Evolution and ion kinetics of a XUV-induced nanoplasma in ammonia clusters

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Abstract

High-intensity extreme ultraviolet (XUV) pulses from a free-electron laser can be used to create a nanoplasma in clusters. In reference Michiels et al (2020 Phys. Chem. Chem. Phys. 22 7828–34) we investigated the formation of excited states in an XUV-induced nanoplasma in ammonia clusters. In the present article we expand our previous study with a detailed analysis of the nanoplasma evolution and ion kinetics. We use a time-delayed UV laser as probe to ionize excited states of H and H2+ in the XUV-induced plasma. Employing covariance mapping techniques, we show that the correlated emission of protons plays an important role in the plasma dynamics. The time-dependent kinetic energy of the ions created by the probe laser is measured, revealing the charge neutralization of the cluster happens on a sub-picosecond timescale. Furthermore, we observe ro-vibrationally excited molecular hydrogen ions H2++ being ejected from the clusters. We rationalize our data through a qualitative model of a finite-size non-thermal plasma.

Keywords: ammonia clusters, nanoplasma, femtosecond pump-probe, photoion spectroscopy, photoelectron spectroscopy, covariance mapping

(Some figures may appear in colour only in the online journal)

1. Introduction

Laser-induced nanoplasmas have been an active field of research in recent years and combine high energy physics [1] with atomic and molecular quantum dynamics on a nanoscale [2–4]. Research has been fueled by the necessity to understand radiation damage and plasma formation in the single-shot imaging of nanoparticles [5]. Many intriguing physical processes have been discovered: for example, nanoplasmas have been investigated as sources of high-energy particles [6–9] and coherent radiation [10, 11]. Nanoplasma research in rare-gas clusters led to the discovery of enhanced absorption by collective quasi-particle resonance of the electrons [12]. Many interesting properties of nanoplasmas come from the enhancement of recombination processes due to the large number of confined electrons and positive ions [13, 14]. The decay of a nanoplasma is governed by a complex interplay of Coulomb explosion and hydrodynamic forces leading to shock shells
molecular clusters, when compared to atomic C in the outer Debye layer [15–18]. Single-shot x-ray diffraction imaging of the nanoplasma evolution in rare-gas clusters revealed sub-picosecond dynamics [19], and a core–shell structure [20]. Nanoplasmas in clusters can be induced using long wavelength radiation (infrared (IR), or near infrared (NIR)) on one hand, or, on the other hand, short-wavelength extreme ultraviolet (XUV) radiation. The two regimes can be distinguished using the Keldysh parameter $\gamma \propto 1/\lambda$, where $\lambda$ is the wavelength of the radiation [21–23]. A Keldysh parameter of $\gamma \ll 1$ denotes the strong-field regime where photoionization proceeds via tunneling processes [24]. On the other hand, the regime where $\gamma \gg 1$ is called the weak-field regime. Photoionization in the weak-field regime proceeds via single- or multi-photon interactions [24]. This applies to XUV-induced nanoplasmas [25], where as a first approximation, electrons are created with kinetic energies $E_{\text{ele}} = h\nu - \text{IP}$. Here, $h\nu$ is the energy of a single XUV photon and IP is the first ionization potential of the species.

Concerning plasma dynamics, molecular clusters with multiple components differ significantly from homogeneous atomic clusters. When hydrogen is among the constituents, the nuclear dynamics speeds up and the lightweight protons offer an efficient pathway for cooling and charge neutralization [26–28]. Calculations predict the inner charge state and temperature in the plasma core to be much lower in (CH$_4$)$_n$ molecular clusters, when compared to atomic C$_n$ clusters [28]. Nanoplasmas in molecular clusters have been studied previously with tabletop [29–32] as well as free-electron lasers [7, 33], showing significant fragmentation of the molecules and the generation of high-energy ions. Previous pump-probe experiments with nanosecond lasers were unable to resolve the fast plasma dynamics happening on a sub picosecond timescale. In a recent femtosecond XUV-pump UV-probe time-resolved experiment on ammonia clusters, we investigated the dynamics of molecular and atomic states upon nanoplasma formation [33].

In the present work, we extend these studies with a detailed analysis of the nanoplasma evolution and ion kinetics. First, we will address the kinetic energy of H$^+$ emitted from the nanoplasma and use covariance analysis to compare how energy is dissipated from the clusters via high-kinetic-energy protons. Secondly, the photoionization of H$^+$ and the photo-dissociation of H$_2^+$ by the probe laser are discussed. Finally, we analyze time-dependent kinetic energy distributions of H$^+$ and photoelectrons upon UV-ionization of H$^+$($n = 2$). We discuss how the observations allow conclusions concerning the lifetime of the Coulomb potential at the cluster surface.

### 2. Experimental setup and methods

The experiment was performed at the low density matter (LDM) endstation [34] at the seeded FEL FERMI in Trieste, Italy [35]. Details on the experimental setup can be found in reference [33] and reference [34]. A jet of neutral ammonia clusters was created via supersonic expansion using a home-built pulsed nozzle. The mean cluster size was $\langle N \rangle = 2000$ molecules and the cluster size distribution is assumed to be a broad log-normal distribution [36]. The cluster jet was crossed perpendicularly with the XUV laser and a 261 nm UV laser. The interaction region was in the focus of a combined velocity-map-imaging (VMI) and Wiley–McLaren [37] type ion time-of-flight (ToF) spectrometer. High intensity XUV pulses were used to multiply ionize the ammonia clusters. The FEL pulse intensity in the interaction region was $I_{\text{XUV}} \approx 3 \times 10^{12}$ W cm$^{-2}$; at 14.3 eV photon energy, $I_{\text{XUV}} \approx 1 \times 10^{13}$ W cm$^{-2}$; at 19.2 eV, 23.8 eV, 28.6 eV and 33.4 eV photon energy and $I_{\text{XUV}} \approx 2 \times 10^{13}$ W cm$^{-2}$; at 42.9 eV photon energy. The UV probe pulse had an intensity of $I_{\text{UV}} \approx 2 \times 10^{12}$ W cm$^{-2}$. Correlation maps were calculated as Pearson’s correlation coefficient [38, 39]. We included partial covariances [40] to compensate for the target density fluctuations and FEL pulse energy fluctuations using the sum intensity of the ion-ToF sig-
nal per FEL shot as a control variable. The correlation map was calculated using 30,000 FEL shots. The shot-to-shot standard deviation of the FEL pulse energy, as well as the standard deviation of the total ion-ToF sum was $\approx 6\%$. We confirm the statistical significance of the correlations with a null hypothesis test using the $p$-value. The $p$-value shows the probability of finding an equal or stronger correlation if the correlation were in fact zero.

In the measured ion-ToF spectrum, the high-kinetic-energy protons create distinct forward and backward peaks corresponding to protons arriving earlier and later than those initially at rest. This process is illustrated in figure 1(a). The flight trajectories of three different protons originating from a highly charged cluster are sketched. The protons created with initial momentum in the forward and backward direction of the spectrometer create separate peaks in the ion-ToF spectrum. Ions with initial velocity components perpendicular to the extraction direction may not be detected. Using ion trajectory simulations and the known geometry of the ToF spectrometer, the initial kinetic energy of the ions can be deduced from their arrival time. An example of the forward and backward peak can be seen in the ToF distribution simulated for protons with 19 eV kinetic energy, which is shown as the black curve in the figure 1(b).

The method used to calculate kinetic energies from the ion-ToF spectra was adapted from reference [41]. We carried out ion-ToF trajectory simulations for discrete integer steps in proton kinetic energy ($E_{\text{h}^+} = 0, 1, 2, 3, \ldots \text{ eV}$) using Simion®. The experimental spectra were fitted to a linear combination of the simulated spectra using least-squares fitting. An example ToF spectrum (red) and the resulting fit (blue) are shown in figure 1(b). To account for the detector resolution, the simulated ion flight times were convoluted with a Gaussian function. The detector resolution was determined experimentally to be 18 ns full width at half maximum (FWHM). From the fit parameters, we calculated the mean kinetic energy of the protons. Because the detection efficiency for protons depends on the kinetic energy, we take it into account when calculating the mean kinetic energy. The detection efficiency was obtained from the Simion simulations by calculating the portion of protons arriving at the detector. Only about 10% of isotropically emitted protons with 10 eV kinetic energy are detected.

### 3. Results and discussion

#### 3.1. Correlated emission of protons from the nanoplasma

A comparison of experimental ToF spectra for protons created with high-intensity XUV pulses (red) and only the UV pulse (blue) is shown in figure 2(a). The red curve is broader than the blue curve, and the shoulders indicate a large contribution from high-kinetic-energy protons. In our experiment, ions are regarded as having a high kinetic energy if they are created with kinetic energies $\geq 4 \text{ eV}$. For both curves, the maximum of the intensity is at the center of the peak, which corresponds to the ToF for protons created with negligible kinetic energy. Taking the reduced detection probability for high-kinetic-energy protons into account, we can deduce that the red curve in the upper left half of the map, we see a region of strong positive correlations centered at $[t_1, t_2] = [1460 \text{ ns}, 1520 \text{ ns}]$, the same flight time as observed for the protons with excess kinetic energy in the top panel. This shows a positive correlation between high-kinetic-energy protons emitted in the forward and backward direction. The positive correlation

![Figure 2](image-url)
Figure 3. (a) Proton signal as a function of FEL intensity for three different proton kinetic energies ($E_{k\nu}$). (b) Proton intensity as a function of kinetic energy (blue curve) and the power exponent $k(E_{k\nu})$ (red curve) indicating the pump-power dependence of the proton intensity for different kinetic energies. Data was taken using a pump laser with 28.6 eV photon energy.

arises for protons with equal kinetic energy and extends from $E_{\text{hi}}^+ \geq 4$ eV to $E_{\text{hi}}^+ \approx 20$ eV. The single-pixel p-value of this correlation feature is $p \leq 1 \times 10^{-4}$. Thus, the correlation is significant even for kinetic energies where the collection efficiency is $< 10\%$. Additionally, there are spots of negative correlation centered at $[t_1, t_2] = [1520 \text{ ns}, 1560 \text{ ns}]$ and $[t_1, t_2] = [1440 \text{ ns}, 1470 \text{ ns}]$. These peaks show an anti-correlation of high-kinetic-energy protons with low-kinetic-energy protons. In the center of the image, the correlation is substantially lower than for the main peaks, converging to zero until overwhelmed by the auto-correlation.

Correlated emission of protons is only possible if there is a common source. To interpret the data, we will look at it in an event-based picture. The correlations observed in the experiment show that there is an underlying event A, which creates high-kinetic-energy protons with equal kinetic energy. Additionally, this event A is negatively correlated to an event B, which creates low kinetic energy protons. Event A needs to be a multi-body fragmentation, either of a doubly charged molecule in the gas phase, or a highly charged cluster. Despite the fact that there are significant amounts of residual non-condensed molecules present in the cluster jet, the correlation observed in figure 2(b) cannot be caused by gas-phase molecules. This is primarily because both detected particles are protons and the parent molecule is ammonia. The only possible fragmentation channel leading to two correlated protons is $\text{NH}_2^+ \rightarrow \text{NH} + \text{H}^+ + \text{H}^+$, a channel which has so far not been observed in photoion–photoion coincidence measurements [42, 43], or ion-impact dissociation [44] on doubly-charged ammonia. Therefore, we assume that event A is associated a single cluster being multiply ionized by absorption of $n \geq 2$ photons.

The shape of the positive correlation feature in figure 2(b) resembles a concerted explosion of a multiply charged object [45]. This is best explained with a core–shell picture of the highly ionized cluster. The protons in the Debye layer of the cluster are ejected by the Coulomb forces, giving them a radially isotropic momentum distribution. Because the absolute kinetic energy of the protons ejected in one shell is approximately equal, the observed correlation feature follows. The core–shell interpretation can be backed-up by the observation that the kinetic energy of the positively correlated protons is in good agreement with the expected plasma potential $hv - IP$ [25], where IP is the first ionization potential of the ammonia cluster ($9.4 \text{ eV}$ [46]). In general, charge ejection out of multiply charged clusters happens either as a concerted Coulomb explosion, or as sequential emission of positively charged ions. In a sequential emission process, each cluster gradually cools and creates positive correlation between all kinetic energies of the cooling cascade, including positive correlations between high- and low-kinetic-energy protons. In contrast, we observe a very pronounced correlation between protons of equal kinetic energy, clearly pointing to a concerted ejection of charges. Consequently, the experimental correlations we observe provide strong evidence for a pronounced core–shell nature of the Coulomb explosion of an XUV-induced nanoplasma in ammonia clusters. In this core–shell explosion, a significant amount of charge and energy is taken away by protons with one specific kinetic energy.

We now take a closer look at how the XUV pulse power influences the kinetic energy of the detected protons. Figure 3(a) shows a typical XUV-induced proton ToF peak with strong broadening (blue curve). Additionally, the power exponent $k$ from a power fit: $A(I_{\text{XUV}}) = A_0I_{\text{XUV}}^k$ is shown for the different areas of the peak (red curve). Here, $A$ is the ion intensity and $I_{\text{XUV}}$ is the XUV pulse intensity. The scaling constant $A_0$ is a free parameter of the fit. Examples of the power fitting for different kinetic energies are given in figure 3(b).

Vertical dashed lines in figure 3(a) mark the area where the time of flight corresponds to a proton kinetic energy of $E_{\text{hi}}^+ = hv - IP$. The power coefficient is close to one and approximately constant for all protons with $E_{\text{hi}}^+ \leq hv - IP$. A power coefficient of one shows a linear relation between the number of photons in the XUV pulse and the number of protons detected. We conclude that the number of protons emitted from the nanoplasma rises linearly as a function of XUV pulse intensity. This linear relation is surprising considering the multi-photon nature of the nanoplasma ignition. However, it can be explained with the core–shell structure of the nanoplasma. An increase in XUV pulse intensity cannot
increase the plasma potential beyond full frustration. Nonetheless, each additional photon absorbed in the cluster will supply an energy of $hν$ to the nanoplasma. A part of this additional energy is dissipated by emitting more protons, and the experimental data show that the relation between the photon flux and the number of protons emitted is linear. Furthermore, we observe that emission of protons with $E_{\text{H}^+} \geq hν - IP$ is highly non-linear in XUV pulse intensity. From this we conclude that a large increase in XUV pulse intensity is required to create a plasma potential that is larger than $hν - IP$.

3.2. Probing the nanoplasma with UV laser radiation

The previous section discussed the kinetics of protons emitted from highly ionized ammonia clusters. In the following, we will look at the difference in the ion spectrum for pump-probe (XUV + UV pulse) and pump only (XUV pulse). The difference between pump-probe ions and pump-only ions will be called probe ion yield. The ions that are created by the pump alone will be called pump ion yield. The UV laser has a large single-photon ionization cross section for the excited states of molecules and atoms. Absorption of multiple photons is required to ionize the electronic ground states of all involved atoms and molecules. Thus, the effect of the UV probe laser on the neutral molecules is negligibly small when compared to the ionization out of excited states. In the nanoplasma, excited states are created via recombination of free electrons and ions, or by electron-impact excitation [25, 33].

We will focus on $\text{H}^+$ and $\text{H}_2^+$ probe ion yields, starting with the asymptotically converged spectrum at a pump-probe delay of 18 ps. The corresponding probe ion-ToF peaks are shown in figures 4(a) and (c), respectively. We observe that the UV pulse can have two different effects on the ion yield, specifically, producing additional $\text{H}^+$ ions, while decreasing the $\text{H}_2^+$ ion yield. We observe that the $\text{H}_2^+$ ToF spectrum has a double peak structure. In contrast, the $\text{H}^+$ ion peak is not significantly broadened. The double peak structure for the $\text{H}_2^+$ probe ion yield is particularly pronounced, forming a local minimum in the center, a feature that is not seen in the pump $\text{H}_2^+$ spectrum (see figure 4(d)). In other words, there is more loss of $\text{H}_2^+$ signal away from the centre of the peak. The decrease of $\text{H}_2^+$ ion yield in the asymptotic difference can only be due to dissociation of $\text{H}_2^+$ by the probe laser. We conclude that, despite the fact that $\text{H}_2^+$ with low kinetic energy is created by the XUV pump laser, the probe laser primarily dissociates the high-kinetic-energy $\text{H}_2^+$.

We will now look at the photodissociation cross section of $\text{H}_2^+$ in order to explain how the probe laser selectively dissociates $\text{H}_2^+$ with kinetic energy. The dissociation probability in our experiment varies between 10% at the center of the $\text{H}_2^+$ ToF peak, and 50% on the outer flanks (cf figures 4(c) and (d)). To explain a dissociation probability of 10% with the used probe laser intensity, a photodissociation cross section of $\sigma \approx 0.3$ Mb is required. Similarly, in order to achieve 50% dissociation probability, a photodissociation cross section of $\sigma \approx 1.5$ Mb is required. Previous research has shown that the photodissociation cross section of $\text{H}_2^+$ has a strong dependence on the ro-vibrational quantum state of the molecule [47]. The cross sections required for the experimentally observed dissociation probabilities of 10% and 50% correspond to a ro-vibrational energy of 2500 K and 8400 K, respectively [47]. From this selectivity, we can draw two conclusions: first, a large part of the $\text{H}_2^+$ created in the ammonia nanoplasma, particularly the $\text{H}_2^+$ with low kinetic energy, has ro-vibrational energies lower than 2500 K. Consequently, the ion temperature in the core of the cluster does not surpass this value. This shows that the ions and electrons in the nanoplasma are not in thermal equilibrium with each other. Second, the $\text{H}_2^+$ ions that emerge from the nanoplasma with significant kinetic energy do also have a larger ro-vibrational energy.

![Figure 4](image-url)

**Figure 4.** (a) Proton-ToF peak created through nanoplasma ignition using different XUV pump laser wavelengths. (b) Corresponding probe proton-ToF peak. (c) $\text{H}_2^+$-ToF peak created through nanoplasma ignition using different XUV pump laser wavelengths. Note that the small peaks at the center are an artifact from the subtraction of background-gas contributions. (d) Corresponding probe $\text{H}_2^+$-ToF peak. (e) Mean kinetic energy difference of the $\text{H}^+$ and $\text{H}_2^+$ ions as a function of the XUV photon energy used for nanoplasma ignition. The values are displayed as differences compared to the mean kinetic energy obtained by UV-ionizing the clusters. This is done in order to distinguish between broadening effects from the cluster environment and kinetic energy release from Coulomb explosion. Red and blue: probe ions. Green: pump $\text{H}^+$ ions. All data shown were obtained at a pump-probe delay of 18 ps.
H$^+_2$ ions are shown in figure 4(e). The values are displayed as $\Delta E_{H^+}$, the difference compared to the mean kinetic energy of protons in a spectrum obtained by UV-ionizing the clusters. This is done in order to distinguish between broadening effects from the cluster environment and kinetic energy release from Coulomb explosion. On the x-axis, the photon energy of the XUV pump is varied in the range of 14.3 eV to 42.9 eV. The kinetic energy of the probe H$^+_2$ ions (blue circles) ranges from 1 eV (12000 K) to 9 eV (100000 K), and is larger than the dissociation energy of H$^+_2$. Furthermore, the mean kinetic energy of probe H$^+_2$ (blue circles) is significantly larger when compared to that of the pump H$^+_2$ (green triangles). The mean kinetic energy of probe and pump H$^+_2$ rises as a function of $h\nu$. The XUV photon energy dependence of the mean kinetic energy of H$^+_2$ shows that a higher XUV photon energy generates a deeper plasma potential. However, the change in mean kinetic energy does not directly correspond to the change in photon energy. This is not surprising, since the total photon flux and the ionization cross section are peaked at 19 eV. From the difference in the mean kinetic energy of probe and pump H$^+_2$ we conclude that the ro-vibrationally excited H$^+_2$ is primarily created in the Debye layer of highly ionized clusters where the electric field is the strongest. On the other hand, the H$^+_2$ with lower ro-vibrational energy are emitted during the later stages of the nanoplasma evolution.

No significant kinetic energy release can be seen in the probe H$^+$ (red rectangles in figure 4(e)). The primary portion of the probe H$^+$ signal is due to UV ionization out of excited states of atomic hydrogen [33, 48]. These excited hydrogen atoms are formed according to the reactions [33]:

$$H^+ + e^- \rightarrow H^+ \quad \text{and} \quad \text{(1)}$$

$$\text{NH}_3^+ \rightarrow \text{NH}_2^+ + H^+. \quad \text{(2)}$$

From the negligible kinetic energy, we deduce that H$^+$ is not subject to a significant plasma potential at the moment of ionization, i.e. at a pump-probe delay of 18 ps. High-kinetic-energy protons that are created through the dissociation of H$^+_2$ contribute only a minor part to the probe H$^+$ yield, as can be seen by comparing the absolute scales of figures 4(a) and (c).

3.3. Time-resolved studies on Coulomb explosion

In the spectra shown in figures 4(a) and (c) we observed that H$^+$ ions created by the probe laser at a pump-probe delay of 18 ps do not have significant kinetic energy. For the case that H$^+$ in the cluster is ionized while the nanoplasma is still active, there are two different options. Either the excited hydrogen atom was formed in the bulk of the plasma, in which case the proton will remain inside the cluster; or, the excited hydrogen was formed in the Debye layer of the plasma, in which case the proton will acquire kinetic energy proportional to the plasma potential at the time of ionization. This allows us to use the time-dependent kinetic energy of the probe protons as a probe for the plasma potential. Figure 5(a) shows the mean kinetic energy of the probe H$^+$ as a function of pump-probe delay. The kinetic energy has a maximum in the vicinity of $t_0$. Afterwards, it decays rapidly and converges at 400 fs pump-probe delay. The decay time-constant is universal for all pump photon energies and only marginally larger than the temporal pulse overlap of the two laser pulses (depicted as the grey shaded area in figure 5(a)). Using our ToF spectrometer, the kinetic energy
of the protons could only be determined for values larger than 500 meV.

Complementary to the ions, we use the VMI spectrometer to detect photoelectrons from the probe ionization of H⁺. The most abundantly populated H⁺ state is the n = 2 state [33], yielding electrons with energy \( E_{\text{ele}} = 1.35 \text{ eV} \) when ionized with the 4.75 eV UV photons. If there is a Coulomb potential present, we observe a shift in the kinetic energy of the photoelectrons. The pump-probe delay dependent shift in the vertical binding energy (VBE) of the electrons (\( \Delta E_{\text{ele}} \)) is displayed in figure 5(b). The inset shows an example of the different peak positions at pump-probe delays of 0.5 ps and 1 ps, respectively. A determination of the peak position was only possible for \( \Delta E_{\text{ele}} < -400 \text{ meV} \). For larger shifts, the peak is strongly broadened and could not be distinguished from the overlapping low energy electrons emitted by the nanoplasma. The VBE shift at a pump-probe delay of 300 fs is roughly –300 meV and converges to zero with a half-lifetime of (200 ± 30) fs.

In section 3.1 we showed that a large number of protons with several eV of kinetic energy are emitted from the nanoplasma. Combining this observation with the dynamics shown in figure 5 leads us to the conclusion that we observe a high-energy Coulomb explosion which governs the charge equalization of the cluster. A proton that is accelerated by a Coulomb potential of 10 eV at the surface of a cluster with a radius of 1.5 nm traverses a distance of \( \approx 1.5 \text{ nm} \), within 100 fs, consistent with the fast timescales. After the Coulomb explosion, the clusters are only mildly charged and hydrodynamic forces are dominant, explaining the slower decay of the remaining –300 meV plasma potential. At a pump-probe delay of 1 ps, the plasma potential is completely neutralized. This two-phase expansion of the cluster is illustrated in figure 5(d).

4. Conclusion

We induced a nanoplasma in ammonia clusters using high power XUV radiation from the FERMI free-electron laser. Using simultaneous photoelectron and ion detection we have shown that emission of high-kinetic-energy ions plays a significant role in the evolution of XUV-induced nanoplasmas. Using shot-to-shot covariance mapping, we show that protons with kinetic energy \( 4 \text{ eV} \leq E_{\text{H}^+} \leq 20 \text{ eV} \) are emitted from the nanoplasma in a correlated way. Furthermore, we use a delayed UV laser as probe to ionize excited states of H and \( \text{H}_2^+ \) in the plasma. From the pump-probe dependent mean kinetic energy of the probe-laser induced \( \text{H}^+ \) ions, we get information on the lifetime of the plasma confining potential. We found that the nanoplasma decays in a two-stage process. In the first stage, the potential reduces drastically through the concerted expansion of protons. These observations show that the highly-ionized ammonia cluster acts as a core–shell system where the Debye layer of the nanoplasma dissipates a large fraction of the energy contained in the nanoplasma through a concerted Coulomb explosion of protons and other light, positively-charged ions. After the Coulomb explosion, the cluster expansion slows down and hydrodynamic forces become dominant. Using UV-laser-induced dissociation of \( \text{H}_2^+ \), we show that a major fraction of the \( \text{H}_2^+ \) molecules in the plasma core has an internal energy that is less than 2500 K.

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