POST-IRRADIATION EXAMINATION
OF THE FUEL ROD CART-C1 N° 4

by

A. FRIGO, R. KLERSY, K.H. SCHRADER, A. SCHÜRENKÄMPER
(EURATOM)
A. BENVENUTI, G. CAMONA and F. MANTEGA
(CISE)

1972

Joint Nuclear Research Centre
Ispra Establishment - Italy
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Post-irradiation examinations on the fuel rod have been carried out. The results are presented in the report. Non destructive gamma-scanning analysis shows a migration of Cs inside the fuel and a more or less homogeneous distribution of the fission products Zr, Nb, Ce-Pr and Ru. Metallographic studies show zirconium hydrides over the whole thickness and oxide layers on the surfaces of the cladding. The welding zone appears perfect. The alteration of the microstructure of the fuel due to the irradiation is shown.
COMMISSION OF THE EUROPEAN COMMUNITIES

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ABSTRACT

Post-irradiation examinations on the fuel rod have been carried out. The results are presented in the report. Non destructive gamma-scanning analysis shows a migration of Cs inside the fuel and a more or less homogeneous distribution of the fission products Zr, Nb, Ce-Pr and Ru. Metallographic studies show zirconium hydrides over the whole thickness and oxide layers on the surfaces of the cladding. The welding zone appears perfect. The alteration of the microstructure of the fuel due to the irradiation is shown.

KEYWORDS

FUEL RODS                   CERIUM 144
RADIATION EFFECTS           PRAESODYMIUM 144
URANIUM DIOXIDE             ZIRCONIUM 95
GAMMA FUEL SCANNING         NIOBIUM 95
FISSION PRODUCTS            MICROSTRUCTURE
DIFFUSION                   FUEL CANS
DISTRIBUTION                ZIRCALOY
CESIUM 134                  ZIRCONIUM HYDRIDES
CESIUM 137                  ZIRCONIUM OXIDES
RUTHENIUM 106               LAYERS
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*) Manuscript received on May 23, 1972"
1. **Fuel rod characteristics and irradiation conditions.**

This report deals with postirradiation examinations of rod No. 4 (peripherical) of CART C-1 fuel element central bundle.

Some data of general interest are listed below:

- **Rod characteristics**

  **Cladding**
  - material: Zircaloy-2 reactor grade
  - supplier: CEFILAC (Paris)
  - nom. size: O.D.: 20 mm
  - thickness: 1 mm

  **Fuel**
  - material: natural UO$_2$ in synthesized pellets
  - manufacturer: FCEC Laboratories (pellets) CNEN Saluggia
  - nom. size: O.D.: 17.80 mm
  - height: 26.00 mm
  - dish depth: 1 mm
  - plug material: Zircaloy-2 reactor grade
  - filling atmosphere: A 90% - He 10%
  - manufacturer: FCEC Laboratories CNEN Saluggia

- **Operative data of the rod**

  - $\bar{\phi}d$ average on the rod: 51.5 W/cm (calculated) with temporal peak (about 1 hour) at 58 W/cm
  - average specific power: nominal 26.7 W/g-UO$_2$; max. temporal 30.0 W/g-UO$_2$
  - average burn-up: 1685 MWD/t U
- cooling: water-steam mixture
- coolant pressure and temperature: 50 kg/cm²; 260°C
- sheath temperature: 270°C
- UO₂ central temperature (calculated): 2000°C
- number of days in power (higher than 50% of the nominal power): 80
- number of days in temperature: 110
2. Non destructive analysis.


The used apparatus consists of a mechanical part installed inside the hot cell (Fig. 1), a collimator system mounted in the concrete wall of the cell and the equipment for gamma-ray analysis outside the cell. An electronic control unit allows to run measurements automatically following a preselected program for the displacement of the fuel rod and for counting time. In the described experiments the fuel rod was rotating during counting but in a fixed position with regard to the axial direction. A cylindrical collimator with a diameter of 1 mm was used. The total gamma-activity in the energy range from 100 to 2,500 KeV has been measured in steps of 0.5 mm using a NaI-crystal. The gamma-spectre have been measured each 20 mm using a Ge-Li crystal.

2.1.1. Total gamma-activity measurements.

Fig. 2 shows the distribution of the total gamma-activity along the fuel rod. The position of the different fuel pellets inside the rod is indicated. The most significant result is the appearance of significant activity peaks. These peaks correspond to the interfaces of fuel pellets. It must be mentioned that the pellets had a dishing. Very often such a dishing leads to pronounced minima of the measured total gamma-activity. In the present case the contrary has been observed. This is an indication that a transport of fission products from the fuel to the interfaces of the pellets have taken place during irradiation.

The mean value of the gamma-activity shows a slightly increase going from the top to the bottom of the rod. This corresponds to the flux distribution during irradiation. The relatively high scattering of the activity is indicating a cracked fuel.
2.1.2. Gamma-spectrometry.

A spectra of the fission products of the irradiated fuel is shown in fig. 3. The isotopes which have been detected in a significant concentration are given in the table.

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Gamma-ray energy (KeV)</th>
<th>Half life (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ru 106</td>
<td>512 622</td>
<td>372.5</td>
</tr>
<tr>
<td>Cs 134</td>
<td>605 796</td>
<td>749</td>
</tr>
<tr>
<td>Cs 137</td>
<td>662</td>
<td>10.990</td>
</tr>
<tr>
<td>Ce-Pr 144</td>
<td>696</td>
<td>285</td>
</tr>
<tr>
<td>Zr 95</td>
<td>724 756</td>
<td>65</td>
</tr>
<tr>
<td>Nb 95</td>
<td>766</td>
<td>35.1</td>
</tr>
</tbody>
</table>

The surface area of an observed peak is roughly proportional to the quantity of the corresponding isotope, which is present in the section under examination. This is only an approximation because different radial distribution of the fission product could lead to different surface areas due to the self adsorption of the X-rays inside the fuel rod itself.

The spectra of the fig. 3 has been measured in the center of a fuel pellet. Fig. 4 shows a spectra measured in a position corresponding to the interface of two pellets. (This position is indicated by A in the figures 5 and 6). It can be seen, that the activity of the Cs 137 and of the Cs 134 is increased in comparison with the other isotopes at the interface of two pellets. In the figures 5 and 6 the distribution of the concentration of the measured isotopes in the fuel rod is plotted together with the total activity. As already mentioned for the total gamma-activity the mean values of the activities of the single
isotopes show also a little increase going from the top to the bottom of the rod. For the isotopes Cs 137 and Cs 134 one observes a high scattering of the measured activities and especially a significant increase of these activities at the interfaces of the fuel pellets. This is not the case for the other fission products.

2.2. Dimensional measurements.

The maximum and minimum diameter of the fuel rod has been measured in three axial positions: in the middle of the rod and at 50 mm distance from the end plugs. A micrometer was used for these measurements. The results are listed in the table below. The corresponding values before irradiation are written in brackets.

<table>
<thead>
<tr>
<th>POSITION 1 (50 mm from the marked plug)</th>
<th>POSITION 2 (center of the rod)</th>
<th>POSITION 3 (50 mm from the non marked plug)</th>
</tr>
</thead>
<tbody>
<tr>
<td>min</td>
<td>max</td>
<td>min</td>
</tr>
<tr>
<td>mm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20,100</td>
<td>20,120</td>
<td>20,075</td>
</tr>
</tbody>
</table>

The length of the rod, measured between the plane surfaces of the two plugs, was $487.56 \pm 0.02$ mm.
3. **Destructive analysis.**

3.1. Preparation of samples for metallographic studies.

Metallographic analysis has been carried out on 7 samples. The location of these samples in the fuel rod is shown in Fig. 7. For the examination of the cladding (samples 5-6-7) the uranium dioxide has been removed mechanically. The samples were then embedded in copper bakelite mixed resin and submitted to metallographic operations (grinding, polishing, chemical etching) to make visible the zirconium oxide layer, the zircaloy morphology and the presence and the orientation of zirconium hydrides. The same metallographic operations have been carried out with the plug (sample 2) of the fuel rod, in order to investigate in addition to the zircaloy properties the welding zone.

The samples 1,3,4 assigned for the fuel examination have been impregnated superficially with epoxy resin to avoid the loss of fuel fragments during metallographic handling. From the samples 8,9,10 the uranium dioxide has been removed. Using the remaining rings, the hydrogen content of the cladding material has been determined.

3.2. Cladding examinations.

3.2.1. Zircaloy.

The microstructure of the Zircaloy has been observed on samples 5,6,7 on the whole cladding thickness and on the whole circumference. The figures 8,9,10 show the typical microstructure of the external, the middle and the internal zones of the cladding. Excluding the regions extremely close to the surfaces an average grain size of 8.5 - 9 ASTM has been estimated. The microstructure of the middle region is slightly different from the microstructure of the adjacent regions. This was observed for each of the three specimens.
Especially the external zones (Fig. 8a, 9a, 10a) show highly deformed grains with considerable twinning and internal stresses due to the mechanical working. The middle zone (Fig. 8b, 9b, 10b) shows a more homogeneously arranged grain structure with appearance of some twins. Comparing equal zones of different samples one can observe that the grains structure is better developed in the samples 5 and 6 than in the sample 7. This is demonstrated if one compares fig. 8c with fig. 10c or fig. 9b with fig. 10b. There is no evidence that these differences are due to the cladding fabrication.

3.2.2. Zirconium hydrides.

The examination of the zirconium hydrides has been carried out on samples 5, 6, 7 after polishing and specific etching. The figures 11, 12 and 13 show for each of the three samples two complete radial zones of the cladding. The angle between the two zones is about 90°. The micrographs are representative for the overall structure of each sample. The distribution of the hydride is not uniform. There is a high concentration on the external surface. The concentration is decreasing up to a deepness of 85 - 100 μ and is rather constant in the remaining region. The hydrides have generally the form of needles, which are long and thick in the external region and somewhat smaller than in the other regions of the samples. Considering the needles with a length of more than 5 μ, the following numbers per mm² have been measured:

- sample 5 : 3470
- sample 6 : 2760
- sample 7 : 2950

Furthermore the number of needles with an inclination smaller than 40° relative to the radius has been determined in order to calculate the orientation factor \( f_n \).
The results are given in the table below.

<table>
<thead>
<tr>
<th>Sample 5</th>
<th>Sample 6</th>
<th>Sample 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>( f_n )</td>
<td>( f_n )</td>
<td>( f_n )</td>
</tr>
<tr>
<td>Fig. 11a</td>
<td>Fig. 12a</td>
<td>Fig. 13a</td>
</tr>
<tr>
<td>0.44</td>
<td>0.30</td>
<td>0.43</td>
</tr>
<tr>
<td>Fig. 11b</td>
<td>Fig. 12b</td>
<td>Fig. 13b</td>
</tr>
<tr>
<td>0.31</td>
<td>0.31</td>
<td>0.27</td>
</tr>
</tbody>
</table>

The values are especially high in all examined regions. It should be noted that for one sample, that means in one cross section, different values were obtained for different radial zones of the cladding.

3.2.3. Zirconium oxide.

A zirconium oxide layer has been observed on the external surface of the cladding. (fig. 14 and 15). This band of oxide was found on the whole periphery and its thickness is between 4.5\( \mu \) and 8\( \mu \). The most frequent value measured was about 6\( \mu \) and from a large number of measurements an average value of 6.1\( \mu \) has been determined for both samples 6 and 7. On both samples some oxide layers have been observed on the inner surface of the cladding (fig. 16 and 17). About 20% of the circumference was covered with oxide. The average value of the thickness was 3.5\( \mu \) with extreme values of 2.5\( \mu \) and 4.5\( \mu \).

3.2.4. Chemical determination of hydrogen content.

The hydrogen content of the cladding has been determined using the samples 8, 9, 10. Two measurements have been carried out on each sample.
The mean values are given below:

<table>
<thead>
<tr>
<th>Ring</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>54.6 H₂ (ppm)</td>
</tr>
<tr>
<td>9</td>
<td>73.3 &quot;</td>
</tr>
<tr>
<td>10</td>
<td>61.8 &quot;</td>
</tr>
</tbody>
</table>

3.3. Plug examination.

The region of the plug has been examined in order to check the plug-cladding junction and to get information on the microstructure of the region near the welding zone.

Fig. 18a shows the welding zone of the plug in polarized light. The junction appears perfect. The interface of the two original pieces can be recognized due to the presence of small needle shaped grains. In bright field observation no interface at all could be detected (fig. 18b).

The variation of the microstructure of the welding zone can be considered as normal. The interface between the normal grain structure of the cladding and the Widmannstätten structure is rather sharp (fig. 19a). Going to the region of the highest temperature grain size increases more continuously (figures 19b, 19c).

3.4. Fuel examination.

3.4.1. Grain size.

Four typical regions could be found on all samples (1,3,4). Starting from the periphery and going towards the centre of the fuel rod, we observe:

- a first region where the annealing during irradiation led to a sintering without secondary recrystallisation;
- a second region with grain growth, which occurred in a more or less orientated way;
- a third region with secondary recrystallisation, which led to large columnar equi-orientated grains;
- a fourth region with large columnar grains, which are radially orientated.

Differences between different samples consist only in the depth of the typical regions. The mean values are given in the following table, indicating also the reference figures, which show the microstructure.

<table>
<thead>
<tr>
<th></th>
<th>Sample 1</th>
<th></th>
<th>Sample 3</th>
<th></th>
<th>Sample 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (mm)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. zone</td>
<td>2.9</td>
<td>2.8</td>
<td>3.25</td>
<td>0.2</td>
<td>0.5</td>
</tr>
<tr>
<td>2. zone</td>
<td>0.2</td>
<td>0.5</td>
<td>0.15</td>
<td>0.35</td>
<td>0.6</td>
</tr>
<tr>
<td>3. zone</td>
<td>0.35</td>
<td>0.6</td>
<td>0.11</td>
<td>5.6</td>
<td>5.1</td>
</tr>
<tr>
<td>4. zone</td>
<td>5.6</td>
<td>-</td>
<td>5.2</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Fig. 23 shows the microstructure of the sample 3 over the whole section. From the table one can see that the diameter of the region with the columnar grain is decreasing going from sample 1 to sample 3 and 4.

It should be mentioned that only the external zone of the first region shows a microstructure which has practically the same characteristics as the original sintered material (figures 20a, 22a), whilst the zone near the interface with the second region shows a well arranged and defined grain structure (20b, 21a, 22b).
3.4.2. Porosity.

The porosity looks different for the different regions of the fuel section, but there is no difference between the three samples. In the external zone of the first region the fuel shows macroporosity and microporosity, as demonstrated in the figures 20a and 22a. Some of the pores have a size equal to the grain size, others are fixed and limited to grain boundaries (fig. 24a, 24b). In the adjacent zone of the first region macroporosity decreases in quantity, whereas the porosity on the grain boundary is still present. In the second region pores appear rather spherical, mainly fixed on grain boundaries (fig. 24d). Only a few pores have been detected inside the grains. In the third and fourth regions pores, definitly spherical, are more and more aligned at grain boundaries as the fuel center is approached (fig. 22c, 23).

3.4.3. Cracks.

Fig. 25 shows a typical example of the cracked fuel. In the external region one observes radial and circumferential cracks. In the adjacent region up to the fuel center, micro- and macrocracks appear. They are partially intergranular and show always a radial direction.
ACKNOWLEDGEMENTS

The Authors wish to thank Mr. Buscaglia G., Mr. Ghezzi E., Mr. Parisotto A., for their help in samples preparation and metallographic examinations.
Fig. 1 : Gamma-scanning equipment.

Fig. 2 : Total gamma activity.

Fig. 3 : Typical gamma spectrum of the fuel.

Fig. 4 : Gamma spectrum measured on the interface between two pellets.

Fig. 5 : Distribution of Cs 134, Ce-Pr 144 and Ru 106 in the fuel rod.

Fig. 6 : Distribution of Zr 95 and Cs 137 in the fuel rod.

Fig. 7 : Location of samples for post-irradiation examination.

Fig. 8 : Zircaloy morphology, sample 5.

Fig. 9 : Zircaloy morphology, sample 6.

Fig. 10 : Zircaloy morphology, sample 7.

Fig. 11(a,b) : Zirconium hydride in two radial zones (sample 5).

Fig. 12(a,b) : Zirconium hydride in two radial zones (sample 6).

Fig. 13(a,b) : Zirconium hydride in two radial zones (sample 7).

Fig. 14(a,b) : Zirconium oxide on the external surface (sample 6).

Fig. 15(a,b) : Zirconium oxide on the external surface (sample 7).

Fig. 16 : Zirconium oxide on the inner surface (sample 6).

Fig. 17 : Zirconium oxide on the inner surface (sample 7).
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Fig. 18(a,b) : Welding zone of the plug
   a : polarized light
   b : bright field

Fig. 19(a,b,c) : Heated zone of the plug
                  (polarized light).

Fig. 20(a,b,c) : Microstructure of the fuel
                  (sample 1).

Fig. 21(a,b,c,d) : Microstructure of the fuel
                   (sample 3).

Fig. 22(a,b,c,d) : Microstructure of the fuel
                   (sample 4).

Fig. 23 : Metallographic structure of fuel rod
          section.

Fig. 24(a,b,c,d) : Porosity of UO₂.

Fig. 25 : Cracked fuel.
Fig. 1 - Gamma-scanning equipment

A: fuel element
B: collimator
Fig. 3 - Typical $\gamma$-spectrum of the fuel.
Fig. 4 - $\gamma$-spectrum measured on the interface of two pellets.
Fig. 6 - Distribution of fission product activities.
Fig. 7 - Location of samples for post-irradiation examination
FIG. 8 - Zircaloy morphology (sample 5)
polarized light  200 x
FIG. 9 - Zircaloy morphology (sample 6)
polarized light  200 x
FIG. 10 - Zircaloy morphology (sample 7)
polarized light  200 x
FIG. 11 - Zirconium hydride in two radial zones (sample 5) 250 x
FIG. 12 - Zirconium hydride in two radial zones (sample 6) 250 x
FIG. 13 - Zirconium hydride in two radial zones. (sample 7) 250 x
FIG. 14 - Zirconium oxide on the external surface. (sample 6) 460 x

FIG. 15 - Zirconium oxide on the external surface. (sample 7) 460 x
FIG. 16 - Zirconium oxide on the inner surface. (sample 6) 460 x

FIG. 17 - Zirconium oxide on the inner surface. (sample 7) 460 x
FIG. 18 - Welding zone of the plug.
a: polarized light
b: bright field
FIG. 19 - Heated zone of the plug.
(polarized light)
200 x
FIG. 20 - Microstructure of the fuel (sample 1) 128 x
FIG. 21 - Microstructure of the fuel (sample 3) 128 x
FIG. 22 - Microstructure of the fuel.
(sample 4) 128 x
FIG. 23—Metallographic structure of fuel rod section.

32 x
FIG. 24 - Porosity of UO$_2$.

625 x
FIG. 25 - Cracked fuel.
To disseminate knowledge is to disseminate prosperity — I mean general prosperity and not individual riches — and with prosperity disappears the greater part of the evil which is our heritage from darker times.

Alfred Nobel
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