

DE LA RECHERCHE À L'INDUSTRIE



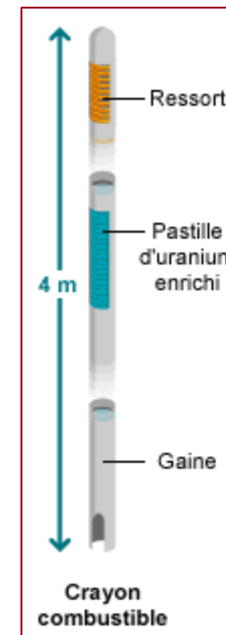
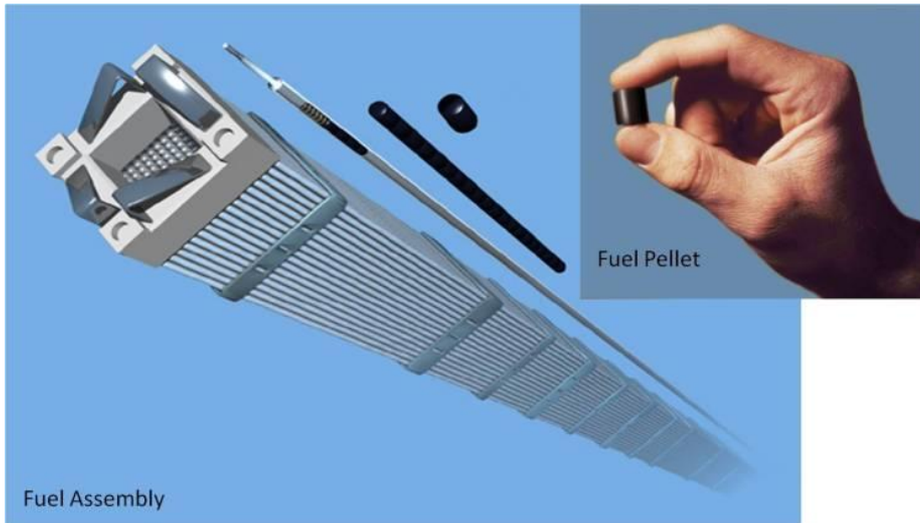
A RENEWED APPROACH OF UO_2 NUCLEAR FUEL USING PDF ANALYSIS

MRS EN 17 | L. Desgranges,¹ Y. Ma¹, Ph. Garcia¹, G. Baldinozzi¹, D. Siméone¹, H.E. Fischer²

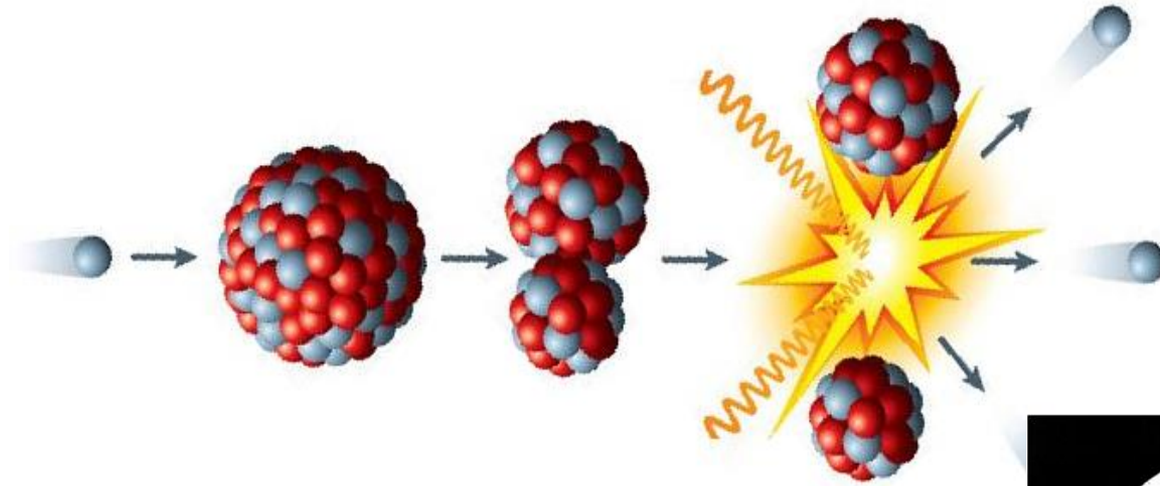
¹CEA, DEN France

²ILL, France

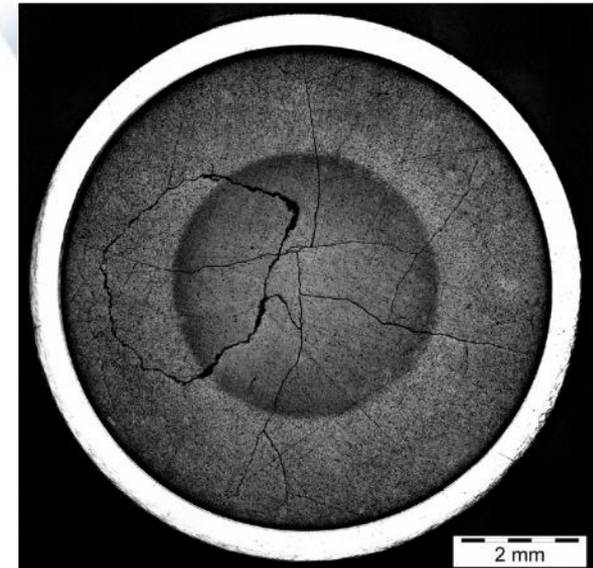
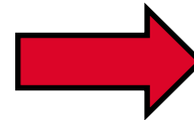
Uranium dioxide is the main component in present-day nuclear fuel reactors



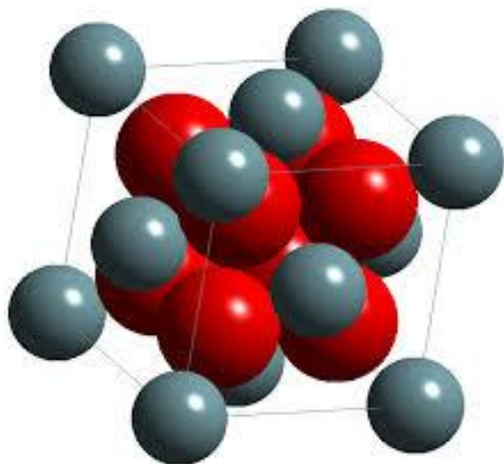
Energy production by Fission



Thermal gradient
Oxygen transfer
Irradiation damage
Fission product incorporation

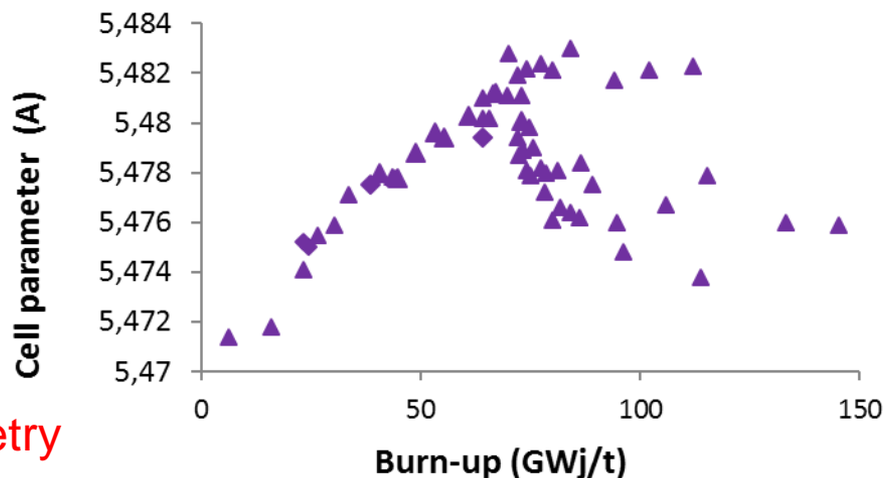


BUT A STABLE CRYSTALLINE STRUCTURE



Fm-3m symmetry
is kept

With slight changes of cell parameter



A simple picture of UO_2 (used in thermodynamic modelling J. Nucl. Mater.419 (2011) 145-167) :

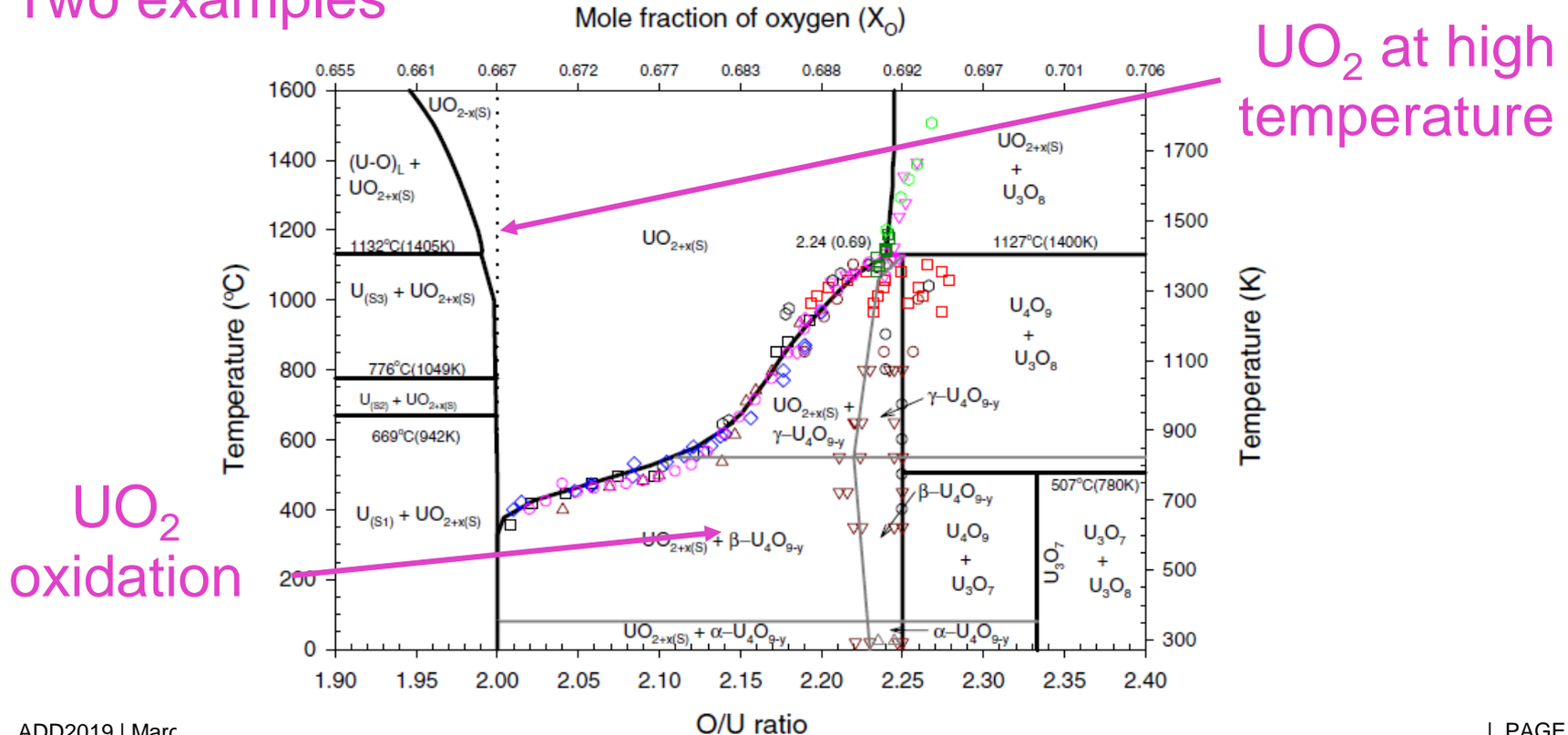
- One crystalline structure
- Three sites : 1 cationic, 1 anionic, 1 interstitial
- Different occupation fractions on each site

A TOO SIMPLIFIED PICTURE?

In the last 15 years, PDF analysis of UO_2 was used to improve the understanding of nuclear fuel behavior.

The obtained results contributed in proving that the ideal solid solution picture was oversimplified.

Two examples



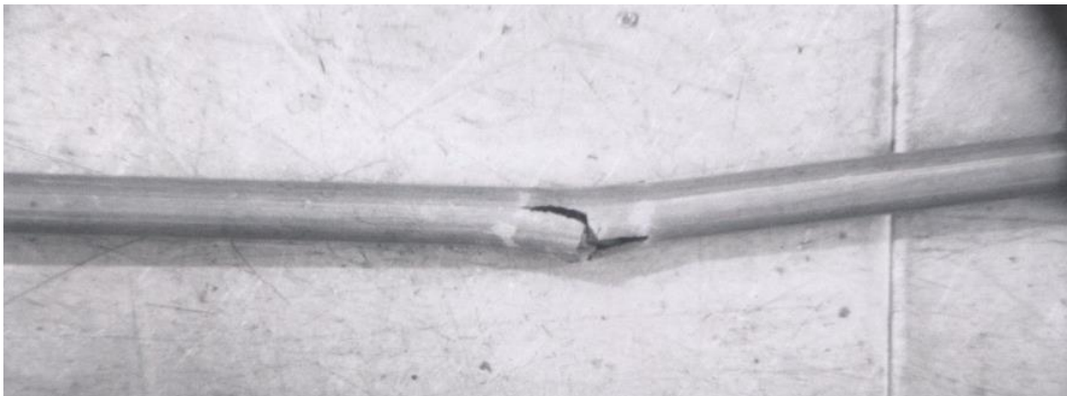
Example 1

UO₂ oxidation

In an accidental scenario, a defect in the cladding might put UO₂ irradiated fuel in contact with atmosphere (T<400°C)



The oxidation of irradiated UO₂ could lead to the ruin of the fuel rod



Need for a better understanding of UO₂ oxidation



cea U₄O₉ CRYSTALLINE STRUCTURE : AN OLD BRAIN TEASER!

- 1961 : first U₄O₉ observation using X-Ray Diffraction
- 1964 : U₄O₉ observation using neutron diffraction
- 1968 : U₄O₉ observation using electron diffraction
- 1960-1980 : evidence of α - β transition at 50°C
- First simple structural models simples : I-43d (1961) I4₁32 (1974)
- 1980 structural models using clusters of of oxygen atoms
- 1989 : determination β phase space group, I-43d
- 2004 : first structural refinement of β -U₄O₉ using Rietveld analysis
- 2005 : EXAFS measurements of α -U₄O₉
- 2006 : First PDF measurement of α -U₄O₉
- 2011 : first structural refinement of α -U₄O₉ using PDF analysis
- 2016 : first structural refinement of γ -U₄O₉ using PDF and Rietveld analysis

why so many difficulties?

DIFFICULTIES FOR SOLVING U_4O_9 STRUCTURE

An heavy cation near to a light anion

Atom	$\sigma_{X\text{-Ray}}$	σ_{neutron}
O	0.64	4.232
U	84.85	8.906

- ▶ neutron diffraction mandatory to evidence O atoms

More than 100 parameters to be refined

β - U_4O_9 is a UO_2 superstructure $4 \times 4 \times 4$ ($a = 21.766 \text{ \AA}$), 21 independent atoms

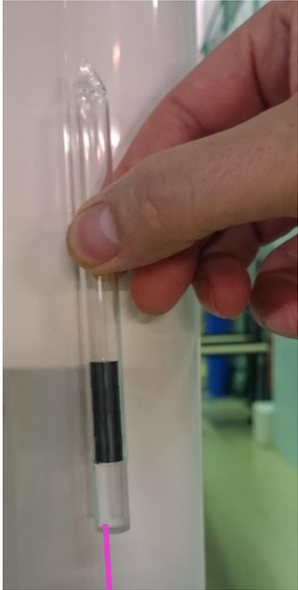
- ▶ a large data set mandatory with numerous diffraction lines

Need for local details

U_4O_9 contains clusters of oxygen interstitial

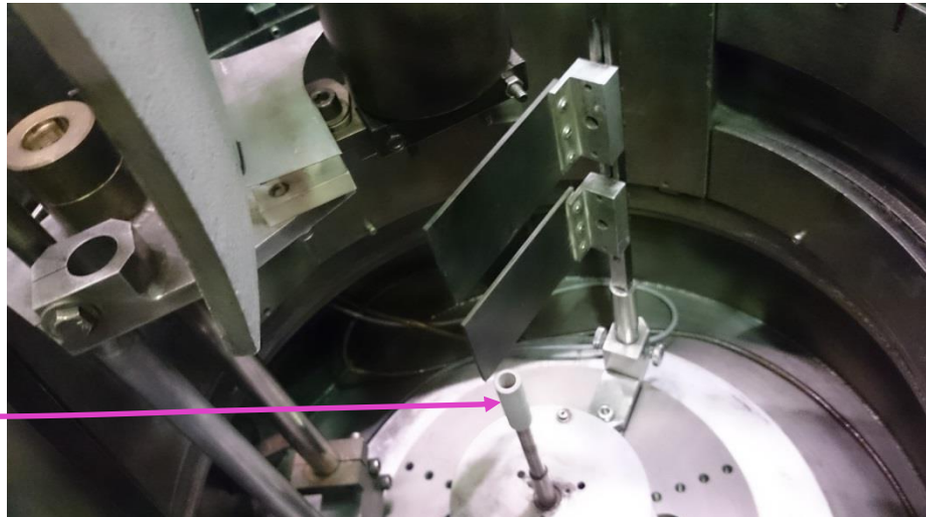
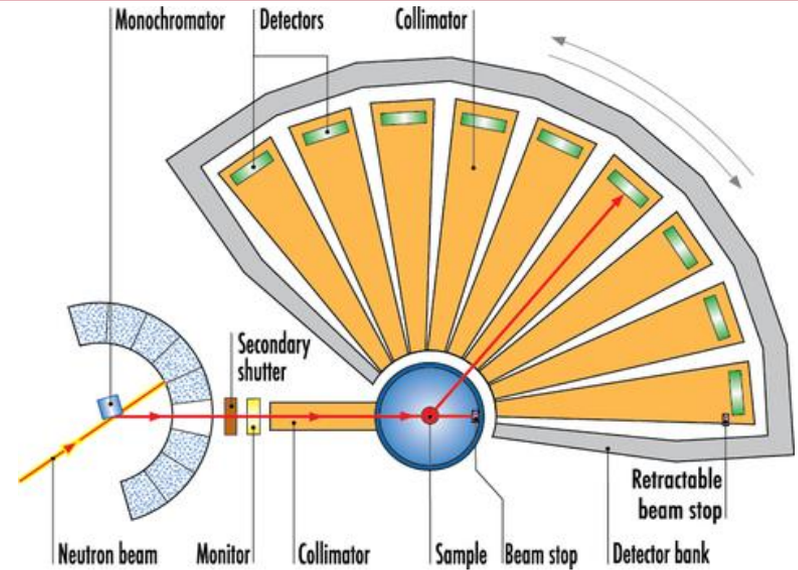
- ▶ PDF analysis mandatory for the short interatomic distance determination

cea NEUTRON DIFFRACTION WITH URANIUM OXIDES AT D4, ILL



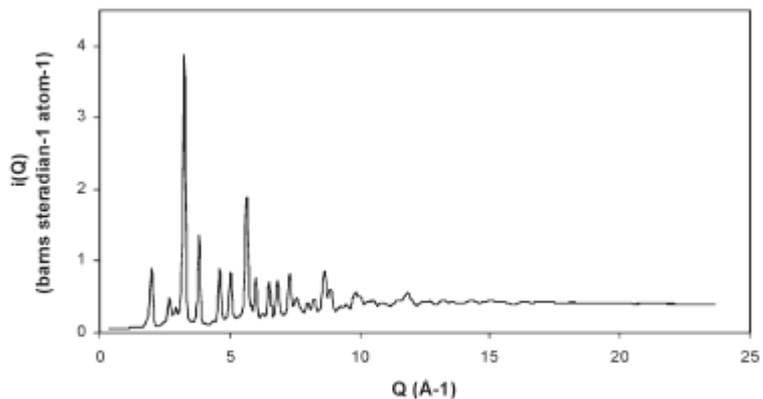
A two UO_2 pellet stack set in a airtight glass tube

Installed inside the sample holder of D4 furnace



Data analyzed in reciprocal and real space

DATA ANALYSIS



Fourier transform
➔

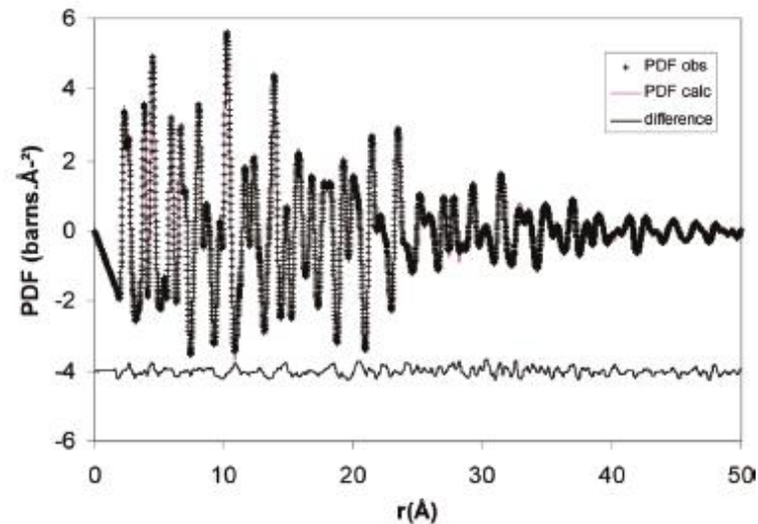
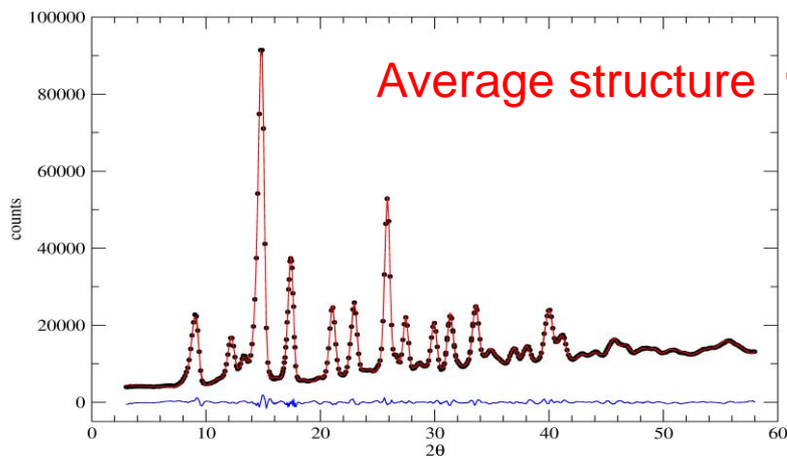
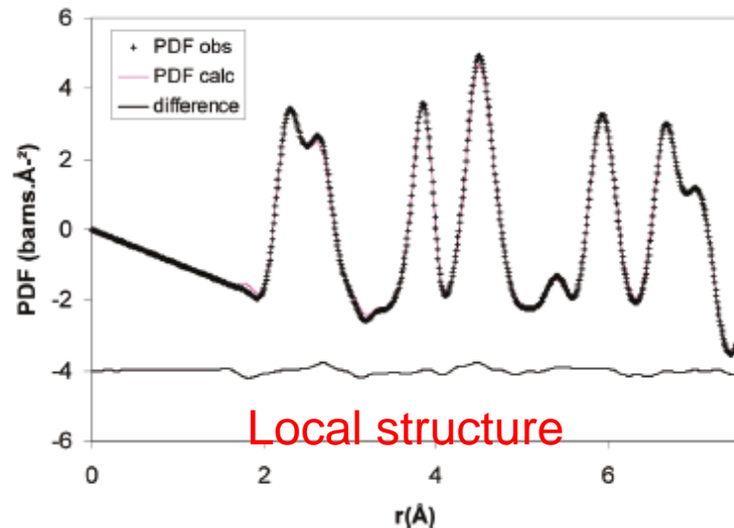


Figure 2. α - U_4O_9 neutron diffraction pattern corrected from air and sample holder diffusion.

↓ Rietvelt refinement (JANA)

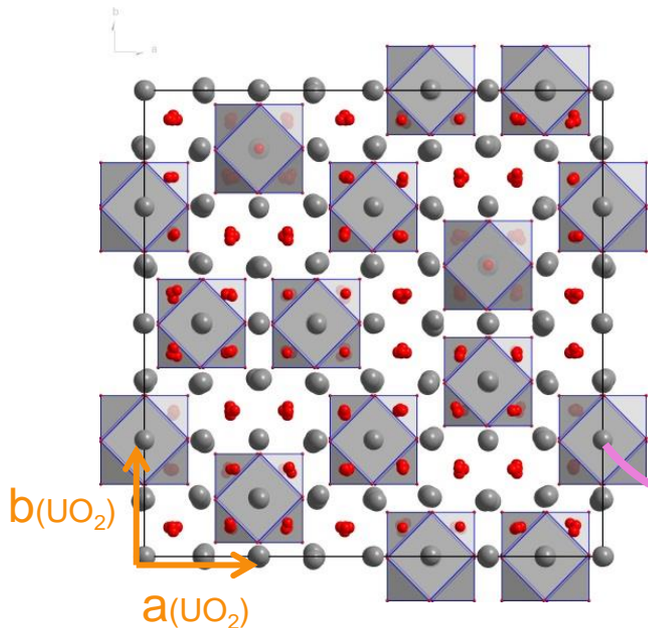


PDF refinement (PDFGUI) ↓

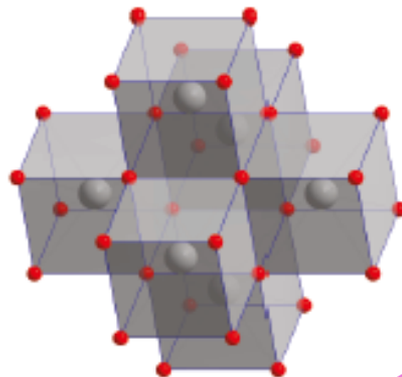


U₄O₉ AVERAGE STRUCTURE

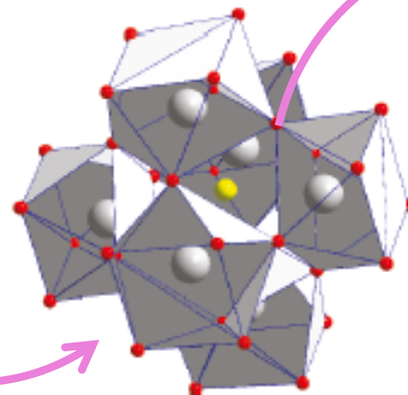
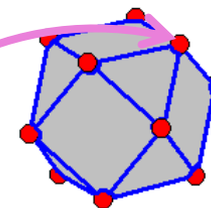
An array of oxygen interstitial clusters in a UO₂ matrix



U₆O₃₂ in UO₂



cuboctahedron

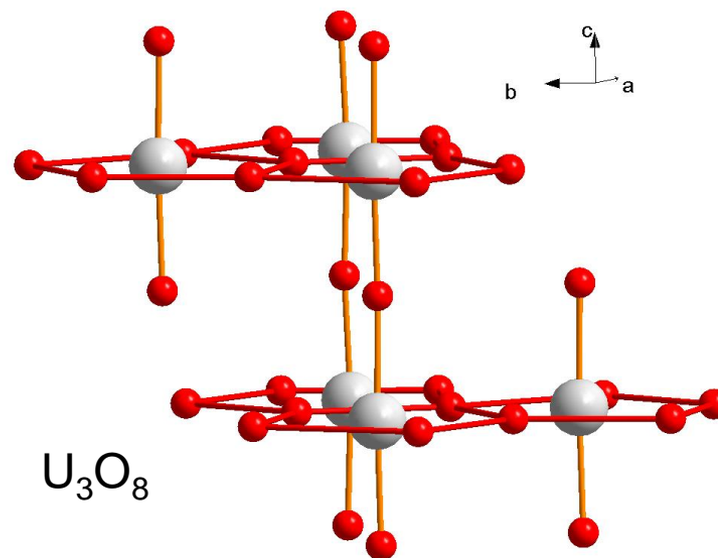
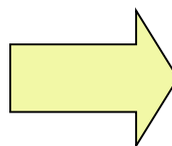
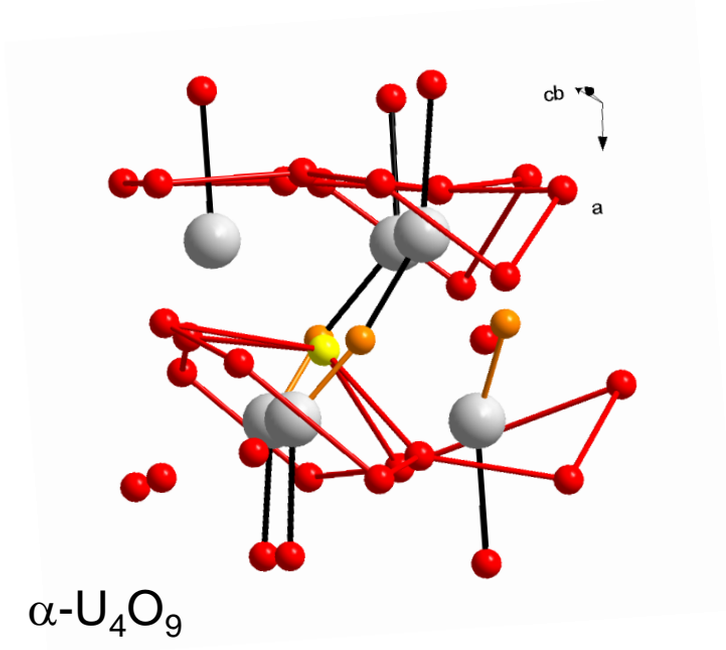
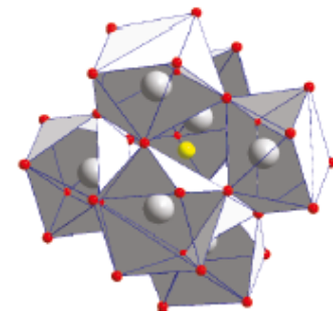


U₆O₃₆₊₁

A more precise description of the cluster was achieved using PDF refinement

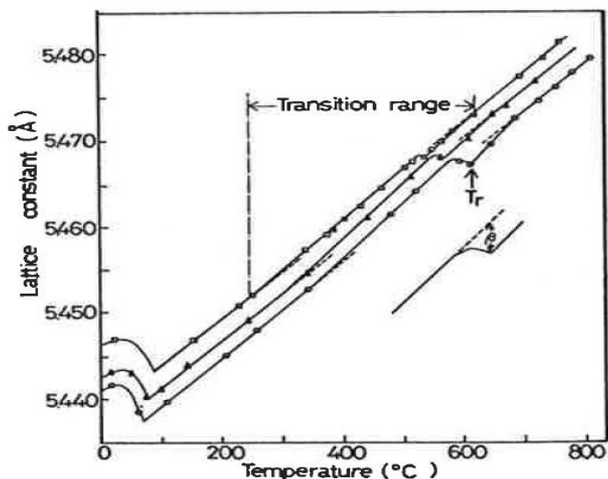
PDF refinement :

- Confirmed the existence of a the central oxygen
- Evidence the cuboctahedron as seed for higher oxide formation

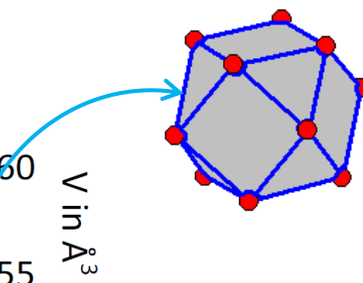
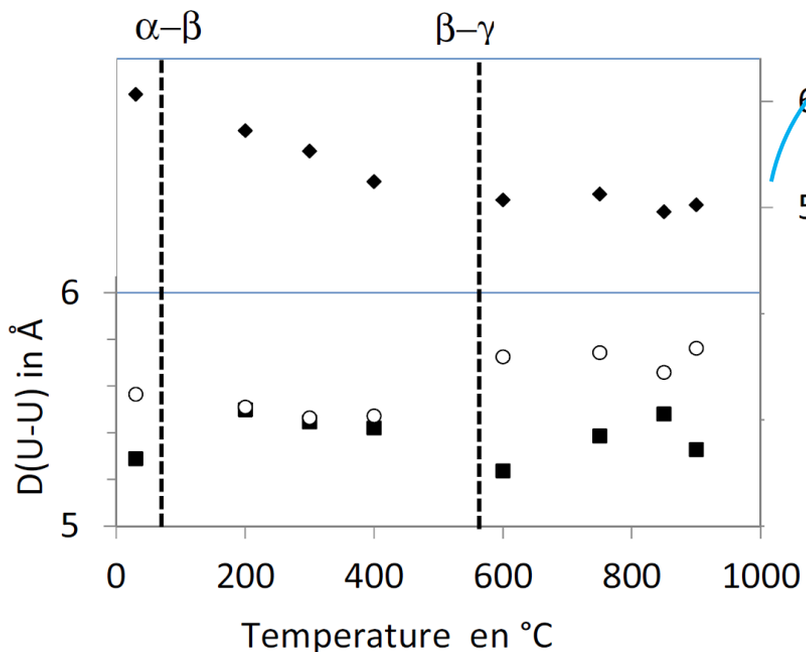


Inorg. Chem. 50 (2011) 6146-6151

PHASE TRANSITION IN U_4O_9

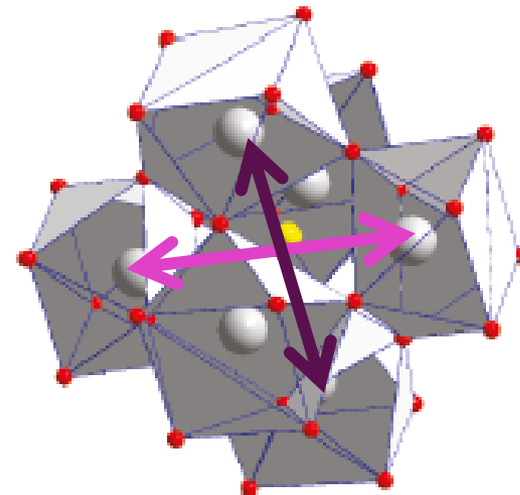


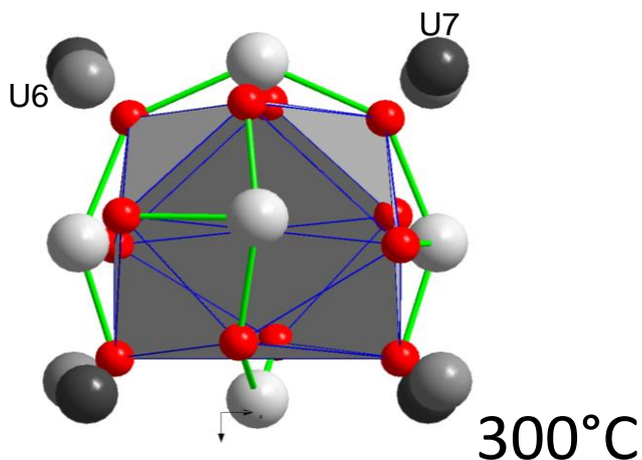
Evidence of changes in the cuboctahedron shape at the phase transition
 β - γ isostructural transition



Shape changes

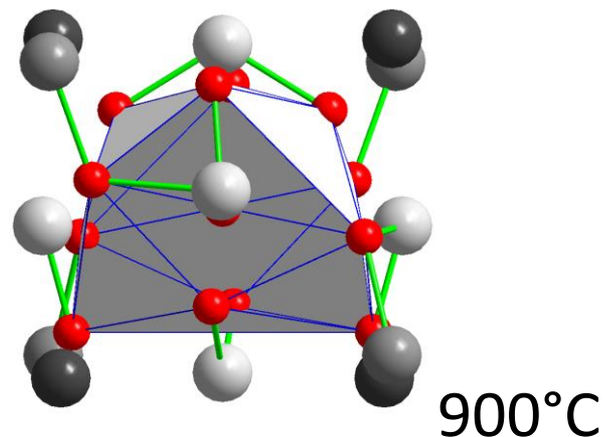
Volume decrease





$\beta\text{-U}_4\text{O}_9$:

- A nearly symmetric packing
- No difference between U6 and U7
- Steric interactions are dominant

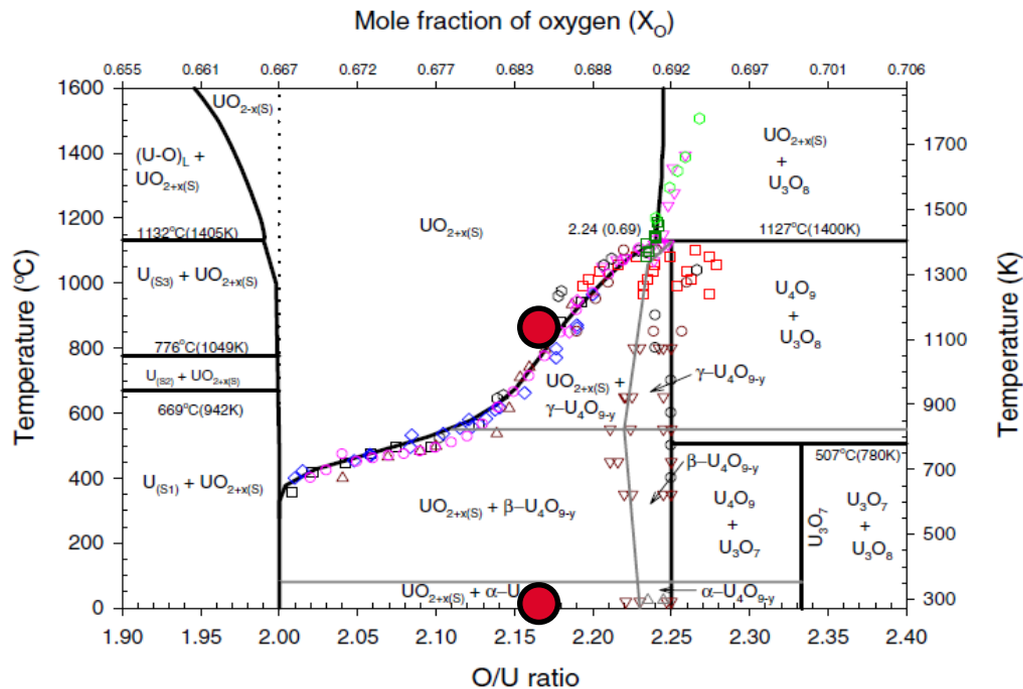


$\gamma\text{-U}_4\text{O}_9$:

- Asymmetric packing
- U atoms with no UO bond shorter than 2.2Å, (U^{4+})
- U atoms with UO bond shorter than 2.2Å, (U^{5+})
- U6 ($5+$) are connected to the cuboctahedron by short bonds (in green)
- Coordination interactions are dominant

Oxygen incorporation in UO_2 proceeds by the formation of oxygen interstitial clusters

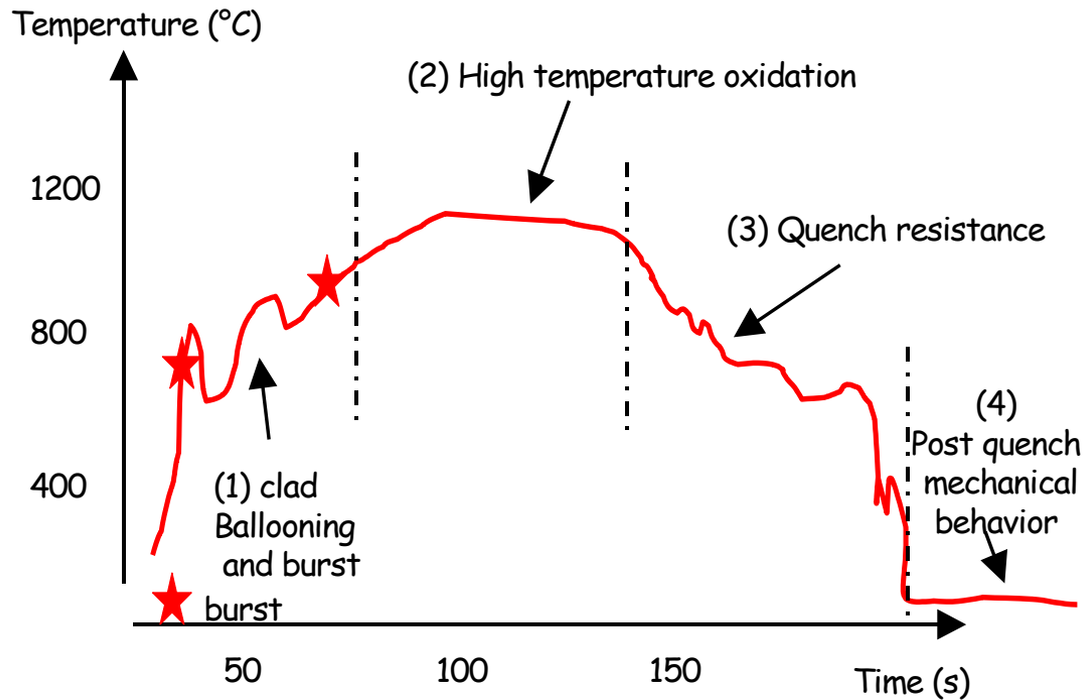
This conclusion is valid for a wide range of U-O phase diagram



Ma et al, Inorg, Chem. 57 (2018) 7064-7076

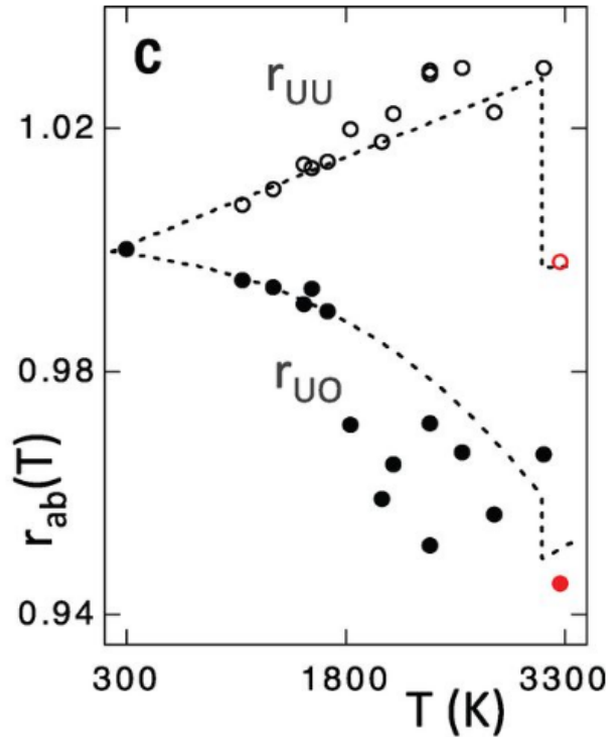
Example 2

UO₂ at high temperature



Temperature change in the case of a LOCA scenario as a function of time
The behavior of UO_2 at high temperature has to be known

No change in UO_2 crystal symmetry is expected



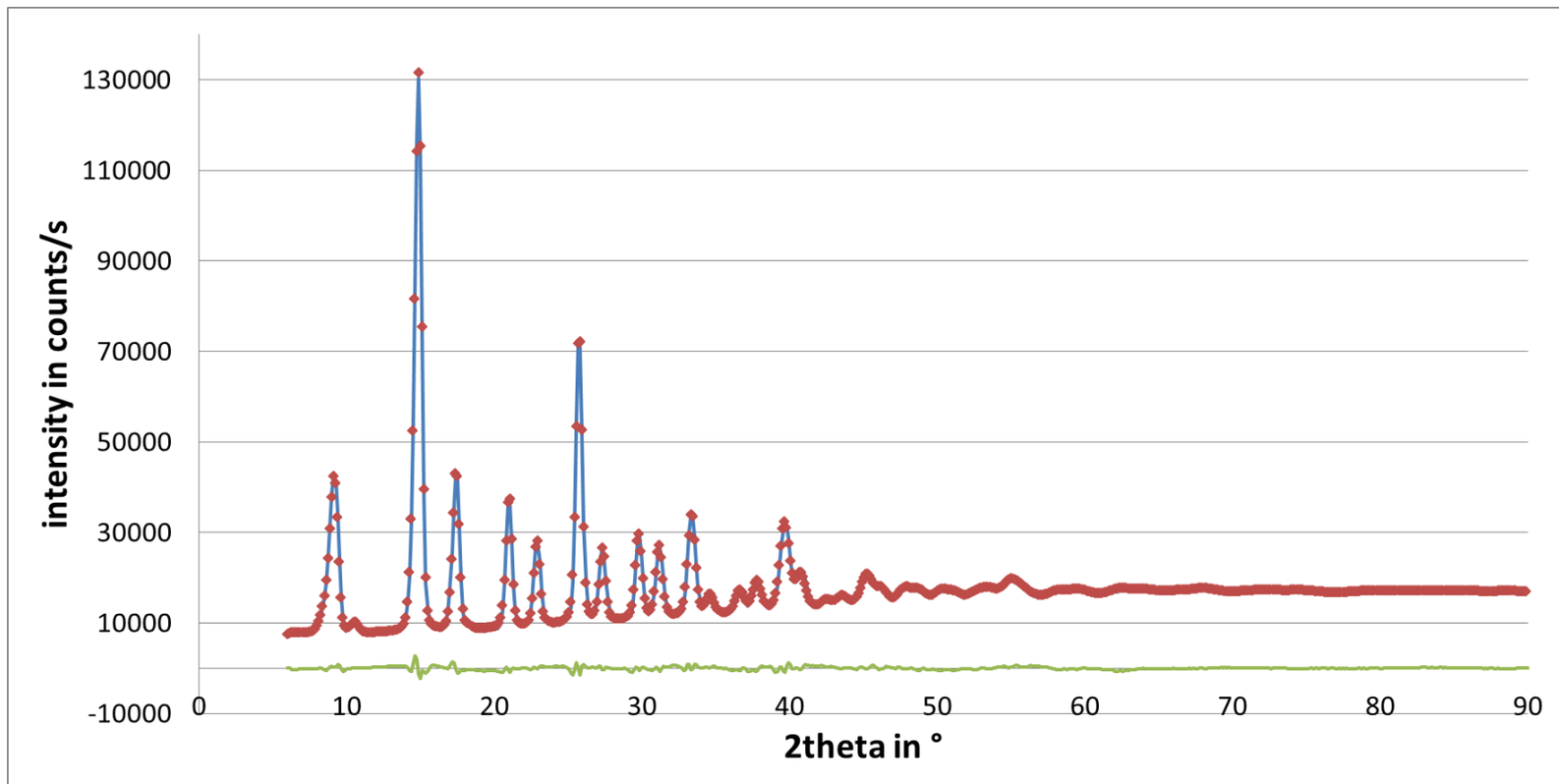
The U-O distance in UO_2 measured by X-Ray diffraction and PDF analysis is not consistent with Fm-3m crystalline structure

Skinner et al. [Science](#). 2014 Nov 21;346(6212):984-7

A change in UO_2 local symmetry at high temperature ???

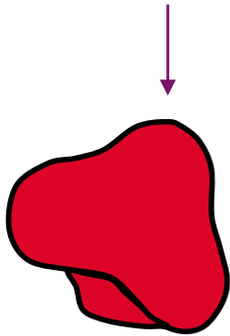
⇒ New experiments using neutron diffraction in order to probe O-O distance that are not visible using X-Ray diffraction

Several diffraction patterns of UO_2 pellets measured from RT to 1000°C

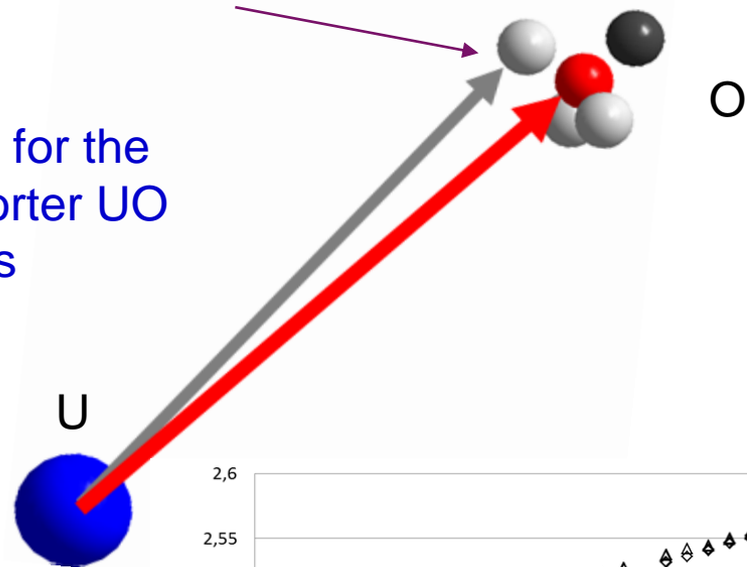


EVIDENCE OF OXYGEN DISORDER

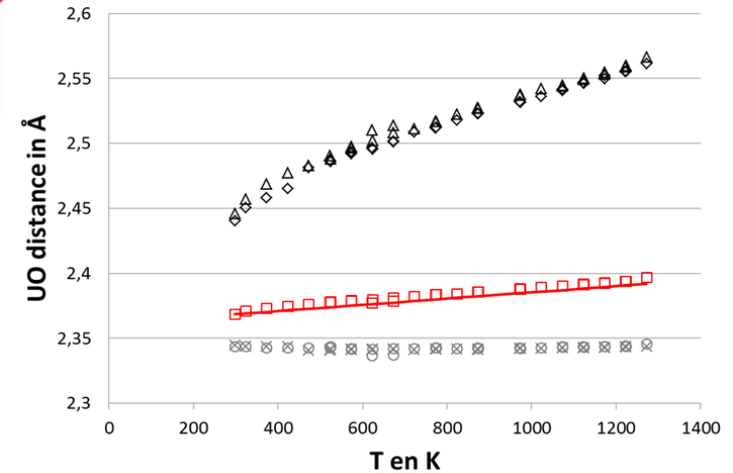
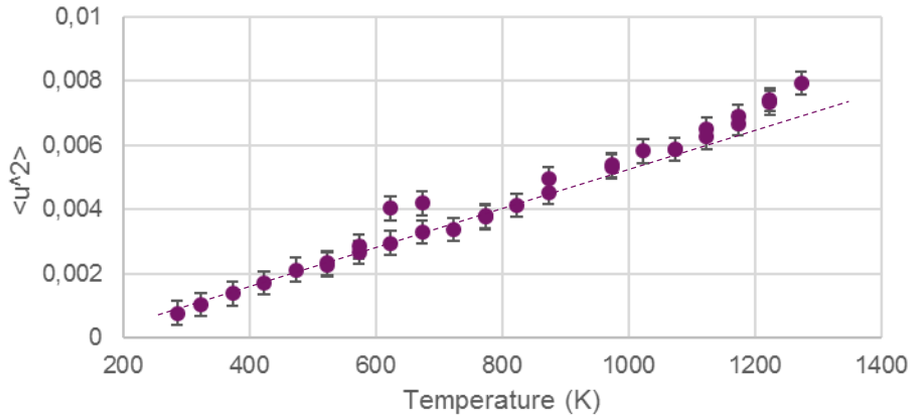
Described by Rietveld analysis,
either with anharmonic thermal motion or with a 4-site model



An explanation for the
existence of shorter UO
distances

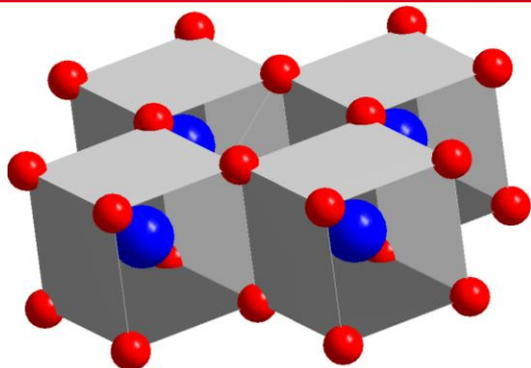


Anharmonic term of Oxygen atom thermal motion



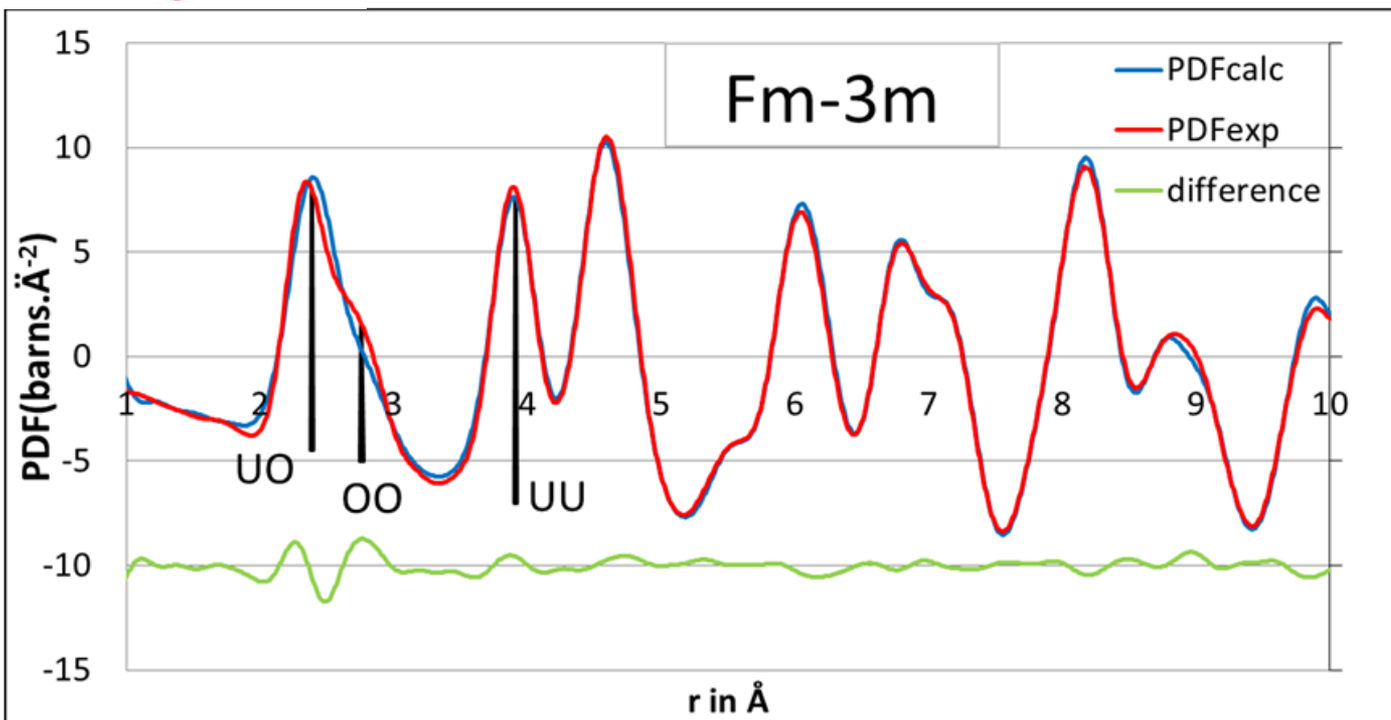
But only an average model

WHAT DOES PDF SAYS

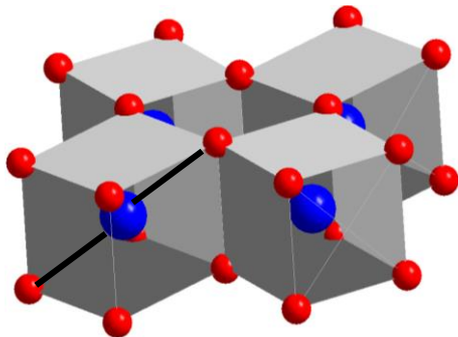


Regular Fm-3m structure does not reproduce well short distance distribution

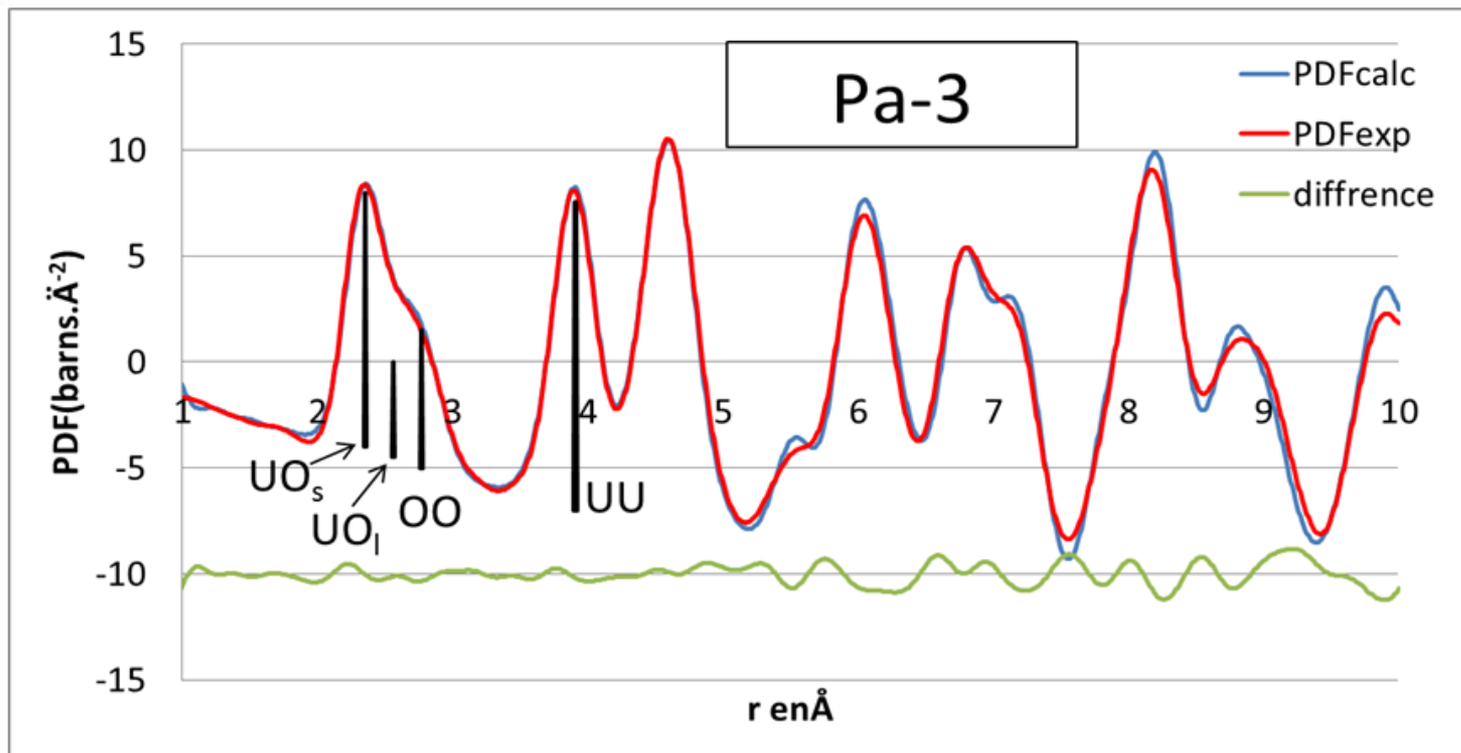
Consistent with oxygen disorder



A FIRST ATTEMPT TO DESCRIBE LOCAL ORDER



A lower symmetry structure, Pa-3, a Fm-3m subgroup having 2 short, 6 long UO and a single OO bond lengths, fits local order better, but no more long range order

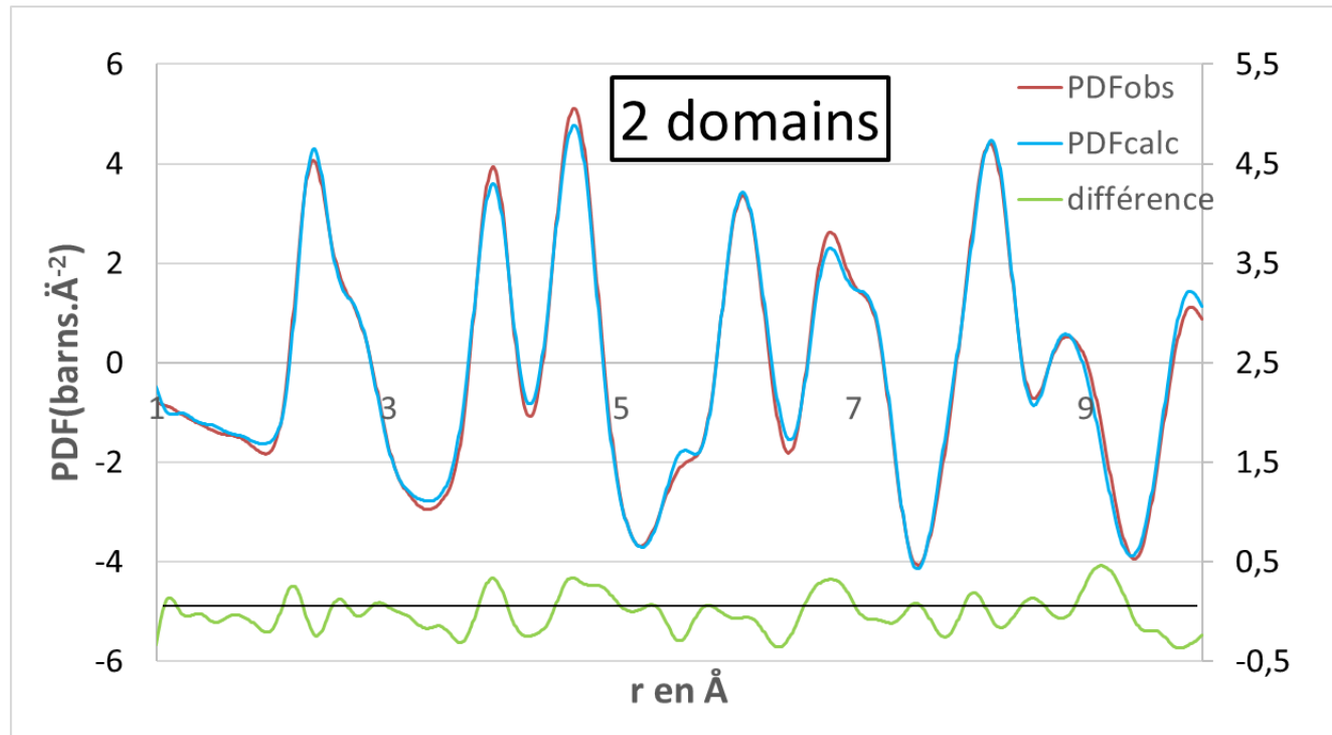


HOW TO LINK LOCAL AND LONG RANGE ORDER?

Putting domains having different orientations the ones next to the others is not enough.

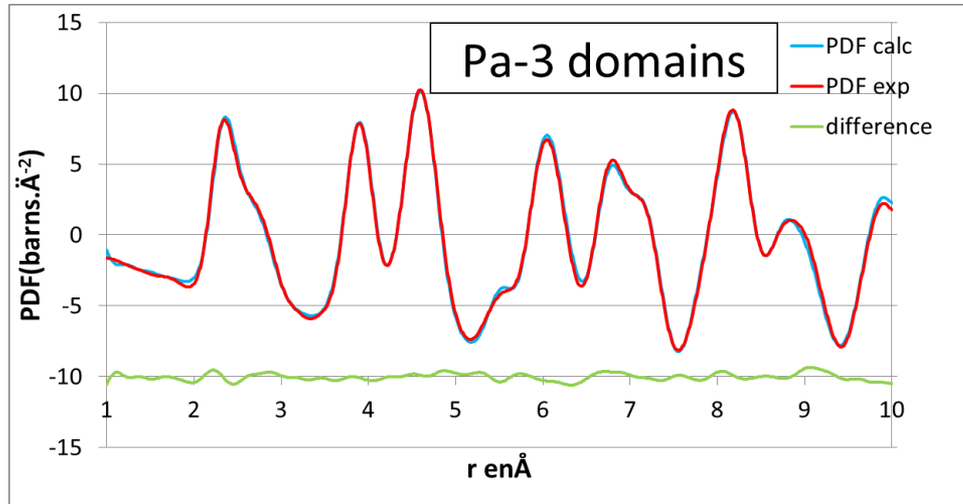


Modeling box with 5 UO_2 cells and 2 domains



Adding more domains does not improve the fit

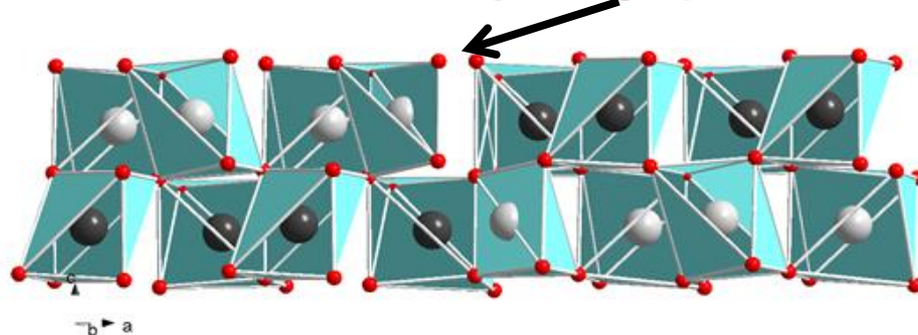
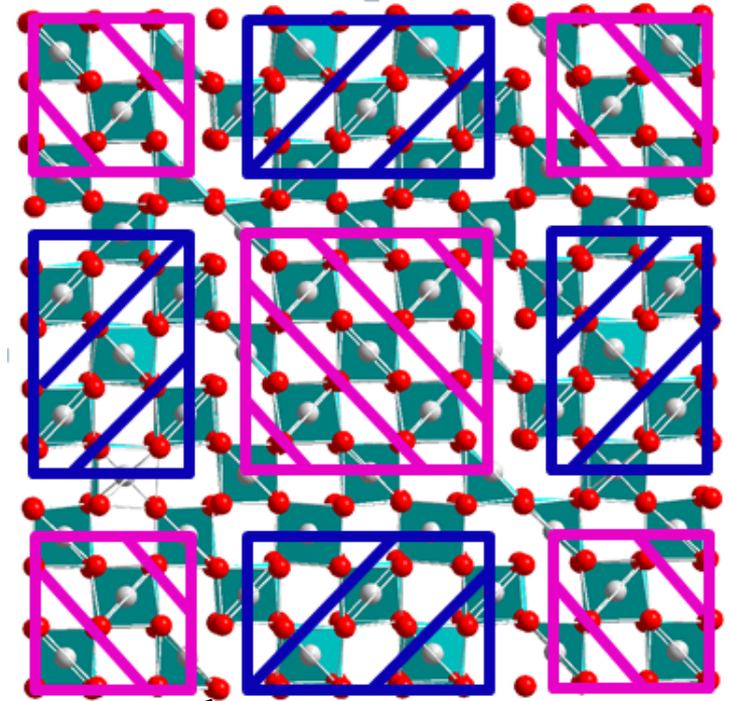
SYMMETRY CONSTRAINTS ARE NEEDED !!



Chem. Eur. J. 2018, 24, 2085 – 2088

Best fit with Pa-3 nm size domain separated by coherent interfaces (compatible with average Fm-3m)

Consistent with the absence of additional diffraction lines



Pyrite type stacking

Marcasite type slab

Pyrite type stacking

Rietveld or PDF fits are only mathematical descriptions of the actual atomic distribution. More physics is needed for interpretation.

A static or dynamic disorder ?

- Diffraction (whether Rietveld refinement or PDF-analysis) cannot distinguish between static versus dynamic disorder, unless data are taken as a function of temperature.

Domains or somethings else ?

- Moving interfaces are also possible

The link with kinetic data would be helpful:

- What are the dynamics inside UO_2 at high temperature that could be related to oxygen motion?

At least dynamically, at high temperature, the local symmetry of UO_2 , Pa-3, is lower than the regular Fm-3m regular symmetry of UO_2

Shorter UO bond length that are observed by EXAFS, X-ray and neutron diffraction
Is consistent with this lower local symmetry

For UO_2 , the assumption : “ one crystalline structure, three sites : 1 cationic, 1 anionic, 1 interstitial, and different occupation fractions on each site” is oversimplified.

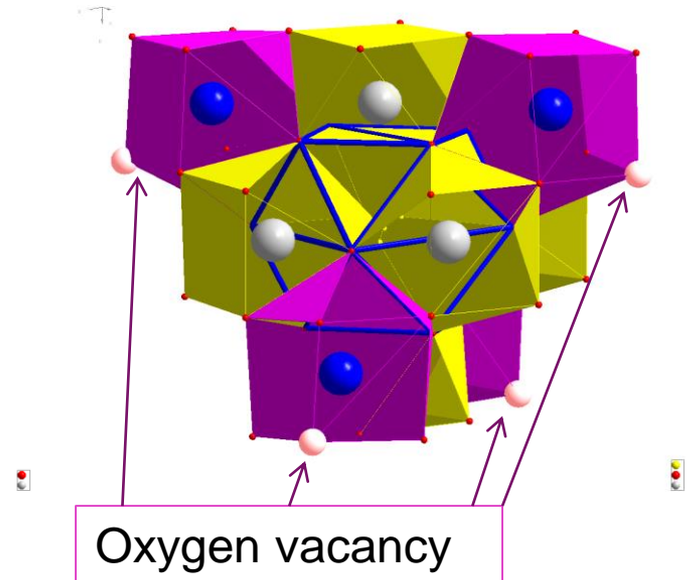
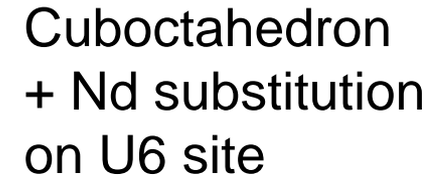
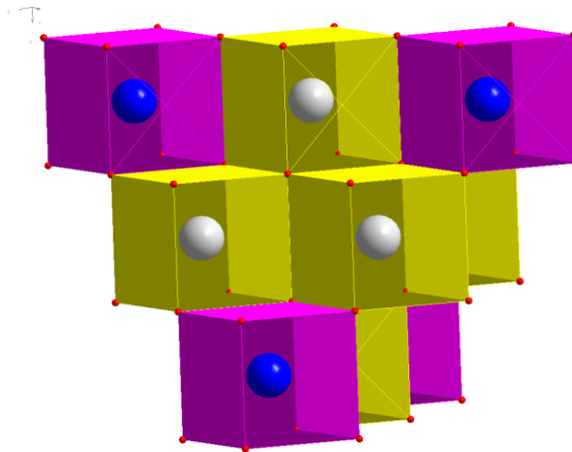
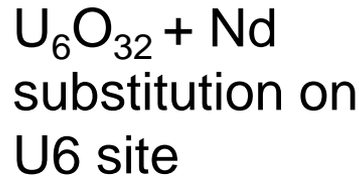
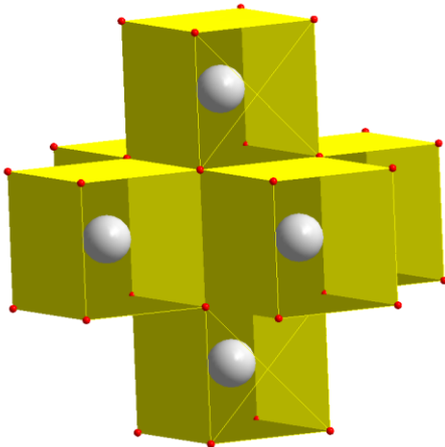
The understanding of UO_2 modification by irradiation requires a description at the local scale

- 2 perspectives:

- A new approach for the understanding of irradiated UO_2
- A more general approach of « simple » oxides

A NEW APPROACH FOR THE UNDERSTANDING OF IRRADIATED UO_2

What about the incorporation of fission products ?



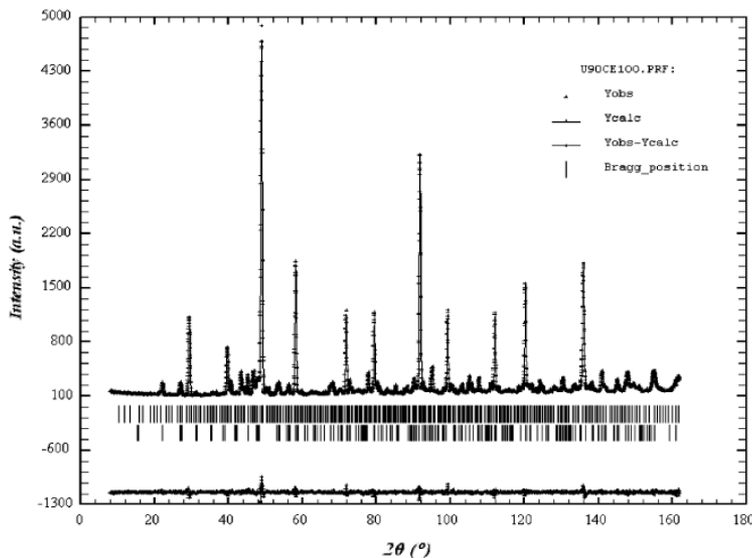
Nd induced cuboctahedron formation :

- Consistent with U5+ formation in $(\text{U},\text{Nd})\text{O}_2$ (J. Nucl. Mater. 507 (2018) 145-150)

SOME EXPERIMENTAL CONFIRMATION

Neutron diffraction on $(U_{0,9}Ce_{0,1})_4O_{9-\delta}$

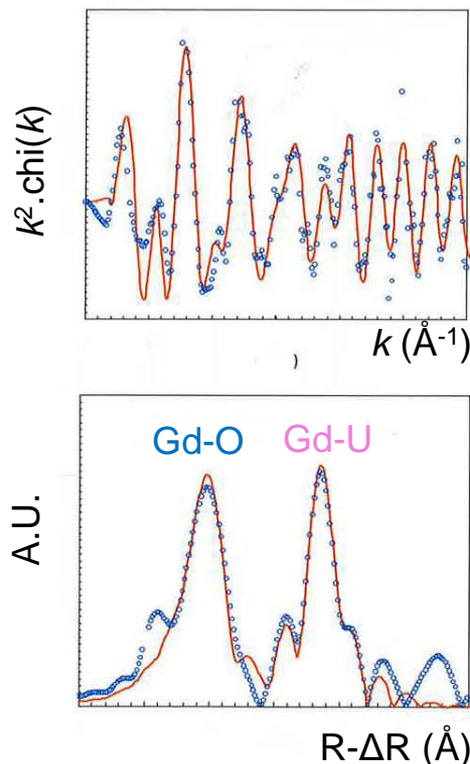
C. Rocanière et al. | *Journal of Solid State Chemistry* 177 (2004) 1758–1767



Rietveld refinement evidenced that Ce lied on U6-U7 site

EXAFS on $(U,Gd)O_2$

P. Martin, private communication



EXAFS Refinement at Gd L3 edge (FEFF8.4) evidenced that Gd has 9.4 (5) oxygen atoms as first neighbours instead of 8 in regular fluorite

Consistent with U6-U7 site

consistent with 3+ cation substitution on U6 site and cuboctahedron formation presented before

Many “simple” oxides evidence lower local symmetry:

- Pyrochlores, Spinel
- Also showing a Fm-3m disordered phase

Does the concept of symmetry coherent interface apply?

- The group subgroup relationships derived in the case of pyrochlores (Sci. Rep. 7 (2017) 3727)
- Go further ?

What are the consequence of a local disorder ?

- Better irradiation resistance ?
- What about transport properties?
- Which thermodynamic description?

Thank You for your attention

Commissariat à l'énergie atomique et aux énergies alternatives
Centre de Cadarache | 13108 Saint-Paul lez Durance Cedex

DEN
DEC
SA3E

Etablissement public à caractère industriel et commercial | R.C.S Paris B 775 685 019