The stopping power of hydrogen in hafnium and the importance of relativistic 4f electrons

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The stopping power of protons through Hf foil has been studied both experimentally and theoretically. The measurements were performed at the Laboratory of Accelerators and X-Ray Diffraction in Lisbon by using the transmission method on self-supporting stopping material. The overall uncertainty of around 5% was established over the protons energy range (0.6-2.5) MeV. The theoretical developments involved fully relativistic atomic structure calculations for Hf, which required the solution of the Dirac equation. The shell-wise local plasma approximation (SLPA) was used to describe the energy transferred to the bound 1s-4f electrons, and the outer four electrons were considered as a free electron gas (FEG). We found the relativistic description of the 4f-shell and the screening between 4f and 5p electrons to be decisive around the stopping maximum. Present theoretical and experimental results are in excellent agreement in the energy region of the new measurements. However, our theoretical stopping cross sections show substantial differences with the most used semi-empirical models (SRIM2013 and ICRU-49) at intermediate to low energies. Our calculations suggest the stopping maximum to be higher and shifted to lower energies than these previous predictions. Future measurements below 100 keV would be necessary for a better understanding of the stopping power around the maximum and below.

I. INTRODUCTION

For impact energies above a few keV/amu, mono-27 energetic charged particles penetrating a foil of any ma-28 terial lose their energy through a series of consecutive in-29 elastic collisions, mainly with target electrons [1, 2]. The ³⁰ information given by the energy loss process is essential 31 not only to have a better comprehension of the physics 32 behind the fundamental interactions but also because it 33 plays a vital role in many applied fields such as materials 34 science, nuclear physics, ionic implantation, and radio-35 therapy [2, 3]. Experimental data on ion mean energy 36 loss per unit path S(E) is of crucial relevance to check ₃₇ the reliability of semi-empirical models and to determine 38 some key parameters [4–6]. The experimental data avail- 39 able is often rather scarce, which is troublesome when the $_{\rm 40}$ material under study corresponds to an element of low $_{41}$ occurrence on the Earth's upper crust, such as hafnium. 42 So far, only one experimental work has been pub- $_{43}$ lished regarding the stopping power cross section of pure 44 hafnium for protons [7], while more attention has been 45 recently given to studies involving hafnium oxide due to 46 its practical use [8–11]. It is well known that signifi-47 cant attention has been paid in recent years to transition $_{48}$ metal-oxides such as HfO₂ because of their potential as

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alternative gate dielectrics to replace SiO_2 for the future generation of nano-electronics with less than 45 nm gate length [12, 13]. Some important physical properties of the above mentioned metal-oxide films depend on their thickness, which is often measured by using Rutherford Backscattering Spectrometry [14, 15]. This method relies heavily on the determination of the stopping power of ion beams in the material of interest.

In this study, we report experimental stopping power cross sections over the incident energy range (0.6-2.5)MeV for protons crossing self-supported Hf thin-film by using the transmission method. We aim not only to upgrade stopping power data compilations [16, 17] but also to provide useful information about the processes governing the slowing down of protons in multi-electronic targets. In the rare earth metals, the 4f electrons play an essential role in the stopping power since they belong to the first shell of bound electrons below the conduction band. As already noted [18], the free electron gas (FEG) shows unexpected behavior in these elements, which casts doubts on its proper description. In the case of Hf, we found the contribution of the 4f-shell to be decisive even at impact energies around the stopping maximum, as will be shown later.

The theoretical approach implemented in this work uses the shell-wise local plasma approximation (SLPA) [19] to describe the energy transferred to the bound 1s-4f electrons and two different models for the

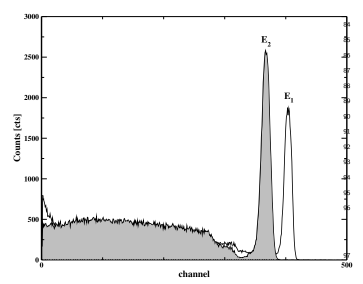


FIG. 1. RBS spectrum for $E_{\text{avg}} = 921.1$ keV protons on $_{99}$ hafnium sample which is subsequently used to determine the $_{100}$ energy loss in the foil.

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FEG; in low energy region, the screened potential with cusp condition model (SPCC) [20], which is non-linear binary formalism, and the Mermin-Lindhard dielectric formalism (ML) [21] for energies around the stopping maximum and above. Our model requires the relativistic wave functions and binding energies of Hf and considers four electrons per atom in the FEG [22]. Hafnium is particularly interesting since the filled 4f-subshell (with 14 electrons) is the main contributor below the FEG, causing the stopping cross sections to be very sensitive to a good description of this shell. The screening among the 4f and 5p electrons has been considered and found to 111 play a significant role within the SLPA calculations.

The experimental details and data are given in Sec-¹¹³ tion II, while the theoretical method is explained in¹¹⁴ Section III. Present theoretical and experimental val-¹¹⁵ ues are finally compared to the only experimental values¹¹⁶ measured by Sirotinin *et al.* [7] with the backscattering¹¹⁷ method, the theoretical results by Grande and Schiwi-¹¹⁸ etz [23, 24], and by Sigmund and Schinner [25], and also¹¹⁹ with the semi-empirical values from the SRIM-2013 pack-¹²⁰ age [26] and the ICRU-49 tabulation [27]. Conclusions¹²¹ and discussions are given in Section V. All the present¹²² data can be found at the Zenodo platform [28].

II. EXPERIMENTAL ARRANGEMENTS

A. Accelerator and scattering Chamber

The procedure used in this work to obtain stop-¹²⁹ ping power data is essentially the same as described in Ref. [29]. The present measurements were made at₁₃₀ the IST/LATR (Laboratory of Accelerators and X-Ray₁₃₁ Diffraction) in Lisbon. This facility uses a 2.5 MV Van₁₃₂

de Graaff accelerator to deliver $^1\mathrm{H}^+$ primary ion beams through a series of electrostatic lenses and collimators onto a thin $\mathrm{Au/SiO_2}$ sample, which is used as a scattering center. This sample was placed in the center of an RBS/C scattering chamber, where a high vacuum (pressure of $\sim 10^{-6}$ Torr) was maintained during the measurements. The beam current on the sample was kept at around 5.0 nA to attain sufficient statistics in each particle spectrum. By using a beam spot of about 1.0 mm in diameter, a solid angle of 11.4 msr was attained. The overall energy resolution (FWHM) of the detection system was about 15 keV relative to 5.486 MeV alpha particles from a $^{241}\mathrm{Am}$ source.

B. Target

The stopping material under analysis was a hafnium foil with a nominal thickness of 1.0 μ m and 99.95% purity, which was supplied by Lebow Company [30]. However, a more precise thickness value was achieved by measuring the energy loss of alpha particles coming from a calibrated (239 Pu, 241 Am, 244 Cm) source. From the alpha spectra with and without the Hf foil interposed, the characteristic energy shift δ E was measured and then combined with the stopping power for 5.486 MeV alphas on hafnium ($55.69 \text{ eV}/10^{15} \text{ at/cm}^2$) found in Ref. [26] to obtain an areal density of $(4.13\pm0.21)\times10^{19} \text{ at/cm}^2$, which corresponds to a thickness of $0.920\pm0.046\mu\text{m}$.

C. Energy loss measurement

Once the beam impinges on the Au/SiO₂ sample, protons are backscattered towards a Si surface barrier detector located at 140° relative to the initial beam direction. Fig. 1 shows two particle spectra, where the ion energies E_2 and E_1 are associated with a placed and removed hafnium sample, respectively. Both energy distributions were fitted by Gaussian functions to obtain the mean energy and width (FWHM) of the peaks [31], and from the difference between these two peak positions in the spectrum, the total energy loss $\Delta E = (E_1 - E_2)$ in the foil was calculated. As established in previous studies [5, 29], the experimental stopping power cross sections $\varepsilon(E)$ are determined at some mean energy E_{avg} by measuring the ion energy losses ΔE within the investigated Hf foil, which has a mean thickness denoted by Δx . In this way, only when the energy loss fraction $\Delta E/E_{\rm avg}$ across the Hf foil is not exceeding 20%, it is possible to define the stopping cross section by [32, 33]:

$$\varepsilon(E) = \frac{S(E)}{N} = -\frac{dE}{N dx} \approx -\frac{\Delta E}{N \Delta x},$$
 (1)

where N denotes the atomic number density (atoms cm⁻³) of the material under study. When this condition was not fulfilled, a small correction to the mean energies

 E_{avg} was applied in order to account for the non-linear dependence on ion energy of stopping powers [34, 35]. 189

III. THEORETICAL METHOD

The energy loss of ions in metal targets responds to ¹⁹⁴ different physical mechanisms, depending on the impact ¹⁹⁵ ion velocity. At low velocities, the binary collisions are ¹⁹⁶ responsible for the loss of energy by the ion. The main ¹⁹⁷ contribution is the ionization of electrons of the metal ¹⁹⁸ conduction band, which is well approximated by a free ¹⁹⁹ electron gas (FEG) of Fermi velocity v_F . Above a partic-²⁰⁰ ular velocity value (i.e., $v \ge 1.5 v_F$), not only binary but²⁰¹ also collective excitations (plasmons) occur [20]. More-²⁰² over, at high energies, also the bound electrons contribute²⁰³ to the stopping power. The method used in this work²⁰⁴ combines a FEG description for the interaction with the²⁰⁵ valence (or conduction) electrons and a different one for²⁰⁶ the interaction with the bound electrons.

We used the SPCC model [20] to describe the stop-208 ping power of low velocity charged particles in the FEG.²⁰⁹ It is a non-perturbative binary collisional approximation,²¹⁰ thus valid at energies below that of plasmon excitations.²¹¹ The SPCC [20] is based on a screened central potential²¹² with cusp condition of the electronic density close to the213 projectile. This model proved to give a good descrip-214 tion of the induced electron density even for negative215 projectiles [20] and reproduces the low velocity proton-216 antiproton differences in the stopping power (Barkas ef-217 fect). The SPCC formalism only depends on the Wigner-218 Seitz radio, r_S , which is a measure of the electronic den-219 sity of the FEG. For metals of well-known r_S , the SPCC₂₂₀ describes the low energy experimental stopping data cor-221 rectly [20], agreeing with the DFT results by Echenique₂₂₂ and coworkers [36, 37] at v = 0.

Hafnium (Z=72, [Xe] $4f^{14} 6s^2 5d^1_{3/2} 5d^1_{5/2}$) belongs to₂₂₄ the first groups of transition metals, with four electrons²²⁵ as FEG ($r_S=2.14$ a.u.) and 1s-4f electrons bound.²²⁶ We compared the computed r_S with the experimental²²⁷ value obtained from the measured energy loss function²²⁸ by Lynch et al. [38]. The experimental plasmon energy²²⁹ of Hf is $\hbar\omega_P\approx 15.8$ eV, with a width at half maximum²³⁰ $\delta\approx 4.4$ eV, and $r_S\approx 2.07$ a.u. [38]. The difference of less²³¹ than 5% between theoretical and experimental r_S assess²³² Hf as a canonical target [20].

Above certain impact velocity, the plasmon contribu-234 tion is essential (i.e., around and above the stopping-235 maximum). A value of interest for our analysis is the-236 minimum impact velocity to excite plasmons, v_P . In-237 the dielectric formalism, this value can be obtained as-238 $v_P \approx v_F [1 + (3\pi v_F)^{-1/2}]$ [39]. To describe the energy-239 loss considering collective and binary excitation, we re-240 sort to the ML dielectric formalism [21], which is a linear-241 response, perturbative approximation, so it depends on-242 the square of the ion charge. In this formalism, the re-243 sponse of target electrons to the ion passage is described-244 through the quantum dielectric function, which depends-245

on the characteristic r_S and δ parameters of the FEG.

For the stopping power due to bound electrons, the SLPA [19, 20] is employed. It is worth mentioning that the only inputs for the SLPA are the space-dependent densities of each shell in the ground state, and their binding energies. Collective processes and screening among electrons are included. Since hafnium is a relativistic target, the wave functions and binding energies must be obtained by solving the many-electron Dirac Hamiltonian. Details of these calculations and a table of binding energies have been published in Ref. [22], while Slater-type orbital expansions are given in Ref. [40].

To assess the importance of a fully relativistic description of bound electrons, Fig. 2 (a) shows our binding energies, $E_{nl\pm}$, with $\pm = j \pm 1/2$; non-relativistic values [41]; and experimental data on solid-state Hf [42], which is available only for 1s to $4f_{\pm}$ subshells, as expected. We notice that not only the most inner shells require relativistic calculations, but also the outer 5p and 4f shells. Furthermore, this figure shows very clearly the disability of non-relativistic calculations to describe the experimental data, which surprisingly worsens from the inner to the outer shells.

From the comparison with the experimental values in Fig. 2 (a), it can be noted that the sign of the binding energy deviations is inverted for the outer 5s and 4f electrons, with the experimental binding energies being less bounded than our theoretical ones. Small differences for the outer shells are expected since the experimental values correspond to hafnium in solid-state, while our theoretical calculations correspond to the element in the gas phase.

More detail of the theoretical binding energies is given in Fig. 2 (b), where relative errors with respect to the experimental values are shown. This figure shows clearly that the relativistic corrections are critical to describing the atomic structure of hafnium, even for the outer shells. It turns out that the errors committed in the non-relativistic calculations of the inner shell orbitals propagate, through the Hartree-Fock approximation, to the outer shells. The importance of fully relativistic calculations for the outer shells has already been noted for Au, Pb, Bi, and W [43].

For the contribution of bound electrons to the total stopping cross sections, the SLPA considers independent contributions of each subshell. Our relativistic binding energies present spin-orbit split. However, in total stopping power, where the initial state of the excited electron is not measured, the quantum uncertainty in energy ΔE melts this split. The criterion $\Delta E \Delta t \geq \hbar/2$ merges the energies $E_{nl+} - E_{nl-}$ for sufficiently small values of Δt (the collisional mean-time). In fact, at sufficiently high impact velocity, we can expect all target electrons to respond together to the ion passage [44, 45]. Following previous works [43], the collisional time is estimated as $\Delta t \approx \langle r_i \rangle/v$, with $\langle r_i \rangle$ and v being the orbital mean radio and impact velocity, respectively. In the case of hafnium, we found that for every sub-shell of electrons,

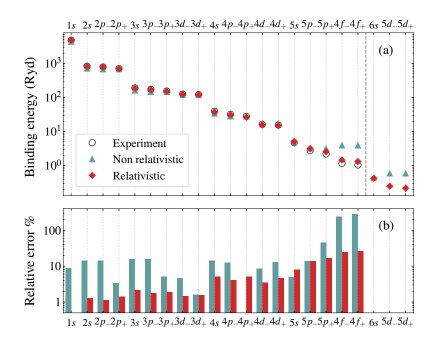


FIG. 2. (a) Binding energies of Hf. Present relativistic and available non-relativistic [41] values are given with filled symbols. Experimental measurements for solids [42] are depicted with open circles. (b) Corresponding relative errors respect to experimental data.

the spin-orbit split is unresolved in the energy region this 277 sub-shell contributes. Therefore, the nl-electrons should₂₇₈ be considered together, responding to the ion passage₂₇₉ as a single gas of electrons with density $\delta_{nl}(r)$ and mean₂₈₀ binding energy E_{nl} . This feature is vital within the SLPA₂₈₁ calculations because it accounts for the screening among₂₈₂ electrons of the same binding energy. For example, the 283 $4f_{-}$ and $4f_{+}$ of Hf can only be resolved for impact en-284 ergies E < 0.05 keV, but the contribution of 4f to the 285 total stopping is negligible for E < 40 keV. Moreover, 286 the 5p and 4f electrons of Hf are very close in energy₂₈₇ $(\Delta E_{5p-4f} \approx 1 \text{ a.u. } [22])$, and they react together at im-288 pact energies E > 40 keV (inter-shell screening). As al-289 ready mentioned, at higher energies, inter-shell screening290 is possible for other subshells (i.e., 4p-4d for impact ener-291 gies above 0.9 MeV), but its weight in the total stopping²⁹² is minor for deeper shells.

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Finally, in all our calculations [20], we assumed the projectile to be proton and not neutral hydrogen. When²⁹³ an ion moves inside a metal, the FEG screens the nucleus,²⁹⁴ so the binding energies will be smaller than outside the metal, and this effect is more critical at low impact veloc-²⁹⁵ ities v. In the case of hydrogen, the difference is drastic,²⁹⁶ i.e., for H inside Hf (rs=2.07), the 1s-bound state is al-²⁹⁷ most null at v < 2 [39]. It is worth to mention that this²⁹⁸ assumption agrees with Ziegler SRIM code [26] but dif-²⁹⁹ fers from CasP code [23], that predicts neutral hydrogen³⁰⁰ at very low velocities.

In Fig. 3, we display the present theoretical stopping₃₀₂ cross section of Hf for protons using the relativistic wave₃₀₃ functions and binding energies, but with and without the₃₀₄

5p-4f screening. We show the FEG and bound electron contributions separately and the total stopping as the addition of both of them. The minimum energy for plasmon excitation was estimated at approximately 37 keV. We used the non-perturbative SPCC model for impact energies E < 37 keV, and the perturbative ML calculation above this value. Bound 1s-4f electrons (relativistic wave functions and binding energies) are calculated with the SLPA and shown separately in Fig. 3 with and without the 4f-5p screening. Below ~ 40 keV, the difference between both calculations is negligible. Considering 5p-4felectrons as a single group of 20 electrons with screening among them gives lower stopping values than the addition of the separate 5p and 4f contributions. Notice that this shell correction can be considered naturally within a many-electron model, such as the SLPA.

IV. ANALYSIS OF THE RESULTS AND DISCUSSION

The present data are displayed in Table I. As can be observed, an overall relative uncertainty of around 5% was achieved for the experimental stopping power values, which are mainly due to the uncertainty in the hafnium foil thickness.

Fig. 4 synthesizes the results of the present work. The agreement between the present theoretical results and the new measurements displayed in Table I is excellent. Present measurements using the transmission method are in good agreement with the previous data by Sirotinin [7],

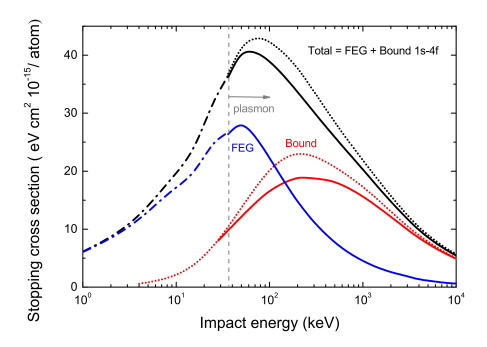


FIG. 3. (colour online) Theoretical stopping cross sections of protons in hafnium. Blue dash-dotted-line, the non-perturbative SPCC for the FEG; blue solid-line, ML results for the FEG (includes plasmon excitation); red solid and dotted-lines, the SLPA results for bound electrons with and without 5p-4f screening, respectively. Black curves, total stopping adding the FEG and bound 1s-4f contributions: dash-dotted-line, SPCC (FEG) + SLPA (bound); solid-line, ML (FEG) + SLPA (bound) with 4f-5p screening; dotted-line, ML (FEG) + SLPA (bound) without 4f-5p screening. The vertical grey dashed-line indicates the energy of 37 keV above which plasmon excitation is possible.

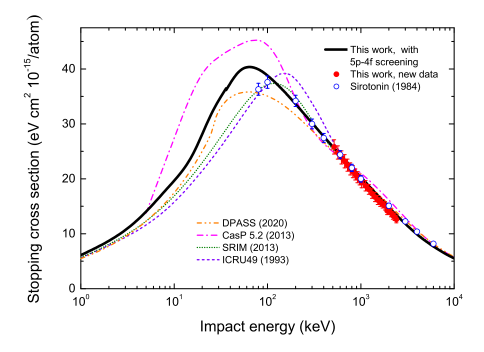


FIG. 4. (colour online) Stopping power cross section of hafnium for protons. Symbols: solid circles, present values; open circles, previous data [7]. Curves: Black solid-line, present full theoretical results with 4f-5p screening; pink dash-dot line, theoretical CasP5.2 [23, 24] values; orange dash-double-dot line, theoretical DPASS [25] results; green dotted-line, semi-empirical SRIM-2013 [26]; violet dashed-line, ICRU49 [27] tabulated values.

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E_{avg} keV	$S_{\rm exp}$ $eV/(10^{15} \ at/cm^2)$	$\Delta \mathrm{E}/\mathrm{E}$	E_{avg} keV	$S_{\rm exp}$ ${\rm eV}/(10^{15}~{\rm at/cm^2})$	$\Delta \mathrm{E}/\mathrm{E}$	E_{avg} keV	$S_{\rm exp}$ $eV/(10^{15} \text{ at/cm}^2)$	$\Delta E/E$
516.6	25.8 ± 1.3	20.5	1170.3	18.25 ± 0.91	6.4	1813.4	15.10 ± 0.76	3.4
567.8	24.8 ± 1.2	17.9	1220.0	18.08 ± 0.90	6.1	1862.7	14.79 ± 0.74	3.3
618.8	23.9 ± 1.2	15.8	1269.6	17.57 ± 0.88	5.7	1912.0	14.21 ± 0.71	3.0
669.6	23.2 ± 1.2	14.2	1319.2	17.32 ± 0.87	5.4	1961.2	14.46 ± 0.72	3.0
720.1	22.5 ± 1.1	12.8	1368.8	17.15 ± 0.86	5.1	2010.4	14.34 ± 0.72	2.9
770.5	21.8 ± 1.1	11.6	1418.3	16.69 ± 0.83	4.8	2059.6	13.76 ± 0.69	2.7
820.8	21.3 ± 1.1	10.7	1467.8	16.43 ± 0.82	4.6	2108.8	13.78 ± 0.69	2.7
871.0	20.8 ± 1.0	9.8	1517.2	16.13 ± 0.81	4.4	2158.0	13.70 ± 0.69	2.6
921.1	20.3 ± 1.0	9.1	1566.7	16.04 ± 0.80	4.2	2206.5	13.33 ± 0.67	2.5
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 15.77 ± 0.79

 15.51 ± 0.78

 15.46 ± 0.77

 14.93 ± 0.75

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TABLE I. Stopping power values S_{exp} of hafnium for protons measured in this work. $\Delta E/E$ values are also shown.

tinin [7], except for the lowest energy measurement at 340 80 keV. We have also included in this figure the theoretical curves from the CasP5.2 code by Grande and $_{_{341}}$ Schiwietz [23, 24] and from the DPASS code by Sigmund and Schinner [25], both available online. Furthermore, 343 we incorporated the semi-empirical results from SRIM- $_{344}$ 2013 [26] and the ICRU49 tables [27]. Interestingly, our 345 full theoretical curve differs from SRIM-2013 for impact $_{346}^{349}$ energies below 100 keV. We obtain a stopping maximum $^{347}_{347}$ of approximately 40×10^{-15} eV cm²/atom at 65 keV. Instead, following the up-to-now only set of data [7], SRIM₃₄₉ 2013 suggests a lower stopping maximum at an impact 350 energy of 115 keV. The stopping maximum is a susceptible region for any full theoretical description without parameters, and this is quite visible in a linear-scale plot $_{_{353}}$ like Fig. 4. However, the impact energy for the maximum seems to agree between our curve and DPASS, although 355 it is 10% below. Instead, CasP a maximum value of the $^{-356}$ stopping power 10% above our value and a completely $_{357}$ different shape at lower energies. It is worth mentioning 358 that our model gives similar results using the experimental value $r_S = 2.07$ a.u. rather than the theoretical one 359 $r_S=2.14$ a.u., with the stopping maximum at the same 360 impact energy but 4% higher. 362

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1764.1

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 19.03 ± 0.95

 18.73 ± 0.94

Future experiments for impact energies below 100 keV^{365} would be essential to complete this study on the stopping³⁶⁶ of Hf for protons. In the energy region 80-500 keV, only³⁶⁷ the measurements by Sirotinin's group [7] from 1984 are³⁶⁸ available. SRIM-2013 includes these values in the code³⁶⁹ fitting, so it does not represent a test for this data. Un-³⁷⁰ doubtedly, new measurements are needed in three spe-³⁷¹ cific energy regions: below 500 keV, around the stopping³⁷² maximum (i.e., 50-200 keV), and at low impact energies.³⁷³

V. CONCLUSION

 13.27 ± 0.66

 13.07 ± 0.65

 12.91 ± 0.65

 12.61 ± 0.63

2256.4

2305.5

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In this work, we have used the transmission method to experimentally determine stopping power cross section values for (0.6-2.5) MeV protons incident on selfsupporting Hf foils with an overall uncertainty of around 5%. Additionally, we calculated values extracted from the theoretical framework that involved the relativistic wave functions and binding energies of Hf and considered four electrons per atom in the free electron gas. The shell-wise local plasma approximation was employed to describe the energy transferred to the bound 1s-4f electrons, and two different models for the FEG: the screened potential with cusp condition (SPCC model) for energies below that of the plasmon excitation, and the Mermin-Lindhard dielectric formalism, for energies around the stopping maximum and above. Present theoretical stopping cross sections cover an extensive energy range from 1 keV/amu to 10 MeV/amu.

At high impact energies, the new stopping measurements are in good agreement with our theoretical results, with previous experimental data and semi-empirical values by SRIM-2013 and ICRU-49. However, we call the attention that around the stopping maximum and at lower impact energies, the difference between our full-theoretical results and SRIM is substantial. We compare our theoretical results with two other models given by the DPASS and CasP5.2 codes. Differences can be noted at intermediate to low impact energies, but they also support a stopping maximum at lower energy than SRIM predictions.

To the best of our knowledge, these are the first theoretical calculations of stopping in Hf to take into account relativistic effects in the atomic structure and screening among electrons in a consistent way, from very low to

high impact energies. Future experiments for impact en-378 ergies around the stopping maximum and in the low energy region would be essential to have a better under-379 standing of the stopping of protons in hafnium.

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