Structural Investigation of Uranium—Neptunium Mixed Oxides Using XRD, XANES, and $^{17}$O MAS NMR

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ABSTRACT: Uranium—neptunium mixed dioxides are considered as fuels and targets for the transmutation of the minor actinides in fast neutron reactors. Hereafter, a local and atomic scale structural analysis was performed on a series of $\text{U}_{1-x}\text{Np}_x\text{O}_2$ ($x = 0.01; 0.05; 0.20; 0.50; 0.75; 0.85$) synthesized by the sol–gel external gelation method, for which longer range structural analysis indicates that the process yields solid solutions. The oxidation state of IV for uranium and neptunium cations was confirmed using $\text{U} \text{L}_{III}$ and $\text{Np} \text{L}_{III}$ edge X-ray absorption near edge structure (XANES). The atomic scale structure was probed with $^{17}$O magic angle spinning nuclear magnetic resonance (MAS NMR) for the anion. Structural distortions due to the substitution of $\text{U}$ by the smaller $\text{Np}$ cation were detected by $^{17}$O MAS NMR.

1. INTRODUCTION

Understanding the incorporation mechanism of minor actinides (MA), that is, Np, Am, Cm, in uranium dioxides and mixed U—Pu oxides (MOX) as solid solutions is essential starting point for the detailed understanding of these materials, which will become particularly relevant, as they are considered as fuels and targets for the transmutation of the minor actinides in fast neutron reactors.1,2 This is an innovative approach to dramatically reduce the long-term radiotoxic impact of these materials and is considered in two modes: the homogeneous MA recycling scenario which relies on the introduction of a few percent of MA in all of the fuel pins of the reactor and, the heterogeneous MA recycling concept which considers deployment of the MA in a UO$_2$ matrix in dedicated fuel rods and assemblies. As such fuels contain no Pu in the fresh fuel, they are often referred as “targets”. Despite being the most abundant minor actinide in spent fuel, targets bearing Np have not been, to our knowledge, the subject of any complete structural study using local structure techniques. We focus our attention here on a most basic solid solution system $\text{U}_{1-x}\text{Np}_x\text{O}_2$. Such Np solid solutions have been investigated by XRD at high temperatures to develop the understanding of their thermodynamic properties in the hyperstoichiometric region of the system.1–3 Further, as these materials possess unusual magnetic ordering at low temperatures significant investigative work have been done.4–9 Neptunium-237 Mössbauer spectroscopy was performed in a series of $\text{U}_{1-x}\text{Np}_x\text{O}_2$ ($x = 0.15; 0.25; 0.5; 0.75$), 1) compounds and demonstrated that Np remains IV valent oxidation state for all compositions.7 Nevertheless, the oxidation state of uranium in these compounds remains uncharacterized, despite its potential importance as U can adopt higher oxidation states than IV, that is, V and VI.

Structural analysis of actinide solid solutions such as Th$_{1-x}$M$_x$O$_2$ ($M = \text{U}, \text{Pu}$)10 and U$_{1-x}$Pu$_x$O$_2$11 were performed mainly using X-ray absorption spectroscopy (XAS). Such studies show more complex local structures than derived from XRD characterization with, for example, a bimodal cation-oxygen distribution. High resolution solid-state MAS NMR should be a very effective tool to probe the atomic scale in such compounds, as it has been extensively utilized for the study of, among other systems, rare-earth solid solutions.12–16 Nevertheless, until recently, its use with radioactive materials has remained elusive, mainly due to strict radiological safety requirements of handling highly radiotoxic materials when high spinning speeds are needed.17 A technological advance has allayed these safety issues and our instrument at JRC-ITU is now available for the characterization of such highly radioactive materials.18 The combination of XAS and MAS NMR will be, then, very powerful for the study of these actinide solid solutions.

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Supporting Information: XAS and $^{17}$O MAS NMR spectra and plots of $^{17}$O chemical shifts vs. $\text{Np}$ concentration for $\text{U}_{1-x}\text{Np}_x\text{O}_2$ x-ray absorption near edge structure (XANES)

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Herein, we present a complete analysis of the $U_{1-x}Np_xO_2$ ($x = 0.01, 0.05, 0.2, 0.5, 0.75, 0.85, 1$) solid solutions prepared by gel-supported precipitation. The stoichiometry of the solid solution was checked by XRD. The oxidation state of both uranium and neptunium was determined using XANES. Finally, high resolution $^{17}$O MAS NMR was used for the first time in such solid solutions to probe the local environment around the oxygen.

2. EXPERIMENTAL SECTION

2.1. Sample Preparation and Chemical Analysis. The $U_{1-x}Np_xO_2$ samples ($x = 0.01, 0.05, 0.20, 0.50, 0.75, 0.85$), denoted hereafter as Np$_x$, have been prepared by gel-supported precipitation (also referred as sol–gel external gelation). This synthesis process avoids powder handling of the highly radiotoxic materials,\textsuperscript{3,19} and one obtains a high composition (metal) homogeneity.\textsuperscript{20} A uranyl nitrate solution was mixed with a nitric acid solution of $^{237}$Np to achieve a molar stoichiometry of $x = \text{Np}/(U + \text{Np})$, as defined above. Before the external gelation step, aliquots of each solution were taken for analysis by inductively coupled plasma–mass spectrometry (ICP-MS) to perform a precise control of the Np/(U + Np) ratio. The results are presented in Table 1 and there is a good agreement between calculated and obtained compositions. For the external gelation step, an organic thickener (Methocel, Dow Chemicals) was added to increase the viscosity of the solutions. These were then dropped into an ammonia bath where the droplet to particle conversion took place due to hydroxide precipitation of the metals inside the polymer backbone of the droplet. The resulting beads were washed, dried, and then calcined at 600 °C in air (2 h) to remove organics and then at 700 °C in Ar/H$_2$ (2 h) to ensure the reduction to U(IV). The samples were enriched at about 30% in $^{17}$O by heating in oxygen-17 enriched gas at 800 °C during 24 h, before being sintered at 1650 °C (4 h) under Ar/H$_2$.

2.2. X-ray Diffraction. The XRD patterns were obtained at room temperature on 20 mg of a powdered sample loaded in an epoxy resin to fix the powders and avoid their dispersion. A Bruker D8 Advance diffractometer (Cu Kα radiation, 40 kV, and 40 mA) with a Bragg–Brentano θ/2θ configuration and implanted in a radioactive glovebox were used for the analysis. This diffractometer is equipped with a curved Ge monochromator (111) and a Lynxeye linear position-sensitive. Instrumental calibration was done using LaB$_6$ SRM660a as external standard. The powder patterns were recorded using a step size of 0.0197° across the angular range $10° \leq 2\theta \leq 120°$. Structural analyses were performed by the Rietveld method using Jana2006 software.\textsuperscript{21} Peak profile fitting was achieved using Pseudo-Voigt functions.

2.3. XANES. The XANES measurements were performed on 5 mg of powdered sample mixed with 55 mg of BN. The data were collected at the Angströmquelle Karlsruhe (ANKA) synchrotron, under dedicated operating conditions (2.5 GeV, 100–160 mA) at the INE-Beamline for actinide science, on which XAS on radioactive materials is licensed.\textsuperscript{22} A Ge [422] double-crystal monochromator coupled to collimating and focusing Rh-coated mirrors was used. The spectra were acquired at room temperature in both transmission and fluorescence modes at U LIII (17166 eV) and Np LIII (17610 eV) edges. In this latter mode, XAS were recorded using a five-element LEGe solid-state detector. The energy calibration was accomplished by measuring the spectrum of an Y (17038 eV) foil located between the second and the third ion chamber filled with Ar at ambient pressure, used to record the transmission.

2.4. $^{17}$O MAS NMR. The NMR experiments were performed on a 9.4 T Bruker spectrometer at the Larmor

![Figure 1](A) XRD pattern of the $U_{1-x}Np_xO_2$ solid solutions and (B) variation of the lattice parameter as a function of composition.

| compound name | Np (mol %) | U (mol %) |
|---------------|------------|-----------|
| Np1           | 1.0(6)     | 99.0(6)   |
| Np5           | 4.7(6)     | 95.3(6)   |
| Np20          | 19.1(3)    | 80.9(3)   |
| Np50          | 51.18(4)   | 48.81(4)  |
| Np75          | 75.8(4)    | 24.2(1)   |
| Np85          | 85.9(5)    | 14.10(7)  |

Figure 1. (A) XRD pattern of the $U_{1-x}Np_xO_2$ solid solutions and (B) variation of the lattice parameter as a function of composition.
frequency of 54.25 MHz. This equipment is combined with an active glovebox dedicated to the handling of radioactive materials, and is described in detail elsewhere. A commercial 1.3 mm probe was used, and the rotors were spun at 55 kHz. The pulse durations were 4 \( \mu s (\pi/2) \) and 8 \( \mu s (\pi) \), respectively, with an echo delay of 18.2 \( \mu s \) (1 rotor period). All spectra were calibrated relative to \( \text{H}_2\text{O} \) enriched in \(^{17}\text{O} \) (0 ppm). The longitudinal relaxation time \((T_L)\) was determined using an inversion−recovery pulse sequence. All the spectra were fitted using the dmfst software and the option “quad first.”

3. RESULTS AND DISCUSSION

3.1. Synthesis and Long-Range Structure. The XRD patterns of the well-crystallized \( \text{U}_{1-x}\text{Np}_x\text{O}_2 \) samples are presented in Figure 1A. These data indicate that all the samples are single phase solid solutions, except for the Np50, for which evidence for two fluorite crystalline phases is found. The presence of two phases in this later sample is most probably due to the experimental conditions during the synthesis and not to a miscibility gap, as this material has already been synthesized as a single phase.\(^9\) Using the Rietveld refinement method, the lattice parameters of each sample were precisely determined, and their variation as a function of composition is presented in Figure 1B. The fluorite \( \text{Fm}3\text{m} \) crystalline structure was found for all the materials. The two crystalline phases identified in Np50 were determined as \( \text{U}_{0.50}\text{Np}_{0.50}\text{O}_2 \) (86\%) and \( \text{U}_{0.25}\text{Np}_{0.75}\text{O}_2 \) (14\%) based on their lattice parameter. As this second phase impurity is very small and given the radiological hazard and our active adherence to the radiological protection of the staff (ALARA = as low as reasonably achievable), the synthesis was not repeated. All the lattice parameters do follow the Vegard’s law exceptionally well, indicating good mixing in the solid solutions. In addition, the diffraction peak broadening was analyzed using Hall−Williamson formula to extract the micro strain in these materials.\(^{25}\) For the Np50 sample, only the strain of the \( \text{U}_2\text{Np}_5\text{O}_{12} \) component has been considered. The results are shown in Figure 2 and indicate a higher strain for the intermediate solid solutions. This trend is readily understood in terms of local distortion created by the coexistence of two cations with different ionic radius in the structure as shown, for example, by Venkateswara Rao et al. in \( \text{Mg}_{5}\text{Ni}_2\text{O}_{26} \) or by Düvel et al. in mixed alkaline-earth fluorine.

Figure 2. Strain variation for the different \( \text{U}_{1-x}\text{Np}_x\text{O}_2 \) samples.

3.2. Oxidation States Determination of the Actinides. The oxidation states have been determined from XANES at U L\(_{III}\) and Np L\(_{III}\) edges. The corresponding spectra are presented in Figure 3 with those of \( \text{U(IV)O}_2 \) and \( \text{Np(IV)O}_2 \) reference materials. Their inflection points and white lines were found at 17170.3 (S) and 17175.5 (S) eV and at 17616.5 (S) and 17621.8 (S) eV, respectively. Within the uncertainty of 0.5 eV, there is no variation of the inflection point and the white lines in the \( \text{U}_{1-x}\text{Np}_x\text{O}_2 \) samples compared to the pure dioxides (Table S1). This shows that uranium and neptunium have an oxidation state of IV for all the compositions. The oxidation state IV of Np in the \( \text{U}_{1-x}\text{Np}_x\text{O}_2 \) is consistent with previous Mössbauer results.\(^7\) Due to the very close UL\(_{III}\) and NpL\(_{III}\) edges, EXAFS spectra cannot be analyzed with this conventional XAS.

3.3. Local Structure Determination by \(^{17}\text{O} \) MAS NMR. The \(^{17}\text{O} \) MAS NMR experiments were performed to probe the local environment around the anions. Due to the presence of unpaired electrons in the cation’s 5f-shell, the NMR resonance of the anion is shifted beyond the normal range found for traditional chemical shifts. This so-called paramagnetic shift is very sensitive to the oxidation states, orbital occupancy, and coordinating geometries of the neighboring cations. In addition, a variation in chemical composition due to the substitution of a cation by another one leads to the apparition of additional peaks on the MAS NMR spectra corresponding to the different local environments around the anion.\(^{25-30}\) This specificity of paramagnetic NMR is therefore very efficient for characterization of such solid solutions. The full spectra, the center band, and the corresponding fits are presented in Figure 4 for \( \text{UO}_2 \), \( \text{NpO}_2 \), and the \( \text{U}_{1-x}\text{Np}_x\text{O}_2 \) under study. Spinning sidebands have been identified for all the spectra and are coming from the satellite transitions. The signals of the two end-members,
Identified at 717 and 475 ppm for UO\textsubscript{2} and NpO\textsubscript{2} respectively,\textsuperscript{18} exhibit a single sharp peak characteristic of O(U)\textsubscript{4} and O(Np)\textsubscript{4} units within the sample. The different oxygen signals observed by increasing Np content are attributed to O(Np)\textsubscript{y}(U)\textsubscript{4-y} (with 0 ≤ y ≤ 4) units. The $^{17}$O MAS spectrum of Np1, characteristic of the lowest Np concentration in our series, is composed of two signals at 718.4 and 636.3 ppm, attributed to (O(U)\textsubscript{4}) and (O(U)\textsubscript{3}(Np)\textsubscript{1}) units. In contrast with XRD where the change of lattice parameter compared to UO\textsubscript{2} is too small to prove clearly the formation of a solid solution,\textsuperscript{17}O MAS spectrum of Np1 gives, here, a clear indication of solid solution formation for such low Np amounts. With an increase of 4 mol % of NpO\textsubscript{2} (Np5 sample), three signals are now visible at 720.9 (O(U)\textsubscript{4}), 639.9 (O(U)\textsubscript{3}(Np)\textsubscript{1}), and 527 ppm (O(U)\textsubscript{2}(Np)\textsubscript{2}). The NMR spectrum of the Np20 sample is composed of four signals between 729.7 ppm (O(U)\textsubscript{4}) and 414.3 ppm (O(U)\textsubscript{1}(Np)\textsubscript{3}). For the Np50 compound, the two crystalline phases observed by XRD cannot be differentiated on the MAS NMR spectrum due to the width of the peaks identified between 749.9 ppm (O(U)\textsubscript{4}) and 337.0 ppm (O(Np)\textsubscript{4}) and a strong overlapping of the two spectra. Nevertheless, as the U\textsubscript{0.50}Np\textsubscript{0.50}O\textsubscript{2} is the dominant composition, the position of the peaks should be characteristic of a sample with this composition. The Np75 sample has an NMR spectrum, which is the least well resolved of all, but, four signals can still be differentiated between 759.8 ppm (O(U)\textsubscript{2}(Np)\textsubscript{2}) and 405.2 ppm (O(Np)\textsubscript{4}). Finally, the spectrum of the Np85 sample presents four well-resolved signals between 759.6 ppm (O(U)\textsubscript{2}(Np)\textsubscript{2}) and 437.8 ppm (O(Np)\textsubscript{4}).

The variation of the paramagnetic shifts of the five different subunits with sample composition is presented in Figure 5. For each identical O(Np)\textsubscript{y}(U)\textsubscript{4-y} (with 0 ≤ y ≤ 4) unit, there is a uniform linear increase of the shifts with increasing Np/(U + Np) content. It is also worth noticing that the slope of the linear fit of the paramagnetic shifts increases with increasing number of Np neighbors. This linear increase of the shifts with increasing Np/(U + Np) content is in agreement with the existence of a solid solution through the entire composition as observed by XRD. In fact, similar variations of the shifts were observed by NMR in alkaline-earth\textsuperscript{27,31,32} and rare-earth solid solutions.\textsuperscript{15} The variation of the shift of each unit with total composition can be thus deemed as an NMR analogue to the Vegard Law’s. The variation of the shifts is more pronounced in our series of solid solution due to the paramagnetic effect, which can be very sensitive to any local distortions.\textsuperscript{13,14} Presence of local distortions were observed on similar, 

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Figure 4. $^{17}$O MAS NMR spectra and fits (red and black curves) of the U\textsubscript{1-x}Np\textsubscript{x}O\textsubscript{2} samples acquired at 55 kHz. The black arrows indicate the spinning sidebands positions.
The width of the individual peak characterized by its full width at half-maximum (fwhm), has been plotted against the Np/(U + Np) ratio in Figure 6. For the same composition, the presence of at least one neptunium neighbor around the O atom leads to an increase of the fwhm and the more surrounding Np atoms the larger the fwhm. In addition, the increase of the fwhm with increasing Np neighbors is more important for the O(Np)₂ units than for the O(U)₁ units. For all the samples, there is an increase of this parameter for each coordination unit with increasing Np content until Np75, then it decreases. The variations of the fwhm can be attributed to both local distortions around the O atoms when Np⁴⁺ replaces U⁴⁺ and transfer of positive electron from the cations.²⁷ Considering the variation of the strain obtained by XRD with a maximum for the U₀.₂₅Np₀.₇₅O₂, additional electronic effects must also contribute to the line broadening for which a maximum occurs for U₀.₂₅Np₀.₇₅O₂. It is worth mentioning that, as¹⁷O is a quadrupolar nucleus (i.e., spin 5/2), presence of local bond angle distortions around the oxygen atoms can create an appearance of quadrupole interaction³³ (i.e., a quadrupolar coupling constant (C_q) not null) being at the origin of the fwhm increase previously observed. To verify if such a distortion exists, nutation experiments that consist of the observation of the center band magnetization variation with increasing 90° pulse lengths (τ_{90°}) have been performed.³⁴–³⁶ In fact, if one compares the nutation curve of a given compound with that of a reference sample, known with C_q = 0 (usually a liquid sample), one can determine if C_q ≠ 0 (i.e., different variation of the nutation curves) and, if so, an estimate of its value is possible by fitting the nutation curve.³⁷,³⁸ The nutation curves obtained at 55 kHz are presented in Figure S1 (2 W of applied power) and Figure S2 (20 W of applied power) for the samples that present the highest fwhm increase Np5, Np20, Np75, and Np85 (Np50 has not been analyzed as two crystalline phases are present). To obtain these nutation curves, the intensities of the main and best resolved peaks have been integrated. In addition, to compare the nutation curves of the different peaks, their intensities have been scaled at the same height. As MAS can influence the nutation curves by reducing the signal intensity at high pulse length,³⁹,⁴⁰ the experiments were also performed at 40 kHz, but no differences were observed between these two spinning rates (not shown). All the nutation curves present similar variation with that of H₂¹⁷O (C_q = 0), as well as between different O(Np)₄/(U)₁−₄⁺ units for the same composition than through the composition range under study. A maximum intensity is detected around 4 μs at 20 W and 12.5 μs at 2 W. These results show that a perfect tetrahedral environment remains around the oxygen atoms even if An–O bond distortions are expected. The broadening observed for the different O unit with increasing Np atoms and content is therefore most probably due to anisotropic interactions created, among others, by the presence of delocalized 5f-electrons and actinide–oxygen bond variations. Considering these results, all the spectra were acquired using the conditions τ_{90°,n.s.} = τ_{90°,s} (with “n.s.” the so-called “nonelective” excitation expected in liquids and for cubic substances and “s” the so-called “selective excitation” expected when C_q ≠ 0³⁴).

The quantitative nature of NMR is one of the key attributes of this technique allowing site population determination for each different oxygen site. The quantitative spectra previously presented were obtained after determination of the 90° pulse excitation but also of the longitudinal relaxation times (T₁) for each O atom in the different environments in all of the samples. The T₁ variations for each different O unit and for all the chemical compositions are presented in Figure 7. For the five types of O subunits, T₁ decreases with the increasing Np content in the sample. This result shows that the addition of Np atom in the vicinity of the O atom leads to a decrease of T₁.
in perfect agreement with faster relaxation time for NpO₂ than UO₂.

In Figure 8, the integrated intensity of each O(U₁₋ₓNpₓ)₁₋ₓ (with \(0 \leq y \leq 4\)) unit obtained experimentally were compared to those expected for a randomly distributed network (RDN). The experimental integrated intensities were obtained by fitting the center band, but also the satellite transition intensities. The chemical shift anisotropy (CSA) values are given in Table 2. The values of the anisotropy of the CSA tensor increase when a cation with a different nature is added to the pure dioxide in agreement with the fwhm evolution. The so-called \(n = 0\) spinning sideband is expected to contribute to the signal intensity of the central transition as it lies below this one. Due to the important spectral broadening, it has unfortunately not been possible to identify properly this later even by changing the spinning rate. The intensity of this spinning sideband has therefore been added in the error bars considering the intensity of the highest spinning sidebands. The theoretical values expected for a RDN were calculated using a binomial distribution function given by

\[
P(n\Omega) = \Omega \times p^n \times (1 - p)^{4-n}
\]

where \(\Omega\) is the number of permutations, \(p = x\) is the probability of finding a Np in U₁₋ₓNpₓO₂, and \(n\) is the number of possible ways Np or U can be arranged around the O atom. As the Np50 sample is composed of two crystalline phases that cannot clearly be differentiated, this sample was excluded from the following analysis. A good agreement is found between theoretical and experimental relative intensities for each solid solution confirming the U/Np random distribution. This later observation do also confirm the previous attributed to all the O(U₁₋ₓNpₓ)₁₋ₓ (with \(0 \leq y \leq 4\)) units.

4. CONCLUSION

A complete analysis of the long-range and atomic scale structure of U₁₋ₓNpₓO₂ samples was performed for the first time using a combination of XRD, XANES, and high resolution solid-state \(^{17}\)O MAS NMR. The robustness of the external gelation synthesis was proven from XRD and Vegard law behavior. In addition, the presence of local distortions due to the coexistence of cations with different ionic radius was observed thanks to the micro strain. XANES has shown that U and Np remain tetravalent for all the compositions. Using \(^{17}\)O MAS NMR, the different oxygen environments were detected thanks to their different paramagnetic shifts. A linear increase of the shifts with decreasing Np content was observed for each O(U₁₋ₓNpₓ)₁₋ₓ (with \(0 \leq y \leq 4\)) unit consistent with the existence of a solid solution. For the same type of unit, the slope of this linear fit increases as the O is surrounded by increasing numbers of Np neighbors. Local distortions around the oxygen atom due to the substitution of \(^{4+}\)U by the smaller \(^{4+}\)Np and transfer of positive electron from the cation lead to a variation of the fwhm of the individual O(U₁₋ₓNpₓ)₁₋ₓ components. Nutation experiments show that no bond angle distortions exist around the oxygen atoms with a perfect tetrahedral environment maintained. Longitudinal relaxation times studies show a decrease of \(T₁\) with increasing Np content in agreement with a larger \(T₁\) for UO₂ than for NpO₂. Finally, the comparison of the intensities in a randomly distributed network and those obtained based on the relative experimental area of the peaks shows a good agreement. The combination of all these techniques and, particularly, the use of solid-state high resolution NMR open new routes for the fine characterization of radioactive materials. In addition, the NMR data should be a strong basis for development of future solid-state NMR calculations, which are not yet possible for 5f-electrons due to, among others, the lack of experimental results.

Caution! As these uranium—neptunium mixed oxides present considerable radiotoxicity hazards, they were all handled under carefully controlled dedicated laboratories at the Institute for Transuranium Elements in Karlsruhe. All steps were performed in hermetically sealed gloveboxes maintained at a relative negative pressure with respect to the laboratory.
The Journal of Physical Chemistry C

**Table 2. Values of Anisotropy of the Chemical Shift Anisotropy (CSA) Tensor (ΔCSA) and Asymmetry Parameters of the CSA Tensor (ηCSA) for the Six U1−xNp,O2 Compounds**

| compound name                  | O(U)x(Np)y units | Np1   | Np5   | Np20  | Np50  | Np75  | Np85  |
|--------------------------------|------------------|-------|-------|-------|-------|-------|-------|
| ΔCSA                           | O(U)x            |       |       |       |       |       |       |
| O(U)x                           | 388.53           | 510   | 493   | 610   |       |       |       |
| O(U)x(Np)y                      | 1440.48          | 1484  | 1314  | 1441  | 1411.40 | 1451  |       |
| O(U)x(Np)y                      | 1100             | 1497  | 1435  | 1492.82 | 1343  |       |       |
| O(U)x(Np)y                      |                   | 1169  | 999.42 | −1344 | −521  |       |       |
| O(Np)y                          | 1098             | −640.74 |       |       |       |       |       |
| ηCSA                           | O(U)x            | 0     | 0     | 0     | 0.1   |       |       |
| O(U)x(Np)y                      | 0.3              | 0.1   | 0     | 0.1   | 0.83  | 0.6   |       |
| O(U)x(Np)y                      | 0.2              | 0.5   | 0.11  | 0.8   |       |       |       |
| O(U)x(Np)y                      |                   | 0.8   | 0.57  | 0.1   |       |       |       |
| O(Np)y                          | 0.4              | 0.88  | 0.6   |       |       |       |       |

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