momentum transfer for a time where many runs at different angles were taken with the other spectrometers, i.e., for one set of runs. This spectrometer thus monitors the relative luminosity. In the course of the measurements at one energy, only a few changes of the monitor angle are necessary to ensure that its event rate is high enough. Each measurement of the luminosity monitor is analyzed in the same way as the normal cross-section measurements, that is, the normal procedure of background subtraction, dead time correction, and normalization to the estimated luminosity is performed.

From the $n$ individual luminosity results in a set of runs, the average cross section is calculated and the cross-section values $\sigma_{\text{exp},i}$ measured with the other spectrometers, are now normalized:

$$\sigma_{\text{exp, norm},i} = \frac{\sigma_{\text{exp},i}}{\sigma_{\text{lum, avg}}} \sigma_{\text{lum},i}. \quad (28)$$

Hence, the common factors in the calculation of $\sigma_{\text{exp},i}$ and $\sigma_{\text{lum},i}$, i.e., beam current, target density, and target length, cancel out and uncertainties in their determination play no role apart from the set-to-set normalization taken as a fit factor in the final analysis.

In this procedure, the statistical error of the normalized data is enhanced by the statistical error of the luminosity measurement,

$$\frac{\Delta \sigma_{\text{exp, norm},i}}{\sigma_{\text{exp, norm},i}} = \sqrt{\left(\frac{\Delta \sigma_{\text{exp},i}}{\sigma_{\text{exp},i}}\right)^2 + \left(\frac{\Delta \sigma_{\text{lum},i}}{\sigma_{\text{lum},i}}\right)^2}. \quad (29)$$

This method provides a stable and precise determination of the relative normalization. It is therefore applied to all data points with exception of the 315-MeV data set. In the analysis of this subset, an error was found in the setup of spectrometer C, which was used as the luminosity monitor for most of these data. Therefore, for this subset we rely on the luminosity provided by the pA meter, as described in Sec. II D 1. This leads to slightly larger uncertainties (cf. Sec. V C 2). Compared to the pA-meter-based luminosity, we observe changes with a one-$\sigma$ width of about 0.1–0.2% for energies of 450 MeV and higher. We find a similar width for the higher currents at 180 MeV. For the lowest currents, the pA-meter measurement and the beam current itself are more noisy and the corrections have a width of about 0.6%.

As described in Sec. IV A, the absolute normalization will be determined by fitting normalization constants to groups of cross sections, determined in a first step with the measured beam currents and target thickness and corrected by the relative luminosity measurements. It is reassuring, that the normalizations of the different sets of measurements are in the expected range of $\pm 3.5\%$ and that they do not depend on the form-factor model used in the fit in any significant way. The largest model spread of the normalization constants is 0.26% for the flexible models discussed below and 0.74%, including the Friedrich-Walcher model. This large difference occurs for cross sections with the highest $Q^2$ and is caused by the different high-$Q^2$ behavior of this model (see Sec. V D 3). The average standard deviation is 0.074%.

5. Further corrections and anomalies

The statistical precision achieved in this experiment together with the conceptual design of overlapping acceptances made several anomalies apparent, which would have been missed in a traditional type of experiment. In the course of the analysis it was found that the acceptances of spectrometers A and C are not completely given by the sheer geometry of the collimators; instead, they depend to some extent on the vertex position. A corresponding correction has been developed and implemented in the simulation software.

Furthermore, it was found that the stray magnetic field of spectrometer C influences the measurement with spectrometer B when the spectrometers are close to each other. In a dedicated beam time, a correction formula has been determined which has been applied to the data. Further details can be found in Ref. [42].

V. DATA ANALYSIS

A. World data basis

In order to provide a coherent analysis and fit of all world data based on the methods developed by this work we included all readily available data related to the determination of the proton form factors known up to today. The inclusion of experiments with polarized electrons [16,17,43–55] allows
In this analysis, to extract a phenomenological two-photon effect. To this end we simply add the published form-factor ratios to the database, with systematic and statistical errors added linearly. Some of the listed references are reanalysis of older measurements—we then only include the updated values.

For Rosenbluth-type measurements [56–70] we include the cross sections instead of extracted form factors in order to make full use of the available information without any bias. We use the quoted statistical errors and let the normalization of the different data sets float. These data sets were taken over the time period of several decades and naturally include a diverse set of radiative corrections. We divided the old corrections out and applied the updated corrections as far as possible to a common standard, following the Maximon-Tjon prescription [31], with the extension to muon and tau loops and exponentiation of the correction to account for higher orders, matching the treatment of the new data as close as possible.

In Refs. [56–61], the radiative corrections are based on Ref. [71], in different variations, and they are straightforward to update. Reference [62] also belongs to this group. For this data set, we include two normalization constants; the data were taken with two different spectrometers and the two subsets show a clear normalization mismatch in the overlap.

The corrections in Refs. [63–67] are based on Ref. [72]. For the two Borkowski et al. data sets [63,64] we assume independent normalizations and include a normalization parameter for each data set.

For Refs. [68–70] the radiative corrections could not be updated since no details on the applied corrections could be obtained.

### B. Form-factor models

For a direct fit of the measured cross sections, an ansatz has to be made for the description of the form factors. Since the true form-factor functional form is unknown, we have to rely on a subjective choice, possibly introducing a bias. However, we can reduce the impact of this by employing a wide variety of models. The model bias can then be judged in this context.

In the frequentist picture, each of these models, together with a specific choice of the parameters, constitutes a hypothesis we can test against the data. For each model we choose the parameter set with the highest $p$-value via a least-squares fitter. From the goodness of fit of the different models, we can rule out some of them; however, this actually leaves the domain of the strict frequentist view. In the Bayesian picture, a selection of a model constitutes a prior. From the infinite function space, we reduce our selection to those which are representable by the model, attributing zero probability to all the others. Additionally, the fit essentially assumes a flat prior for the probability distribution for the parameters.

In the following the models used in this work will be discussed. For the magnetic form factor, $G_M$, the factor $\mu_p$ has been suppressed to improve readability. All models are normalized to 1 at $Q^2 = 0$. This will be used by the fit to fix the global normalization.

#### 1. Dipole

The designation “standard dipole,”

$$G_{\text{standard dipole}}(Q^2) = \left(1 + \frac{Q^2}{0.71 \text{ GeV}^2}\right)^{-2}, \quad (30)$$

was coined by Hand et al. [73]. For a long time it was the accepted form for the electric form factor of the proton and—scaled with $\mu_p/n$—also for the magnetic form factor of both the proton (“scaling relation”) and the neutron, and it is today found in many textbooks (e.g., Ref. [74]). While the choice of the dipole form was originally purely phenomenological, the related exponential falloff in $r$-space comes about as the probability function of a quantum mechanical particle trapped in a narrow potential well.

In the present analysis, the scaling relation is not enforced. Instead, different parameters for the electric and magnetic form factor are used,

$$G_{\text{dipole}}^{E,M}(Q^2) = \left(1 + \frac{Q^2}{a_{E,M}^2}\right)^{-2}. \quad (31)$$

With only two free parameters, $a_E$ and $a_M$, this model is very rigid, and it will be seen that it is not able to describe the data of this experiment, as was the case already for earlier data (e.g., Simon et al. [60]).

#### 2. Double dipole

A somewhat more flexible ansatz consists of the sum of two dipoles,

$$G_{\text{double dipole}}^{E,M}(Q^2) = a_0^{E,M} \left(1 + \frac{Q^2}{a_1^{E,M}}\right)^{-2} + \left(1 - a_0^{E,M}\right) \left(1 + \frac{Q^2}{a_2^{E,M}}\right)^{-2}. \quad (32)$$

#### 3. Polynomials

- **a. Simple polynomial.** A polynomial is a simple model without theoretical idea of the nature of the form factors except some level of continuity or smoothness. The constant term is fixed to 1 by the normalization. With a polynomial of the order $n$, the form factors are parameterized as follows:

$$G_{\text{polynomial},n}^{E,M}(Q^2) = 1 + \sum_{i=1}^{n} a_i^{E,M} Q^{2i}. \quad (33)$$

Since the form factors drop rapidly with $Q^2$, high orders are needed to describe them adequately over a larger $Q^2$ range.

- **b. Polynomial × dipole.** In order to free the polynomial from the necessity to describe the gross behavior of the form factors, the latter may be accounted for by multiplying the polynomial by the standard dipole as follows:

$$G_{\text{polynomial} \times \text{dipole},n}^{E,M}(Q^2) = G_{\text{dipole}}^{E,M}(Q^2) \left(1 + \sum_{i=1}^{n} a_i^{E,M} Q^{2i}\right). \quad (34)$$
In principle, it would be possible to optimize also the parameter of the dipole. It was found, however, that this additional freedom does not improve the fits and has a high computational cost.

c. Polynomial + dipole. A variation of the aforementioned splitting off of the gross behavior of the form factors is the sum of a polynomial and the standard dipole instead of the product,

\[ G_{\text{polynomial+dipole}}^{E,M}(Q^2) = G_{\text{standard dipole}}(Q^2) + \left( \sum_{i=1}^{n} a_i^{E,M} Q^{2i} \right). \]  

While the multiplication parameterizes the relative deviation from the standard dipole, the sum parametrizes the absolute deviation.

d. Inverse polynomial. A variation of the polynomial model is the inverse polynomial ansatz as in Ref. [3],

\[ G_{\text{inv. poly.}}^{E,M}(Q^2) = \frac{1}{1 + \sum_{i=1}^{n} a_i^{E,M} Q^{2i}}. \]  

4. Splines

In all other models described in this section, the behavior of the function in different \( Q^2 \) regions is highly correlated. Therefore, possible shortcomings in the description of the data in one \( Q^2 \) region may influence negatively the description in other regions. Functions that decouple the behavior in different \( Q^2 \) regions to a great extent are splines.

A spline ansatz has multiple advantages. Depending on the number of knots, a spline can be very flexible. Nevertheless, the fit converges even for a large number of knots quickly since the fit converges even for a large number of knots quickly since it had the best results in Ref. [80] for existing proton data. While this was studied in [75] for a model-independent analysis of nuclear charge distributions in 3 space. The bump contribution is hence described by the following:

\[ G_b(Q^2, Q_b, \sigma_b) = e^{-\frac{1}{2} \left( \frac{Q^2}{\sigma_b^2} \right)^2} + e^{-\frac{1}{2} \left( \frac{Q^2}{\sigma_b^2} \right)^2}. \]  

To attribute the full normalization to the smooth part, the bump contribution is multiplied by \( Q^2 \). The complete model is, therefore,

\[ G^{E,M}_{\text{Friedrich-Walcher}}(Q^2) = G_s(Q^2, a_{0,1,2}) + a_b^{E/M} Q^2 G_b(Q^2, \sigma_b^{E/M}). \]  

6. Continued fraction

A popular model introduced by Sick [76] is the continued fraction ansatz. However, it turns out that this model produces slowly converging fits and the results are difficult to control due to poles in the denominator. While this was studied in some detail, it was not included into the final analysis.

7. Extended Gari-Krümpelmann model

While all previous models are just mathematical procedures with no physical meaning for the description of the data, the extended Gari-Krümpelmann model [77–80]—actually a group of models which differ in their details—is based on physical considerations. In this work, the version called DR-GK(1) [78] (respectively, GKex(01) [79,80]) is selected, since it had the best results in Ref. [80] for existing proton form-factor data when the normalization of the data sets is not varied.

Under the assumption that QCD is the fundamental theory of the strong interaction, the \( Q^2 \) dependence of the electromagnetic form factors can be calculated in perturbative QCD.
(pQCD) for very high momentum transfers. For small momentum transfers, the confinement property of QCD leads to an effective hadronic description with vector meson dominance (VMD), the coupling of a photon to a vector meson which itself couples to the nucleon. Earlier models that were based solely on VMD introduced multiple, phenomenological poles of higher mass in addition to the $\rho$, $\omega$, and $\phi$ poles. Gari and Krümpelmann limit the VMD contributions to these three poles but enforce the asymptotic $Q^2$ behavior dictated by the scaling behavior of pQCD by additional terms.

In the model used here, the dispersion integral approximation of the $\rho$-meson contribution is replaced by an analytical form. The model was extended to include the $\rho'(1450)$ pole (for details see Refs. [77,78]).

As mentioned in Sec. III A, the Sachs form factors can be rewritten in terms of the Dirac and Pauli form factors $F_i$ and $F_\pi$, which are preferred by VMD models. Those can be divided into an isoscalar and an isovector component as follows:

$$2F^p_{i,2} = F^{iv}_{i,2} + F^{iv}_{-i,2}, \quad 2F^n_{i,2} = F^{is}_{i,2} - F^{is}_{-i,2}. \quad (44)$$

The model GKE(01) is formulated in terms of these four form factors with the poles for $\rho$, $\rho'$, $\omega$, $\omega'$, and $\phi$ mesons,

$$F^{iv}_{i}(Q^2) = \frac{N}{2} \left[ 1.0317 + 0.0875 \left( 1 + \frac{Q^2}{0.3176 \text{GeV}^2} \right) \right]^{-1} F^i(0^2)$$

$$+ \frac{\frac{g_{\rho',\rho}}{f_{\rho',\rho}}}{m_{\rho'}^2 + Q^2} F^i_{\rho'}(Q^2)$$

$$+ \left( 1 - 1.1192 \frac{N}{2} \frac{g_{\rho',\rho}}{f_{\rho',\rho}} \right) F^i_{\rho'}(Q^2),$$

$$F^{iv}_{i}(Q^2) = \frac{N}{2} \left[ 5.7824 + 0.3907 \left( 1 + \frac{Q^2}{0.1422 \text{GeV}^2} \right) \right]^{-1} F^i_{\omega}(Q^2)$$

$$+ \frac{\frac{g_{\omega',\omega}}{f_{\omega',\omega}}}{m_{\omega'}^2 + Q^2} F^i_{\omega'}(Q^2)$$

$$+ \left( 1 - 6.1731 \frac{N}{2} \frac{g_{\omega',\omega}}{f_{\omega',\omega}} \right) F^i_{\omega'}(Q^2), \quad (45)$$

$$F^{is}_{i}(Q^2) = \sum_{\alpha=\rho,\omega,\rho',\omega'} \frac{f_{\alpha}}{m_{\alpha}^2} m_{\alpha}^2 + Q^2 F^i_{\alpha}(Q^2)$$

$$+ \left( 1 - \frac{g_{\alpha}}{f_{\alpha}} - \frac{g_{\alpha}}{f_{\omega}} \right) F^D(0^2),$$

$$F^{is}_{i}(Q^2) = \sum_{\alpha=\rho,\omega,\rho',\omega'} \frac{\kappa_{\alpha}}{f_{\alpha}} m_{\alpha}^2 + Q^2 F^i_{\alpha}(Q^2)$$

$$+ \left( 1 - \frac{g_{\alpha}}{f_{\alpha}} - \kappa_{\omega} \frac{g_{\omega}}{f_{\omega}} - \kappa_{\phi} \frac{g_{\phi}}{f_{\phi}} \right) F^D(0^2).$$

In this model, the form factors $F^\alpha_i$ ($\alpha = \rho$, $\omega$, $\rho'$, $\omega'$, $\phi$, meson-nucleon) and $F^D_i$ (quark-nucleon) are parameterized as follows:

$$F^\alpha_i(Q^2) = \frac{\lambda^2_{\alpha,D}}{\lambda^2_{\alpha,D} + Q^2} \lambda^2_{\rho} + Q^2,$$

$$F^\rho_i(Q^2) = \frac{\lambda^2_{\rho,D}}{\lambda^2_{\rho,D} + Q^2} \lambda^2_{\rho} + Q^2,$$

$$F^\phi_i(Q^2) = \frac{\lambda^2_{\phi} Q^2}{\lambda^2_{\phi} + Q^2}, \quad (46)$$

$$F^\phi_i(Q^2) = \frac{\lambda^2_{\phi} Q^2}{\lambda^2_{\phi} + Q^2}, \quad (46)$$

$$\tilde{Q}^2 = Q^2 \ln \frac{\left( \lambda^2_{\rho} + Q^2 / \lambda^2_{\rho,\omega} \right)}{\ln \left( \lambda^2_{\rho} / \lambda^2_{\rho,\omega} \right)} \quad (47)$$

The parametrization fulfills the normalization constraint for $Q^2 = 0$. The constants $\lambda_{\rho}$ and $\lambda_{\phi}$ and the masses $m_{\rho}$, $m_{\omega}$, $m_{\phi}$, $m_{\rho'}$, and $m_{\omega'}$ are taken as $\lambda_{\rho(1450)} = 3.706$, $\lambda_{\phi} = -1.12$, $m_{\rho(1731)} = 0.776 \text{GeV}$, $m_{\omega(1270)} = 0.784 \text{GeV}$, $m_{\phi} = 1.019 \text{GeV}$, $m_{\rho'} = 1.45 \text{GeV}$, and $m_{\omega'} = 1.419 \text{GeV}$.

There remain at most 14 free parameters: eight couplings (four $g_\alpha/f_\alpha$, four $\kappa$); four cut-off masses ($\lambda_1$, $\lambda_2$, $\lambda_D$, and $\mu_{\phi}$); the mass $\lambda_{QCD}$, which gives the size of the logarithmic $Q^2$ behavior; and the normalization parameter $N$ for the dispersion relation part of the $\rho$ meson.

In Ref. [79], at most 12 of these parameters were varied, since either the $\omega'$-meson contribution was neglected or $\lambda_{QCD}$ and $\lambda_{QCD}$ were fixed to $N = 1$ and $\lambda_{QCD} = 0.150 \text{GeV}$, the physical value. The latter constraints are also used in the present work. Still, the fit shows slow convergence and a high time complexity because of the type of mathematical operations used and the mathematical properties of the formulas.

8. Other models not described in this paper

The new high-precision Mainz data were used in several fits not described in this paper, including simultaneous fits to both proton and neutron form factors. Bauer et al. [81] calculate the electromagnetic form factors of the nucleon to third chiral order in manifestly Lorentz-invariant effective field theory and fit to the Mainz data for the proton and the world data for the neutron. Lorenz et al. [82] use a dispersion relation approach to analyze the Mainz data.

C. Fits to cross sections and polarization measurements

1. Fit strategy

The experimentally determined cross sections are analyzed, performing a direct fit of the different models for the form factors.

For the new Mainz data, the fit minimizes the sum of the weighted deviations squared,

$$M^2 = \sum_i \left[ \frac{\int_{\Delta r_i} \frac{d^2 \sigma}{d^2 \Omega}}{\int_{\Delta r_i} \frac{d^2 \sigma}{d^2 \Omega \text{ std. dipole}}} \right] \frac{d\Omega}{d\Omega} \sum_i \left[ \frac{\sigma_i}{\sigma_{\text{std. dipole}}} - 1 \right]^2 \Delta r_i^2. \quad (48)$$
By dividing the measured count rate by the simulated count rate, we have extracted the cross sections from the measurement as ratios $r_j$ to the cross sections for the standard dipole. This is compared to the ratio of $(\frac{d\sigma}{d\Omega})_{\text{model}}$ and $(\frac{d\sigma}{d\Omega})_{\text{std. dipole}}$, the cross sections calculated with the fit model and with the standard dipole, respectively, individually integrated over the acceptance $A_i$ of run $i$. $\Delta r_j$ is the uncertainty of $r_j$ and $\Pi$, the product of normalization parameters to be discussed further below. As discussed, it is not possible to determine the normalization of the different measurements, i.e., runs, to much better than 2%, an order of magnitude larger than the statistical errors. However, the overall normalization can be determined from the measured data themselves due to the knowledge of the cross section at $Q^2 = 0$. Furthermore, the relative normalization of sets of runs, grouped by the relative luminosity determination method described above, are well constrained through the kinematical overlaps of the different sets and can be easily determined as free parameters in the fit. Overall, 31 normalization constants $n_j$ are used as free parameters in addition to the model parameters in the fit to a total of 1422 measurements from this experiment (not yet including the external world data). These parameters model the overall normalization difference between spectrometers and between different sets of runs. Each individual measurement by a spectrometer has an individual combination, i.e., a product $\Pi$, of a subset of these normalization constants. For example, to a measurement of spectrometer B might belong the product of the overall efficiency of spectrometer B compared to A, and a constant for the normalization of the set this measurement belongs to. All normalization parameters are used in different combinations for a large number of individual measurements.

$M^2$ can be identified with $\chi^2$ only when we compare with a known true theory curve and when the deviations from the true theory curve are Gaussian distributed with a true variance matching our estimated errors $\Pi, \Delta r_j$, which one cannot prove. For the problem at hand the true theoretical curve is not known and it has to be estimated by the best fit curve which is nonlinear in the fit parameters by construction. As is so often the case in experimental physics, the frequentist interpretation of $M^2$ is problematic and the identification $\chi^2 = M^2$ can be only approximate. It is, however, customary to use $\chi^2 \approx M^2_{\text{min}}$ or, better, the reduced $\chi^2$, $\chi^2_{\text{red}} = \chi^2/(N_{\text{data points}} - N_{\text{parameters}})$, as an approximate measure of the quality of the different fit models.

The external cross-section data are included in a similar way. In addition, for each data set, we add a term $[(1 - n_{\text{ext},j})/\Delta n_{\text{ext},j}]^2$ with the normalization fit parameter $n_{\text{ext},j}$ and the normalization uncertainty quoted in the corresponding paper, $\Delta n_{\text{ext},j}$. For the form-factor ratio results from experiments using polarization techniques, we use terms of the form $[(G_E/G_M)_{\text{model}}(Q^2_i) - R_i]/\Delta R_i]^2$, where $R_i$ is one of the externally measured ratios and $\Delta R_i$ the quoted error, with systematic and statistic errors added linearly.

### 2. Point-to-point errors beyond counting statistics

Besides the errors from counting statistics, different additional effects contribute to the point-to-point error of the cross sections. These include the dead-time estimation, the uncertainty of the current measurement for the 315-MeV data (see Sec. IV C 4), the uncertainty of the background estimation, and undetected slight variations of the detector and accelerator performance. At this level of precision these effects are hard to quantify, even with direct measurements. In order to estimate these effects we group the data by incident beam energy and by the spectrometer with which the data are measured. We then inspect for each group the distribution of the deviation of the data points from the fit divided by the error from the counting statistics. These distributions follow a bell-shape curve and it is therefore safe to assume that the point-to-point errors are, to a large extent, also normally distributed. We therefore scale the statistical errors of the different data groups by the width of the bell curve to account for the additional point-to-point uncertainty. This effectively assumes that the additional error contribution scales with the statistics of the individual measurement, which should be true for the dominant source, the background estimation, and also for the dead-time estimation. Iterating the fit with updated scaling factors leads to a meta-stable situation, in which the scaling factors oscillate between two solutions, without significant changes in the fit function. We suppress these oscillations by hand and find a set of factors which, when iterated, are stable but result in a $\chi^2$ slightly larger than 1. In order to achieve a $\chi^2_{\text{red}}$ of unity for the best models the statistical errors would need to be increased further by less than 7%. In view of the smallness of this change, we choose not to apply such a further modification. The factors are determined using the spline fit, which gives the best $\chi^2_{\text{red}}$ without any scaling, and the same set of factors is used for all models.

The scaling factors of the point-to-point errors lie in the range between 1.07 and 1.8 with exception of the 315-MeV data. For these the luminosity could only be determined by the less precise measurement of the beam current and target density, hence the errors had to be scaled by 1.7 to 2.3.

It has to be noted that an overall scaling of the errors by a common factor does not change the best fit. However, since the factors differ for each group, we observe that cross-section values calculated from the fits with and without these error renormalizations change at most by 0.3%. In any case, it is possible to rank the fits by the $\chi^2_{\text{red}}$ when the same set of fixed scaling factors are used for all fits. This ranking is almost independent of the chosen set of scaling factors—only widely varying scaling factors from data set to data set can change the rank of a fit. Such a large variation in the point-to-point uncertainty can be ruled out from the data.

### 3. Fits without external data

The spline and polynomial models allow for a varying number of parameters. For the determination of the optimal number, one encounters the basic fact that it is not possible to determine simultaneously which model describes the data and how statistically pure a data sample is. In the extreme case, a model goes through all data points, i.e., it interpolates the data. The choice of the number of parameters is therefore a trade-off: With too few parameters, the model cannot describe the gross behavior of the data and the deduced quantities cannot be trusted; on the other hand, a fit with too many parameters
The extended Gari-Krümpelmann model achieves a $\chi^2$ of 1759, which is only slightly better than the double dipole. This fit is rather unstable and it seems that there are ambiguities in the solutions. Since the calculation and convergence is slow due to the large number of logarithms and the numerical properties of the model, it was not possible to perform a full study of this model. Such a study would require varying the starting conditions and constraints. For a reliable fit of this model, it may be necessary to fix the 31 normalization parameters obtained with one of the flexible models beforehand. Subsequently, the data could be used for a fit with this model. This has not been pursued further in this work.

Figure 9 shows the measured cross section divided by the cross section calculated with the standard dipole and with the scaled statistical errors, compared to the different fits. The measured cross sections are normalized with the normalization parameters extracted with the spline fit. The precision is better than 0.4% (average) per data point. It is noted that the normalization parameters depend slightly on the model. Therefore, for a comparison of the data to a fitted model, the normalization found in the fit of that model should be used. However, the models that achieve a small $\chi^2$ yield very similar normalizations, so it is reasonable to present the data normalized to only the spline model which has the smallest $\chi^2_{\text{red}}$. For the flexible models the maximum difference in a normalization parameter is 0.26% and the average standard deviation is 0.073%. The largest difference for the “good” models occurs for the 855-MeV data, where the (less flexible) Friedrich-Walcher model shifts the data slightly upwards by 0.7% for the data measured with spectrometer C.

For the models that do not yield a $\chi^2 < 1600$, i.e., the double-dipole and the extended Gari-Krümpelmann model, the differences in the normalizations are larger (up to 1.6% in the case of the double-dipole fit). Both models would shift the cross sections down, therefore both fit curves are below the data with normalizations from the spline fit (see Fig. 9).

The analyses with the “good” models yield cross sections which differ by less than 1% for almost all of the $Q^2$ range of the data. In the high-$Q^2$ range the fits start to diverge. Above 0.55 GeV$^2$ only data from 720 and 855 MeV contribute. Therefore, the separation into $G_E$ and $G_M$ is not well determined. In the $Q^2$ region covered only by 855-MeV data, the allocation of the cross-section strength to the electric or magnetic part is undetermined, giving rise to the larger spread of the models.

### 4. Fits including the world cross-section and polarization data and possible two-photon-exchange effect

The addition of the world data extends the range in $Q^2$ considerably, with data points reaching above 30 GeV$^2$. However, their uncertainty and density vary widely. Fits with high-order polynomials are therefore problematic and spline fits with a constant knot spacing small enough to accommodate the low-$Q^2$ data are impossible. We therefore extended the spline × dipole model to nonconstant knot spacing and placed

<table>
<thead>
<tr>
<th>Model</th>
<th>$\chi^2$</th>
<th>Number of parameters</th>
<th>$\chi^2_{\text{red}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single dipole</td>
<td>3422</td>
<td>2 $\times$ 1 + 31</td>
<td>2.4635</td>
</tr>
<tr>
<td>Double dipole</td>
<td>1786</td>
<td>2 $\times$ 3 + 31</td>
<td>1.2893</td>
</tr>
<tr>
<td>Polynomial</td>
<td>1563</td>
<td>2 $\times$ 10 + 31</td>
<td>1.1399</td>
</tr>
<tr>
<td>Poly. + std. dipole</td>
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<td>2 $\times$ 10 + 31</td>
<td>1.1400</td>
</tr>
<tr>
<td>Poly. × std. dipole</td>
<td>1572</td>
<td>2 $\times$ 8 + 31</td>
<td>1.1436</td>
</tr>
<tr>
<td>Inv. poly.</td>
<td>1571</td>
<td>2 $\times$ 7 + 31</td>
<td>1.1406</td>
</tr>
<tr>
<td>Spline</td>
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<td>2 $\times$ 8 + 31</td>
<td>1.1385</td>
</tr>
<tr>
<td>Spline × std. dipole</td>
<td>1570</td>
<td>2 $\times$ 7 + 31</td>
<td>1.1403</td>
</tr>
<tr>
<td>Friedrich-Walcher</td>
<td>1598</td>
<td>2 $\times$ 7 + 31</td>
<td>1.1588</td>
</tr>
<tr>
<td>Ext. Gari-Krümpelmann</td>
<td>1759</td>
<td>14 + 31</td>
<td>1.2777</td>
</tr>
</tbody>
</table>
knots roughly according to the data point density at 0, 0.25, 0.5, 0.75, 1, 1.5, 3, 5, 10, and 40 GeV$^2$.

A fit including only external Rosenbluth data in addition to the new Mainz data results in $\chi^2_{\text{red}} = 2074.64/1810 = 1.146$, well comparable to the numbers above.

Including all available data, i.e., also polarization data on the form-factor ratio, raises this to $\chi^2_{\text{red}} = 2282.24/1868 = 1.222$, a rather large increase in $\chi^2$ of 207.6 for only 58 additional data points. This demonstrates that the difference between the Rosenbluth and polarization methods seen at higher $Q^2$ does not vanish with our floating normalization of the cross-section data.

The most likely explanation for this discrepancy is the effect of hard two-photon exchange which is believed to have a larger effect on the Rosenbluth separation than on the ratio determined from polarization measurements [11]. The data basis is not broad enough to disentangle the contributing amplitudes over the whole $Q^2$ range as has been done in Refs. [83,84] for a single $Q^2$ point. In fact, the experimental information is just enough to constrain rather simple models. Therefore, we assume a simple linear dependence on $\varepsilon$ which vanishes at $\varepsilon = 1$ and a logarithmic dependence on $Q^2$, similarly to Ref. [85], as an additional multiplicative term $(1 + \delta_{\text{TPE}})$ on top of Feshbach’s Coulomb correction Eq. (20),

$$\delta_{\text{TPE}} = -(1 - \varepsilon) a \ln(b Q^2 + 1),$$

where $a$ and $b$ are fit parameters. The global fit of Alberico et al. [7] uses a similar approach, with a two-parameter model introduced by Chen et al. [86]. In contrast to our approach, their model assumes a given $Q^2$ dependence but gives more freedom in the epsilon dependence. The fit including our TPE parametrization to all data gives $\chi^2_{\text{red}} = 2151.72/1866 = 1.153$ now as good as the “good” fits above.

We also performed fits of all data with a low-order Padé model,

$$G_{\text{Pade}}^{E,M}(Q^2) = \frac{1 + \sum_{i=1}^{3} a_i^{E,M} Q^{2i}}{1 + \sum_{j=1}^{5} b_j^{E,M} Q^{2j}},$$

i.e., the same parametrization as in Ref. [4]. This model has only 8 instead of 11 parameters per form factor and therefore has much lower flexibility and the achieved $\chi^2$s are considerably higher. Table V gives an overview over the achieved $\chi^2$s.

D. Statistical significance and model dependence

Our model for uncertainties divides up all sources into three parts: the first part is the global component, i.e., uncertainties which affect all points in a normalization group the same way. These are automatically corrected for by the floating normalization. The next component affects each measurement individually—we treat these via the scaling of the counting statistics errors described above and in the following subsection. The third, and most critical, component covers effects which change the cross sections in a systematic way. We describe this in Sec. V D 2. A given source of uncertainty does not necessarily fall into only one of these categories. For example, the detector efficiencies bulk contribution is a global effect, but we also treat this source in the third category. While
TABLE V. The $\chi^2_{\text{red}}$ for several fits to (1) all cross-section data, i.e., to the new data of this paper and previous cross-section world data; (2) all cross-section and polarization results without any TPE model; and (3) same as (2) with the simple TPE model. All fits were performed for the variable spline model and the Padé model.

<table>
<thead>
<tr>
<th>Data base</th>
<th>Model</th>
<th>$\chi^2_{\text{red}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cross sections</td>
<td>Spline with var. knots</td>
<td>$2074.64$ (\text{TM}^5)</td>
</tr>
<tr>
<td>c.s. + pol.</td>
<td>Spline with var. knots</td>
<td>$2282.76$ (\text{TM}^6)</td>
</tr>
<tr>
<td>c.s. + pol.</td>
<td>” + TPE</td>
<td>$2131.72$ (\text{TM}^6)</td>
</tr>
<tr>
<td>Cross sections</td>
<td>Padé</td>
<td>$2280.14$ (\text{TM}^5)</td>
</tr>
<tr>
<td>c.s. + pol.</td>
<td>Padé</td>
<td>$2465.51$ (\text{TM}^7)</td>
</tr>
<tr>
<td>c.s. + pol.</td>
<td>Padé + TPE</td>
<td>$2360.81$ (\text{TM}^7)</td>
</tr>
</tbody>
</table>

point-to-point efficiency differences are less probable, they are taken care of by the error scaling.

A source of systematic error is the selection of models used in the fit which needs special attention. We describe our approach in Sec. V D 3.

1. Point-to-point uncertainties

To express the uncertainty in the result due to uncertainties in the data and extraction procedure we must construct a confidence region around the best fit. However, the meaning of this region is a delicate point.

As described above the uncertainties from counting statistic are scaled to account for point-to-point uncertainties from other sources like the luminosity determination and detector efficiency fluctuations. Classically, such point-to-point uncertainties are treated by standard error propagation, which assumes a linear approximation. In order to circumvent this limitation, we calculate the confidence bands using a Monte Carlo technique: From the best fit, we generate a large number of pseudo data sets, with data varied according to the individual uncertainties. We additionally vary the normalization of the data groups with a ±5% uncertainty. Each of these data sets is now fitted, and from the resulting fit ensemble one can construct envelopes as the confidence bands to a selected confidence level.

For the interpretation of this band, it has to be noted that it is dependent on the model function. One uses an implicit prior assumption, namely that the true curve can actually be expressed by the model function. From this, it is clear that a less flexible model will have a smaller uncertainty; the assumption is then a stronger statement and therefore reduces the uncertainty. It is also clear that it is impossible to define a model-independent band: Without any prior, that is, allowing any function (or distribution), the uncertainty at positions off the exact points of the input data is infinite.

Per se, standard error propagation and the Monte Carlo method construct the pointwise confidence bands, that is, one expects the value derived from the experiment to be inside the confidence band around the true value at a given $Q^2$ with the specified confidence level without any limitation on the behavior at a different $Q^2$. In the linear approximation, this is commonly reformulated in the not-quite-precise inversion: One expects that the unknown true value is inside the confidence band around the experimentally determined value. In a nonlinear model, this inversion is even less accurate, a fact which cannot be cured without the introduction of further prior assumptions.

However, with our method, it is possible to overcome the other limitation, i.e., the reduction to single $Q^2$ points. We can construct simultaneous confidence bands, i.e., the statement is extended to express that, with the chosen confidence level, the true function is inside the band for a chosen $Q^2$ range and not just at a single point. These bands are therefore strictly wider than the point-to-point bands of the same confidence level. Assuming the same shape of the bands, the Monte Carlo technique allows us to find scaling factors: for the models at hand, a 68% simultaneous band is about 2.3 times wider than the pointwise band with the same confidence level. To achieve 95%, one has to scale the 68% pointwise bands by a factor of around 3.

In this paper, we present pointwise bands with a confidence level of 68%, the usual “one-σ errors.”

2. No point-to-point uncertainties

Besides statistical errors, one has to take into account uncertainties which affect several measurements in a systematic way. Most of these are irrelevant as they affect either all points or all points of a set in the same way. For example, an error in the target density or thickness will shift all points up or down. Due to the way the fits are constructed these shifts are subsumed in the fitted normalization constants. A ±5% uncertainty of this normalization scaling is included in the simulation and therefore also in the confidence bands as described above. What remains are slow drifts over time or scattering angle which may affect the outcome of the fits. We identified several experimental sources:

(i) Energy cut in the elastic tail. This error can be estimated by varying the cut-off energy. It changes the form-factor results by at most 0.2% for high $Q^2$ and by less than 0.1% for $Q^2 < 0.55$ GeV².

(ii) Drift of the normalization. This error might occur due to unaccounted dead-time effects in the detectors or electronics when the event rate changes. From the long-term experience with the detector setup, this error on the cross sections is estimated to be below 0.05%.

(iii) Efficiency change due to different positions of the elastic peak on the focal plane. The detector efficiency is position dependent because of different wire tension, missing wires, or quality of the scintillators. By adjusting the central momentum, the position of the electron trajectories in the focal plane was almost constant. This effect on the cross sections is estimated to be at most 0.05%.

(iv) The vertex-dependent acceptance correction for spectrometers A and C. A comparison of the 720-MeV data, measured with the long and short target cells, leads to a cross-section uncertainty below 0.1%.

(v) The influence of spectrometer C on the measurement with spectrometer B. We split this uncertainty in a part
which is effectively point to point, reflected by the error scaling, and a part which behaves systematically as a function of the angle. The latter is estimated to be below 0.1%.

(vi) The background estimation. Depending on the size of the background below the elastic hydrogen peak this error is estimated to be between 0.1% and 0.5%.

While the first point can be tested directly by fitting data with varied cut-off energy, the other uncertainties have to be treated by hand. To this end the cross sections are grouped by the energy and by the spectrometer with which they are measured. For each group, we define a linear function \( c(\theta) = a(\theta - \theta_{\text{min}}) \) interpolating from 0 for the smallest scattering angle to the full estimated uncertainty at the maximum angle of the group. The cross sections are then multiplied by \( 1 + c(\theta) \). The sign of \( a \) was kept constant for all energies. The so-modified cross sections were then refitted with the form-factor models. In order to determine an upper and a lower bound the fits were repeated with negated \( a \). The uncertainties found in this way are added quadratically to the uncertainties from the radiative tail cutoff. The choice of a linear function in \( \theta \) is certainly arbitrary, but we checked several different reasonable functional dependencies on \( \theta \) and \( Q^2 \), e.g., imitating the effect of a spectrometer angle offset or target position offset. They all produced similar results. The so-determined uncertainties are reflected by the experimental systematic confidence bands presented in this paper.

A possible source of uncertainty not from data but from theory are the radiative corrections. The absolute value of the radiative corrections should already be correct to better than 1% and a constant error in the correction will be absorbed in the normalization. Any slope introduced as a function of \( \theta \) or \( Q^2 \) by the radiation correction will be contained in the slope-uncertainty discussed above up to a negligible residual; it is therefore not considered.

In order to evaluate the influence of the applied Coulomb correction, the amplitude of the correction was varied by \( \pm 50\% \). The so-modified cross sections are refitted with the different models. The differences of the extracted form factors to the results for the data with the unmodified correction are shown as a band in Fig. 10.

Except for the phenomenological TPE model included in the fit to the full data set, we do not include any theoretical correction of the hard two-photon exchange to the cross sections in our analysis but apply Feshbach’s Coulomb correction. Published Rosenbluth data normally do not include a Coulomb correction. This has to be considered for comparisons of our fits with old Rosenbluth separations.

3. Model dependence

An important issue is the question of whether the form-factor functions are sufficiently flexible to be a suitable estimator for the unknown true curve or whether they introduce any bias, especially in the extraction of the radius. We have studied this problem in two ways.

First, we used a Monte Carlo technique similar to the method described in Sec. V D 1. We analyzed Monte Carlo data sets produced at the kinematics of the data of the present experiment with a series of published form factors: the standard dipole, the Padé and polynomial descriptions of...
models have a very small spread between themselves, at least in the region where a reliable disentanglement of the form factors is possible. The less flexible fits exhibit larger deviations, especially above 0.5 GeV$^2$. The Friedrich-Walcher parametrization has good agreement at lower $Q^2$, similarly to the flexible models, but exhibits the same bias as the other less flexible fits at higher $Q^2$.

For the fits which include the world data we use the mentioned variation of the spline model. The additional freedom of picking the location of the knots gives local flexibility of the model function. This variation provides an alternative handle on testing model dependencies since the flexibility of the model can be varied in an almost continuous way. For the purpose of this paper, we kept the number of knots constant and only varied the positions with a Monte Carlo approach. We select for each knot a random position between half the distance to the previous and half the distance to the next knot with a uniform distribution and refit. The distribution of $M^2$'s of this ensemble of models is presented in Fig. 12. It has to be stressed here that, besides the points we raised earlier, this is not a $\chi^2$–like distribution—it is not a distribution of $M^2$ fitting an ensemble of repeated experiments with the same model, but the distribution of $M^2$ fitting the same data with an ensemble of models. As can be seen from Fig. 12 the original choice of the knot positions (“nominal knots”) was already close to optimal. We construct a confidence band by taking the envelope for the 68% best models. The result is displayed in Figs. 13 and 14 together with the other contributions to the uncertainty.

VI. RESULTS FOR THE FORM FACTORS

A. Fits to the new Mainz data alone

In Fig. 10 we present the results for $G_E$ and $G_M$ and of their ratio for fits of the spline model to the new data without additional external data. In the same figure we show previous measurements and fits to old data. It has to be noted that the previous measurements are plotted as given in the original publication without the update to the radiative corrections described above. The error bars of the previous

<table>
<thead>
<tr>
<th>Fit model</th>
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<tbody>
<tr>
<td></td>
<td>Std. dip.</td>
</tr>
<tr>
<td>Single dipole</td>
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<tr>
<td>Double dipole</td>
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<td>Polynomial</td>
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<tr>
<td>Poly. + dipole</td>
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<tr>
<td>Poly. × dipole</td>
<td>1.000</td>
</tr>
<tr>
<td>Inv. poly.</td>
<td>1.000</td>
</tr>
<tr>
<td>Spline</td>
<td>1.000</td>
</tr>
<tr>
<td>Spline × dipole</td>
<td>1.000</td>
</tr>
<tr>
<td>Friedrich-Walcher</td>
<td>1.005</td>
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</table>

One way. For the purpose of this paper, we kept the number of knots constant and only varied the positions with a Monte Carlo approach. We select for each knot a random position between half the distance to the previous and half the distance to the next knot with a uniform distribution and refit. The distribution of $M^2$'s of this ensemble of models is presented in Fig. 12. It has to be stressed here that, besides the points we raised earlier, this is not a $\chi^2$–like distribution—it is not a distribution of $M^2$ fitting an ensemble of repeated experiments with the same model, but the distribution of $M^2$ fitting the same data with an ensemble of models. As can be seen from Fig. 12 the original choice of the knot positions (“nominal knots”) was already close to optimal. We construct a confidence band by taking the envelope for the 68% best models. The result is displayed in Figs. 13 and 14 together with the other contributions to the uncertainty.

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<td>Single dipole</td>
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<tr>
<td>Double dipole</td>
<td>1.002</td>
</tr>
<tr>
<td>Polynomial</td>
<td>1.000</td>
</tr>
<tr>
<td>Poly. + dipole</td>
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</tr>
<tr>
<td>Poly. × dipole</td>
<td>1.000</td>
</tr>
<tr>
<td>Inv. poly.</td>
<td>1.000</td>
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<tr>
<td>Spline</td>
<td>1.000</td>
</tr>
<tr>
<td>Spline × dipole</td>
<td>1.000</td>
</tr>
<tr>
<td>Friedrich-Walcher</td>
<td>1.005</td>
</tr>
</tbody>
</table>
The results for $G_E$ exhibit some peculiar structure for small $Q^2$, therefore we show this form factor also with an extended scale [Fig. 10 (a)]. First, $G_E$ exhibits a significant negative slope relative to the standard dipole at $Q^2 \approx 0$, giving rise to the larger electric radius. After the slope levels out around 0.1 GeV$^2$, there is an indication of a bump around 0.15 GeV$^2$, however, at the limit of significance. Further, the ratio to the standard dipole remains constant up to 0.55 GeV$^2$ when the slope again becomes larger. In that region, however, only measurements at large scattering angles for only two beam energies contribute so the fit becomes less reliable and more sensitive to systematic errors such as the neglect of TPE. For even higher $Q^2$ measurements have been taken only at one energy and a separation of $G_E$ and $G_M$ is not possible.

The magnetic form factor $G_M$ deviates from earlier measurements. We relate this to the normalization depending on the extrapolation with an assumed analytical form which—in previous analyses—does not include the wiggle first seen by this experiment. The specifics of the maximum and the minimum of the wiggle structure depend, of course, on the functional form one divides by—in our case, the standard dipole.

The structure at small $Q^2$ seen in both form factors corresponds to the scale of the pion of about $Q^2 \approx m_\pi^2 \approx 0.02$ GeV$^2$ and may be indicative of the influence of the pion cloud [1].

While the deviation of $G_M$ from previous measurements seems surprising at first glance, it reconciles the form-factor ratios from experiments with unpolarized electrons, like this one, with those found with polarized electrons, especially with the high-precision measurements in Refs. [16,54,55].

Due to the deviation of the results of $G_M$ from most of the previous determinations the geometric reliability of the spectrometer motion has been questioned by some experimenters after the publication of Ref. [8]. The verification of the rotational axes of the spectrometers in 2013 found them to be within the assumed limits and is far of from explaining the change in $G_M$ form previous measurements.

A possible general concern with fits is the question of convergence. In the time-like region, the form factors have poles which limit the convergence of a polynomial expansion

<table>
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<tbody>
<tr>
<td></td>
<td>Std. dip.</td>
</tr>
<tr>
<td>Single dipole</td>
<td>0 ± 0.3</td>
</tr>
<tr>
<td>Double dip.</td>
<td>0 ± 1</td>
</tr>
<tr>
<td>Polynomial</td>
<td>−1 ± 18</td>
</tr>
<tr>
<td>Poly. + dip.</td>
<td>0 ± 15</td>
</tr>
<tr>
<td>Poly. × dip.</td>
<td>−1 ± 14</td>
</tr>
<tr>
<td>Inv. poly.</td>
<td>0 ± 13</td>
</tr>
<tr>
<td>Spline</td>
<td>1 ± 7</td>
</tr>
<tr>
<td>Spline × dip.</td>
<td>0 ± 6</td>
</tr>
<tr>
<td>F.-W.</td>
<td>0 ± 2</td>
</tr>
</tbody>
</table>

FIG. 11. (Color) Relative deviation of the different models to the spline fit.
around 0 to $|Q^2| < 4m^2$. To test this, we modified the spline model; we add to the spline model a calculation of the nonanalytic terms [93]. In effect, the splines will then only have to fit the remaining, analytical part. The result is almost indistinguishable from the spline fit without this addition with a relative change of below $6 \times 10^{-4}$.

B. Form factors via Rosenbluth separation

The classical way of determining $G_E$ and $G_M$ is the Rosenbluth separation of cross sections measured at fixed $Q^2$ for different polarization $\varepsilon$. Rewriting the cross section [Eq. (7)] as

$$
\sigma_{\text{red}} = \varepsilon (1 + \tau) \left( \frac{d\sigma}{d\Omega} \right)_{0} / \left( \frac{d\sigma}{d\Omega} \right)_{\text{Mot}} = \left[ \varepsilon G_E^2(Q^2) + \tau G_M^2(Q^2) \right]
$$

(51)

makes it obvious that, for constant $Q^2$, the reduced cross section $\sigma_{\text{red}}$ depends linearly on $\varepsilon$ with $G_E^2(Q^2)$ as slope and $\tau G_M^2(Q^2)$ as ordinate intercept. Hence, a linear fit can be used to extract $G_E$ and $G_M$. We have discussed in Sec. I the advantages of extracting the form factors through a global fit to the cross sections. Nevertheless, we also perform a classic Rosenbluth separation of our data in order to reconcile our analysis with the expectation the community might have.

One of the problems with a direct Rosenbluth separation of measured cross sections is a coherent inclusion of a normalization of the data. We handle this by first fixing the cross section with the normalizations extracted by the spline fit.

Another problem is the necessity of several data points at constant $Q^2$ but varied $\varepsilon$. Due to the large number of measurements with overlapping acceptances, it is possible to find a set of 77 narrowly spaced $Q^2_i$ with measurements at least three different $\varepsilon$, so the linearity can be tested. Obviously, not all of the measured data are being used in this case—especially unfortunate is the loss of information on the lower end of $Q^2$: The lowest point is 4 times larger than what is available in the data. To project the cross section, which has been averaged over a finite-size $Q^2$ range given by the spectrometer acceptance, to the nearest $Q^2_i$ point, we divide by the numerically integrated result of the Monte Carlo simulation with the standard dipole for $G_E$ and $G_M$ and multiply by the differential cross section evaluated with the same form factors at the given $Q^2_i$ point.

This procedure implies an error which, to first order, depends on the difference of the curvature of the true cross
FIG. 14. As in Fig. 13 but for fits to the data of this work and the external data from unpolarized and polarized measurements under the inclusion of the TPE parametrization Eq. (49).

section and the one used in the Monte Carlo calculation, multiplied by the square of the acceptance [94], when the cross section is attributed to the central $Q^2_{\text{central}}$; this error is found to be negligible for our measurements. Attributing the resulting cross section to an off-central value, say $Q^2_i$, results in an additional error proportional to the difference in the slopes of the true and the reference cross sections and on the projection distance $Q^2_i - Q^2_{\text{central}}$. Our measurements are so narrowly spaced in $Q^2$ that this uncertainty is below 0.15% for the highest $Q^2_i$ presented here and considerably less for lower $Q^2_i$.

The Rosenbluth-separated form factors are shown in Fig. 15, together with the result of the global fit (spline model). For the lowest $Q^2$ points, $G_M/(\mu_p G_{\text{std dipole}})$ was not determined by the Rosenbluth fit but, for each point, set once to 1 and once to 1.05, as one would expect it to be in that range and not larger. For each point, the difference in $G_E$ of the two constraint fits and the errors of the individual fits give the error of $G_E$ shown in Fig. 15. The points from the unconstrained fits are presented in gray for reference. The use of the prior knowledge that the magnetic form factor cannot differ too much from the standard dipole for $Q^2 \sim 0$ helps to reduce the error bars on $G_E$ for low $Q^2$ considerably.

The agreement of the Rosenbluth-separated form factors with those from the global fit has been tested by calculating a reduced $\chi^2$ from the differences of the Rosenbluth data points to the spline fit. The rather large value of 2.2 is found, with similar numbers for a comparison of $G_E$ or $G_M$ alone. Fits of polynomials (order 10) to $G_E$ and $G_M$ from the Rosenbluth separation yield also $\chi^2_{\text{red}}$ values above 2. In order to put these numbers into perspective, one has to note that the $\chi^2_{\text{red}}$ distribution is much wider for the fit to the Rosenbluth-separated form factors, due to the lower number of degrees of freedom. In fact, interpreting the deviation of the flexible fits from the expectancy value 1 as purely statistical, $\chi^2_{\text{red}}$ values up to 1.7 for the fit to the Rosenbluth-separated form factors would have the same probability as $\chi^2_{\text{red}}$ values up to 1.14 for the global fits.

While the “ingredients” of the global fit and the Rosenbluth separation are in principle similar, the explicit Rosenbluth separation differs fundamentally from the global fit since it (a) has to contract the large acceptance of the measurements to single $Q^2$ points and (b) acts on subsamples of the complete...
data set. In this way a large part of the dependence on the primary kinematic variables, the scattering angle and the incident energy, disappears and the information is lost. This is exacerbated by the fact that the set of \( Q^2 \) values has to be the same for all energies. The consequences of these differences have not been studied fully, however, the robustness (see Sec. 8.7 of Ref. [95]), i.e., the insensitivity to unaccounted non-Gaussian errors of the input data, has been tested for both estimators, i.e., the form factors determined via the global fit and via the Rosenbluth separation. To this end, statistically pure pseudo data are generated from the spline fit and then perturbed with systematic errors. Unperturbed, both methods result in an average \( \chi^2 \) of 1. Adding systematic shifts, however, increased the \( \chi^2 \) for the Rosenbluth method more than for the global fit method.

In fact, if we shift 5% of the data points by a systematic “not normal distributed” shift of 0.5%, the fits yield a difference in the \( \chi^2 \) increase comparable to the difference seen for the measured data. We therefore conclude that the global fit is a much more robust estimator of the form factors with respect to non-normal errors in the measured cross sections.

As to other systematic insufficiencies in the measurement, the Rosenbluth separation is also prone to errors in theoretical corrections of the cross sections. Such insufficiencies might show up as deviations of the reduced cross section from Eq. (51), i.e., from a straight line in the Rosenbluth plot. Figure 16 shows the relative deviations of the measured \( \sigma_{\text{red}} \) from the Rosenbluth fit for 3 of the 77 \( Q^2 \) values. At the (quite high) level of statistical accuracy of this experiment no systematic deviations from the straight line are visible. This was tested further for all 77 points by fitting polynomials of second order, where the coefficient of the quadratic term was found to be compatible with zero.

C. Fits including external data

1. Unpolarized elastic scattering only

For global fits with the addition of external data we have to adapt the models to be usable with the drastic change in data point density over \( Q^2 \) as already described. We use a flexible spline \( \times \) dipole model with variable knots described above and the less flexible Padé model given by Eq. (50). Figure 13 shows both form factors and the form-factor ratio from both models.

The overall behavior of the Padé model is quite similar to the spline model up to 5 GeV\(^2\), without following the small-scale wiggles of the spline fit.

Differences in the gross behavior appear in \( G_E \) for \( Q^2 \geq 5 \) GeV\(^2\), where the existing data start to determine \( G_E \) badly. Here the Padé model does not exhibit the downward bend, however, without leaving the model confidence band of the spline model. For \( G_M \), there is a distinct knee between 1 and 2 GeV\(^2\) for the spline model, which arises slightly washed out but clearly visible also in the Padé model. The high-\( Q^2 \) behavior of both models is closer together in trend in \( G_M \) than in \( G_E \); however, the confidence band of the spline fit does not overlap the Padé fit over the complete range.

The large model-dependency estimate for larger \( Q^2 \) illustrates a point which is often underestimated: The standard error propagation used for the construction of the confidence bands gives an estimate of the statistical error for the chosen model only.

2. Unpolarized and polarized elastic scattering

Adding additional information in the form of form-factor ratio data from polarized experiments can help to reduce the uncertainty in the form-factor separation, especially affecting the uncertainty of \( G_E \) at large \( Q^2 \). As discussed in Sec. V C 4, we need to introduce additional parameters for a TPE parametrization given by Eq. (49) to reconcile the two measurement methods. Figure 14 shows the results of these fits, again for both models. While \( G_E/G_{\text{std.dip}} \) now decreases more or less linearly, \( G_M \) is shifted upward for \( Q^2 > 1 \) GeV\(^2\). As a result, the ratio also decreases almost linearly. This behavior is similar for both the spline as well as the Padé model. In spite of the added parameters, the widths of the confidence bands are reduced.

Figure 17 shows the contribution from our TPE parametrization as a function of \( Q^2 \) at \( \varepsilon = 0 \) where the
correction is maximal. This contribution is similar for both models. Table IX lists the fit result for the parameters \(a\) and \(b\).

It is clear that the TPE model used is simple and only weakly motivated by theoretical considerations. However, it gives much better fits for the wide \(Q^2\) region in which the ratio \(G_E / G_M\) derived without accounting for TPE from the unpolarized scattering and from the polarized measurements differs significantly. While the total \(\chi^2\) increases by 207 when adding 58 polarization data points without any TPE parametrization taken into account, the increase is reduced to 77 by the inclusion of only two free parameters (for the detailed numbers, cf. Table V).

Figure 18 shows the TPE correction as a function of \(\varepsilon\) for selected \(Q^2\) values in comparison with the calculation of Borisyuk et al. [96] and the calculation by Blunden et al. [97–99]. Both calculations are valid in the low-\(Q^2\) region. Our simple parametrization can evidently not reproduce the strong curvature of these particular calculations in the low-\(\varepsilon\) region. Most of the data, however, are situated in the midrange of \(\varepsilon\), where the calculations are almost linear and our parametrization gives a similar slope but a different overall normalization of about 0.5%, growing slightly with \(Q^2\). Such a correction linear in \(\varepsilon\) cannot be tested for with the Rosenbluth separation as it is indistinguishable from a change in \(G_E\). The change of the overall normalization is mostly absorbed in our floating normalization of the fits.

An indication for such a normalization effect can be seen by a comparison of the normalization factors of the external data found in the fits: In Fig. 19 the shift of the cross-section normalization \(n_{\text{ext},j} - 1\) of the external data is displayed. Figure 19(a) shows the result when only the external unpolarized cross-section data are added to the data set used for the fit. As can be seen, all shifts are positive, i.e., the actual cross sections as reconstructed by the fit are larger than the values quoted in the original publication. The spread of the shifts is quite large. However, a fit to just the previous data sets without the new Mainz data shows a similar spread of almost the same shifts for the low-\(Q^2\) data sets and shifts smaller by about 4% absolute for the large-\(Q^2\) data. While it may look strange that all shifts are positive, the mean of the normalization falls together with the shift of the oldest measurement [57], for which the absolute normalization was certainly not better than a few percent, and the other older measurements may have checked their normalization with regard to Ref. [57].

Figure 19(b) shows the normalization when, in addition, the polarization data and the TPE parametrization are taken into account. Figure 19(c) shows the difference in the normalization introduced by this. While there is virtually no change in the normalization with respect to the analysis without the polarization data for the data sets at \(Q^2\) below 0.2 GeV\(^2\), we find a shift of about 1% for the large-\(Q^2\) data sets, similarly to the spread of the curves in Fig. 18.

Figure 20 shows the low-\(Q^2\) region of the spline fit. We find similar features as in the fits to the Mainz data alone which did not include a TPE correction: Both the change in slope of \(G_E\) and the bump-dip structure in \(G_M\) are visible.

The figure also shows a fit of the data without our empirical TPE-correction model but with the Feshbach correction replaced by a TPE calculation by Blunden et al. [97,98], who provided us with a numeric evaluation [99]. Since this calculation is valid only up to around 5 (GeV/c)\(^2\), we fit data only up to this value. The correction reduces the disagreement between unpolarized and polarized data: the \(\chi^2\) reduces from 2232 for a fit with the Feshbach correction to 2142 with the Blunden correction, equal to an increase of about 1.85 for each data point from polarization experiments. Our full fit achieves 2107, i.e., 1.22 for each data point from polarization. Fits up to 3 (GeV/c)\(^2\) produce similar results.

The fit for \(G_E\) with the Blunden TPE correction lies higher at around \(Q^2 = 0.2\) (GeV/c)\(^2\), making the change in slope even more pronounced. On the other hand, the bump in \(G_M\) is

---

**FIG. 17.** The TPE contribution to the cross section (excluding the Feshbach term) as determined by the fit to the complete world data set as a function of \(Q^2\) for \(\varepsilon = 0\). Same nomenclature as in Fig. 13.

**FIG. 18.** Comparison of the complete TPE correction \(\delta_{\text{TPE}}\) (without the soft part included in Maximon-Tjon) for four different \(Q^2\) values. Solid: Fit to data; dashed: calculation with the approximation of Ref. [96]; dotted: TPE corrections from Refs. [97–99].

**FIG. 19.** The TPE contribution to the cross section (excluding the Feshbach term) as determined by the fit to the complete world data set as a function of \(Q^2\) for \(\varepsilon = 0\). Same nomenclature as in Fig. 13.

**TABLE IX.** The values of \(a\) and \(b\) of the TPE parametrization, extracted from the fits to all data.

<table>
<thead>
<tr>
<th>Model</th>
<th>(a)</th>
<th>(b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spline with var. knots</td>
<td>0.069</td>
<td>0.394 GeV(^{-2})</td>
</tr>
<tr>
<td>Padé</td>
<td>0.104</td>
<td>0.188 GeV(^{-2})</td>
</tr>
</tbody>
</table>
much reduced. However, the fit still indicates that the following dip compared to the dipole does not start at $Q^2 = 0$, like previous fits extracted, but at slightly higher $Q^2$ values.

FIG. 19. Shifts of the normalization $n_{ext,j} - 1$ of the external cross-section data found by the spline fits. (a) Fits to the data of this work and external data from unpolarized scattering. (b) Data from polarized measurements are additionally included and the fit is extended with the TPE parametrization. (c) The difference of the normalization shift $[b] - [a]$. The numbers in the legend indicate the normalization uncertainty we assumed in the fit.

FIG. 20. The spline fit with variable knots from Fig. 14, zoomed in to $Q^2 < 1$ (GeV/c)^2. The dashed-dotted line represents a fit of all data up to 5(GeV/c)^2, with TPE corrections according to Refs. [97–99] applied.

VII. ELECTRIC AND MAGNETIC RADII

A. Extraction method and model dependence

According to Eq. (9), the electric and magnetic rms radii, $r_E = \sqrt{\langle r^2_E \rangle}$ and $r_M = \sqrt{\langle r^2_M \rangle}$, are given by the slopes of the corresponding form factors at $Q^2 = 0$. Therefore the accuracy with which they are determined by the measurement is given by the accuracy of the data in particular at low $Q^2$. Since the accuracy of $G_E$ is high at low $Q^2$, $r_E$ will be determined with good precision, while $G_M$ and therefore $r_M$ is less well determined due to its small contribution to the cross section at low $Q^2$. In any case, the determination of $r_E$ and $r_M$ corresponds to an extrapolation of $G_E$ and $G_M$ to $Q^2 = 0$, and one has to ask the question to which extent the result depends on the ansatz for the fit model, in particular on its flexibility which depends on functional form and number of parameters, $N_p$. For too low flexibility, the resulting radii are
not trustworthy since the data are not correctly reproduced. If the flexibility is too high, the fit can follow the smallest statistical fluctuations, which results in erratic determinations of the radii. In fact, one has to compromise between these two extremes, as we did for the choice of the number of parameters \( N_p \) by looking at the development of \( \chi^2 \) as a function of \( N_p \).

Here, we also inspect the resulting values of the radii and search for the range of \( N_p \) in which the extracted radii are stable.

For both the electric and the magnetic radius the polynomial and the polynomial + dipole model produce a stable result for \( N_p > 9 \). The polynomial × dipole model works comparably well for the electric radius for \( N_p > 8 \) but shows erratic behavior for the magnetic radius already for \( N_p > 9 \). The inverse polynomial, which has a quicker convergence to the \( \chi^2 \) plateau, also deteriorates faster into such erratic behavior for \( r_M \). Nevertheless, these models agree quite well for both radii when one confines oneself to \( N_p \) at the beginning of the plateau.

The erratic behavior of the magnetic radius stems from the less stringent determination of the magnetic form factor at low \( Q^2 \), where the magnetic contribution is small and where, with enough flexibility (large \( N_p \)), the fit follows smallest statistical deviations, resulting in larger uncertainties. Low \( N_p \) give the fit enough stability to extrapolate \( G_M \) from higher \( Q^2 \) values, where the magnetic contribution is sizable, down to \( Q^2 = 0 \), but may not be flexible enough to capture the true behavior. It has to be noted that in previous determinations of \( r_M \) only models with much less flexibility have been used and that the data had been taken at \( Q^2 > 0.0053 \text{ GeV}^2 \) only and they had significantly larger errors.

The spline fits based on polynomials of third order tend to give a smaller electric radius than the rest of the models; they additionally exhibit a depression in the value of the radius of about 0.015 fm, that is, about 2.5 times the statistical uncertainty, for \( N_p \) around 10. This difference between the result from the splines and the polynomials was further investigated, but no conclusive cause was found. The spline model’s tendency to underpredict, albeit less pronounced, was already seen in the model dependency analysis described in Sec. V D 3. The curvature of the spline models is limited by the order of the base polynomial. To test for a possible bias, we also use splines based on polynomials of fourth and fifth order. They produce progressively larger radii.

Focusing on the \( \chi^2 \) of points below \( Q^2 = 0.06 \text{ GeV}^2 \) (543 data points) the spline fits yield a \( \chi^2 \) around 581 while the rest of the models give around 576. While this might indicate a worse fit of the low-\( Q^2 \) region by the spline models, the \( \Delta \chi^2 \) of 5 is small compared to the 1σ width of the \( \chi^2 \) distribution \( \sqrt{\langle \chi^2 \rangle_{\text{dof}} \approx 543} \approx 33 \).

In order to estimate the model dependency for the extracted radii, the radii are determined with all models described before and for some variation in \( N_p \). The results are shown in Fig. 21.

The results for the charge radius fall somewhat apart into two groups according to the model of the analysis, namely those determined with the spline-based models and those from the polynomial-based models. For each group, the electric and magnetic radius and their statistical and systematic error have been determined as the weighted average over the results from the single fits, where the weights are the linear sum of statistical and systematical error. The model error has been calculated from the weighted variance of the values.

The final result is the (unweighted) arithmetic average of the values of the two groups. An additional error (labeled “group”), accounting for the difference of the two groups, is attributed to the result. Since it cannot be assumed that this error is normal distributed, it is taken as half of the difference of the two groups.

For most of the results below we applied the radiative corrections described above, that is, with the Feshbach Coulomb correction. In order to get a feeling for the effect of different corrections, we repeated the fits with a TPE calculation using the approximation of Ref. [96], yielding the values already published in Ref. [19] and also using the calculation from Refs. [97–99].

### B. Electric radius

As the average of the flexible models, we obtain

\[
r_E = 0.879(5)_{\text{stat}}(4)_{\text{syst}}(2)_{\text{model}}(4)_{\text{group}}.
\]

This value is in complete agreement with the CODATA06 [13] value of 0.8768(69) fm based mostly on atomic measurements. It is also in complete accord with the old Mainz result [60] when the Coulomb corrections [37,76] are applied. However, the results from recent Lamb shift measurements on muonic hydrogen [12,14] are 0.04 fm smaller, i.e., 5 standard deviations from our result (quadratically added errors) and almost 8 from the updated CODATA value from 2010 [100] which combines our data and earlier scattering and atomic level measurements. Since this difference is unexplained yet, despite a multitude of efforts, we looked whether other ways of analysis of the scattering data would yield different results. The Friedrich-Walcher model gives a slightly larger, but fully compatible, radius.

For small \( Q^2 \), the contribution of the magnetic form factor to the cross section is so small that one can adopt one parametrization for \( G_M \); subtract the magnetic contribution from the cross section, and then fit the resulting \( G_E \) at low \( Q^2 \) only using a simple model like a low-order polynomial. This technique is similar to the method employed by Simon et al. [60], where \( G_M \) was set to \( \mu_p G_E \) (scaling relation). In the present work, we apply it to the 180-MeV data alone, using different parametrizations, different cut-off values in \( Q^2 \), and different \( G_M \) models. The normalization was left floating, but the fit recovered the normalization given by the global fit on the 0.1% level. This approach yields radii between 0.870 and 0.895 fm, with most values close to 0.880 fm, thus in excellent agreement with the final result of the global fit.

The two different TPE calculations we applied change the result by significantly less than the statistical uncertainty alone. In Ref. [101], Arrington compares several recent calculations and finds all of them in good agreement for low-\( Q^2 \) and high \( \varepsilon \). A full evaluation of the effect of all available calculations on the fit and the extracted radii is beyond the scope of this paper.

The inclusion of external data changes the value marginally for the spline model with variable knot positions. The Padé
model is not sufficiently flexible to achieve a comparably small $\chi^2_{\text{red}}$. Hence, its extracted radius is unreliable and we refrain from quoting its result.

Table X summarizes the electric radii determined with the different approaches together with the final result Eq. (52). Despite all these efforts, we do not see a way to reconcile our result with those from muonic hydrogen. We do not expect that a future calculation of TPE corrections can reconcile our result with the muonic measurement completely, but we cannot rule out that such calculations may reduce the discrepancy. We want to note, however, that a large shift in the radius from TPE would in turn create tension with atomic measurements with electric hydrogen, albeit probably with less significance.

C. Magnetic radius

Table XI gives an overview of the results for the magnetic radius of the proton. The statistical and systematic uncertainties are larger, since the radius is determined as an extrapolation for $Q^2 \to 0$ where the cross section is less sensitive to magnetic scattering. Interestingly, the difference between splines and polynomials is much smaller than for

![Graph showing electric and magnetic rms radii as extracted with different models. Gray: Statistical error; black: Linearly added systematic error.](image-url)

**FIG. 21.** The electric and magnetic rms radii as extracted with the different models. Gray: Statistical error; black: Linearly added systematic error.

Table X. Results for the electric radius.

<table>
<thead>
<tr>
<th>Method</th>
<th>Electric radius $r_E$ in fm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spline models (1)</td>
<td>0.875(5)<em>{\text{stat}} (4)</em>{\text{sys}} (2)_{\text{model}}</td>
</tr>
<tr>
<td>Polynomial models (2)</td>
<td>0.883(5)<em>{\text{stat}} (3)</em>{\text{sys}} (2)_{\text{model}}</td>
</tr>
<tr>
<td>Friedrich-Walcher (a)</td>
<td>0.884(7)<em>{\text{stat}} (5)</em>{\text{sys}} (4)_{\text{model}}</td>
</tr>
<tr>
<td>Spline with variable knots</td>
<td></td>
</tr>
<tr>
<td>+ external data</td>
<td></td>
</tr>
<tr>
<td>+ Rosenbluth data</td>
<td>0.878</td>
</tr>
<tr>
<td>+ all external data</td>
<td>0.878</td>
</tr>
<tr>
<td>Average of (1) and (2)</td>
<td>0.879(5)<em>{\text{stat}} (4)</em>{\text{sys}} (2)<em>{\text{model}} (4)</em>{\text{group}}</td>
</tr>
<tr>
<td>With TPE from Ref. [96]</td>
<td>0.876(5)<em>{\text{stat}} (4)</em>{\text{sys}} (2)<em>{\text{model}} (5)</em>{\text{group}}</td>
</tr>
<tr>
<td>With TPE from Refs. [97–99]</td>
<td>0.875(5)<em>{\text{stat}} (4)</em>{\text{sys}} (2)<em>{\text{model}} (5)</em>{\text{group}}</td>
</tr>
</tbody>
</table>

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The hyperfine splitting in electric hydrogen represents an alternative method to determine $r_M$. The radius $r_M$ enters in the hyperfine splitting via the Zemach radius $r_Z$. For the extraction, one has to make an ansatz for the electric and magnetic form-factor shape. The corresponding analysis in Ref. [15] was performed with the standard dipole with $r_E = 0.8750(68)$ fm and $r_M$ was left as a fit parameter. The measured value is $r_Z = 1.045(16)$ fm and the variation resulted in $r_M = 0.778(29)$ fm in complete agreement with the value we obtain. However, Carlson et al. [107] elaborated an update of this analysis by including better polarization corrections, i.e., TPE effects, and more recent form-factor parametrizations. These parametrizations yield $r_Z = 1.069$ fm [6], $1.091$ fm [108], and $1.089$ fm [4]. It is somewhat model dependent to convert these $r_Z$ to $r_M$, but one gets as an indication $r_M = 0.82(1)$ fm, $0.86(1)$ fm, and $0.86(1)$ fm for the $r_{ZS}$ above, respectively. Their analysis did not yet take our new data into account and, hence, has to be taken with the poor knowledge of $G_{Mq}$ at small $Q^2$ from the previous measurements and, consequently, also of $r_Z$ in mind.

The most recent determination of the magnetic radius stems from the laser spectroscopy of muonic hydrogen [14]. From the measurement of two transition frequencies the Zemach radius $r_{Z} = 1.082(37)$ fm has been extracted and the charge radius has been reevaluated as $r_{E} = 0.84087(39)$ fm. Using the analytical ansatz with the standard dipole the authors determine $r_{M} = 0.87(6)$ fm and claim consistency with electron scattering. However, assuming the dipole parametrization they used for the extraction, these values predict a form-factor ratio at odds with that of polarization experiments (e.g., Ref. [16]). One would need to shift $r_{M}$ down to resolve this discrepancy, and then be at odds with the scattering experiments they compared to, or assume a different form-factor shape, which could in turn invalidate the extraction. Electron scattering provides more information than just the radii, and only if simultaneous agreement in additional moments of the charge and magnetization distribution is reached, one can claim consistency.

The Zemach radius derived from the form factors presented in this paper is $r_{Z} = 1.045(4)$ fm [109], i.e., within the error margin of the laser spectroscopy of muonic hydrogen and identical to the value from normal hydrogen. Putting this Zemach radius into the calculation of Carlson et al., one gets $r_{M} = 0.777(10)$ fm, in perfect agreement with the result from the elastic electron scattering of this work.

### VIII. CONCLUSIONS

This paper presents the details of the highly precise measurement of elastic electron scattering off the proton performed at MAMI, the first results of which have already been published in Ref. [8]. The analysis differs in some respects from the customary approach as follows:

<table>
<thead>
<tr>
<th>Method</th>
<th>Magnetic radius $r_M$ in fm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spline with variable knots + external data:</td>
<td>0.775(12)$<em>{lat}(9)$</em>{syst}(4)$_{model}$</td>
</tr>
<tr>
<td>+ Rosenbluth data</td>
<td>0.777(13)$<em>{lat}(9)$</em>{syst}(5)$<em>{model(2)}$</em>{group}$</td>
</tr>
<tr>
<td>+ all external data</td>
<td>0.778(14)$<em>{lat}(10)$</em>{syst}(6)$_{model}$</td>
</tr>
</tbody>
</table>

The values for $G_E$ of Ref. [63] indicate a $\sim 1\%$ deviation in the normalization of the form factors—the extrapolation to $Q^2$ aims at 0.99. Applying the same shift to the $G_M$ data of that paper, one might even recognize the wiggle in this old data set.

The electric form factor. This gives some confidence that the wiggle in $G_M$ at small $Q^2$ (see Fig. 10) is not an artifact of the fitted models. The only experimental reason for this wiggle, which we could think of and not rule out directly, would be a systematic error with the angular rotation of the spectrometers around the target. From the precision of the setup of the spectrometer turntable [9], which we have rechecked, and the checks with the overlapping angular settings of the spectrometer described above, in particular the measurements on the left and right side, we can exclude such an explanation.

On the other hand, the change is relatively large when applying the two different TPE corrections. This may be due to the somewhat unexpectedly large deviations of the TPE calculations from a linear behavior at small $\epsilon$, i.e., larger scattering angles. The Coulomb calculation from Refs. [18,102] does not show this behavior at small $\epsilon$ and it has been argued in Ref. [101] that calculations beyond second Born might be needed, reflecting the uncertainty in current TPE calculations. The phenomenological determination of TPE effects in this work may serve as a help for a better theoretical description.

Future results from direct measurements of the TPE effect are expected from several experiments [103–105] for large $Q^2$ and from Ref. [106] for low $Q^2$. They will help resolve this issue.

We observe that larger parameter numbers tend to produce larger magnetic radii. We do not believe that this stems from insufficient flexibility for the lower $N_p$, as a plateau in $\chi^2_{red}$ is clearly already reached for the smaller parameter numbers. However, more flexible fits are more susceptible to follow statistical fluctuations. These possibly less reliable fits have larger uncertainties than fits with smaller parameter numbers. Hence, with the chosen weighted averaging, the impact of these fits is lessened. However, this choice and the choice of the parameter number are somewhat subjective. For comparison, an unweighted average would enlarge the radius by 0.008 fm.

Previous determinations from elastic electron scattering give a significantly larger magnetic radius (see Ref. [16] and references therein). However, since available data for $Q^2 < 0.2$ GeV$^2$ had large error bars and could not resolve the structure the new data indicates, the validity of the extrapolation of these previous determinations is questionable.
The Rosenbluth separation and classic error propagation are given up as they are unnecessary steps limiting the precision and the amount of information extracted from the data. Instead, we perform a direct fit of sufficiently flexible models for the form factors to the whole body of measurements, avoiding the unnecessary limitation for the Rosenbluth separation and the badly controlled correlation in the resulting form factors. However, we also show the consistency of the two methods. We extracted sensible confidence bands without any approximation to a linear behavior using Monte Carlo techniques and performed an extensive study of model dependency.

The calibration problem present in any determination of absolute cross sections has been overcome by fitting the normalization of sets of cross sections of which the relative measurement-to-measurement normalization was well under control by an explicit luminosity measurement using an extra spectrometer. The absolute normalization was fixed by the well-known form-factor values at \( Q^2 = 0 \). This procedure only needs a weak assumption on the smoothness of the form factors for the \( Q^2 = 0 \) limit. Our data has the furthest reach toward lowest \( Q^2 \) to date, minimizing the impact of this assumption. The use of a spectrometer as luminosity monitor to fix the relative normalization between individual measurements and the large overlap between our data sets makes a precise determination of the absolute normalization for all measurements possible.

We have only applied the standard radiative corrections but not the hitherto debated two-photon-exchange contributions. However, an empirical form has been derived from the inconsistency of the \( G_E \) and \( G_M \) data, extracted from measurements with polarized and unpolarized electrons, respectively, which may be interpreted by radiative corrections as TPE or other physics (see also Ref. [19]).

The new method has been also applied to the world data set together with the data of this paper. The analysis represents a coherent summary of the present knowledge of the form factor of the proton.

From the slopes of the form factors for \( Q^2 \to 0 \) we determined the electric and magnetic radii of the proton. The values extracted here from the whole body of electron-scattering data are at variance with those determined recently with very high precision from muonic hydrogen. In spite of a multitude of efforts, there is no generally accepted explanation yet.

The discrepancy for the magnetic radius determined with the different methods is somewhat less dramatic. While its determination from the hyperfine splitting in electric hydrogen requires the knowledge of the Zemach radius (which, in turn, needs the information from the electric and magnetic form factors), the determination from electron scattering is hampered by the limited sensitivity of the cross section to magnetic scattering at low \( Q^2 \). In contrast to some (re)analyses of the hyperfine splitting and to a new measurement on muonic hydrogen, we determine a magnetic radius for the proton which is substantially smaller than the electric radius. While this result fits well to the direct measurement of the ratio \( G_E/G_M \) using polarized degrees of freedom at quite small \( Q^2 \), the muonic result does not.

The larger slope (with respect to the dipole) observed in \( G_E \) gives rise to a larger charge radius, compatible with older extractions. While the data clearly exhibit this feature, the conflict with the muonic Lamb-shift measurements certainly warrants further study. The wiggles in \( G_M \) are at the limit of significance and further measurements are needed for an independent verification. If the results of this paper would be confirmed and if these structures would survive other efforts for an explanation like the application of TPE, they would hint at the existence of effective degrees of freedom which may be a yardstick feature to be replicated by theory, for example, in lattice calculations (we refer to Refs. [110–112] for current results).

ACKNOWLEDGMENTS

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[99] P. G. Blunden (private communication).