Current research activities in the field of multilayers for soft X-rays in Japan

T. Namioka

To cite this version:


HAL Id: jpa-00245999
https://hal.archives-ouvertes.fr/jpa-00245999
Submitted on 1 Jan 1988

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Current research activities in the field of multilayers for soft X-rays in Japan

T. Namioka

Research Institute for Scientific Measurements, Tohoku University, Sendai 980, Japan

(Reçu le 29 septembre 1987, accepté le 8 février 1988)

Résumé. — Les programmes en cours de développement au Japon sont passés brièvement en revue. Les thèmes d'études examinés portent sur les principes des optiques envisagées et sur quelques applications, les études de constantes optiques X-UV des substrats et des films minces ainsi que sur les techniques d'évaporation.

Abstract. — The present status of studies on soft X-ray multilayers in Japan is briefly reviewed. This includes the design concepts, optical constants of substrates and thin films, fabrication techniques, evaluation methods, and some applications.

1. Introduction.

Studies of multilayers in Japan are relatively new in the field of soft X-ray optics, in striking contrast to extensive research in the field of semiconductors. Our current efforts have thus been directed more to basic research, both academic and technical, than applications, though those researches are all aimed at specific applications.

This paper describes, as a representative example, the current activities of several Japanese research groups in the field of multilayers for soft X-ray optics. This includes various design concepts, examples of designed multilayers, measurement of optical constants, fabrication of substrates and multilayers, evaluation of roughness, entrapped Ar, crystal structure, and reflectance, and applications to X-ray lithography and X-ray telescopes. The groups participating in this paper are listed below.

Tohoku group: T. Namioka, M. Yamamoto, M. Yanagihara, and A. Arai, Research Institute for Scientific Measurements, Tohoku University (Katahira, Sendai 980);
Photon Factory (PF) group: T. Matsushita, A. Iida, and T. Ishikawa, Photon Factory, National Laboratory for High Energy Physics (Oho, Tsukuba-shi, Ibaraki 305);
Yoshida Nano-Mechanism Project (YNMP) group: S. Yoshida, H. Nagata, and Y. Suzuki, JRDC (2-5-2 Marunouchi, Chiyoda-ku, Tokyo 100);
NTT group: Y. Ishii, H. Takenaka, and H. Takaoka, NTT Electrical Communication Laboratory (3-9-11 Midori-machi, Musashino-shi, Tokyo 180);
H. Kinoshita, T. Kaneko, H. Takei, N. Takeuchi, and T. Ishihara, NTT Atsugi Electrical Communication Laboratory (1839 Ono, Atsugi-shi, Kanagawa 243-01);
Canon group: S. Ogura and Y. Watanabe, Canon Research Center, Canon Inc. (5 Morinosato-Wakamiya, Atsugi-shi, Kanagawa 243-01);
Nagoya group: F. Nagase, H. Kunieda, Y. Tawara, and T. Kii, Dept. of Astrophysics, Nagoya University (Chikusa-ku, Nagoya 464);
Chubu group: Y. Namba, Chubu University (Kasugai, Aichi 487);
Kyoto group: T. Shinjo, N. Nakayama, and I. Moritani, Institute for Chemical Research, Kyoto University (Uji, Kyoto 611);
Osaka group I: K. Yamashita, H. Tsunemi, S. Kitamoto, I. Hatsuakde, A. Miyake, and Y. Ueno, Dept. of Physics, Osaka University (Toyonaka, Osaka 560);
Osaka group II: Y. Kato, H. Shiraga, K. Shihoyama, and T. Endoh, Institute of Laser Engineering, Osaka University (Suita, Osaka 565);
2. Design of multilayers.

2.1 PERIODIC MULTILAYERS. — In designing multilayers the first thing we have to do is to choose a proper pair of materials for a specific purpose. YNMP group, who is interested in developing X-ray microscopes and X-ray lithography devices for wavelengths of 10-50 Å, calculated the saturated reflectance at normal incidence of periodic multilayers, consisting of various pair materials A and B, by using formulas based on Fresnel equations [1-3] with the optical constants derived from the data of Henke et al. [4]. The result obtained for a wavelength of 25.2 Å is summarized in Table I.

Table I shows that combinations of any material in the group of Ni, Co, Cu, Fe, Re, Zn, W, Mn, Ta and Au with any one in the group of Be, Mg, Sn, Sb, V and Te would provide relatively high reflectance and that among them the Ni/Be combination would give the highest reflectance of 44.5%. It is interesting to note in the table that the multilayers containing V (L absorption edge at 24.03 Å), Sn (M edge at 24.03 Å), or Sb (M edge at 23.44 Å) show relatively high reflectances at 25.2 Å, even though their atomic numbers are not small. This is due to the effect of anomalous dispersion, which causes a large difference in the refractive indices of the two materials and reduces the amount of absorption.

Figure 1 shows some examples of the enhancement in reflectances due to this effect. The spectral reflectances of all the multilayers at normal incidence were calculated by keeping the total thickness of the multilayers to a constant value of 3 660 Å. This clearly illustrates the usefulness of a material whose absorption edge lies at a wavelength slightly shorter than the design wavelength. An experimental proof of this idea is given in section 5.2.

With a pair of materials thus chosen, a further increase in the reflectance would be possible by controlling the density of the materials. Figure 2 compares the calculated reflectances at the 45-deg. angle of incidence of a 128-pair W/C multilayer with and without density control in the C layers. The carbon density is assumed to be 80% of the bulk value in the controlled case. As is seen in the figure, the peak reflectance of the density-controlled multilayer is 1.3 times higher than that of the non-controlled one. Similar results were obtained for multilayers of other combinations in different wavelength ranges, suggesting the possibility of increasing the reflectance by means of density control.

Table I. — Saturated reflectances (%) of multilayers composed of two materials A and B. Reflectances are computed for a wavelength of 25.2 Å (courtesy of YNMP group).

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mn</td>
<td>Fe</td>
<td>Co</td>
<td>Ni</td>
<td>Cu</td>
<td>Zn</td>
<td>Ga</td>
<td>As</td>
<td>Nb</td>
<td>Mo</td>
<td>Tc</td>
<td>Ru</td>
<td>Pd</td>
<td>Hf</td>
<td>Ta</td>
<td>W</td>
<td>Re</td>
</tr>
<tr>
<td>Be</td>
<td>31.9</td>
<td>37.1</td>
<td>42.5</td>
<td>44.5</td>
<td>40.4</td>
<td>34.0</td>
<td>25.4</td>
<td>22.7</td>
<td>24.1</td>
<td>25.8</td>
<td>28.1</td>
<td>26.6</td>
<td>24.8</td>
<td>26.7</td>
<td>31.3</td>
<td>33.9</td>
<td>35.5</td>
</tr>
<tr>
<td>V</td>
<td>28.9</td>
<td>32.3</td>
<td>36.2</td>
<td>37.7</td>
<td>33.7</td>
<td>28.6</td>
<td>22.0</td>
<td>19.2</td>
<td>17.6</td>
<td>18.6</td>
<td>20.2</td>
<td>18.5</td>
<td>16.9</td>
<td>19.9</td>
<td>23.5</td>
<td>25.7</td>
<td>27.1</td>
</tr>
<tr>
<td>Sr</td>
<td>20.7</td>
<td>22.4</td>
<td>19.1</td>
<td>14.4</td>
<td>9.2</td>
<td>7.5</td>
<td>8.0</td>
<td>9.1</td>
<td>10.7</td>
<td>9.7</td>
<td>8.5</td>
<td>9.6</td>
<td>12.9</td>
<td>14.9</td>
<td>16.2</td>
<td>12.9</td>
<td>6.8</td>
</tr>
<tr>
<td>Sn</td>
<td>34.9</td>
<td>36.6</td>
<td>38.8</td>
<td>39.6</td>
<td>35.8</td>
<td>31.9</td>
<td>27.1</td>
<td>23.7</td>
<td>18.5</td>
<td>18.6</td>
<td>19.3</td>
<td>17.2</td>
<td>15.9</td>
<td>20.5</td>
<td>23.2</td>
<td>24.7</td>
<td>25.9</td>
</tr>
<tr>
<td>Sb</td>
<td>30.7</td>
<td>33.1</td>
<td>36.0</td>
<td>37.1</td>
<td>33.3</td>
<td>28.8</td>
<td>23.3</td>
<td>20.3</td>
<td>16.8</td>
<td>17.3</td>
<td>18.4</td>
<td>16.5</td>
<td>15.2</td>
<td>18.9</td>
<td>21.9</td>
<td>23.8</td>
<td>25.1</td>
</tr>
<tr>
<td>Te</td>
<td>28.3</td>
<td>31.4</td>
<td>34.4</td>
<td>35.7</td>
<td>31.8</td>
<td>27.1</td>
<td>21.3</td>
<td>18.4</td>
<td>15.9</td>
<td>16.6</td>
<td>17.9</td>
<td>16.2</td>
<td>14.8</td>
<td>18.0</td>
<td>21.3</td>
<td>23.2</td>
<td>24.6</td>
</tr>
<tr>
<td>La</td>
<td>13.7</td>
<td>18.9</td>
<td>21.4</td>
<td>18.0</td>
<td>12.5</td>
<td>6.5</td>
<td>5.5</td>
<td>8.3</td>
<td>10.0</td>
<td>12.1</td>
<td>11.3</td>
<td>10.0</td>
<td>10.0</td>
<td>13.8</td>
<td>16.3</td>
<td>17.7</td>
<td>14.3</td>
</tr>
</tbody>
</table>

© N° 10 REVUE DE PHYSIQUE APPLIQUÉE
2.2 APERIODIC MULTILAYERS. — Tohoku group has developed a new design method of multilayers utilizing a Gaussian-plane plot of the complex amplitude reflectance [5]. Reformulating a well known recurrence formula [1], they derived the following formula for the complex amplitude reflectance $R_m$ of a $m$-layer system in a vacuum:

$$R_m = \frac{r_m(1 - r_m R_{m-1}) + (R_{m-1} - r_m) \exp(-i \delta_m)}{1 - r_m R_{m-1} + r_m(R_{m-1} - r_m) \exp(-i \delta_m)},$$

where

$$\delta_m = 4 \pi n_m d_m \cos \phi_m / \lambda,$$

$$R_0 = r_0 \text{ (for the substrate)},$$

$r_m$ is the Fresnel reflection coefficient of the $m$-th layer material with respect to the vacuum, $n_m = n_m - i k_m$ the complex refractive index of the $m$-th layer material, $d_m$ the thickness of the $m$-th layer, $\phi_m$ the angle of refraction into the $m$-th layer, and $\lambda$ the wavelength of incident light.

A graphical representation of equation (1) enables one to visualize the changes in the amplitude reflectance with the growth of layered structure, giving a clear insight into the multilayer design. A typical example of such a representation in the complex plane is given in figure 3. Points $r_s$, $r_A$, and $r_B$ represent the Fresnel reflection coefficients of the substrate, material A, and material B with respect to the vacuum respectively. Suppose we start, in figure 3, deposition of A on the substrate. As the deposition proceeds, i.e., as the layer thickness $d_i$ increases, the amplitude reflectance $R_1(d_1)$ starts to move from $r_S$ and spirals in toward $r_A$, producing...
the reflectance maxima and minima alternately. To increase the reflectance it is necessary to switch the deposition of A over to that of B at a certain stage of growth. If this is done, then the amplitude reflectance \( R_2(d_2) \) of the bi-layer spirals in toward \( r_B \) as the thickness \( d_2 \) of the second layer increases, and its course depends solely on its starting point, at which the deposition of A was switched over to that of B (see Fig. 3). In view of this, one can see that a highly reflective multilayer can be realized effectively by choosing switching points properly so as to make a series of amplitude reflectance curves to expand outward most efficiently as the multilayer structure grows. It can be shown that the optimum multilayer structure that gives the highest reflectance for a given pair of materials is realized by switching the deposition of materials at the points where the successive amplitude reflectance curves connect smoothly.

Table II compares, for a pair of Re and Al, the reflectance of an optimum aperiodic multilayer with that of an optimized periodic multilayer as a function of the number of layers. The reflectance increases much faster for the aperiodic structure as compared with the periodic one, especially for the first 11 layers or so. The difference, however, becomes smaller as the number of layer increases.

Table II. — Comparison of calculated reflectances of an optimum aperiodic Re/Al multilayer with those of an optimized periodic one.

<table>
<thead>
<tr>
<th>m</th>
<th>( d (\text{Å}) ) Re</th>
<th>( R(% ) )</th>
<th>( d (\text{Å}) ) Al, Re</th>
<th>( R(% ) )</th>
<th>( \Delta R )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>28.3</td>
<td>6.01</td>
<td>72.4</td>
<td>12.68</td>
<td>6.97</td>
</tr>
<tr>
<td>2</td>
<td>28.3</td>
<td>6.01</td>
<td>72.4</td>
<td>12.68</td>
<td>6.97</td>
</tr>
<tr>
<td>3</td>
<td>85.6, 28.3</td>
<td>17.17</td>
<td>78.5, 72.4</td>
<td>29.79</td>
<td>12.62</td>
</tr>
<tr>
<td>4</td>
<td>ditto</td>
<td>17.17</td>
<td>78.5, 72.4</td>
<td>29.79</td>
<td>12.62</td>
</tr>
<tr>
<td>5</td>
<td>28.5</td>
<td>6.01</td>
<td>72.4</td>
<td>12.68</td>
<td>6.97</td>
</tr>
<tr>
<td>6</td>
<td>ditto</td>
<td>17.17</td>
<td>78.5, 72.4</td>
<td>29.79</td>
<td>12.62</td>
</tr>
<tr>
<td>7</td>
<td>37.9</td>
<td>45.27</td>
<td>76.1, 37.3</td>
<td>54.10</td>
<td>8.83</td>
</tr>
<tr>
<td>8</td>
<td>72.7</td>
<td>45.27</td>
<td>76.1, 37.3</td>
<td>54.10</td>
<td>8.83</td>
</tr>
<tr>
<td>9</td>
<td>45.27</td>
<td>76.1, 37.3</td>
<td>54.10</td>
<td>8.83</td>
<td></td>
</tr>
</tbody>
</table>


It is extremely important in the design of multilayers to have accurate optical constants of materials in the soft X-ray region. An effort has, therefore, been made by Tohoku group to measure the optical constants of substrate materials and of thin films, using synchrotron radiation at the Photon Factory.

3.1 Optical constants of substrate materials. — Yanagihara et al. [6] designed and installed a high-precision UHV reflectometer on beamline 11A at the Photon Factory. Figure 4 is a schematic of the reflectometer. The reflectometer is constructed in such a way that alignment of the rotation axes of the sample and the detector remains unaffected when it is evacuated and/or baked. The accuracy of the sample rotation and detector rotation was found to be better than 30 s of arc. Reflectance measurement can be done at any angle of incidence in a range 9.6-89.4 deg.

Using the reflectometer together with a 2-m grasshopper monochromator, Yanagihara et al. measured the reflectances of polished CVD-SiC (samples I and II), CVD-TiC, and sintered-WC mirrors at photon energies of 80-1 200 eV. In the analysis the Debye-Waller factor was taken into consideration. In figure 5 is shown a typical example of \( R-\theta \) plots \( (R : \text{reflectance}, \theta : \text{angle of incidence}) \) obtained for the CVD-SiC mirror (sample I). Errors in the observed reflectances are less than 3 %. The optical constants \( \delta (= 1 - n) \) and \( k \) were determined by means of curve fitting to the measured reflectance curves (see solid lines in Fig. 5). Their values are listed in table III together with the rms surface roughness \( \sigma \) in the Debye-Waller factor. These \( \sigma \)
Fig. 5. — Measured reflectance as a function of incidence angle for polished CVD-SiC (sample I) at $h\nu = 1000$ eV ($\bullet$), 700 eV ($\circ$), 500 eV ($\Delta$), and 350 eV (+). Solid lines are calculated values (after Ref. [6]).

Table III. — Optical constants of SiC in the soft X-ray region. $\sigma$ is the rms roughness (after Ref. [6]).

<table>
<thead>
<tr>
<th>$E$ (eV)</th>
<th>$\delta = 1 - n$</th>
<th>$k$</th>
<th>$\sigma$ (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample I</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>80</td>
<td>0.0337</td>
<td>1.28 E-2</td>
<td>33.8</td>
</tr>
<tr>
<td>100</td>
<td>0.00718</td>
<td>4.09 E-3</td>
<td>18.1</td>
</tr>
<tr>
<td>120</td>
<td>0.00753</td>
<td>1.58 E-2</td>
<td>28.3</td>
</tr>
<tr>
<td>150</td>
<td>0.0116</td>
<td>1.39 E-2</td>
<td>28.0</td>
</tr>
<tr>
<td>200</td>
<td>0.00934</td>
<td>6.47 E-3</td>
<td>27.5</td>
</tr>
<tr>
<td>250</td>
<td>0.00729</td>
<td>5.03 E-3</td>
<td>29.6</td>
</tr>
<tr>
<td>Sample II</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>350</td>
<td>0.00488</td>
<td>2.33 E-3</td>
<td>22.9</td>
</tr>
<tr>
<td>500</td>
<td>0.00253</td>
<td>6.45 E-4</td>
<td>18.0</td>
</tr>
<tr>
<td>700</td>
<td>0.00129</td>
<td>2.06 E-4</td>
<td>15.0</td>
</tr>
<tr>
<td>1000</td>
<td>0.00062</td>
<td>4.30 E-5</td>
<td>17.2</td>
</tr>
</tbody>
</table>

values obtained at different photon energies are consistent with one another for the same sample and the averaged value of 28 Å for sample II is in good agreement with a value of 23 Å measured with a WYKO profiler. This result, together with those for the CVD-TiC and WC mirrors, suggests that the simple diffractive scattering model is applicable even to the case of specular reflections of soft X-rays from rough surfaces that violate the assumption $\lambda \gg \sigma$. The reason for this is not clear at the present.

Tohoku group [7] determined also the optical constants of polished BK7 glass in a wavelength range 120-200 Å using the same reflectometer.

3.2 Optical constants of thin films. — The optical constants of various materials in the form of thin film ($\sim 300$ Å) are of great importance not only in the design of multilayers, but also in the study of the dependence of the optical constants on deposition conditions and methods and in the study of roughness, interdiffusion, etc.

Figure 6 shows the optical constants of Mo film prepared by Tohoku group using an ion-beam sputtering system. The reflectance data used for the analysis were taken for s-polarization at the Photon Factory. In the analysis, the measured data of reflectance vs. wavelength were compiled in the form of reflectance vs. angle of incidence for every wavelength sampled. The optical constants were, then, determined by means of least-squares curve fitting by taking multiple reflections into account. The film thickness measured by the Tolansky method was used in the initial phase of analysis, and then the film thickness was finely adjusted, within the possible errors in the thickness measurement, until it gave the best fit for all the data available to the analysis.

In figure 6, two published data points, one at 124 Å (100 eV) by Henke et al. [4] and the other at 210 Å by Lynch and Hunter [8], are shown by solid circles. The optical constants used in the simulation of reflectance curves by Barbee et al. [9] are also shown at a wavelength of 170 Å. A weak structure is clearly seen at about 140 Å, whose existence was not known to date. Tohoku group has so far accumulated reflectance data at 9.6-88 deg. angles of incidence over a wavelength range of 225-43 Å for single layers of Mo, C, Al, Ni, Rh, Ru, LiF, and B prepared by UHV electron-beam deposition, and of Re, Mo, Si, C, Au, Ni, Rh, Ru, and BN by ion-beam sputtering. Analyses of their optical constants and surface roughness are now under way.

Fig. 6. — Optical constants ($n - ik$) of Mo film in the XUV region. The solid circles show the values given in references [8] and [9].
4. Fabrication of multilayer mirrors.

4.1 Polishing of substrates. — It is needless to say that low-scatter supersmooth optical surfaces are essential in the fabrication of multilayer mirrors for soft X-rays. Bowl feed polishing [10] is used by Tohoku group, and surface finish of about 5 Å rms roughness on BK7 glass has been obtained [11]. Namba of Chubu group invented a better polishing technique called float polishing [12, 13] and constructed special machines [14, 15].

The essential parts of the new float-polishing machine are schematically shown in figure 7. The machine is capable of polishing samples of up to 180 mm in diameter. While the machine is running, the polishing fluid rushes through the small gap between the lap, made of tin, and the workpiece, producing polishing action. The polishing fluid consists of 2 wt. % of 70 Å-diam fumed silica powder (Aerosil 300, Degussa Corp.) in filtered deionized- or distilled-water. The key to successful float polishing is the flatness of the tin lap. Since the flatness of the lap is transferred on the surface of the workpiece, the lap should be flat to about 1 μm over its entire 46-cm diameter. The micro-grooves on the lap surface also help the polishing fluid to come in contact with the workpiece.

Figure 8 shows the change in flatness of a 100-mm diam Zerodur sample before, during, and after float polishing. This figure illustrates the effectiveness of float polishing. The flatness was measured with a Zygo Mark III interferometer. The surface roughness of a number of samples thus polished has been measured with a Nomarski interference microscope, optical heterodyne profilometer [16], digital profiler [17] (WYKO TOPO-2D), Talystep, laser scatterometer, and X-ray reflectometer [11]. The results of all the measurements, which are based on different principles, are in good agreement with one another, giving the order of 1-2 Å rms surface roughness. Figure 9 shows an example of the surface roughness of a float-polished glass-ceramic sample as measured with a Talystep [18]. An rms roughness of 1.9 Å over a distance of 0.6 mm is obtained, with an instrumental noise of about 1 Å rms. The best rms roughness so far obtained is 0.8 Å rms over a distance of 1 mm on a sapphire single crystal [15].

4.2 Fabrication of multilayer mirrors. — UHV electron-beam deposition systems and sputtering systems are currently in use in Japan for the preparation of multilayers. The former is used by Kyoto, Osaka I, and Canon groups and the latter by NTT and Tohoku groups.

Fig. 7. — Principal structure of float polishing machine (after Ref. [14]).

Fig. 8. — Flatness change of a 100-mm Zerodur sample in the float polishing process (after Ref. [14]).

Fig. 9. — Surface profile of a float-polished glass-ceramic sample, measured with a Talystep (after Ref. [14]).
Kwansei-Gakuin group is looking into the possibility of epitaxial growth of metallic multilayers by molecular beam epitaxy (MBE), in spite of practical difficulties associated with MBE such as substrate heating that might deform the surface of superpolished substrates, a narrow choice in depositing material pairs and substrates that fulfill epitaxial relations. They have prepared by MBE Nb(20 Å)/Ta(30 Å) multilayers with 60 periods on several kinds of sapphire substrates.

YNMP group is setting up a magnetron sputtering and a photo-CVD system.

A) Electron-beam deposition.

Kyoto group has prepared many metallic multilayers containing 3d transition metals, such as Fe/Sb, Fe/V, Fe/Mg, Co/Sb, Fe/C, Mn/Sb, Fe/Mn, Fe/Dy, Ag/V, V/Si, etc. [19], by using a UHV electron-beam deposition system [20] (see Fig. 10). Multilayers were formed on cooled substrates (−50 °C), made of glass, mylar, etc., in the ultra-high vacuum (~10−9 Torr). The samples deposited on glass, mylar, and triacetyle cellulose were used for measurements of X-ray diffraction and reflection, Mössbauer absorption and ferromagnetic resonance, and electron microscopy, respectively. Programmable quartz-oscillator thickness-monitors were employed to monitor the layer thickness and to control two pneumatic shutters. The values of layer thickness observed by the monitor are in good agreement with the values determined from X-ray diffraction measurements. As an example, the periods determined by the two methods are compared in figure 11 for the case of Fe/Mg multilayers [21]. Difficulty was encountered, however, in depositing carbon. The quartz oscillator was heated up by thermal radiation, and this introduced an error in the thickness determination. The thickness of carbon layers had to be estimated after closing the shutter and waiting for a while until the temperature returned to the room temperature. A typical deposition rate of Fe is ~0.2 Å/s. A quadrupole mass analyser and an auger electron spectrometer are also incorporated in the deposition system for analysing the residual gases and the surface contamination.

Deposition systems quite similar to the Kyoto group’s are used by Osaka and Canon groups. Osaka group 1 has made Mo/Si, Mo/C, and Ni/C multilayers on float glass and float-polished glass [12, 13] substrates of 10 x 10 cm² with a rms roughness of 2-3 Å [22]. In their system a movable mask system is mounted immediately below the sample holder so as to permit the preparation of several different multilayers on one substrate without breaking vacuum.

Canon group has prepared Ru/Si multilayers on Si substrates of 2" (diam) x 10 mm (thick) having a peak-to-valley roughness of ~30 Å.

Tohoku group is testing a UHV electron-beam deposition system equipped with an in situ ellipsometric monitor.

B) Magnetron sputtering.

NTT group constructed a magnetron sputtering system, which is capable of coating several substrates of the size 4" in diameter or 70 x 60 mm² and prepared W/C multilayers of Si wafers and Pyrex and quartz substrates. The sputtering system is schematically shown in figure 12. The substrates mounted on the rotating table are alternately exposed to individual well-isolated magnetron sources. The individual sputtering rates and the table rotation speed can be accurately controlled. This system offers the flexibility necessary for the preparation of
multilayers having various layer thicknesses. The individual layer thicknesses are controlled by the deposition rate of the sputtered material, the speed of the rotating table, the source-to-substrate distance, and the open duration of individual shutters. Shield masks are placed between the source and the substrates in order to achieve the uniformity in the layer thickness. The thickness uniformity of a W/C multilayer formed on a 4" Si wafer was examined by a surface profiler along the direction that coincides with a diametrical direction of the rotating table. The thickness was found to be uniform within ±0.4% along the direction measured.

C) Ion beam sputtering.
Tohoku group has prepared Au/C, Mo/Si, Ni/C, Re/C, Ru/C, and Rh/C on polished BK7 glass substrates of 40-mm diam by using an ion beam sputtering system equipped with an in situ ellipsometric monitor [23]. The sputtering system is schematically shown in figure 13. Argon ions produced in the electron cyclotron resonance gun chamber are accelerated by a grid system and form a uniform ion-shower of 100 mm in diameter. The maximum acceleration voltage and current density are 2 kV and 1 mA/cm², respectively. Ion shower is stopped down to a size to sputter only the target held at an angle of 45 deg. to the beam. Three targets can be mounted on a water-cooled prism-shaped holder. Although deposition rates are very slow (<1 Å/min), the system has the advantage over other sputtering systems: the sample chamber is well isolated from the gun housing and is kept at a pressure of ~3 × 10⁻⁴ Torr.

The in situ ellipsometric monitor consists of a 1-mW He-Ne laser, a Glan-Thompson polarizer (P), two vacuum windows (V.W.) on the sample chamber, a compensator (C), a Glan-Thompson analyser (A), and a photodetector (see Fig. 14). Azimuthal angles of P, C, and A are detected by rotary encoders to 0.005 deg. The in situ extinction measurement at the zone 3 can be made automatically by computer-assisted driving and data-acquisi-
tion systems. The thickness sensitivity of the ellipsometer was experimentally found to be much better than 1 Å: e.g., 0.1 Å for Au, Pd, and Mo, 0.2 Å for W, and 0.8 Å for Ta [23].

The ellipsometer is a sensitive tool for monitoring the thickness and uniformity of layers during the deposition. The solid curve in figure 15 shows changes in ρ, the ratio of the complex amplitude reflectance for the p-component to that for the s-component, which were observed by the ellipsometer during the initial phase of W deposition on a polished BK7 glass substrate. A simulation curve (dashed line) is also drawn in the figure, for comparison. The numerals beside the open circles are the layer thicknesses at the respective points. As is clearly seen in the figure, the simulation curve fits very well to the observed curve down to the point corresponding to ~30-Å thick W layer. This lower-limit thickness d₁ is a good indicator for the formation of a layer which satisfies the assumption of isotropic and parallel-faced layer model. The values of d₁ were found to be 50, 35, 30, 20, and ~0 Å for Au, Pd, W, Mo, and Ta [23]. Figure 16 shows the observed ρ's (solid line) during the formation of a 7-layer Au/C mirror (layer thicknesses of Au and C are 50 Å) together with the simulation curve (dashed line). As is expected from a large value of d₁ for Au, the observed ρ's for the Au layers deviate by a large amount from the simulation curve at the beginning and then approach very close to the simulation points as the thickness of Au comes closer to 50 Å. This indicates the usefulness of ellipsometry in monitoring the formation of multilayers.

5. Evaluation of multilayers.

The overall performance of multilayers for soft X-rays can best be evaluated by their spectral reflectances at the wavelengths of interest. However, spectral reflectance measurements alone do not furnish detailed information on the periodicity, interdiffusion at the interfaces, crystal structure, entrapped gases, surface roughness, interface roughness, etc. In the following subsections are described various methods and techniques employed by the Japanese groups for the characterization of multilayers.

5.1 STRUCTURAL CHARACTERIZATION.

A) X-ray diffraction and scattering.

X-ray diffraction is the most widely used technique in the evaluation of multilayer structure, such as surface roughness, interface roughness, periodicity,
etc. We, therefore, restrict ourselves to describe
here some methods and results of interest only.
To examine the establishment of periodic structure
in a super-thin multilayer, Kyoto group measured X-
ray diffraction patterns of Fe(8 Å)/Mg(24 Å),
Fe(2 Å)/Mg(16 Å), and Fe(4 Å)/Mg(4 Å) mul-
tilayers, using Cu Kα line (1.54 Å) [24]. The result
shown in figure 17 confirms the existence of periodic
structure even in these multilayers, whose Fe layers
are of super-thin. The periods estimated from the
diffraction-peak angles agree well with the designed
periods.

Kwansei-Gakuin group measured X-ray diffrac-
tion patterns of single crystals of a Nb(20 Å)/
Ta(30 Å) multilayer grown by MBE and determined
the epitaxial relations between the component ma-
terials and several kinds of sapphire substrates with
Mo Kα line [25]. Figure 18 shows the intensity
contours with Cu Kα line around the (002) Bragg
reflection from the Nb/Ta multilayer formed on a
sapphire-R (1102) substrate. As is clearly seen in
the figure, the satellite reflections line up in the
direction tilted from the [001] axis, implying that the
Nb and Ta layers have a terraced structure in the
interface region. The interdiffusion region at the
interfaces is estimated to be about 6 Å from the
width of the satellite reflections. The results indicate
that a fairly good metallic multilayer is obtained by
MBE, in spite of fairly large lattice mismatch
between the component materials and the substrate,
as is seen in the figure.

NTT group designed a new triple-axis goniometer
for large area X-ray diffraction topography capable
of utilizing synchrotron X-rays of 3 × 50 mm² and
installed it on the beamline 15B at the Photon
Factory [26]. The precision of the third rotation axis
is better than 0.01 s of arc, which is sufficient for
most plane-wave work. With this goniometer they
carried out preliminary experiments to characterize
the structural uniformity of a 70 × 60-mm²
W(16 Å)/C (24 Å) multilayer with 40 periods,
which was formed on an optically flat quartz sub-
strate by means of magnetron sputtering. The exper-
imental arrangement used is schematically shown
in figure 19. Using a monochromatic X-ray beam of
2 Å, they observed a rocking curve of the sample at
the first-order diffraction. A half width of 135 s of

Fig. 17. — X-ray diffraction patterns in a small angle
region for Fe/Mg multilayers. The layer thicknesses of Fe
and Mg are denoted in the figure (after Ref. [24]).

Fig. 18. — Intensity contours of scattered X-ray around
the Nb-Ta (002) Bragg reflection. The intensity is
expressed in counts. The solid and the dashed line
represent the [001] axis of Nb and Ta and the direction
parallel to the sapphire [1102] axis, respectively (after
Ref. [25]).

Fig. 19. — Schematic of the triple-axis goniometer system
(after Ref. [26]).
arc was obtained. This shows that the period of the multilayer is uniform within ±2.3% over the measured area of 70 × 50 mm².

The measurement of X-ray scattering has been proved to be a good method for evaluating an X-ray mirror surface [27]. Nagoya group extended the method and succeeded not only in a quantitative evaluation of the rms height as a measure of the surface roughness, but also in a measurement of the power spectral density function, which represents the characteristics of surface wavings [11]. They measured the scattering profiles of Al-Kα (8.34 Å) X-rays for plate glass, float glass, polished BK7 glass, and metal films coated on float glass with a 12-m X-ray beam facility at Nagoya University. A schematic diagram of the facility is shown in figure 20. The angular resolution in the scattered beam experiment was 44-s of arc with the second slit of 0.1 mm wide. The power spectral density functions of the surface height distribution for these materials, except for the Au-coated sample, were represented by the power-law spectra with power indices ranging from −1 to −2. The rms heights were found to be 1.8–8.3 Å for the surface wavings with period lengths of 10–100 μm. The results thus obtained were consistent with those measured with a WYKO optical profiler (NCP-1 000). This method was successfully applied also to UPILEX (polyimide produced by Ube Industries, Inc.), Al foil replica backed by epoxy, and Mo/C and Mo/Si multilayers [22] to estimate their surface roughness.

B) Transmission electron microscopy. Transmission electron microscopy has been utilized by NTT and Canon groups, to a varying degree, for the direct characterization of multilayers.

NTT group uses a 400-kV transmission electron microscope to characterize the periodicity and the interfacial structure [28]. Cross-sectional specimens were prepared by means of Ar-ion etching. For precise determination of the periodic length, the specimen was observed by varying its tilt with respect to the electron beam. Electron micrographs thus taken were traced with a microphotometer. The accuracy of period determinations with this method was found to be ±0.15 Å. Figure 21 shows an electron micrograph of a W/C multilayer prepared by magnetron sputtering and its microphotometer trace. From this a mean period of 88.6 Å was obtained with a fluctuation of ±1.9 Å. This value agrees quite well with the designed value of 90 Å.

A detailed observation of the interfacial structure was carried out using thin (~100 Å) specimens. Figure 22 shows a cross-sectional electron micrograph of a thinned W(100 Å)/C(60 Å) multilayer. Fluctuations were observed at the interfaces of Si substrate and C layer and of W and C layers. The average fluctuations were ±1.3 Å with a period of 50 Å at the Si-C interface and ±2.3 Å with a period of 50 Å at the W-C interfaces. Fluctuations at the Si-C interface is probably due to the surface roughness of the Si substrate.

Crystallization of the component layers was further investigated by observing lattice images. Crystallization was not observed in the C layers of 20–1000 Å thick. Crystal grains were observed in the W layers of 100–1000 Å thick. Electron diffraction patterns of these W layers exhibited the same characteristics as those of bulk tungsten. In W layers of 30–60 Å thick, however, no crystal grain was observed. Electron diffraction patterns of these layers showed crystalline patterns, suggesting the existence of fine crystalline particles in these thin layers.

Fig. 20. — Schematic of the 12-m X-ray beam facility at Nagoya University (after Ref. [11]).
It was found that with a further decrease in the layer thickness tungsten became amorphous and fine crystalline particles were dispersed in the amorphous matrix.

Osaka group II observed, in collaboration with Canon group, the existence of diffusion layers of ~10 Å thick at the Mo-Si interfaces of a Mo/Si multilayer, which was fabricated by ECD, Inc. It was found from electron diffraction patterns that the Si layers were amorphous, whereas the Mo layers were almost completely crystallized.

C) X-ray standing waves.

X-ray standing wave field is excited in a multilayer as a consequence of interference between the incident and Bragg-reflected beams. The anti-nodal planes lie in between the diffracting planes when the glancing angle of the X-ray beam is slightly below the exact Bragg angle, and continuously move onto the diffracting planes as the glancing angle is scanned to just above the Bragg angle. Fluorescent X-rays become maximum (or minimum) when the anti-nodal (or nodal) planes coincide with the positions of atoms in the multilayer. Measuring the intensity of fluorescence excited by the standing wave and comparing it with the calculated values, one can determine the position of impurity atoms.

PF group [29] used this method to probe the interlayer distribution of entrapped Ar atoms in a W(12 Å)/Si(18 Å) multilayer with 48 layer-pairs on a silicon wafer. The sample was prepared by ECD, Inc. with magnetron sputtering in argon. The period of the multilayer was measured to be 29.56 Å by the X-ray diffraction measurement. They used synchrotron X-ray beam monochromated by a silicon (111) channel-cut crystal to excite fluorescence [30]. Figure 23 shows the angle dependence of the reflectance and intensity of Ar Kα(2.96 keV) excited in the multilayer by the standing waves of 8.41 keV. Comparison of the experimental and calculated curves suggests that the density of Ar atoms in the Si layers is 2-4 times higher than that in the W layers. The calculation was based on the treatment by Parratt [31]. The averaged density of Ar atoms was estimated to be in the order of 0.1%.

D) Mössbauer spectroscopy.

Kyoto group has been using Mössbauer spectroscopy, in combination with X-ray diffraction, etc., for the structural characterization of Fe/Mg...
Fig. 23. — Intensity of Ar Kα fluorescence X-rays emitted from entrapped Ar atoms in a W/Si multilayer. Reflectance is also shown. Thick and thin solid lines are experimental and calculated curves, respectively. α is a parameter for showing that the density of Ar atoms in the Si layers is α / (1 - α) times higher than that in the W layers (after Ref. [29]).

and Fe/C multilayers, which are of interest in soft X-ray use. It was found from Mössbauer spectra that a structural change occurs in the Fe layers when their thickness is thinner than a critical thickness.

In the case of Fe/Mg multilayers, the Fe layers thicker than 15 Å have bcc structure, whereas the Fe layers thinner than 15 Å have amorphous-like structure [32]. This structural change can be clearly seen in the 57Fe Mössbauer absorption spectra shown in figure 24. The spectrum for Fe(15 Å) shows a sharp six-line pattern similar to the bulk spectrum. This indicates that most part of 15-Å Fe layers has bcc structure. When the Fe layer thickness is less than 15 Å, the hyperfine field disappears at 300 K, and the spectrum shows a broad split six-line pattern at 4.2 K indicating the presence of a distributed hyperfine field. The monolayer samples, Fe(2 Å) and Fe(1 Å), show similar spectra with hyperfine splitting, and no non-magnetic component is observable. These results suggest that the diffusion or mixing of Fe atoms into the Mg layer is negligible and that Fe(< 15 Å) layers have amorphous-like structure.

In the case of Fe/C multilayers, the Fe(30 Å) layers have bcc structure, whereas the Fe(15 Å) and Fe(8 Å) layers have amorphous-like structure [33, 34]. Mössbauer spectra in figure 25 show this structural change, which is quite similar to those of the Fe/Mg multilayers. The spectra of Fe(15 Å) and Fe(8 Å) indicate a structural change of the Fe layers from bcc-Fe. This change was confirmed also by X-ray diffraction patterns and electron micrographs [33]. If the observed small hyperfine field is caused by bcc-Fe particles of small size, Mössbauer spectra at 4.2 K should show magnetic splitting corresponding to that of bcc-Fe. It was found experimentally that this is not the case. The detailed study of Mössbauer spectra of various samples shows that amorphous-like carbide layers or ultra-fine carbide particles are formed in the Fe(15 Å) and Fe(8 Å) layers and that amorphous carbide layers of ~ 5 Å thick are formed at each Fe-C interface [34].
5.2 REFLECTANCE MEASUREMENT USING SYNCHROTRON RADIATION. — The overall performance of multilayers can be evaluated by measuring their reflectances as a function of wavelength and angle of incidence. Use of a high-precision reflectometer with synchrotron radiation is best suited to such measurements. At present, such facilities are available at the Photon Factory and the UVSOR.

The Photon Factory provides an UHV reflectometer of high precision with a 2-m grasshopper monochromator (refer to Sect. 3.1 for the detail). The shortest wavelength available for the measurement is 10 Å. Unfortunately, this reflectometer cannot be rotated about the incident beam axis. A new reflectometer which has a provision for the rotation is now under adjustment and will be put in service shortly. A reflectometer at the UVSOR can be rotated by 90 deg. and permits measurements down to 75 Å with a plane grating monochromator.

A few examples of reflectance measurements made at the UVSOR and the Photon Factory are shown in figures 26 and 27. Figure 26 shows the peak reflectance vs. the angle of incidence measured by Osaka group I at the UVSOR for a multilayer of Mo (28.5 Å)/Si (66.5 Å) (10 pairs) overcoated with a carbon layer of 100 Å thick [35]. The reflectances of the s- and the p-component show a distinct difference at about 45 deg. angle of incidence, indicating a possible application of multilayers to polarizers.

Figure 27 shows the intensity variation of the 1st order reflection from a Fe (15 Å)/Mg (50 Å) multilayer [36]. The measurement was carried out by Kyoto group at the Photon Factory with X-rays of 1.25-2.3 Å so as to cover the K-absorption edges of Fe (1.7432 Å) and Mn (1.8959 Å). As is clearly seen in the figure, the intensity abruptly increases at about 1.90 Å, which corresponds to the Mn K-edge, and then decreases nearly exponentially at longer wavelengths. No anomalous behaviour was, however, observed at the Fe K-edge. This is quite interesting, because Fe and Mn have nearly equal atomic numbers and thus Fe/Mn multilayers are expected to have a very low reflectance. The reason for the observed enhancement is the effect of anomalous dispersion in the slightly longer wavelength region than the Mn K-edge. Nonappearance of anomaly at the Fe K-edge is due to large absorption by both Mn and Fe atoms. A similar intensity enhancement was observed for a Fe (15 Å)/V (30 Å) multilayer at the V K-edge (2.269 Å). These observations confirm the idea of YNMP group and suggest the use of the absorption edge of lower atomic number component of a multilayer to enhance the reflectance.

6. Applications.

Various applications are being considered by the research groups: X-ray telescopes by Osaka group I and Nagoya group, optical cavities for XUV lasers by Osaka group II, spectroscopic devices by Tohoku group, for example.

NTT group has made a preliminary experiment on X-ray lithography with a Schwarzschild demagnifying projection optics, whose components were coated with W/C multilayer [37]. The multilayer was designed to have a high reflectance at around a
wavelength of 110 Å for ease of experiment and prepared by means of ion-beam sputtering. The Schwarzschild optics was tested using synchrotron radiation on the beamline 15B at the Photon Factory. The experimental arrangement is illustrated in figure 28. The resist pattern of a wire mesh of 20 μm wide thus obtained is shown in figure 29, as an example. The pattern recorded in PMMA (0.4 μm thick) is the image of the mesh demagnified to 1/5 and shows a 4-μm grid. Although the wavelength used was not appropriate for X-ray lithography, the result was quite encouraging.

Nagoya group [38] has been developing a grazing incidence nested thin-foil X-ray telescope, which was originally proposed by Serlemitsos [39], to propose for a future mission of the Japanese X-ray astronomical satellite. The effective area of a grazing incidence telescope is limited by the small critical angle. They considered the use of multilayers as a means of obtaining a large effective area at the energy range of 6-8 keV. They made simulations on the performance of the telescope for the two cases: one for mirrors coated with Au single layer and the other for mirrors with Au/C bi-layer. Besides these simulations, they measured the reflectance of bi-layers, the surface roughness of various thin-foil mirror materials, the accuracy of a pre-engineering model structure for thin-foil nests, etc. Preliminary studies show that multilayers and single layers have their own merit and demerit. Further careful assessments are, therefore, necessary to draw any definite conclusion.

Acknowledgments.

The author would like to express his sincere thanks to all the members of the groups cited in Introduction for their cooperations in preparing the present article.

Fig. 28. — Schematic of the Schwarzschild demagnifying projection optics for X-ray lithography experiment (after Ref. [37]).

Fig. 29. — Resist pattern of a 20-μm wide wire mesh recorded with the arrangement of figure 28 (after Ref. [37]).

References

[34] NAKAYAMA, N., KATAMOTO, T., SHINJO, T., in preparation.