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Primary Radicals Production from Water Fragmentation by Heavy Ions

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The importance of fragmentation of water molecules by heavy ions is discussed for several different physical environments, including tumor treatment, planetary sciences and fission reactors. The characteristics of the fragmentation yields by electrons, protons and photons are presented and compared with fragmentation yields of water molecules by C^{3+} and C^{5+} ions at energies corresponding to the Bragg peak. It is shown that for low energy electron loss and high energy electron capture, the water molecules essentially blow-out, releasing a much larger fraction of C^{9+} ions as compared with electrons, protons and photons.

I. INTRODUCTION

Water is the main component of our bodies, covers most of Earth surface, is found in several moons of our planetary system and has been continuously brought to our vicinities by comets, revealing part the distant history of our solar system formation. In all these scenarios the water molecules can be found bombarded by photons as well as by a variety of particles coming from several different sources: the Sun, planetary magnetospheres, radioactive sources, particle accelerators or nuclear power plants, for example. These ionizing radiations have the power to transform the benefic water molecule in quite reactive pieces. When water molecules are splitted due to the action of these particles, very reactive fragments are produced which strongly interfere on the nearby environment. Different particles produce different fragmentation yields which must be known to model the complex chain of reactions which follow the primary products.

The splitting of water by photons is a much more familiar phenomenon than by heavy particles. Available ionizing heavy particles are by far less abundant than photons but there are several circumstances where they play an essential role. However, there are very few studies on the dynamics of water fragmentation by heavy ions, from both the experimental and theoretical points of view. In case of heavy particles, the collision with water molecules involves collision channels which are not present in the photon or electron cases and whose dynamics are not well described by existing theoretical models, making the experimental data essential to understand and quantify the effects of the ionizing radiation.

This paper has three purposes. The first is to give to the reader a general view of some of the environments where the interaction between heavy ions and water occurs. Thus, in the first sections of this paper, we describe the role played by heavy ions in splliting water in some important scenarios of four branches of sciences: planetary sciences, Precambrian research, cometary atmospheres and tumor therapy. The second is to describe the experimental procedures and results of recent measurements performed by the authors with C³⁺ and O⁵⁺ ions on water molecules in the vapor phase. These are

the first measurements to cover a broad range of fragition yields and which have opened the perspective of ide ing common signatures of water fragmentation by participation types, energies and charge states. Finally, the is to show that water fragmentation, either by photons trons, protons or heavy particles presents universal particles is a key result to organize several apparently dissocollision regimes allowing a better modeling of the interbetween heavy particles and water.

II. WATER RADIOLYSIS

Before we describe some of the environments where ions act as important ionizing agents of water molecu general, we briefly describe how these ions can affect chemical composition by looking at the important case uid water. Swift ions passing through liquid water re electrons from the water molecules leaving a track of ic species. The final inventory of the new species generated lated to a complex sequence of events which are usually sified as physical, physicochemical and chemical stages ing the first 10^{-15} s from the primary electron removal the molecular ion relaxes generating ionic fragments and trons which can produce further ionizations, giving a small regions of high ionization densities commonly l as spurs. Between 10^{-15} s and 10^{-12} s the electrons ever termalize becoming solvated (e_{aq}^-) , the H_2O^+ ions diffus domly, the spur expands, and some fast ionic reactions as $OH^+ + H_2O \rightarrow H^+ + H_2O_2$ occur. Between 10^{-12} 10^{-6} s radicals such as OH, H, O as well as e_{aa}^- specigrate randomly and can react with each other through ety of reactions such as $OH + OH \rightarrow H_2O_2$, $O + H_2O$ + H₂ resulting, in the end, on the removal of most chen reactive species [1, 2]. Because, as we will show later, ions can produce a much larger variety of primary fra ions as compared with photons, electrons or protons, so the final products can be effectively enhanced, as for ple, the production of O2 [3]. The same sort of enhance occurs in bombardment of ice, where much larger amou charged H, O, H₂, and O₂ species are sputtered by high izing 252 Cf fission fragments as compared with weakly ionizing 0.85 MeV N^+ ions [4].

III. EUROPA

The Jupiter moon Europa is about 670,000 km from Jupiter, has roughly the size of our moon, is smoothly covered with ice and has a very tenuous oxygen atmosphere. Unlike our Earth, the source of oxygen in Europa's atmosphere is not likely to be biologically produced by plants and bacteria. It comes from the bombardment of the surface by H, O and S ions from the Jupiter magnetosphere which erodes the surface by sputtering, splitting water molecules and producing H_2 and O_2 [5]. The hydrogen escapes from the atmosphere while the oxygen is left behind. A further and interesting consequence of this bombardment is the change of the surface reflectance for UV radiation [6] as observed by the Galileo spacecraft. Water radiolysis can produce, for example H + OH radicals which, after diffusion, can lead to a chemical combination of two OH radicals yielding H₂O₂. Laboratory experiments [6] show that the reflectance of water ice with just 0.16 % of H₂O₂ results in a decrease of 20 % of the surface reflectance at wavelengths of 210 nm as compared with pure water ices. As stressed in Ref. [6], the abundance of H_2O_2 and the existence of O_2 in the atmosphere demonstrate that the surface chemistry on Europa is dominated by water radiolysis induced by energetic heavy particle bombardment.

IV. OKLO NATURAL FISSION REACTOR

In the beginning of 1970s, a uranium shipment to France from a mine near the Oklo river, central-east Gabon, Africa, was found tinily depleted from the usual 0.7202 % ²³⁵U fissionable isotope abundance. The reason for this depletion was found to have its origin in the Precambrian period, about 2 billion years ago. By that time, the evolution of plants and algae started the increase of atmospheric oxygen. Uranium has a peculiar chemistry which makes it insoluble in water under oxygen-free conditions but soluble in its oxidized form, UO₂. With the increase of oxygen in the atmosphere as well as in the ground water, the UO2 began to form and dissolve, becoming present as trace elements in flowing waters. In the Oklo region there are evidences that some quite peculiar microorganism has the ability to collect and concentrate uranium [7]. The U started then to be deposited at higher concentrations at that locations. The differences in the life-time of ²³⁵U and ²³⁸U is such that, by that time, the fraction of ²³⁵U was around 3% and, when an enough amount of U was accumulated by these strange bacteria, a self-sustained chain reaction started, lasting around 1 million years [8, 9]. At critical operation, fission fragments from ²³⁵U could directly interact with water, producing further amounts of O2 as by-products. This extra O₂ not only favored the solubility of fission products but also modified the near-field redox conditions making new oxidized species such as, for example Pb₃O₄. On the other hand, in zones rich in organic matter it could favor the oxygen consumption resulting in a reducing environment [9]. The radiolysis by heavy ion fission fragments thus plays an itant role in the understanding of this natural reactor.

V. COMETARY X-RAYS

The Sun emits a continuous stream of particles who jor constituents are electrons and protons, and minor are C, N, O, Ne, Mg, Si, etc. The heavy ions are in several charge states and their interactions with pla and cometary atmospheres are responsible for a broad v of phenomena such as auroras, x-ray emission, atmos chemistry, etc. The interaction of highly-charged ion planetary and cometary gases such as hydrogen, methan water not only produces ionization and dissociation of atoms and molecules but also changes the ion charge through electron capture and stripping. Electron capture cited states can result in a subsequent decay through em of photons in the ultra-violet or x-ray regions, which of the major sources of such photons in very cold atmosp In 1996 the observation of x-ray emission from the Hyakutake was reported [10]. Being icy bodies, the sion of UV and X-rays from such objects was unexp The origin of such emission is presently considered to b sult of charge exchange processes of highly-charged o ions from the solar wind with water [10–12]. For the age velocity of the solar wind ions, the electron capture ions occurs into states with high quantum numbers. Th tured electron subsequently decays by emitting energeti tons. In addition, the chemical reactions involving the fragment ions produced by the collisions with the solar should be added to those resulting from the photofragr tion process, in order to model the net ionic composit the ion tail of comets. Due to the lack of information of mary water fragmentation by the solar wind, the magnit its effect on the comet ion tail cannot be properly evalu

VI. C-ION THERAPY

For about 30 % of cancer patients the disease is well ized within a specific region of the body when it is diag and good chances of cure are obtained through chem apy, surgery or radiation therapy. In the latter, the c aim of maximizing damage in the tumor sites while min ing damages to healthy surrounding tissue has lead to t of protons and, more recently, carbon ions to provide spatial-specific damage profiles [13-18]. In radiation of ticle therapy there is a possibility that the particle intera rectly with critical components of the cell or with wate latter case is known as indirect action and the damage is by producing free radicals resulting from the interaction bombarding particles with water molecules. These ra can diffuse and make subsequent damage to the mite dria, enzymes or DNA. It is estimated that 75 % of the damage in mammalian cells are due to the action of the radical. As these radicals are extremely reactive, the dif

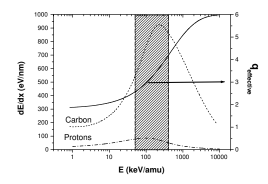


FIG. 1: Energy loss of protons and C-ions in water [19]. The right abscissa gives the C ion effective charge [21], indicated by the dashed curve. The energy region covered by the present measurements is indicated by the dashed area.

length must be within about 20 Å from the target. Recently, good effective radiotoxic responses from C-ion tumor therapy was reported [14–18]. The high ionization densities within the Bragg peak make normal DNA enzymatic repair difficult in contrast to low LET (Linear Energy Transfer) electron, proton and photon irradiation [13]. In spite of C- ion therapy being in current use there are no previous measurements, in the Bragg peak region, of water fragmentation by C ions.

The Bragg peak is where most of the energy of the ionizing particle is deposited and the behavior of several physical systems under the effect of heavy particle irradiation depends on that information. For carbon ions, the energy of the ions at the top of the Bragg peak is nearly 250 KeV/amu [19, 20], with an average charge state of approximately q=3 [21]. Here, we report the general behavior of the branching ratios for positive ions production in collisions of C^{3+} ions on water molecules at energies around the Bragg peak. The region of the Bragg peak covered in this work is shown in Fig. 1.

VII. PREVIOUS DATA OF ELECTRONS, PHOTONS, PROTONS AND HEAVY IONS

Previous measurements of the absolute yields for fragmentation of water through electron removal - essentially ionization - have been carried out by photons, electrons, protons and Xe^{44} ions. These measurements are displayed in Fig. 2, where the ternary graph shows the normalized fractions of H_2O^+ , OH^+ and Σ_q O^{q+} cross sections by proton ionization of Ref. [22] in the 100 - 400 keV range, electron ionization from Ref. [23] in the 40 - 1000 eV range, 6.7 MeV/amu Xe^{44+} ionization from Ref. [2], and photofragmentation results from Ref. [24] in the 30 - 60 eV range. The sum of the three fractions was made equal to one. It is clear that the photon, electron and proton data tends to cluster with the same Σ_q O^{q+} and slightly different H_2O^+ , OH^+ fragmentation fractions. It is also noteworthy that, for each of these clusters, the fragmen-

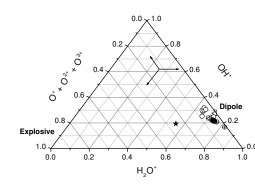


FIG. 2: Ternary plot of probabilities for H_2O^+ , OH^+ , and formation by proton ionization (open circles), (Ref. [22]); e ionization (crossed circles) (Ref. [23]); 6.7 MeV/amu Xe^{44+} tion (star), Ref. [2]; and photofragmentation results from Re (side open triangles). The three diverging arrows in the upper the graph indicate how a particular data point is connected wi of the three axes. The different data belonging to the same far symbols corrrespond to different projectile energies.

tation fractions are quite independent of the particle e On the other hand, the 6.7 MeV/amu Xe⁴⁴⁺ fragmer fractions are clearly different from the others, with a amount of the Σ_q O^{q+} fractions.

The results for protons, electrons and photons displa Fig. 2 clearly show that the fragments follow a sequenti tribution with $H_2O^+ > OH^+ > \Sigma_q O^{q+}$. This distribu characteristic of the dipole ionization of the water mol [24] and express the decay probabilities following the i tion of one of its four valence molecular orbitals 1b₁, 3a and $2a_1$, with ionization energies of 12.6, 14.7, 18.5 an eV, respectively [24]. This dipole, sequential fragmer pattern, has been always assumed to be the ionization tern, independently of the particle type and energy, in radiolysis modeling [1]. The former particles never real explosive regime, when the Σ_q O^{q+} become the major mentation products. In fact, not even the Xe⁴⁴⁺ data rea explosive regime, when the above sequence is inverte the Σ_q O^{q+} fraction becomes the major fragmentation uct. In the following sections we show that collision ch other than ionization can perform collisions which are enough to reach the explosive regime.

VIII. EXPERIMENT

Mono-energetic O⁺ and C⁺ beams are delivered 4MV Van de Graaff accelerator of the Catholic Univer Rio de Janeiro. A 90⁰ magnet analyzes the primary making its charge/mass and energy selection. Subsequence the beam passes through a striping gas cell producing a vof charged daughter beams. The desired charge state of

nent is then selected and directed onto the experimental line by a switching magnet. After collimation, the beam is further cleaned from spurious components just before the collision chamber by a third small magnet.

The target cell is placed inside the collision chamber and connected to an effusive water vapor set-up. The set-up is composed by a manifold connecting a needle valve to a Pyrex bottle containing 10 ml of de-ionized water, which is first pumped by an rough pump. When the pressure inside the bottle decreases down to a critical value, the water changes phase into ice. The pump is then valved off and the bottle naturally warm up . The process is repeated several times to drive out any remaining dissolved gas, in order to assure a pure $\rm H_2O$ target. The water is then allowed to sublimate into the cell through a needle valve.

A capacitive manometer (MKS-Baratron) provides an absolute measurement of the pressure inside the gas cell. Determination of the gas pressure for water vapor is more difficult compared to more standard molecular gases. Working with $\rm H_2O$ requires longer times for the system to reach the equilibrium (order of a few minutes) and a significant drift of the zero reading of the manometer can occur. We overcame this difficulty by doing sets of short runs and by checking the zero reading of the gauge each run. Typical working pressures employed during the measurements were kept between 0.5 to 1 mbar, always under single collision conditions.

After interacting with the target, the projectile beam leaves the collision chamber and an analyzing magnet separates the projectile beam from its products. The final projectile charge state will define the collision process related to the molecular ionization/break-up. For example, the O⁵⁺ projectile can pick up one electron from the H₂O target (electron capture process) leading to a final projectile charge state of O4+. All the beam components are split and detected together by a position sensitive detector placed at the end of the beam line, 4 m downstream. Micro-channel plates detectors usually present a loss of local efficiency due to saturation effects when the number of particles/channel is increased. In this work, as the beam image is very important to separate the different collision channels, it was necessary to work with small counting rates for which the local efficiency was found to be constant. For O⁵⁺ projectiles, for example, we have found 500 particles/sec as being our upper limit. Therefore, 250 to 350 particle/sec were used throughout the measurements.

The charged target products, H^+ , O^{q+} , OH^+ and H_2O^+ resulting from the interaction with the incident projectile beam are collected by a transverse electric field. The use of a strong electric field (960 V/cm) assures the maximum collecting efficiency for all target products. These slow ions pass the electrodes, enter into a field free region, and finally go onto a microchannel plate detector. The target products are separated and analyzed by a standard time-of-flight technique in coincidence with the position-sensitive projectile detector output. The recoil-ion detection efficiencies are obtained using the same procedure described by Santos et al. [25, 26]. In Fig. 3 we have a typical time-of-flight spectrum for 2.0 MeV O^{5+} on H_2O . The H_2O^+ production for the electron capture channel is clearly reduced when compared with the ionization channel.

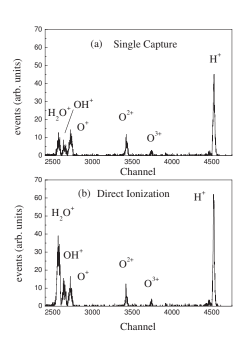


FIG. 3: Time-of-flight spectra for electron capture and direct tion by 2.0 MeV O⁵⁺ on water vapor molecules.

IX. EVIDENCE OF EXPLOSIVE FRAGMENTATION

The experiment described above measured the fragr tion of water due to three different collision channels: i tion, electron capture and electron loss. These three ch have different dynamics and energy transfers. For exa for C³⁺ at 3.5 MeV the energy transfer in the ionization nel is \cong 60 eV while for the electron capture channel is eV. These large differences in the energy transfer as v the much smaller impact parameter in which capture at this energy compared to ionization give rise to large ences in the fragmentation pattern. This is indeed show 4, where the present data is added to the data shown in 1 The possible fragmentation yields are now scattered a broad range of Σ_q O^{q+}fractions, ranging from the dipo ues, around 5 %, to a really explosive dynamics, with O^{q+} fraction lying around 80 %. The data correspond the most explosive outcome is associated to processes i ing high energy transfers and/or close collisions: high-e electron capture and low-energy electron loss. As the co becomes more distant and involving smaller energy tran the fragmentation pattern goes towards the dipole chara tics. For example, the most explosive channels and en shown in Fig. 4 are electron capture (closed triangles) a velocities and electron loss (closed circles) at low velo For electron capture, the collision becomes softer as t locity decreases and this family of data points moves,

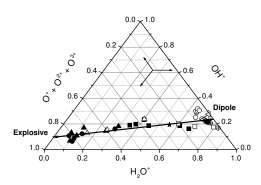


FIG. 4: Same as Fig. 2 including fragment formation by C^{3+} ionization (closed squares), capture (closed triangles) and loss (closed circles), O^{5+} ionization (open squares) and capture (open triangles). The line is a fit through the data. For the C and O data shown, the energy increases along this line from right to left for the ionization and capture channels and from left to right for the loss. The explosive regime is reached for high energy electron capture and low energy electron loss.

the indicated line, from the explosive region (higher velocities) towards the dipole region (lower velocities) of the graph. For electron loss the reverse behavior occurs.

The coalescence of the data on the line shown in Fig. 4 is also remarkable. This coalescence is pretty much insensitive to the collision energy, collision channel, particle type and charge state. It can be shown that this coalescence is a related to the constraint that all fragment products are due to either one- or two-electron removal cross sections plus a decay scheme which is independent of the collision dynamics [27]. This last point is an indication that the removal and the decay processes can be well separated in time.

X. CONSEQUENCES OF EXPLOSIVE PRIMARY FRAGMENTATION

From Fig. 4 it is clear that the O^{q+} production by heavy ions for energies around the Bragg peak is much larger than

the mere 5% production by electrons, photons and profound any energy. This finding can explain the observed incre O_2 by high LET particles. After being produced by the mary collision, the O ion rapidly neutralizes through electrons of the $O(^3P)$ states. The singlet $O(^1D)$ reacts rapidly with through the reactions $O + H_2O \rightarrow H_2O_2$ or $O + H_2O \rightarrow H_2O_2$. On the other hand, the ground state $O(^3P)$ is reactly inert in water and can survive enough time to ever react with OH through the reaction $O(^3P) + OH \rightarrow HO_2$ ducing HO_2 . The suggestion of Ref. [3] that the increasing production by high-LET particles is due to an increprimary O production is supported by our findings are consequence of the increasing importance of the double tron removal at the Bragg peak region [27].

XI. CONCLUSIONS

Here we find evidence that in C and O ions interactio water, a large fraction of molecules completely blows or ducing large abundances of H⁺ and O⁺ fragments in co ison to proton, electron and photon irradiation, or even ions with energies outside the Bragg peak region. Indee cross sections for O+ production by C3+ ions at the peak are about one order of magnitude larger as con with those for the above mentioned projectiles at any e [23, 24, 27]. Our findings have straightforward implic in the subsequent fast chemistry at the ionization site, s the production of the highly reactive OH radicals, hyd peroxide, H₂O₂, as well as O₂, showing that the market ferences in primary O production between high-LET pa and low-LET electrons, protons and photons can be the ble source of the observed differences in the final produ ionizing particles.

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