An electron gun is used with a mirror electrostatic field to produce zero or near zero velocity electrons by forming a turning point in their trajectories. A gas capable of attaching zero or near zero velocity is introduced at this turning point, and negative ions are produced by the attachment or dissociative attachment process. Operation may be continuous or pulsed. Ions thus formed are extracted by a simple lens system and suitable biasing of grids.
FIG. 1

multichannel analyzer

channel electron multiplier

quadrupole mass filter

ion lenses

FIG. 2

electrons on off off on

reversal field

ions to mass analyzer

electrode C₁

ring R

mirror M

extractor G₁

time, μs
FIG. 3
GENERATION OF INTENSE NEGATIVE ION BEAMS

ORIGIN OF INVENTION

The invention described herein was made in the performance of work under a NASA contract, and is subject to the provisions of Public Law 96-517 (35 USC 202) in which the Contractor has elected not to retain title.

BACKGROUND OF THE INVENTION

This invention relates to a method and apparatus for the generation of atomic and molecular negative ion beams, either pulsed or continuous beams.

The production of beams of atomic and molecular negative ions is of considerable interest in diverse areas of atomic, molecular, and plasma physics. Such beams are required for fusion plasma heating, heavy ion inertial-confinement fusion, and in basic atomic and molecular scattering studies. The types of sources for ion production are many and varied, involving both plasma and surface-plasma interactions.

SUMMARY OF THE INVENTION

In accordance with the present invention, an ion source utilizes a beam of electrons and target molecules. The source includes an electrode which reverses the electron beam, producing electrons at their turning point having a distribution of velocities centered at zero velocity. A gas which attaches zero velocity electrons or some near-zero velocity electrons is introduced at or near this turning point. Negative ions of the gas are produced by an attachment, or a dissociative attachment, process. For many of the thermal electron-attaching molecules, the attachment cross section at zero energy can be quite large, varying as (electron energy)^{-1}, or just the s-wave threshold law.

The novel features of the invention are set forth with particularity in the appended claims. The invention will best be understood from the following description when read in conjunction with the drawings.

BRIEF DESCRIPTION OF DRAWINGS

FIG. 1 is a schematic diagram of the apparatus and method for generation of intense beams of negative ions by zero-energy electron attachment, or dissociative attachment, in accordance with the present invention.

FIG. 2 illustrates the grid pulsing sequence for pulsed operation of the source shown in FIG. 1.

FIG. 3 illustrates the relative Cl^+ yields for continuous operation of the source shown in FIG. 1 as a function of the ring potential V_R, and at electron energies E_e of 20 eV and 30 eV.

FIG. 4 illustrates the relative Cl^+ yields for continuous operation at 40, 50, 60, and 80 eV electron energies.

FIG. 5 illustrates the distribution of Cl^- pulses during several pulsing cycles of the ring electrode voltage V_{Rmax}.

DESCRIPTION OF PREFERRED EMBODIMENTS

In the following description, the cross sections referred to are "attachment cross sections" defined in the usual sense of the probability per molecule that an electron traversing a gas will undergo an attaching collision with the gas molecules.

Many molecules, such as SF_6, CFCl_3, perfluorinated carbon compounds, and chlorocarbon compounds, have extremely large cross sections for attachment of zero-energy electrons to form negative ions such as F^-, Cl^-, Br^-, and I-. In conventional ion sources, electron energies are too high (of the order 2-3 eV or greater) to attach to these molecules. In the apparatus of the present invention, "zero energy" electrons are generated momentarily by pulsing a reversing electrode R negatively with respect to an electron-gun cathode F,P. In this preferred embodiment, the reversing electrode is comprised of a conductive ring R and a conductive mirror plate M. These electrons can then attach to molecules, such as CFCl_3, which effuse from a 1 mm diameter stainless-steel tube B placed at the center of the electrostatic reversal field. This field is produced by applying appropriate negative potential on the ring R and mirror M. The location of the electron turning point is governed by the magnitude of potentials on the electrodes R and M. Applying larger negative potentials E_R and E_M has the effect of moving the turning point to the left in FIG. 1, and vice versa. The "mirror" effect produced by the ring R and mirror M can be either concave, convex or planar, depending upon the potential ratio. However, the invention is not limited to such an arrangement of these electrodes R and M as the "reversing electrode" for production of zero energy electrons.

The present invention takes advantage of the extremely large, threshold (zero-energy) electron attachment cross sections in several molecules to generate, for example, intense beams of F^-, Cl^-, Br^-, and I^- ions. Thus, an ion source is provided in which an electrode or grid, which reverses electron trajectories, is placed just beyond an ion extraction aperture comprised of ion lenses L_1 and L_2. When the reversing electrodes are pulsed negatively, this pulse voltage reverses the direction of electrons momentarily, giving the electrons zero kinetic energy. At this zero-energy turning point the electrons attach to the ambient gas, and generate negative ions as the product of the attachment process. The method described herein as an example is applied to the generation of Cl^- ions from CFCl_3. Other ions from effusing molecules may be formed and extracted through the aperture formed by the ion lenses L_1 and L_2 in the same manner.

Recent experimental evidence has shown that the zero-energy cross section to CFCl_3, SF_6, CCl_4, and other molecules is much larger than previously expected. Thus, the probability of electron attachment during the field-reversal pulse will be much larger, making this apparatus an even more efficient ion source. Generally, the production yield of negative ions is the product of the electron energy distribution function and the dissociative attachment cross section, integrated over all electron energies. The reversing electrode momentarily creates a maximum in the distribution function at the maximum of cross section (zero eV), thus maximizing the integral and therefore the yield of negative ions.

The electrons in the turning point region B will have a peak at about zero eV in their energy distribution. These electrons can then attach to an admixed gas having a peak in its dissociative attachment (DA) cross section at zero eV. Cl^- was generated as an example from CFCl_3 for several reasons: the behavior of the cross section for Cl^- production at zero eV (greater than 10^{-13} cm^2) is now well known, and is many orders
of magnitude greater than for H− production from H2 at higher energies, an effect due to the divergent nature of the s-wave attachment cross section; Cl− is by far the major ion produced in CFC13 at any electron energy; and CFC13 is inexpensive and inert.

Other examples of the effect of reversal of electron trajectories is seen in a high-pressure electron swarm and the diffuse discharge plasmas. Electrons in a swarm, for example are backscattered by elastic and inelastic gas collisions. These electrons move against the drift field until their trajectories are reversed, and are accelerated along the field lines again. At their turning point, they have essentially zero kinetic energy, and can be removed from the swarm by attachment to SF6, 12, or an admixed perfluorocarbon compound.

The overall process can be described, for molecules AB and electron energy E, as

\[
\text{AB}^− (\text{attachment}) \rightarrow \text{A}^+ + \text{B}^− (\text{dissociative attachment})
\]

From the Wigner threshold law, the cross section to form the atomic ions Cl− or F−, or the corresponding parent negative ions, behaves as \(E^{−4}\) in the limit \(E→0\), thus providing an extremely efficient path for negative ion formation. Use of CFC13 had the additional virtue that the cross section for Cl− production greatly exceeded that for production of other possible ions (F−, Cl2−, and CCl3−) at energies below 3 eV. This situation arises in other molecular targets as well, and makes for an efficient source with minimal interference from other species during extraction and acceleration in an ion beam transport system.

The theoretical calculation of Henkelman and Ottensmeyer, J. Phys. E7, 176 (1974), is used to fix the placements of the C3, M, and R electrodes, and to determine the aperture diameters in C3 and R. The distance between C3 and R was 9.8 mm in our experiments.

Assuming the origin as the center of the aperture in C3 and neglecting aberrations in the reversal field, the longitudinal energy \(E_L\) and transverse energy \(E_T\) are

\[
E_L = E_0 - e|V_R|
\]

\[
E_T = E_0e
\]

(2)

where \(e\) is the electric field intensity, and \(E_0\) and \(E_0\) refer to launched or reverse increasingly energetic electron (where \(e\) is the magnitude of electron charge) and the potential \(V_R\) on R was obtained from the calculations of Henkelman and Ottensmeyer, supra for the ratio \(V_R/V_M\).

Results of Cl− signal vs \(V_R\) for continuous-mode operation are shown in Figs. 3 and 4 at the indicated \(E_0\). It is clear from these spectra that, as expected, increasingly greater ring voltages \(V_R\) are required to reverse increasingly energetic electron (higher \(E_0\)). Also, the highest signal count rates and narrowest widths were encountered as \(E_0\) above 30 eV. While this effect of width is not clearly understood, several effects which could give rise to this behavior with \(E_0\) are: optimum electron-molecule spatial overlap (e.g., focusing at region B the smaller “disk of least confusion” rather than the Gaussian image), a minimal transverse energy \(E_T\), thus bringing more electrons into the peak of the attachment cross section at zero electron energy (see Eq. 2), minimal aberrations in the reversal field, or optimal extraction and focusing efficiency by G1, L1, and L2.

All count rates in Figs. 3 and 4 are given relative to the 30 eV count rate, so that relative efficiencies at the different energies can be compared. In the following table we list at each energy the peak counting rate.
For pulsed operation of the source described herein—before with reference to FIG. 2 there is shown in FIG. 5 an oscilloscope image of the distribution of $C^-$ pulses during several cycles of the ring voltage $V_{R_{max}}$. The electron energy is 40 eV and the value $V_{R_{max}}/V_M$ is 0.75. The maximum yield and peak-integrated yield for this pulsed operation is found to be about ten times greater than for continuous operation, even after a 30% duty cycle is taken into account. This increase is almost certainly due to the fact that $C^-$ ions are now being extracted from a field-free region, and not from the field of the ring and mirror which tends to drive the ions from the turning point region back toward $C_3$.

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Efficiency

The production rate $I$ of $C^-$ ions can be written as

$$I(C^-) = N \phi_{(E)} \sigma(E) \epsilon,$$

where $N$ (cm$^{-3}$) is the CFCl$_3$ target density, $\psi$ (cm$^3$) is the overlap volume between the electron and target beams, $\phi$ (s$^{-1}$ cm$^{-2}$) is the incident electron flux, and $\sigma(E)$ (cm$^2$) is the attachment cross section at $E=0$ energy. Estimates of the quantities in Eq. (3) are as follows: A pressure of CFCl$_3$ in the beam was taken as 0.133 Pa, or $N = 3.3 \times 10^{17}$ cm$^{-3}$. The interaction volume is taken as that of a cylindrical electron beam of 0.2 cm diameter intersecting a target beam of 0.2 cm diameter, or $V = 6.5 \times 10^{-3}$ cm$^3$. The electron flux is taken as a 1.5 $\mu$A electron beam in a cylinder of 0.2 cm diameter or $3.0 \times 10^{14}$ s$^{-1}$ cm$^{-2}$, and an average attachment cross section is taken as $2 \times 10^{-14}$ cm$^2$ for $E$ less than approximately 0.1 eV. Assuming that the entire 1.5 $\mu$A incident current is reversed to give a sum of longitudinal and transverse energies $E$ of less than about 0.1 eV, then the production rate $I(C^-)$ is $1.2 \times 10^{12}$ s$^{-1}$, or $0.2$ $\mu$A. Thus, it appears that 13% of the incident electron beam can be converted to $C^-$. This corresponds to a current density $\rho$ of $0.2 \mu$A/$\sqrt{3} \times 10^{-2} = 6.7$ $\mu$A/cm$^2$ for emission from a 0.2 cm diam region of the target. This density can be increased by several orders of magnitude through use of (a) more intense electron beams (milli-ampere or amperes as opposed to microampere currently used) with a correspondingly larger beam size, and (b) denser targets.

It should also be possible to extend the use of the source to the generation of negative ions through attachment resonances located at nonzero electron energies, since a continuous distribution of electron energies exists between $C_3$ and $R$. While for the present demonstration of the source we have chosen $E_p=0$, one can choose (see Eq. 2) any other energy $E_p$ located at the same $y$, but at a value of

\[
\epsilon' = \frac{(E_p - E_p)}{E_p}
\]

This energy could correspond, for example, to the 3.7 eV$^2$ resonance in H$_2$ to produce $H^-$. Moreover, any given energy $E_p$ is attained twice: once when $\epsilon'$ and $y$ are parallel (electrons decelerating towards the turning point), and once when antiparallel (electrons accelerating away from the turning point). Thus, the electron beam is "used" twice.

What is claimed is:

1. A method for generating negative ions using a beam of electrons comprising the steps of:
   - producing an electric field in the path of said electron beam for reversal of said electron beam to produce at the turning point electrons having a distribution of energies centered at zero;
   - introducing a gas containing thermal electron-attaching molecules at a point in the path of said electron beam where said electrons have a low energy and a high probability for attachment to said molecules, thereby efficiently generating ions; and
   - directing said ions in a focused beam to a utilization device.

2. A method as defined in claim 1 for generating negative ions from thermal electron-attaching molecules having an extremely large attachment cross section at zero electron energy wherein said gas of said molecules is introduced at said turning point.

3. A method as defined in claim 1 for generating negative ions from thermal electron attaching molecules having an attachment cross section at a level near zero electron energy wherein said gas of said molecules is introduced at a point in the path of said electron beam ahead of said turning point, whereby said electrons of the appropriate energy make two passes through said gas of said molecules for enriched ion generation.

4. A method as defined in claim 1 wherein said electric field is controlled to be constant for continuous generation of ions.

5. A method as defined in claim 1 wherein said electric field is controlled for pulsed generation of ions.

6. A method as defined in claim 5 wherein said control of said electric field for pulsed generation of ions includes pulsed gating of said beam of electrons to said turning point and simultaneously pulsing said reversal of said electron beam while said electron beam is gated on, and directing said ions to said utilization device while reversal of said electron beam, and the generation of said electron beam, are pulsed off.

7. A negative ion source comprising:
   - means for generating a beam of electrons, electrodes in the direct path of said beam;
   - means for biasing said electrodes for reversal of said electron beam to produce at the turning point electrons having a distribution of energies centered at zero;
   - means for introducing a gas containing thermal electron-attaching molecules at a point in the path of said electron beam where said electrons have low energy and a high probability for attachment to said molecules, thereby efficiently generating ions; and
   - means for directing said ions in a focused beam to a utilization device.

8. Apparatus as defined in claim 7 for generating negative ions from thermal electron-attaching molecules having an extremely large attachment cross sec-
7. Apparatus as defined in claim 7 for generating negative ions from thermal electron-attaching molecules having an attachment cross section at a level near zero wherein said gas of said molecules is introduced at a point in the path of said electron beam between said electron gun and said turning point, whereby said electrons of the appropriate energy make two passes through said gas of said molecules for enriched negative ion generation.

8. Apparatus as defined in claim 7 wherein said biasing means is controlled with constant voltages for continuous generation of ions.

9. Apparatus as defined in claim 7 wherein said biasing means is controlled for pulsed generation of ions.

10. Apparatus as defined in claim 11 wherein said control of said biasing means for pulsed generation of ions is comprised of means for pulsed gating of electrons to said turning point and simultaneously pulsing said electrodes for reversal of said electron beam while said electron beam is turned on, and turning on said means for directing said ions to said utilization device while said electrodes for reversal of said electron beam, and said electron beam, are pulsed off.