REALIZATION OF THICK HYBRID TYPE CARBON STRIPPER FOILS WITH HIGH DURABILITY AT 1800K FOR RCS OF J-PARC

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Abstract

The Japan-Proton Accelerator Research Complexes (J-PARC) requires thick carbon stripper foils (250-500 μg/cm²) to strip electrons from the H- beam supplied by the linac before injection into the RCS (Rapid Cycling Synchrotron). The 200 MeV H- beam from the linac has a pulse length of 0.5 ms with a repetition rate of 25 Hz and an average beam current of 335 μA. For this high-energy and high-intensity beam, conventional carbon stripper foils will break in a very short time and even a diamond foil will be ruptured at around 1800 K by the MW class accelerator. Thus, thick carbon stripper foils with high durability at 1800 K produced by energy deposition in the foil are indispensable for this accelerator. For this purpose, we have been developing carbon stripper foils of 350 μg/cm² by means of both the controlled DC and AC/DC arc-discharge methods. Recently, we have successfully developed hybrid type thick boron doped carbon stripper foils, which showed a drastic improvement not only with respect to the lifetime, but also with respect to thickness reduction and shrinkage at high temperature during long beam irradiation. In this report the preparation procedure and lifetime measurements with a 3.2MeV, Ne+ beam are presented.

INTRODUCTION

Figure 1 shows the layout of the RCS with the injection and extraction beam transport line. A 200 MeV (first stage) H- beam from the J-PARC linac is stripped directly to H+ by a 300±50 μg/cm² thick carbon foil before injection into the RCS, in which the H- beam is post-stripped by carbon foils placed at three different positions. At the 1st position, the primary H- beams are converted to H+ with an efficiency of about 99.6 - 99.7%, and the unconverted fraction of the beam can be converted H+ at the 2nd and the 3rd position and transported to the beam dump. Finally unconverted H- beams are also transported to the beam dump by a magnet. The lifetime of the carbon stripper foils strongly depends on the beam current intensity and density. Due to the energy loss of the H- beam in the carbon foil at the injection stage and due to collisions with the circulating bunched beam, which is much more intense than the H- injection beam, the temperature of the stripper foils

Fig.1 Schematic layout of the accelerator complex of J-PARC

The properties of the injection beam for the 3 GeV ring of the J-PARC are listed in Table 1. These data are relevant for the development of desired carbon stripper foils.

For comparison, we calculated the energy deposition as a function of foil thickness for a low energy Ne++ DC beam of 3.2MeV and high-energy proton beams of 200 and 400MeV. The results are shown in Fig. 2.

Table 1: Injection Beam and Stripper Foil Properties of

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
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<tbody>
<tr>
<td>Kinetic energy</td>
<td>200 MeV (first stage) 400 MeV (second stage)</td>
</tr>
<tr>
<td>Beam pulse length</td>
<td>0.5 ms</td>
</tr>
<tr>
<td>Repetition rate</td>
<td>25 Hz</td>
</tr>
<tr>
<td>Average beam current</td>
<td>0.335 mA</td>
</tr>
<tr>
<td>Beam size</td>
<td>10 mm x 10 mm</td>
</tr>
<tr>
<td>Foil thickness</td>
<td>250 - 500 μg/cm²</td>
</tr>
<tr>
<td>Foil peak temperature</td>
<td>&gt; 1800 K</td>
</tr>
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</table>

Figure 2: Energy deposition as a function of foil thickness for 200, 400 MeV and Ne+ DC beam.
The energy deposition of the H\textsuperscript{-} injected beam is only about 0.5 W/cm\textsuperscript{2}. Hence, most of energy deposition is originating from the circulating beams. As can be deduced from Fig. 2, the thermal power of the 3.2 MeV Ne\textsuperscript{+} DC beam is nearly same as those by 200 and 400 MeV proton beams. Thus it seems justified to use a 3.2 MeV Ne\textsuperscript{+} DC beam in the lifetime measurements of the stripper foils.

During irradiation of carbon stripper foils with pulsed 200 and 400 MeV H\textsuperscript{-} ions of high current density, the peak temperature of the foils with 250-500 μg/cm\textsuperscript{2} thickness may rise above 1800 K and the limiting effect on the lifetime might be thinning due to evaporation until burning of a hole. In order to solve these issues, there are mainly two ways. The one is to use a diamond foil [1], and the other is to apply the new laser stripping technique [2]. The diamond foil might be a very attractive candidate for ion stripping due to its drastically high thermal conductivity (five times higher, compared to copper). In our investigations, however, the foil ruptured, may be, by changing into graphite structure above 1800±100 K [3]. The laser stripping technique is also an attractive method. It is, however, presently in an early stage of development. Hence, the development of long-lived type of carbon stripper foil with high durability at 1800 K is one of the key technologies for the 3 GeV proton beam at the RCS of J-PARC.

**EXPERIMENTAL RESULTS**

For this purpose, we have prepared cluster carbon stripper foils of 130 μg/cm\textsuperscript{2} thickness by means of the controlled AC/DC arc-discharge (CADAD) method, as a test preparation [4]. Although the foils had a very long lifetime, we found that the foil thickness reduced to about 25% of the original thickness and pinholes were observed at the irradiated area. In addition, the maximum accessible foil thickness turned out to be 170 ± 30 μg/cm\textsuperscript{2} due to lack of adhesion to the substrate. To overcome these problems, we succeeded in developing a boron mixed type of carbon stripper foil of 250-380 μg/cm\textsuperscript{2} accessible thickness. Comparative measurements of the lifetime and other properties of this hybrid type stripper foils as well as of the diamond foils and of the best available commercial foils were performed with a 3.2 MeV Ne\textsuperscript{+} DC ion beam. This carbon discharge-arc evaporation source was installed in a new vacuum chamber (EBX-2000C) of dimensions 750 x 750 x 900 mm\textsuperscript{3}. It is equipped with a cryopump of 5000 l/s pumping speed for N\textsubscript{2}, which produces a basic vacuum of 4·10\textsuperscript{-6} Pa in the empty chamber. The distance between the evaporation source and the substrate was 180 mm. The principal axis of both electrodes was tilted by 45° against the plane of the substrate holder. As mentioned in ref. [4-6], the lifetime of the foil was found to be correlated to the expression R=\(W_{c}(W_{c}+W_{a})^{-1}\) calculated in %, where \(W_{c}\) and \(W_{a}\) are the carbon source weight losses due to ablation from the cathode and the anode electrodes, respectively. In the present preparation, the ratio R was kept between 60 and 70 %. The layer thickness was controlled with a quartz thickness monitor and measured after deposition. For carbon layer deposition, the pressure in the vacuum chamber was 8·10\textsuperscript{-5} to 2·10\textsuperscript{-4} Pa with DC arc power on, when the arc was off, the pressure was 3·10\textsuperscript{-5} Pa. Stripper foils prepared by this method are referred to as HBC-foils (Hybrid Boron mixed Carbon stripper foils).

Fig.3 shows photographs of two different carbon deposits made by the CADAD method. Fig. 3(a) represents a HBC-foil of 320 μg/cm\textsuperscript{2}, while Fig. 3(b) depicts a cluster foil of approximately 180 μg/cm\textsuperscript{2} with R=75%.

The foil in Fig. 3 (a) shows strong adherence to the substrate. It was deposited on the blank substrate without using a release agent, because HBC-foils deposited on glass substrates coated with a release agent tend to peel off at a thickness of about 50μg/cm\textsuperscript{2}. The foil in Fig. 3(b), however, apparently shows bad adherence to the substrate.

Fig.3 Photographs of carbon layers on slide glasses: (a) represents an HBC-foil of 320 μg/cm\textsuperscript{2} and (b) is for a Cluster –foil of 180 μg/cm\textsuperscript{2} made by the CADAD method.

Self-supported foils were obtained by applying an annealing technique. Here, a carbon layer deposited on a glass slide of 26 x 74 mm\textsuperscript{2} was fixed to a SUS-304 target holder which was placed at a distance of 30 mm from a Ta heater source. Subsequently it was heated to a temperature of about 680 K for 7 h in a vacuum of approximately 10\textsuperscript{-5} Pa. During this annealing, the foils were separated from the substrate, keeping a flat shape. The lifetime measurements were performed with a 3.2 MeV Ne\textsuperscript{+} DC beam of 2.5±0.5 μA and 3.5 mm beam spot diameter, supplied from the Van de Graff accelerator at the Tokyo Institute of Technology. In this case, the lifetime was determined as a total integrated irradiation dose ( mC/cm\textsuperscript{2}) until foil rupture occurred assuming a constant current during irradiation. We investigated four foils of the HBC type of 200-380 μg/cm\textsuperscript{2}, three diamond foils (DM-foils) of 350 μg/cm\textsuperscript{2}, two diamond foils of 680 and 780 μg/cm\textsuperscript{2}, and, for comparison, four commercially available thick foils (CM-foils) of 200-400 μg/cm\textsuperscript{2}. The diamond film was deposited by Kobe-Steel company and Sumitomo Electric Industries, ltd. in Japan on a 0.5 mm thick Si substrate. It had to be prepared self supporting by chemically etching the Si substrate except for an approximately 18 mm outer circle frame, which survived etching due to coating with a Kapton film (Fig. 4(b)).
Figure 4: Photographs of a HBC-foil (a) (340 μg/cm²) and a diamond foil (b) on its Si ring–frame at different stages of ion irradiation.

In this lifetime measurement, we used the ribbon type foils of 20 mm x 30 mm and the foils were mounted on an Al holder (38 mm x 58 mm), as shown in Fig. 4. We observed the state of shrinkage, brittle and rupture from a viewing port. Table 2 shows the results of the lifetime measurements.

Table 2: Maximum and average lifetime of HBC-foil, DM-foil and CM-foil measured with a 3.3 MeV, Ne⁺ ion beam of 3 μA on a 3.5 mm diameter beam spot

<table>
<thead>
<tr>
<th>Type of foils</th>
<th>HBC-foil</th>
<th>DM-foil</th>
<th>CM-foil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. lifetime (mC/cm²)</td>
<td>9800</td>
<td>104</td>
<td>24</td>
</tr>
<tr>
<td>Average (mC/cm²)</td>
<td>5600</td>
<td>88</td>
<td>20</td>
</tr>
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</table>

From the Fig. 4(a), the HBC-foil did not show any shrinkage even after a long time irradiation of approximately 9800 mC/cm² at temperature of 1600±200 K. The thickness of the foil was reduced, but not by more than 20 % of the original thickness, as measured with an α particle thickness gauge [7]. The tested DM-foil in Fig.4(b) showed a little pale dark color after a few minutes of irradiation, and it became a deeply dark color with more than irradiation time, and then it was broken at approximately 104 mC/cm² at the irradiated area. The temperature of the foil during irradiation was approximately 1700±100 K. In this case, the foil did not show any shrinkage until the foil was broken. The CM-foil showed a strong shrinkage after approximately 20 mC/cm², it continued to shrink, and then the foil broke by making a hole. The maximum temperature was 1600±200 K.

**SUMMARY**

We have successfully developed HBC-foils using the CDAD method, and the HBC-foils showed the following great improvements:

1. Strong adhesion to the substrate and thus the foils could be prepared with thickness up to a maximum of 480 μg/cm².
2. Noticeably less shrinkage than for foils made by the CADAD method, and thickness reduction less than 20 %, even after long-time irradiation at the high temperature of 1600±200K.
3. A maximum lifetime of a HBC-foil of 9800 mC/cm² corresponding to approximately 408 times longer or 94 times longer than those of the best CM foils or high quality diamond (DM) foils, respectively.

In conclusion, this method is a simple and promising one for production of long-lived carbon stripper foils thicker than 200 μg/cm² for use in J-PARC.

**REFERENCES**