The State of Hydrogen in Nb-based Nb–Mo Alloys Analyzed by the Channelling Method

Eiichi YAGI

The Institute of Physical and Chemical Research (RIKEN), Wako, Saitama 351-0198 Japan, also at The School of Science and Engineering, Waseda University, Okubo, Shinjuku-ku, Tokyo 169-8555 Japan.

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In order to locate hydrogen dissolved in metals a channelling method utilizing a nuclear reaction \(^{1}\text{H}(^{11}\text{B}, a)\alpha\alpha\) with a \(^{11}\text{B}\) beam has been developed. This method has been applied to study the effect of metallic solutes on the state of hydrogen in metals. The lattice location of hydrogen in Nb-based Nb–Mo alloys with different concentrations \((C_{\text{Mo}})\) of undersized Mo atoms of 3, 10, 20 and 26 at\% has hitherto been investigated. These results are summarized in this paper and the interaction of hydrogen with metal solutes is discussed. At low Mo concentrations less than 10 at\% hydrogen is trapped by Mo atoms at room temperature due to attractive interaction to be located at sites displaced from tetrahedral \((T)\) sites by about 0.6 Å toward the nearest neighbour Mo atom. In the case of \(C_{\text{Mo}}=3\) at\%, hydrogen is detrapped at 373 K and enter a \(T\) site. The number of H atoms trapped by one Mo atom is limited. It is 0.7–1.0 for \(C_{\text{Mo}}=3\) at\% and decreases with increasing \(C_{\text{Mo}}\). The excess H atoms are located at \(T\) sites. For \(C_{\text{Mo}}=20\) at\% such trapped site occupancy is not observed, and 70–80\% of the H atoms are located at \(T\) sites and the rest are at octahedral \((O)\) sites at low H concentrations. For \(C_{\text{Mo}}=26\) at\%, most of the H atoms are located at \(T\) sites.

KEY WORDS: hydrogen; lattice location; channelling analysis; solute atoms; Nb–Mo alloy.

1. Introduction

In order to elucidate the mechanism of hydrogen embrittlement, knowledge of the fundamental properties of hydrogen such as its lattice location, its interaction with impurities, the behaviour in an atomic scale under stress, etc. is highly required. However, such knowledge is still insufficient, because the experimental methods for hydrogen detection are limited. Information on these properties is also important in the understanding of the hydrogen-related problems in the solid state physics. As to the lattice location of hydrogen isotopes, a number of experiments have been performed on deuterium by the neutron diffraction method and also by the channelling method with a \(^{3}\text{He}\) beam, whereas on hydrogen there had been only a few neutron diffraction experiments because of their experimental difficulties and no channelling experiments despite their usefulness, because the method used for deuterium is ineffective for hydrogen. The behaviour of hydrogen is not necessarily the same as that of deuterium as recognized from, for example, the difference in the phase diagram between \(V–H\) and \(V–D\) systems and the difference in the diffusivity between \(H\) and \(D\). From this point of view, isotope effect experiments on hydrogen are indispensable. Therefore, to study the lattice location of hydrogen a nuclear reaction \(^{1}\text{H}(^{11}\text{B}, a)\alpha\alpha\) with a \(^{11}\text{B}\) beam of about 2 MeV was applied to the channelling method. This method was demonstrated to be very useful and it has become possible to locate hydrogen dissolved in metals.\(^1\) This method has been used to various studies. Hydrogen dissolved in group \(V\) metals in the periodic table (\(V\), Nb and Ta) was observed to be located at tetrahedral \((T)\) sites.\(^2\)–\(^4\) It was demonstrated that hydrogen in \(V\) is extremely sensitive to stress and changes its location reversibly from a \(T\) site toward octahedral \((O)\) sites by 0.44 Å under application and release of stress below the elastic limit.\(^2\)–\(^5\)

In the present paper an interaction of hydrogen with solute atoms in metallic alloys is focused, and the results of studies hitherto made on the change of the lattice location of hydrogen in Nb–Mo alloys with Mo concentration, \(C_{\text{Mo}}\) are reported. This interaction is one of the important factors that influence behaviour and the state of hydrogen in metallic alloys. However, microscopic information on such interactions has been extremely limited. One of the examples is the effect of alloying on the terminal solubility of hydrogen (TSH) in group \(V\) metals (\(V\), Nb and Ta). It has been reported that for undersized metal solutes, \(e.g.\), Nb in Ta, and Mo in Nb, the TSH increases rapidly with solute concentration up to a certain value, whereas for the oversized solutes, \(e.g.\), Ta in Nb, and Nb in \(V\), the TSH increases less rapidly with solute concentration.\(^6\)–\(^11\) In order to understand the mechanism of this effect, the lattice location of hydrogen in such alloy systems is to be investigated. In the present study, an Nb–Mo alloy system has been chosen. This system forms a solid solution over the entire Mo concentration range, and has a bcc crystal structure. The TSH in Nb at room temperature is a few per cent, whereas the TSH in Mo is very low. Therefore, the systematic studies on the \(C_{\text{Mo}}\) dependence of the hydrogen site occupancy are desired. Hence, lattice location of hydrogen was investigated by the channelling method for Nb-based Nb–Mo alloys with various Mo concentrations up to 26 at\% Mo. Even for deuteri-
um, whose lattice location has been investigated extensively by both neutron and channelling methods, there has been only a few studies made in metallic alloys. The present study for Nb–Mo alloys with the same crystal structure as Fe will give helpful information to investigations of the state of hydrogen in iron and its alloys.

2. Experimental Method

The reason why the nuclear reaction $^1$H($^{11}$B, a)aα was applied is as follows. In the detection of impurities by ion beams, usually the Rutherford backscattering spectroscopy (RBS) is used, but this method is not useful to the detection of hydrogen. Therefore, the nuclear reaction was utilized. This reaction exhibits a narrow resonance with a resonance width of about 66 keV at about 1.8 MeV of a $^{11}$B beam. The energy of an incident $^{11}$B beam (~2 MeV) is reduced to about 1.8 MeV at the depth of a few thousand Å in a specimen, and there the resonance reaction takes place effectively to emit α particles. Hydrogen can be detected by measuring these α particles, whose energy ranges from zero to about 5 MeV. To locate hydrogen, this hydrogen detection technique was combined with the channelling method.

Specimens are Nb and Nb–Mo alloy single crystal slices whose largest face is perpendicular to the (110) axis. They were annealed and then doped with hydrogen from the gas phase up to a desired concentration lower than the solubility limit. They were mounted on the specimen holder of the three-axis-goniometer. Channelling analyses were performed at room temperature for the {100} and {110} axial channels, the {100} planar channel, and, if necessary, the {111} planar channel at different spots on the same specimen with a $^{11}$B ion beam of about 2 MeV. The beam was collimated to the divergence of less than 0.076°. The beam spot was 1 mm in diameter and the intensity was 1.5–2.0 nA. The goniometer was tilted on the horizontal plane by 45° with respect to the incident beam. The yield of $^{11}$B ions scattered by host metal atoms and the yield of the emitted α particles were measured as a function of the incident angle $\psi$ with respect to the channel direction in question by tilting the goniometer on the horizontal plane (channelling angular scan). Thus $^{11}$B- and α-angular profiles were obtained. The backscattered $^{11}$B ions were detected at a scattering angle of 160° in the vertical plane including the incident beam direction, while the α particles were detected at about 90° with respect to the incident beam direction on the horizontal plane. In front of the detector for the α particles a 4-μm-thick Mylar foil was placed to eliminate the $^{11}$B ions which were scattered by the host metal atoms in the direction of 90°. The α particles can penetrate this foil to be detected. The experimental set-up is shown in Fig. 1.

If hydrogen is located at a site which is projected at the position $r_i$ in the channel on the plane perpendicular to the channel (a transverse plane) in question, the α-angular profile exhibits peaks at angular positions $\psi_p$, corresponding to the potential at the projected position, $r_i$, due to flux-peak- ing effect in the spatial flux distribution of an incident beam in a channel under the channelling condition. The position of a subpeak $\psi_p$ is approximately given by

$$E\psi_p^2 = U(r_i).$$

$E$ is the incident energy of the beam and $U(r)$, which is the difference from its minimum value in the channel, is the continuum potential experienced by particles travelling in the channel. The origin for the position vector $r$ is taken at the centre of the channel. Therefore, when hydrogen is located at a site that is projected at the centre of the channel, the α-angular profile exhibits a central peak, while for hydrogen projected at off-centre positions, it exhibits off-centre peaks.

3. Experimental Results

3.1. Nb

Figure 2 shows the channelling angular profiles obtained for the Nb specimen which was doped with hydrogen up to the hydrogen-to-metal-atom ratio $C_{H_i}$=0.023. This specimen is in the α phase at room temperature. As the backscattering yields, those by atoms in the depth region from 60 to 180 nm were plotted. This region corresponds to the energy range where the resonance reaction in question takes place effectively. As the α-particle yields, those with energies higher than 0.7 MeV were plotted. (This is also the case for the Nb–Mo alloy specimens.) The angular profiles of the α-particle yield (α-angular profiles) exhibit a sharp central peak for the {100} channel, a broad central peak consisting of three subpeaks at $\psi=0°$ and around $\pm 0.35°$ for the {110} channel, and a small central peak superimposed on a shallow dip for the {100} channel. In the inset the projections of T and O sites on the plane perpendicular to the channel in question are shown with open squares. The angular positions of subpeaks which are expected for the T-site occupancy are indicated by arrows. They are esti-
In the calculation of a continuum potential, the crystal close to the axis or plane will not be able to pass through the centre of the channel before they penetrate a distance of the order of $\lambda = d_\alpha/(2\psi_{1/2})$, where $d_\alpha$ is the planar spacing of the channel and $\psi_{1/2}$ is the half angle of the channelling dip. In the case of the $\{100\}$ planar channel having the smallest $\psi_{1/2}$ among the three channels investigated, $\lambda$ is estimated to be about 19 nm. Since there are a number of ions deflected by less than $\psi_{1/2}$, the equilibrium in the flux distribution will not be fully established until the ion beam penetrates distance much larger than $\lambda$. As the angular profiles shown in Figs. 2, 5 and 7 were obtained from backscattering yields and $\alpha$-particle yields associated with atoms in the region deeper than 60 nm, this condition is considered to be fulfilled.

The normalized $\alpha$-particle yields $\chi_\alpha(\psi)$ were estimated from Eq. (2), considering that they are proportional to $F$. It was assumed that H atoms give the same angular profiles as those for the host metal atoms, i.e., the $^{11}$B-angular profiles, when they are shadowed behind atomic rows or planes in the channel. In this calculation the beam divergence and a change in $E_\perp$ due to scattering by electrons were taken into account, but the effect of lattice vibration was not. The average increase in $E_\perp$, $\delta E_\perp$, was taken to be 1.8 eV. This corresponds to $\delta E_\perp$ during penetration of ions to the depth of 60 nm where the nuclear reaction cross section begins to increase. For comparison of the measured $\alpha$-angular profiles with the calculated ones, the theoretical $\alpha$-angular profiles for the perfect lattice were corrected in the following way by taking account of the random fraction of the incident beam arising from dechanneling during transmission through the surface region of the specimen. The random fraction $f_a(\psi)$ is given by

$$f_a(\psi) = \frac{1}{\psi_{1/2}} \left( \frac{\psi_{1/2}}{\psi_{1/2} - \psi} \right)$$
3.2. 97at%Nb–3at%Mo and 90at%Nb–10at%Mo alloys

In the 3 at% Mo alloys, for $C_H = 0.02$, which is lower than $C_{Mo}$, the $\alpha$-angular profiles obtained for $\langle 100 \rangle$, $\langle 110 \rangle$ and $\langle 100 \rangle$ channels at room temperature were completely different from those for hydrogen in Nb.20) The $\alpha$-angular profiles did not exhibit a peak at $\psi = 0^\circ$ for all the three channels, but exhibited a small peak at $\psi = 1^\circ$ for the $\langle 100 \rangle$ channel, small off-centre peaks at $\psi = \pm 0.2^\circ$ and $\pm 0.6^\circ$ for the $\langle 110 \rangle$ channel, and only a shallow dip for the $\langle 100 \rangle$ channel. Hence, most of the H atoms are considered to be located at sites different from $T$ sites. When the hydrogen concentration was higher than $C_{Mo}$, i.e., $C_H = 0.05$, the $\alpha$-angular profiles obtained at room temperature markedly changed from those for $C_H = 0.02$. The $\langle 100 \rangle$ $\alpha$-angular profile exhibited a higher central peak at $\psi = 0^\circ$. In the $\langle 110 \rangle$ channel, higher subpeaks newly appeared at $\psi = 0^\circ$ and around $\pm 0.35^\circ$, and a small peak at $\psi = -0.6^\circ$ for $C_H = 0.02$ was still present. The $\langle 100 \rangle$ $\alpha$-dip became deeper, and a small central peak appeared to be superimposed on the dip. These results suggest that some portion of the H atoms are located at $T$ sites and the rest at the same sites as those observed for $C_H = 0.02$. After the measurement at room temperature, the specimen with $C_H = 0.05$ was measured at 373 K. In the $\langle 100 \rangle$ $\alpha$-angular profile the central peak became higher, and for the $\langle 100 \rangle$ channel a central peak superimposed on the dip became more conspicuous. These profiles are characteristic of the $T$-site occupancy. These results can be interpreted as follows. In the alloy with $C_H = 0.02$, most of the H atoms are trapped by Mo atoms at room temperature and located at sites different from $T$ sites (trapped sites $T_u$). In the alloy with $C_H = 0.05$, some portion of hydrogen atoms are located at $T_u$ sites and the rest are at $T$ sites at room temperature, while at 373 K the trapped hydrogen atoms are detrapped and enter $T$ sites.

In order to determine the lattice location of the $T_u$ sites, the $\alpha$-angular profiles were calculated, considering five possible kinds of sites and that all H atoms are located at only one kind of these sites. Thus calculated results were compared with the observed ones. The observed $\alpha$-angular pro-

![Graph](image)
files obtained for $C_H=0.02$ were well reproduced by the calculation made for the site displaced from a $T$ site toward the nearest neighbour lattice point by 0.62 Å, which is about 34% of their original separation (Fig. 4). Hence, it is concluded that in the 3 at% Mo alloys with $C_H=0.02$ all H atoms are trapped at room temperature by Mo atoms and are located at sites displaced from $T$ sites by about 0.6 Å toward the Mo atoms at the nearest neighbour substitutional sites, and that at 373 K they are detrapped and enter $T$ sites. Admittedly, there is no way to determine directly the hydrogen position relative to Mo atoms, but, in view of the smaller size of an Mo atom in comparison to an Nb atom, the $T_{O}$ site nearest to an Mo atom is believed to be most probable. The result that only a portion of the H atoms are trapped for $C_H=0.05$ indicates that hydrogen trapping by Mo atoms has already been saturated. From the comparison of $\alpha$-angular profiles for H in Nb and the 3 at% Mo alloys with $C_H=0.02$ and 0.05, the number of hydrogen atoms trapped by one Mo atom, $\gamma$, can be roughly estimated to be 0.67–1.0. Hence, the binding energy between hydrogen and an Mo atom was roughly estimated to be 0.05 eV in reasonable agreement with the previous determination by other technique.6)

In the 10 at% Mo alloys the $\alpha$-angular profiles were similar to those for the 3 at% Mo alloys with $C_H=0.05$. It was concluded that H atoms are also distributed over $T$ and $T_{O}$ sites, but the magnitude of $\gamma$ is reduced to 0.25–0.35.21,22)

3.3. 80 at% Nb–20 at% Mo Alloys

In the 20 at% Mo alloy, channeling experiments were performed at room temperature for three different hydrogen concentrations $C_H=0.026, 0.042$ and 0.08.23) The results are shown in Fig. 5. For $C_H=0.026$ and 0.042, the $\alpha$-angular profiles exhibit a large and sharp central peak superimposed on a shallow dip for the $\{100\}$ channel, a broad central peak consisting of three subpeaks at $\varphi=0^\circ$ and around $\pm 0.35^\circ$ superimposed on a shallow dip for the $\{110\}$ channel, and a small central peak superimposed on a shallow dip for the $\{100\}$ channel. Except for the $\{100\}$ and $\{110\}$ shallow dip, they are similar to those observed for H in Nb (Fig. 2). These results suggest that a large portion of H atoms are located at $T$ sites and some portion of them are at sites which give the shallow dip in the $\{100\}$ and $\{110\}$ $\alpha$-angular profiles, i.e., which are shadowed behind $\{100\}$ and $\{110\}$ atomic rows. These sites are considered to be $O$ sites, because one third of $O$ sites are shadowed behind $\{100\}$ and $\{110\}$ atomic rows (inset of Fig. 5), whereas $T$ sites are not. For $C_H=0.08$, similar peaks are still observed for the three channels, but the shallow dips for the $\{100\}$ and $\{110\}$ channels became much shallower or almost non-existent, and, in addition, for the $\{100\}$ channel the relative depth of the $\alpha$-dip with respect to the $\{\bar{1}1\}0\bar{0}$-dip became smaller than those in the case of $C_H=0.026$ and 0.042. This result suggests that for $C_H=0.08$ the fraction of the H atoms located at $O$ sites is significantly reduced and most of the H atoms are at $T$ sites.

The lattice location of hydrogen and its distribution in the 20 at% Mo alloy are determined by fitting the calculated $\alpha$-angular profiles to the observed ones.23) Examples of $\alpha$-angular profiles calculated for various distributions of H atoms over $T$ and $O$ sites are shown in Fig. 6 together with experimental results. Taking account of the statistical error, for $C_H=0.026$ and 0.042 the observed $\alpha$-angular profiles for the three channels are rather well reproduced by the H distribution of 70–80% at $T$ and 20–30% at $O$ sites. On the other hand, for $C_H=0.08$, they are reproduced by the H distribution of 90–100% at $T$ sites and 0–10% at $O$ sites. It indicates that at high H concentrations the fraction of the $O$-site occupancy markedly decreases.23) The $T_{O}$-site occupancy was not observed, i.e., $\gamma=0$.

3.4. 74 at% Nb–26 at% Mo Alloys

The channeling angular profiles for more concentrated 26 at% Mo alloys with different concentrations of hydrogen are shown in Fig. 7.6) Irrespective of the hydrogen concentration the $\alpha$-angular profiles are very similar to those for H in Nb in Fig. 2. As distinct from the results on H in the 20 at% Mo alloys with low concentration of hydrogen,23) a shallow dip is not observed for the $\{100\}$ and $\{110\}$ channels. The lattice location of hydrogen was determined by comparing the observed $\alpha$-angular profiles with the calculated ones.6) It can be said that as in the case of hydrogen in Nb the observed profiles are well reproduced by the calculated ones for the $T$-site occupancy, although the subpeaks are not observed to be sharply resolved ones. The possible occupancy of 10% H atoms at $O$ sites cannot be excluded, however, because the $\{100\}$ and $\{110\}$ $\alpha$-angular profiles calculated for this distribution do not give a shallow dip so clearly as in the case of 20–30% H atoms at $O$ sites.23) It is, therefore, concluded that in the 26 at% Mo alloys H atoms are mostly located at $T$ sites and, the fraction of the $O$-site occupancy is at most 10%. The $T_{O}$-site occupancy was not observed. The site occupancy of hydrogen that has hitherto been observed in Nb–Mo alloys is summarized in Table 1. The dependence of the number of H atoms trapped per one Mo atom, $\gamma$, on $C_{Mo}$ is shown in Fig. 8.

4. Discussion

Previously it was reported that the halfwidth of X-ray reflection lines in 5 at% Mo and 5 at% V alloys6) and the halfwidth of $^{53}$Nb NMR absorption curves in 5 at% V and 8 at% V alloys24) remarkably decreased with increasing H concentration, when the H concentration was less than the metal solute concentration. This result was interpreted as a result of the relaxation of internal strain induced by the undersized metal solutes, Mo and V, by hydrogen occupying their neighbouring interstitial sites, i.e., the trapping of hydrogen by solute atoms due to elastic interaction between...
The site change of hydrogen in the Nb–Mo alloys containing Mo less than 20 at% has been explained as follows. At low $C_{\text{Mo}}$, H atoms occupy well-defined trapped sites ($T$-sites) near Mo atoms due to the attractive elastic interaction. With increasing $C_{\text{Mo}}$, the interference of the strain field induced by nearby Mo atoms changes the energy of the trapped sites, leading to a distribution of local atomic configurations and concomitant distribution of interstitial site energies. Correspondingly, trapping behaviour will become less well-defined, and, therefore, $\gamma$ becomes smaller. At higher $C_{\text{Mo}}$, 20 at% Mo, the average separation between Mo atoms becomes even smaller, and the effect of strain fields of individual Mo atoms would tend to average out. It is then considered more appropriate to regard this alloy as a pure metal having a homogeneously reduced lattice parameter.

A theoretical calculation on H in group V a metals by Sugimoto and Fukai has shown that as the lattice is elongated along one of its cube axes, e.g., [001] axis, the wave function of hydrogen, originally localized on one of the four $T$-sites around the $O$ site on the (001) plane ($O_z$ site), is gradually displaced toward the $O_z$ site ($d-T$ site) and, after going through a resonant state of these four $d-T$ configurations (4$\pi$), finally attains the $O_z$ site. (The symmetry of the 4$\pi$ configuration allows for the $O_z$-site occupancy as a particular case.) The energy difference between $T$ and 4$\pi$ configurations, $\Delta E_{4\pi-T}$, depends on metal species and decreases

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**Fig. 5.** Channelling angular profiles of $\alpha$-particle yield (●) and the yield of $^{11}\text{B}$ ions backscattered by metal atoms (○), obtained at room temperature for the 20 at% Mo alloys with hydrogen of (a) $C_H=0.026$, (b) 0.042 and (c) 0.08. The full curves and the dashed curves have been drawn to guide the eye. The angular positions of subpeaks which are expected for the $T$-site occupancy are indicated by arrows.
in order of Ta, Nb and V, i.e., with decreasing lattice parameter. The variation of \( \Delta E_{\text{T}T} \) with the relative size of an interstitial with respect to the host lattice arises from the fact that more drastic lowering of the potential energy of an O site with outward displacement of its two nearest neighbour metal atoms occurs in more congested circumstances. Such a site change of H atoms from T to d–T or 4T configurations was observed by a channelling method in V under uniaxial stress,\(^{2,5} \) although the stress level was much lower than that expected from the calculation. Such site changes were not observed in Nb and Ta.\(^{3,4} \)

The appearance of the O-site occupancy in the 20 at% Mo alloys can be explained qualitatively by the size effect described above. On alloying with 20 at% Mo the lattice parameter of Nb is reduced by 0.8%. Hence the energy difference between T and O site becomes smaller, and O sites become more easily available for occupation. The reduction of the fraction of the O-site occupancy, which was observed at a higher hydrogen concentration \( (C_{\text{H}}=0.08) \), is considered to be due to the interaction between H atoms, because the average separation between H atoms becomes smaller, \( \sim 2.2a \) \((a: \text{a lattice constant})\).\(^{23} \)

However, the results on the 26 at% Mo alloys cannot be explained in a similar way by such size effects. Whereas the smaller lattice parameter of the 26 at% Mo alloys suggests the larger fraction of the O-site occupancy in the 26 at% Mo alloys, the experimental results indicate the contrary, which requires some additional considerations. With increasing \( C_{\text{Mo}} \), individual T and O sites have a variety of configurations of the nearest neighbour atoms, i.e., tetrahe-

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Fig. 6. The observed (open circles) \( \chi \)-angular profiles and calculated ones for various hydrogen distributions over T and O sites for the 20 at% Mo alloys with hydrogen of (a) \( C_{\text{H}}=0.026 \), (b) 0.042 and (c) 0.08. Curve 1; 75% at T, 25% at O, curve 2; 60%, 40%, curve 3; 90%, 10%, and curve 4; 100%, 0%. Some of the curves are only partly shown, and the calculated yield for \( \psi=0^\circ \) are indicated on the right hand side of the curves by horizontal lines of the type specific to each curve, because the figures are busy.
dra and octahedra comprising various numbers of Mo atoms. In the 26 at% Mo alloys there exists approximately one Mo atom per two unit cells on average. Therefore, all of the T sites come to be surrounded by tetrahedra including one Mo atom as one of their constituent atoms on average, while each half of the O sites come to be surrounded by octahedra including one Mo atom and two Mo atoms on average, respectively. Generally speaking, the interaction between metal solutes and hydrogen consists of electronic and elastic interactions. The electronic structure of Nb–Mo alloys have been studied by angular correlation of positron annihilation radiation. The two dimensional angular correlation studies by West on Nb, 10 at% Mo, 30 at% Mo alloys and Mo clearly revealed the change of Fermi surface topology with increasing Mo concentration. From the one dimensional angular correlation experiments on 20 at% Mo and 50 at% Mo alloys, Shiotani et al. demonstrated that extra valence electrons from Mo atoms fill the hole region in the momentum space and there exist well-defined Fermi surfaces even in the most concentrated alloy. The result of the present experiments suggests that, in addition to a change in the electronic structure of alloys on alloying with higher concentration of Mo atoms, the change in the distribution of local atomic configurations as described above in-

Fig. 7. Channelling angular profiles of α-particle yield (〇) and the yield of 11B ions backscattered by metal atoms (○), obtained at room temperature for the 26 at% Mo alloys with hydrogen of (a) C_H=0.016, (b) 0.026 and (c) 0.059. The full curves and the dashed curves have been drawn to guide the eye. The angular positions of subpeaks which are expected for the T-site occupancy are indicated by arrows.
alloying with undersized metal solutes in group V a metals, Nb–Mo alloys is under consideration. On the lattice location of hydrogen in more concentrated electronic interaction becomes important. The experiment ascribed to a simple trapping of hydrogen.\(^{28,29}\) Oates and in Nb–V alloys for a wide range of solute concentration pressure) solubility of hydrogen and the hydrogen diffusion observed in the studies on the isopiestic (equal hydrogeners reported that evidence for trapping of H could not be detected in Nb–Mo and NMR experiments on H in Nb-based Nb–Mo and trapping model) has been proposed from X-ray diffraction a model on the trapping of hydrogen by solute atoms (the elastic interaction is more important in the low C\(_{Mo}\) region, while in the high C\(_{Mo}\) region the contribution of electronic interaction becomes important. The experiment on the lattice location of hydrogen in more concentrated Nb–Mo alloys is under consideration.

On the mechanism for the enhancement of the TSH on alloying with undersized metal solutes in group V\(_a\) metals, a model on the trapping of hydrogen by solute atoms (the trapping model) has been proposed from X-ray diffraction and NMR experiments on H in Nb-based Nb–Mo and Nb–V alloys.\(^{6,24}\) On the other hand, Peterson and co-workers reported that evidence for trapping of H could not be observed in the studies on the isopiestic (equal hydrogen pressure) solubility of hydrogen and the hydrogen diffusion in Nb–V alloys for a wide range of solute concentration and, therefore, the enhancement of the TSH could not be ascribed to a simple trapping of hydrogen.\(^{25,29}\) Oates and Flanagan suggested that the enhancement of the TSH on alloying is likely to be fairly general and explained this by considering only the average effect of alloying on macroscopic thermodynamical quantities.\(^{30}\) Therefore, the mechanism of the enhancement of the TSH is still an open question. From the studies on the lattice location of hydrogen, however, it is considered that in the Nb-based Nb–Mo alloy system the trapping of hydrogen by Mo solutes is effective for this enhancement of TSH, at least in the low concentration region of Mo solute atoms.

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