Phase decomposition of γ-U (bcc) in U-10 wt% Mo fuel alloy during hot isostatic pressing of monolithic fuel plate

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Eutectoid decomposition of γ-phase (cI2) into α-phase (oC4) and γ'-phase (tI6) during the hot isostatic pressing (HIP) of the U-10 wt% Mo (U10Mo) alloy was investigated using monolithic fuel plate samples consisting of U10Mo fuel alloy, Zr diffusion barrier and AA6061 cladding. The decomposition of the γ-phase was observed because the HIP process is carried out near the eutectoid temperature, 555 °C. Initially, a cellular structure, consisting of γ-phase surrounded by α-phase, developed from the destabilization of the γ-phase. The cellular structure further developed into an alternating lamellar structure of α- and γ'-phases. Using scanning electron microscopy and transmission electron microscopy, qualitative and quantitative microstructural analyses were carried out to identify the phase constituents, and elucidate the microstructural development based on time-temperature-transformation diagram of the U10Mo alloy. The destabilization of γ-phase into α- and γ'-phases would be minimized when HIP process was carried out with rapid ramping/cooling rate and dwell temperature higher than 560 °C.

1. Introduction

The replacement of high-enriched uranium (HEU) fuel with low-enriched uranium (LEU) fuel is the primary goal of Materials Management and Minimization Reactor Conversion (MMRRC) program, also known as Reduced Enrichment for Research and Test Reactor (RERTR) program [1–4]. Uranium based metallic fuel has garnered heavy consideration based on several advantageous properties, including high fissionable density, high thermal conductivity and good compatibility with cladding materials.

Monolithic design using U-10 wt%Mo (U10Mo) fuel alloy is being developed to achieve a higher uranium density for high flux reactors, because of high fissionable density and excellent irradiation behavior [5–7]. The monolithic fuel plates are fabricated via hot isostatic pressing (HIP) in order to encase the fuel plate foil in an Al-alloy based cladding. A Zr diffusion barrier is typically placed between the U10Mo fuel alloy and Al-alloy cladding to prevent interaction between the fuel and cladding. Previously, microstructure resulting from the interdiffusion and solid-state reaction between different constituents of the fuel plate was reported [8–13]. In addition, within the fuel alloy, Meyer et al. [11] reported the presence of “chemical banding” with inhomogeneous Mo content within the fuel alloy, and Jue et al. [12] identified γ-phase decomposed regions consisting of α- and γ'-phases due to heat treatment near the eutectoid temperature. The low-Mo (~5 wt %) and high-Mo (~13 wt%) phases were assumed as α and γ', respectively [12]. In this study, decomposition of γ-U10Mo (γ) into α-U (α) and γ'-U2Mo (γ') was examined in detail by electron diffraction and quantitative image analysis, because the HIP process is carried out near the eutectoid temperature [14].

2. Background

Pure U exhibits two allotropic transformations, and thus three different crystal structures. Orthorhombic α-, tetragonal β- and body-centered cubic (BCC) γ-phases are stable up to 667 °C, between 667 °C and 771 °C, and above 771 °C, respectively [14–17]. The α-U has undesirable characteristics as a nuclear fuel such as low hardness, rapid oxidation, low corrosion resistance and anisotropic irradiation behavior [18–21]. A single crystal of α shortens in a-axis and extends in b-axis, and there is no significant change in c-axis...
during irradiation under 500 °C [16,22], leading to dimensional instability [23,24].

In contrast, the γ-phase exhibits favorable properties as a nuclear fuel. The bcc γ displays good dimensional stability because of isotropic expansion during irradiation [24]. The γ-phase also has a higher corrosion resistance to water, both liquid and vapor [25]. In order to retain the metastable γ-phase below 560 °C, rapid cooling or alloying additions has been employed [26,27]. It is difficult for pure U to exist as γ-phase even after quenching, because α-phase is thermodynamically favorable at low temperature [24,26,27]. Thus, alloying U with Mo or Zr or Nb has been adopted to stabilized the γ-phase and produce good irradiation behavior [28-30]. Table 1 shows the characteristics of potential alloying elements in U. Mo, Zr and Nb have been considered because of appropriate characteristics such as low neutron absorption, high melting point, high thermal conductivity and good corrosion resistance among all elements. U-Mo has been selected for higher uranium density, stabilization of γ-phase, favorable properties listed in Table 1, despite the higher thermal neutron absorption and thermal conductivity, and excellent irradiation behavior [5,6,24,29,31].

The γ decomposition is affected by the Mo concentration in the alloy. The time-temperature-transformation (T-T-T) diagram presented in Fig. 1 demonstrates that the decomposition is hindered by the Mo addition. With the Mo content 8 wt%, or above, the γ-phase is stabilized even from slow cooling with acceptable uranium density for good irradiation behavior [32].

Despite the sluggish kinetics, according to the equilibrium phase diagram, the high-temperature γ-phase undergoes a eutectoid decomposition upon cooling into low-temperature orthorhombic α- and tetragonal γ-phases as shown in Fig. 2 [14,33-36]. The equilibrium temperature of decomposition is relevant to the high temperature employed during co-rolling and HIP for the monolithic fuel system. Table 2 lists selected and relevant properties of the γ-, α- and γ-phases. Generally, the microstructure related to the decomposition was observed first at γ-phase grain boundaries [28,37].

Fig. 3 presents the T-T diagram that illustrates the time-dependent phase decomposition of the γ-phase in U10Mo alloy. The T-T-T diagrams of Repas et al. [26] and Peterson et al. [27] were constructed based on hardness, dilatometry, XRD and metallographic analyses. Saller et al. [37] also reported measurements of the electrical resistivity and hardness for U-21 at.% Mo alloy, indicating a structural transformation from BCC to an ordered body-centered tetragonal through isothermal transformation study at 500 °C. The transformation began approximately at 27 h and completed around 170 h. It should be noted that a reverse decomposition, i.e., α and γ' into γ was observed during irradiation testing [38,39].

Fig. 4 shows a typical backscatter electron (BSE) micrograph of the HIP fuel plate investigated in this study. The U10Mo fuel alloy was cladded in AA6061 with a Zr diffusion barrier in-between. Microstructure of the U10Mo alloy in HIP fuel plates was characterized by scanning electron microscopy (SEM), transmission electron microscopy (TEM), and a variety of analytical techniques including X-ray energy dispersive spectroscopy (XEDS). Quantitative image analyses were carried out to estimate volume fraction of decomposed regions based on BSE micrographs.

### 3. Experimental details

The U10Mo alloy was fabricated by arc-melting pellets of depleted U and 99.95% pure Mo which were cleaned with 30% nitric-acid to remove oxidation at the surface. The melting process was carried out in a vacuum of 1.3 × 10⁻³ Pa (10⁻⁵ Torr) containing less than 50 ppm oxygen, and the U10Mo alloy was turned and remelted three times in graphite crucible to ensure the homogeneity. The cast U10Mo alloy had an approximate dimension of 88.8 × 48.9 × 3.2 mm and a carbon content of approximately 700 ppm. The ingot was laminated in a carbon steel can using a pure Zr (99.9% pure) foil with a starting thickness of 250 μm on each surface. A ~30% nitric acid solution was used to etch the reactor-grade Zr prior to lamination. The Zr-laminated U10Mo coupon, pre-heated at 650 °C for 30 min in a furnace, was co-rolled 15 times to a thickness of approximately 0.46 mm. Subsequently, a post-rolling heat treatment was performed at 650 °C for 45 min. The hot-rolled and annealed foil was removed from the can, and cold-rolled to a final thickness of 0.33 mm. The final dimension of as-rolled foil was approximately 746 × 65 × 0.33 mm. The foil was exposed for approximately 130 min at 650 °C. The final thickness of the Zr diffusion barriers on each surface was approximately at 25 μm. Each small U-Mo foil sectioned from the as-rolled parent foil was used in this study. In general, each small foil sample exhibited the same microstructure in the co-rolled U10Mo/Zr laminate previously observed by Park et al. [9]. “AR” in this study is defined as the as-co-rolled/sectioned sample prior to the HIP process.

The polished AR sample (each U10Mo/Zr mini foil) was cleaned using a mixture of nitric and hydrofluoric acid (2.5% HF, 35% HNO₃ and 62.5% H₂O), and any residual oxide scale of AA6061 cladding was removed by etching using a 1.85 M NaOH and then a 30% nitric acid solution. After cleaning, the AR samples were rinsed with ethanol and stacked with AA6061 (in wt.% nominal composition of 0.65Si–0.35Fe–0.28Cu–0.08Mn–1.09Mg–0.20Cr–0.13Zn–0.08Ti–AI balance) cladding pieces. The U10Mo/Zr minifoil was then HIPed with the AA6061 at various temperatures (520, 540, 560 and 580 °C) and durations (45, 60, 90 180 and 345 min). The U10Mo/Zr/AA6061 assemblies were placed into stainless steel HIP cans with tool-steel strong-backs that were welded with corners open for degassing in an argon atmosphere glove box. The assemblies were subsequently leak-checked and vacuum degassed at 315 °C for 3 h. After loading into a laboratory-size HIP, the HIP cans were heated with heat-up and cool-down rates (H/C rate) of 35 and 70 °C/h to the target temperature. The isostatic pressure of 103 MPa (15 ksi) for HIP process was applied in an argon environment. The samples characterized as functions of HIP parameters in this study are listed in Table 3.

After fabrication, the fuel plates were cross-sectioned and mounted in epoxy. The mounted fuel plates were polished using silicon carbide papers (240, 600, 800 and 1200 grits) with an ethanol lubricant. Final polishing was carried out utilizing diamond pastes (3 and 1 μm) with an oil lubricant in an Ar glove box to prevent oxidation and contamination. Microstructural features in U10Mo solid solution were examined utilizing a Zeiss ULTRA-55 field-emission SEM with XEDS capabilities. High-contrast in backscatter electron micrographs were utilized to distinguish the different phases and quantitative analyses. Transmission electron microscopy (FEI/Tecnai™ F30) equipped with Fischione™ high angle annular dark-field (HAADF) and XEDS were used to examine the detailed microstructures and identify the phase constituents via selected area electron diffraction (SAED) and high resolution

### Table 1: Elemental characteristics of Mo, Zr and Nb [42-44].

<table>
<thead>
<tr>
<th>Property</th>
<th>Mo</th>
<th>Zr</th>
<th>Nb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermal neutron absorption (barn)</td>
<td>2.48</td>
<td>0.18</td>
<td>1.15</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>10.22</td>
<td>6.49</td>
<td>8.35</td>
</tr>
<tr>
<td>Melting point (°C)</td>
<td>2623</td>
<td>1854</td>
<td>2477</td>
</tr>
<tr>
<td>Thermal expansion coefficient (K⁻¹)</td>
<td>5.35 × 10⁻⁶</td>
<td>5.78 × 10⁻⁶</td>
<td>7.34 × 10⁻⁶</td>
</tr>
<tr>
<td>Thermal Conductivity (W/m·K⁻¹)</td>
<td>138</td>
<td>22.7</td>
<td>54</td>
</tr>
<tr>
<td>Corrosion resistance</td>
<td>Good</td>
<td>Good</td>
<td>Good</td>
</tr>
</tbody>
</table>
Fig. 1. Time-temperature-transformation diagram of U-Mo alloys as a function of Mo content [26].

Fig. 2. (a) Equilibrium phase diagram for the U-Mo system with (b) a magnified view of the γ Û a+γ eutectoid reaction [14].
TEM (HRTEM). The TEM samples were prepared by focused ion beam and an in-situ lift-out process was utilized for sample extraction (FIB-INLO; FEI™ TEM200) using site-specific capability. Typical phase constituents and microstructural developments due to interdiffusion and reaction at the U10Mo/Zr and Zr/AA6061 interfaces in these samples were reported previously [8,9]. This paper reports, in detail, the phase transformation of $\gamma$-phase in the U10Mo fuel alloy as functions of HIP temperature, holding time and ramping/cooling (H/C) rate as listed in Table 3.

### Result and discussion

The AR sample before HIP process exhibited a small amount of cellular structure which consists of $\alpha$, $\gamma$, and $\gamma'$-phases as shown in Fig. 5. The cellular structure is assumed as the initial feature associated with onset of $\gamma'$-formation (i.e., start of the $\alpha + \gamma + \gamma'$ phase region in Fig. 3) prior to development of lamellar structure, and is also assumed to nucleate at heterogeneous sites, such as grain boundaries. Only the cellular structure without any lamellar

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### Table 2

Selected properties of $\gamma$-U (U – 10 wt% Mo), $\alpha$-U and $\gamma'$-U2Mo phases [14,23,32,40,41,45–51,52].

<table>
<thead>
<tr>
<th>Property</th>
<th>$\gamma$-(U10Mo) (at 800 °C)</th>
<th>$\alpha$-U (at 627 °C)</th>
<th>$\gamma'$-(U2Mo) (at 500 °C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mo solubility (wt.%)</td>
<td>10</td>
<td>0.2</td>
<td>16</td>
</tr>
<tr>
<td>Strukturbericht designation</td>
<td>A2</td>
<td>A20</td>
<td>C11b</td>
</tr>
<tr>
<td>Prototype</td>
<td>W</td>
<td>$\alpha$-U</td>
<td>MoSi$_2$</td>
</tr>
<tr>
<td>Crystal structure</td>
<td>Body-centered cubic</td>
<td>C-centered orthorhombic</td>
<td>Body-centered tetragonal</td>
</tr>
<tr>
<td>Lattice parameter (Å)</td>
<td>a 3.474</td>
<td>b 2.854</td>
<td>c 3.427</td>
</tr>
<tr>
<td></td>
<td>b 3.474</td>
<td>5.869</td>
<td>4.955</td>
</tr>
<tr>
<td></td>
<td>c 3.474</td>
<td>3.427</td>
<td>9.834</td>
</tr>
<tr>
<td>Space group</td>
<td>Im3m (229)</td>
<td>Cmc2 (63)</td>
<td>I4/mmm (139)</td>
</tr>
<tr>
<td>Pearson symbol</td>
<td>cl2</td>
<td>oc4</td>
<td>tI6</td>
</tr>
<tr>
<td>Young’s modulus (GPa)</td>
<td>98</td>
<td>117</td>
<td>102</td>
</tr>
<tr>
<td>Hardness (GPa)</td>
<td>3.1</td>
<td>2.3</td>
<td>N/R</td>
</tr>
<tr>
<td>Poisson ratio</td>
<td>0.35</td>
<td>0.26</td>
<td>0.40</td>
</tr>
<tr>
<td>Thermal conductivity (W-m$^{-1}$K$^{-1}$)</td>
<td>3.65</td>
<td>34</td>
<td>N/R</td>
</tr>
<tr>
<td>Thermal expansion coefficient ($10^{-6}$ K$^{-1}$)</td>
<td>19.2</td>
<td>18.0</td>
<td>14.7</td>
</tr>
<tr>
<td>Specific heat (J-K$^{-1}$g$^{-1}$)</td>
<td>0.194</td>
<td>0.192</td>
<td>0.154</td>
</tr>
</tbody>
</table>

### Table 3

Hot isostatic pressing parameters of monolithic fuel plate specimens examined in this study.

<table>
<thead>
<tr>
<th>Sample identification</th>
<th>Temp. (°C)</th>
<th>Hold time (min)</th>
<th>Ramp rate (°C/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>52-90-280</td>
<td>520</td>
<td>90</td>
<td>280</td>
</tr>
<tr>
<td>54-90-280</td>
<td>540</td>
<td>90</td>
<td>280</td>
</tr>
<tr>
<td>56-90-280</td>
<td>560</td>
<td>90</td>
<td>280</td>
</tr>
<tr>
<td>58-90-280</td>
<td>580</td>
<td>90</td>
<td>280</td>
</tr>
<tr>
<td>56-45-280</td>
<td>560</td>
<td>45</td>
<td>280</td>
</tr>
<tr>
<td>56-60-280</td>
<td>560</td>
<td>60</td>
<td>280</td>
</tr>
<tr>
<td>56-180-280</td>
<td>560</td>
<td>180</td>
<td>280</td>
</tr>
<tr>
<td>56-345-280</td>
<td>560</td>
<td>345</td>
<td>280</td>
</tr>
<tr>
<td>56-90-35</td>
<td>560</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>56-90-70</td>
<td>560</td>
<td>70</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 3. Time-temperature-transformation diagram of the U - 10 wt.% Mo alloy [24]. “AR” represents the time-coordinate of the alloy in as-co-rolled condition, based on microstructural analysis.

Fig. 4. Typical backscatter electron micrograph of the U10Mo monolithic fuel encased in AA6061 with Zr diffusion barrier, produced by co-rolling and hot-isostatic pressing (sample 56-345, hot-isostatic pressed at 560 °C for 345 min) [9].

Fig. 5. Backscatter electron micrograph of the cellular structure observed from the decomposition of $\gamma$ into $\alpha$ and $\gamma'$ found in the as-co-rolled U - 10 wt.% Mo alloy prior to hot-isostatic pressing.

U10Mo fuel alloy as functions of HIP temperature, holding time and ramping/cooling (H/C) rate as listed in Table 3.
features were observed in the AR sample. The cellular structure is also characterized by the Mo depleted regions surrounding the Mo-rich nuclei as shown by the inset in Fig. 5. Based on these observations, AR sample is assumed to exhibit features characteristic of an U10Mo alloy corresponding to a time coordinate of 25 h in the U10Mo T-T-T diagram as labeled in Fig. 3. In other words, subsequent HIP and the corresponding characterization results assume that the starting coordinates for U10Mo alloy fuel in T-T-T diagram is at 25 h as labeled in Fig. 3.

Fig. 6 shows microstructure of U10Mo fuel alloy HIP'ed at 560 °C for 90 min with a H/C rate variation of 280, 70 and 35 °C/hour. The micrograph in Fig. 6(a) from the U10Mo alloy with the higher H/C rate of 280 °C/h shows cellular regions along grain boundaries of U10Mo alloy. Cellular structure is better defined than the AR sample, and some of the cellular reaction was observed within the grains of the U10Mo solid solution. For U10Mo alloy HIP'ed with H/C rate of 70 °C/h, a clear development of lamellar structure was observed in addition to the cellular structure as shown in Fig. 6(b). The lamellar structure is assumed to begin from the cellular precipitates along the grain boundaries, and grow into the grain of γ'- Rich
(U10Mo). The largest volume fraction of lamellar structure region was observed in the sample with 35 °C/h H/C rate. The lamellar structure of α and γ is presented in detail with a higher magnification BSE micrograph in Fig. 7. Fig. 8 shows that the volume fraction estimated for the lamellar structure region resulting from decomposition of γ increases as the H/C rate decreases. This observation corresponds to the fact that slow H/C rate exposes the U10Mo alloy in the three-phase (α + γ + γ′) region the longest as illustrated on the T-T-T diagram in Fig. 9.

A selected region highlighted by black rectangle in Fig. 6(c) was prepared by FIB-INLO for TEM analyses presented in Figs. 10 and 11. The SAED patterns presented in Fig. 11 were collected from circled regions in Fig. 10(a). The HAADF and bright-field images shown in Fig. 10 along with SAED patterns presented in Fig. 11 were employed to identify the decomposition of bcc-γ into alternating orthorhombic-α and tetragonal-γ′ lamellar structure. These phases were identified by analyzing the SAED patterns, and the crystallographic characteristics of each phase corresponded well to those reported in Table 2. Variation in Mo composition in γ-phase with ~10 wt% Mo, Mo-depleted α, Mo-enriched γ′ was also confirmed by extensive analyses by XEDS equipped on TEM. Other regions consisted of γ and α, without γ′. Therefore the presence of transition region from α+γ to α+γ+γ′ in the T-T-T diagram appears to be consistent with the current TEM analyses.

Fig. 12 shows microstructural variation in U10Mo fuel alloy as a function of HIP temperature. The time at HIP temperature and H/C rate remained constant for 90 min and 280 °C/hour, respectively, for these samples. In general, the volume fractions of cellular and lamellar structure regions decreased with an increase in HIP temperature as presented in Fig. 13. The HIP temperature variation examined in this study is close to the transition that separates γ and α+γ (and α+γ and γ) on the upper-left-corner of the T-T-T diagram as presented in Fig. 14. Repas et al. [26] postulated a transition between α + γ′ and α + γ at 573 °C (lower critical temperature), and between α + γ′ and γ at 580 °C (upper critical temperature) in the T-T-T diagram. However, these temperatures have been subject to some scrutiny by other investigations [14,29,37,38,40,41]. Table 4
summarizes the critical transition temperatures reported in literature [14,26,39] for the U10 wt\% Mo alloy. In this study, the samples HIP’ed at 520 and 540 °C show the lamellar structure of α and γ0 within the γ matrix, while only cellular structure was observed when the samples were HIP’ed at 560 °C and above. This demonstrates the transition of α and γ’ into α and γ starts above 540 °C and below 560 °C.

Effects of HIP holding time was examined with U10Mo alloys HIP’ed at 560 °C with H/C rate of 280 °C/h as presented in Fig. 15. There was no significant change in the amount of decomposed region because, upon heating, a transformation from $\alpha + \gamma'$ (or $\alpha + \gamma + \gamma'$) into $\alpha + \gamma$ in the temperature range of 555–580 °C is expected as shown in Fig. 16. Given the HIP temperature of 560 °C, the volume fraction of decomposed region showed no significant trend with HIP holding time, up to 345 min, as shown in Fig. 17. However such an observation may change when HIP temperature is selected at 540 °C or below (e.g., 555 °C according to Okamoto) according to the T-T-T diagram. Therefore, in order to minimize the decomposition of the γ-phase into α- and γ’-phases in the U10Mo fuel, HIP

| Repas [26] | 573 °C | 580 °C |
| Kim [39] | N/R | >570 |
| Okamoto [13] | 555 °C | ~580 °C |
utilizing at least 560 °C and the highest H/C rate would be recommended. HIP at 580 °C may allow longer duration without concern for any decomposition.

A related consideration for higher HIP temperature is the interdiffusion and reaction among U10Mo fuel alloy, Zr diffusion barrier, and AA6061 cladding alloy. In our previous study, HIP did not cause any observable growth of the interaction layer at the U10Mo/Zr interface within the temperatures and holding time...
range examined [9]. Development of the interaction layer containing α-U, UZr2, and Mo2Zr at the U10Mo/Zr interface primarily occurred during the co-rolling process (650 °C) prior to HIP. However, the intermetallic (Al2Si)3Zr layer at the AA6061/Zr interface was observed to grow with the HIP duration, and the growth followed the Arrhenius relation with an activation energy of 457.54 ± 27.60 kJ/mol. Fig. 18 illustrates the growth kinetics of (Al2Si)3Zr at the AA6061/Zr interface [9]. Therefore the HIP process for the monolithic fuel system should employ higher temperature (e.g., 560° and 580 °C) to (1) ensure the highest quality of adhesion at the U10Mo/Zr and AA6061/Zr interfaces [9,10], and (2) minimize the γ-phase decomposition, (3) however, without excessive growth of (Al2Si)3Zr layer at the AA6061/Zr.

5. Conclusion

U − 10 wt% Mo alloy for the monolithic fuel plates encased in AA6061 cladding with Zr barrier was characterized as functions of HIP parameters with an emphasis on decomposition of γ-phase into α- and γ′-phases. From TEM SAED, the constituent phases of the U10Mo alloy consisted of γ cI2 Im3m (3.474 × 3.474 × 3.474 Å), α Oc4 Cmcm (2.854 × 5.869 × 4.955 Å) and γ′ t64 I4/mmm (3.427 × 3.427 × 5.834 Å). The α- and γ′-phases formed by decomposition of the γ-phase because the HIP process was carried out near the eutectoid temperature with varying HIP parameters including ramping/cooling rate, temperature, and holding time. Initially, a cellular structure of α and γ′ from the decomposition of γ developed discontinuously on grain boundaries, and evolved into lamellar structures of alternating α and γ′. Volume fraction associated with decomposition (e.g., cellular and lamellar structure) decreased with (1) an increase in ramping/cooling rate, (2) an increase in HIP temperature due to γ-phase stabilization at higher temperature. There was no significant development of decomposed lamellar structure when the HIP was carried out at 560 °C for up to 345 min.

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