Dalton Transactions



PAPER View Journal | View Issue



Cite this: *Dalton Trans.*, 2025, **54**, 6847

Precipitation of morphology-controlled uranium(vi) peroxide in nitric acid media†

The precipitation of U(vi) peroxide (UO₂(O₂)·4H₂O, known as studtite) has been extensively studied in mildly acidic media (i.e. pH > 1). However, only a few studies have investigated the influence of highly acidic conditions (i.e. pH < 1) on the precipitation of studtite, particularly regarding the morphology of the final powder. Therefore, the influence of high acidity (0.1 M < $C(H^+)_{inj}$ < 2.0 M) and the $C(H_2O_2)_{inj}/C(U)_{inj}$ molar ratio (ranging from 10 to 70) on the precipitation kinetics, as well as the morphological and crystallographic properties of U(vi) peroxide, was investigated in this study. Decreasing the acidity and increasing the C(H₂O₂)_{ini}/C(U)_{ini} molar ratio have significantly enhanced both the final precipitation yield and the reaction kinetics. Furthermore, a correlation was found between the initial saturation index of the reaction mixture and the final morphology and crystallite size of the precipitated U(vI) peroxide, independent of the initial precipitation conditions. These results demonstrate that in highly acidic nitric media, it is possible to control the morphology of the precipitated U(vi) peroxide by simply adjusting the initial saturation index. This correlation opens up interesting perspectives for the synthesis of powders with specific morphologies, in particular uranium peroxide or corresponding oxide powders obtained after thermal treatment. This morphology control is of great interest in the field of nuclear fuel fabrication, as the characteristics of oxide powder, such as flowability or sintering reactivity, have a strong impact on the manufacturing process and the properties of the final ceramic.

Received 16th December 2024, Accepted 19th March 2025 DOI: 10.1039/d4dt03467h

rsc.li/dalton

1. Introduction

Uranium(v₁) peroxide tetrahydrate, $UO_2(O_2)(H_2O)_2 \cdot 2H_2O$, commonly called "studtite", from the name of the natural mineral form of this compound, ¹ is ubiquitous in the industrial uranium chemical treatment process. In the initial steps of the nuclear fuel cycle, particularly during the uranium extraction process, uranium(v₁) peroxide can be precipitated and then calcined at low temperature to recover uranium from the *in situ* leaching of uranium ore:^{2,3} it is usually precipitated in sulfuric media at low acidity (pH 3–4).⁴ Peroxide precipitation can also be a purification step prior to isotopic enrichment.⁵ Downstream, in advanced reprocessing of spent nuclear fuel, uranium can be recovered by peroxide precipitation.⁶ In addition, in the context of the direct long-term repository of spent nuclear fuel, the interactions between dissolved uranium and H_2O_2 produced by water radiolysis⁷ likely lead to

the formation of UO₂(O₂)·4H₂O that can passivate the dissolving surface. 8-12

There have been many studies on the precipitation of uranium(vi) peroxide. Some phenomenological ones have demonstrated quantitative precipitation of studtite at low acidity (pH 2–3), 13,14 and then its conversion to pure U_3O_8 after calcination in air. 15 Several reactors technologies have also been studied to precipitate uranium(vi) peroxide, such as continuous reactors 16 or fluidized beds. 17 Nevertheless, only a few studies have dealt with a fundamental approach. For example, Planteur *et al.* have addressed the crystal growth of uranium(vi) peroxide at pH = 3 in sulfuric acid 18,19 while Kim *et al.* 20 and Schwerdt *et al.* 21 have established a relationship between the particle size and initial conditions of precipitation.

Only two papers have measured precipitation yields in acidic media. Kornilov $et\ al.^{22}$ assessed the precipitation yield after 24 hours in 1 to 3 M HNO $_3$ solutions and for different peroxide ion concentrations. The higher the acidity, the lower the yield was for a given initial peroxide concentration. Hibert $et\ al.$ studied the precipitation of U(vI) and demonstrated the co-precipitation of U(vI) and Pu(IV) (potential production of MOx fuel directly from U-Pu flow) working between 0.8 M and 2.0 M HNO $_3.^{23}$

^aCEA, DES, ISEC, DMRC, Univ Montpellier, Marcoule, France. E-mail: paul.estevenon@cea.fr; Tel: +33 4 66 79 65 66

^bICSM, Univ Montpellier, CEA, CNRS, ENSCM, Marcoule, France

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4dt03467h

Paper

Still, there is very little data on the morphology adopted by U(vi) peroxide, especially in very acidic media (0.1 to 2.0 M), despite the fact that $UO_2(O_2)\cdot 4H_2O$ precipitation is a way of conversion and fabrication of nuclear fuel²⁴ with the preservation of the peroxide morphology during the calcination step. With this objective, the shape of the particles is of great importance, as it is the main parameter that impacts the powder flowability²⁵ and sinterability.²⁶

In this work, the precipitation of uranium(vI) peroxide has been studied in nitric acid concentrations between 0.1 M and 2.0 M. To this end, several precipitation reactions were performed with a fixed uranium concentration (0.07 M) over a wide acidity range from 0.1 to 2.0 M H $^+$ and molar ratios of $C(H_2O_2)_{ini}/C(U)_{ini}$ ranging from 10 to 70. The precipitated U(vI) peroxide was characterized by X-ray diffraction and observed using scanning electron microscopy. The evolution of the precipitation yield was also measured under specific conditions.

Materials and methods

2.1. Precipitation of uranium(vi) peroxide

Caution! 238 U is an α -emitter and is considered a health risk. Experiments involving actinides require appropriate facilities and persons trained in the handling of radioactive materials.

Commercial powdered uranyl nitrate, $UO_2(NO_3)_2 \cdot 6H_2O$ (Prolabo, purity 99%), was used as the uranium source for all the experiments. Acidity was fixed using concentrated analytical grade nitric acid solution at 70 wt% from CARLO ERBA. The hydrogen peroxide used was an analytical grade commercial solution at 30 wt%, supplied by CARLO ERBA (the concentration was determined to be $C(H_2O_2) = 10.05$ M by potentiometric titration with Ce(IV)).

1.00 mL of uranium(v1) solution with a uranyl concentration of 0.46 M was prepared in nitric media by dissolving around 235 mg of uranyl nitrate hydrate in diluted nitric acid solution with acidity ranging from 0.1 to 2.0 M. 5.80 mL of hydrogen peroxide solution was prepared by diluting concentrated $\rm H_2O_2$ in nitric acid media with the same acidity as the uranium solution. A uranyl concentration of 0.46 M was arbitrarily set so that the final uranyl concentration after mixing these two solutions was 0.07 M. The concentration of hydrogen peroxide was determined so that the $\rm C(H_2O_2)_{ini}/\rm C(U)_{ini}$ molar ratio was between 10 and 70 (Table ESI 1†) after the mixing of the two solutions.

Uranyl peroxide precipitation was achieved by adding the uranyl solution dropwise to the corresponding hydrogen peroxide solution in 10 mL glass beakers with a diameter of 2.5 cm and stirring with a 1.2 cm magnetic stirring rod at 500 rpm. The resulting mixture was stirred for a few hours, resulting in the formation of a yellow precipitate, with kinetics varying significantly depending on the chemical conditions considered. The progress of the reaction at the end of the experiment was calculated from the ratio of the effective yield to the theoretical yield at equilibrium. The reaction was arbi-

trarily considered to be at equilibrium when its progress reached 95%.

At the end of the precipitation, the suspension was collected and centrifuged at 4500 rpm (*i.e.* $3260 \times g$) for 5 min in 15 mL tubes. The powder and the supernatant were then separated. A washing step was carried out twice with 2 mL of a 50:50 (vol:vol) water/ethanol mixture. The solid was isolated from the supernatant by centrifugation and finally dried overnight at room temperature.

It is worth noting that under the experimental conditions considered, although nitrous acid (HNO_2) was autocatalytically generated in the nitric medium²⁷ and can catalyze the decomposition of hydrogen peroxide, this reaction remains negligible for HNO_2 concentrations below 8 M.²⁸ In addition, since the reactions were carried out at room temperature, the decomposition rate of H_2O_2 was further limited. Additionally, the reaction durations are relatively short compared to the characteristic decomposition kinetics of H_2O_2 . Consequently, the amount of H_2O_2 added can be considered nearly constant throughout the experiment.

2.2. Determination of uranium concentrations

Uranium concentration measurements were performed using an ICP-OES (inductively coupled plasma optical emission spectroscopy) Spectro Arcos apparatus. $\lambda = 367.007$ nm, $\lambda =$ 385.958 nm, $\lambda = 263.553$ nm and $\lambda = 393.203$ nm wavelengths were considered to determine the uranium concentration. Calibration was performed using a series of standard uranium solutions (0.5 to 20 ppm), prepared by dilution of a 1000 ppm standard solution supplied by SCP Science. Uranium concentration was also determined by photon electron rejecting alpha liquid scintillation (PERALS) considering the protocol developed by Dacheux et al.29 The extraction was performed by contacting 6 mL of analyzed aqueous solutions with 1.5 mL of Alphaex™ scintillating cocktail purchased from ORDELA Inc. (Oak Ridge, USA). AlphaexTM is an organic cocktail containing HDEHP (bis-(2-ethylhexyl)phosphoric acid) as the extractant, PBBO (2-(4'-biphenylyl)-6-phenylbenzoxazole), PPO (2,5-diphenyloxazole) and NPO (2-(1-naphthyl)-5-phenyloxazole) as scintillators, naphthalene as the energy transfer agent and toluene as the solvent. The mixture was shaken for approximately 2-3 minutes and then centrifuged at 2000 rpm (i.e. $643 \times g$) for 5 minutes. 1.3 mL of the organic phase was then collected and the ²³⁸U + ²³⁴U activity was measured, leading to uranium concentration in the leachate.

2.3. Characterization of the prepared solids

Solids were analyzed using a Bruker D8 diffractometer equipped with a Lynxeye detector adopting the reflection geometry with a Göbel mirror and using Cu $K\alpha_{1,2}$ radiation ($\lambda K\alpha_1=1.540593$ Å; $\lambda K\alpha_2=1.544427$ Å). For each acquisition, the PXRD (powder X-ray diffraction) pattern was recorded in the $10^\circ < 2\theta < 60^\circ$ range with a step size of 0.027° (2θ) and a counting time of 1 s per step. Powders were prepared on flat sample holders with top-loading, placed in a dome-shaped specimen holder aiming to avoid radioactive contamination (Bruker airtight specimen holder with a knife edge, acting as

Dalton Transactions Paper

an anti-scattering blade). Note that this PMMA (poly(methyl methacrylate)) dome can be the source of a large diffusion peak in the range of $10^{\circ} < 2\theta < 20^{\circ}$. Le Bail refinements³⁰ of these patterns were performed using the FullProf Suite program³¹ with the Cox–Hastings pseudo-Voigt profile function.³² The instrumental function was obtained by refinement of a LaB₆ reference. The backgrounds were manually selected

For SEM characterization, the samples were deposited on carbon adhesive tape and measured directly without any metal coating step. An FEI Quanta 200 scanning electron microscope, equipped with an Everhart–Thornley detector (ETD) and a backscattered electron detector (BSED), was used to record images at an acceleration voltage of 15 kV under high-vacuum conditions (10^{-6} Pa).

Specific surface areas (SSA) were measured on a Micrometric ASAP 2020 apparatus with the BET method (nitrogen adsorption at $77 \, \text{K}$).

$$\varepsilon = \frac{\left(\text{C}(U)_0 + \text{C}(\text{H}_2\text{O}_2)_0 - 4 \times K \times \text{C}(\text{H}^+)_0\right) - \sqrt{\left(\text{C}(U)_0 + \text{C}(\text{H}_2\text{O}_2)_0 - 4 \times K \times \text{C}(\text{H}^+)_0\right)^2 - \left(4 - 16 \times K\right) \times \left(\text{C}(U)_0 + \text{C}(\text{H}_2\text{O}_2)_0 - K \times \text{C}(\text{H}^+)_0^2\right)}{(2 - 8 \times K)}}$$

2.4. Determination of precipitation yield, saturation index and progress of the reaction

The precipitation yield at time t was calculated from the initial and final uranium concentrations in the supernatant, by using eqn (1). The final uranium concentrations were measured either by ICP-OES or PERALS (depending on experimental availability), after a 100-fold dilution to avoid any precipitation in the analysis tube. The initial uranium concentration was calculated from the mass of uranyl nitrate added.

$$\text{Yield}_{t} = \frac{[\text{UO}_{2}^{2+}]_{\text{ini}} - [\text{UO}_{2}^{2+}]_{t}}{[\text{UO}_{2}^{2+}]_{\text{ini}}} \% \tag{1}$$

The initial saturation index (SI_{ini}) was calculated from the U(vi) peroxide precipitation reaction (eqn (2)) according to eqn (3), *i.e.* from the logarithm of the ratio between the reaction quotient out of equilibrium, Q_{ini} (eqn (4)), and the solubility constant, K (eqn (5)). The saturation index is useful here, as it expresses the driving force of the precipitation reaction. Indeed, when SI > 0, the solution is supersaturated with respect to the precipitated phase, whereas it remains undersaturated for SI < 0. In the latter case, no phase precipitation is expected in the solution. For the solubility constant of studtite (eqn (5)), the recommendations of the Nuclear Energy Agency were followed³³ and the value from Kubatko *et al.*³⁴ was selected, *i.e.* $\log (K) = -2.87$.

$$\begin{split} &UO_{2}(O_{2})\cdot 4H_{2}O_{(s)} + 2H_{(aq)}^{+} \\ &\to UO_{2}{}^{2+}_{(aq)} + H_{2}O_{2(aq)} + 4H_{2}O_{(aq)} \end{split} \tag{2}$$

$$SI_{ini} = \log\left(\frac{Q_{ini}}{K_{s}}\right) \tag{3}$$

$$Q_{\rm ini} = \frac{a_{\rm H_2O_2}^{\rm ini} \times a_{\rm UO_2^{2+}}^{\rm ini}}{(a_{\rm tot}^{\rm ini})^2} \tag{4}$$

$$K = \frac{a_{\text{H}_2\text{O}_2}^{\text{eq}} \times a_{\text{UO}_2^{2+}}^{\text{eq}}}{(a_{\text{H}^+}^{\text{eq}})^2}$$
 (5)

To simplify the calculations, the reaction progress has been calculated by taking into account only the $U(v_1)$ peroxide precipitation reaction (eqn (2)). The first step is to calculate the reaction quotient Q_t for a time t (eqn 6).

$$Q_t = \frac{a_{\text{H}_2\text{O}_2}^t \times a_{\text{UO}_2^{2+}}^t}{\left(a_{\text{U}^+}^t\right)^2} \tag{6}$$

Taking into account the initial reactant concentration and the progress ε , Q_t can be expressed as follows (eqn 7):

$$Q_t = \frac{(\mathbf{C}(\mathbf{U})_0 - \varepsilon) \times (\mathbf{C}(\mathbf{H}_2 \mathbf{O}_2)_0 - \varepsilon)}{(C(\mathbf{H}^+)_0 + 2\varepsilon)^2}$$
(7)

The reaction reaches equilibrium when Q_t equals K. The progress of the reaction is thus (eqn 8):

It is then possible to calculate the theoretical yield of precipitation, which would correspond to (eqn 9):

$$Yield_{theo}^{eq} = \frac{\varepsilon}{C(U)_o} \%$$
 (9)

And then to access the rate of progress of the reaction by comparison with the final experimental yield (eqn 10):

$$\tau = \frac{\text{yield}_{\text{exp}}^{\text{final}}}{\text{vield}_{\text{eq}}^{\text{eq}}} \% \tag{10}$$

3. Results

3.1. Kinetic study associated with the precipitation of studtite

Two sets of precipitation experiments were carried out. For the first one, the initial nitric acid concentration was varied from 0.1 to 2.0 M and the $C(H_2O_2)_{\rm ini}/C(U)_{\rm ini}$ molar ratio was set to 10 (Fig. 1a). In contrast, for the second set, $C(H_2O_2)_{\rm ini}/C(U)_{\rm ini}$ was varied from 10 to 30, while the initial nitric acid concentration was set to 0.5 M. For both sets, the evolution of the precipitation yield (eqn (1)) was monitored (Fig. 1b), since it is clear from eqn (2) that the initial concentrations of both nitric acid and peroxide have an effect on the precipitation of $UO_2(O_2)\cdot 4H_2O$.

Simulations were performed using Phreeqc software and the Thermochimie v12 database³⁵ to determine the uranium speciation under these conditions (Fig ESI 1 and Table ESI 3†). These simulations exclusively relied on thermodynamic constants validated by the latest NEA (Nuclear Energy Agency) review.³³ In addition to the various species included in the Thermochimie database, H_2O_2 , HO_2^- , and the complexes $[(UO_2)(O_2)(OH)]^-$ and $[(UO_2)_2(O_2)_2(OH)]^-$ were added.

Paper **Dalton Transactions**

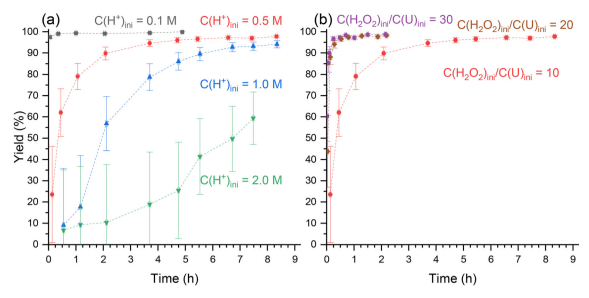


Fig. 1 Evolution of uranium(vi) precipitation yield (a) for various initial nitric acid concentrations (0.1 M in black squares, 0.5 M in red circles, 1.0 M in blue upward triangles, 2.0 M in green downward triangles) and considering $C(H_2O_2)_{ini}/C(U)_{ini} = 10$ and $C(U)_{ini} = 0.07$ M and (b) for various initial $C(H_2O_2)_{inj}/C(U)_{inj}$ molar ratios (10 in red circles, 20 in brown diamonds and 30 in purple hexagons) and with $C(H^+)_{inj} = 0.5$ M and $C(U)_{inj} = 0.07$ M. The dashed line is a guide to the eye and does not represent any mathematical fit.

However, these simulations do not match the experimental data, as the precipitation of U(v1) peroxide is significantly overestimated. This discrepancy is most likely due to the omission of higher-order uranyl nitrate complexes, for which no constants have been validated by the NEA, except for the mononitrate species.

Nevertheless, it remains possible to qualitatively estimate that the only expected solid phase under these conditions is U(vi) peroxide, while the predominant species in solution are uranyl nitrate and the U(v1) hydroxide complex. Regarding peroxo complexes, they only form under highly basic conditions and are therefore not expected in our system.

For both sets of experiments, a plateau seems to be reached more or less quickly depending on the initial conditions. The maximum precipitation yield is different for each precipitation and depends on the acidity or the C(H₂O₂)_{ini}/C(U)_{ini} molar ratio. The higher the initial acidity, the lower the plateau value. From the mass action law, it is clear that the precipitation reaction is favored by increasing the initial uranium or peroxide concentrations and disfavored by increasing the initial acidity. This explains why the plateau reached at the end of the precipitation reaction increases as the initial nitric acid concentration decreases (Fig. 1a) and as the C(H2O2)ini/C (U)ini molar ratio increases (Fig. 1b), which agrees well with Le Chatelier's principle. Kornilov et al. obtained similar results for similar initial acidities of 0.5 and 1.0 M, a molar ratio C(H₂O₂)_{ini}/C(U)_{ini} of 10, and an initial uranium concentration of 0.1 M after 24 hours.²² In 2.0 M H⁺, a yield of 90% was measured after 24 h, while in this work it did not exceed 60% after 7 hours, without reaching equilibrium. This means that the precipitation in 2.0 M H⁺ had not reached equilibrium after 7 hours.

In terms of kinetics, the precipitation rate can be written considering eqn (11), where k_1 and k_{-1} represent the straightforward and the indirect kinetic constants, respectively. α , β , and γ stand for the partial orders related to uranium, hydrogen peroxide and proton concentrations, respectively, and s is the surface of the particles. From this equation, it can be confirmed that the precipitation rate increases with the initial uranium or peroxide concentration (i.e. C(H₂O₂)_{ini}/C(U)_{ini} molar ratio for a given uranium concentration) (Fig. 1b) and decreases with increasing initial proton concentration (Fig. 1a).

$$\nu = \frac{d[U]}{dt} = k_1 \times [U]^{\alpha} \times [H_2 O_2]^{\beta} - k_{-1} \times f(s) \times [H^+]^{2\gamma}$$
 (11)

3.2. Morphological characterization of the studtite precipitates

The influence of the initial nitric acid concentration (i.e. 0.1, 0.5, 1.0 and 2.0 M) and the $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratio (i.e. 10, 20, 40, and 68) on the powder morphology was also studied. SEM images are shown in Fig. 2. It is clear that larger particles are formed when the initial nitric acid concentration is increased for a given C(H₂O₂)_{ini}/C(U)_{ini} molar ratio. A similar effect is also observed when the C(H2O2)ini/C(U)ini molar ratio is decreased for a given initial nitric acid concentration.

As it has already been described for other systems, the morphology of the precipitate can also be correlated with saturation conditions in the solution.36,37 For each sample, the initial saturation index associated with the corresponding precipitation conditions was calculated. A correlation is clearly evidenced, as the particle size decreases with increasing starting SI value. Furthermore, three different types of particle mor-

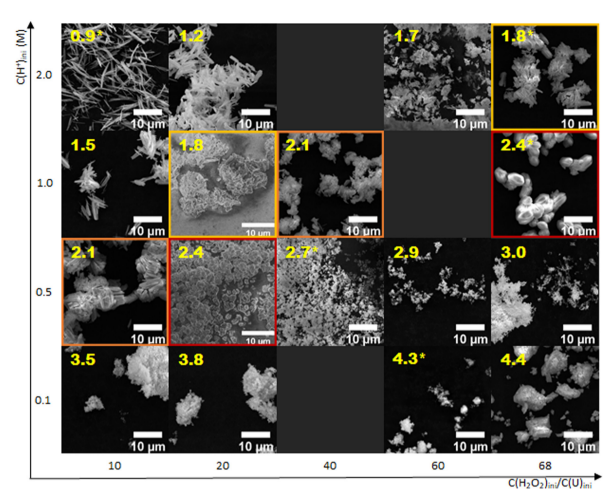


Fig. 2 SEM pictures of uranium(vi) peroxide powders obtained for various initial $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratios and various H^+ concentrations and for $C(U)_{ini} = 0.07M$. The initial saturation index values are shown in yellow. The framed pictures in matching colors correspond to two samples prepared with different initial conditions but with an identical initial saturation index. *The reaction was stopped before reaching 95% progress.

phologies were observed depending on the initial SI values. For an initial $SI_{\rm ini}$ < 2.9, the powder is composed of needle-like crystals less than 1 µm wide and a few micrometers long, which become longer for lower $SI_{\rm ini}$ (around 10 µm for SI=0.9). These needles are agglomerated to form pseudo-spheres for 2.4 < $SI_{\rm ini}$ < 2.9 (from a few micrometers for SI=2.4 and less than 1 µm for SI=2.9). It should be noted that, to our knowledge, these particular shapes have never been observed for $UO_2(O_2)\cdot 4H_2O$. Nanoparticles are obtained for $SI_{\rm ini}$ > 2.9.

The influence of the saturation index on the morphology can be explained by the impact of saturation on the nucleation phenomenon. The higher the saturation index, the more primary nucleation is promoted and the less uranium in solution is available for crystal growth. Consequently, as SI_{ini} increases, the crystal size formed at equilibrium decreases.

With respect to the spherical agglomerates, the morphology of the particles was followed during precipitation with $C(H^+)_{\rm ini}$ = 0.5 M and $C(H_2O_2)_{\rm ini}/C(U)_{\rm ini}$ = 20 by regularly sampling the powder throughout the process (Fig. 3). SEM images show that at the very beginning of the reaction, when only about 20% of the uranium has precipitated, needles of 1–2 µm are formed.

They quickly agglomerate into a parallelepiped shape with nearly aligned needles. The needles and agglomerates stop growing when the yield exceeds 80%. This could mean that the number of agglomerated particles is likely to increase.

3.3. Structural characterization of the studtite precipitates

PXRD patterns were recorded for all the samples prepared. All powders correspond to studtite, $UO_2(O)_2 \cdot 4H_2O$ (monoclinic, space group $C2/c^{38}$). For $C(H_2O_2)_{ini}/C(U)_{ini} = 10$, $UO_2(O)_2 \cdot 4H_2O$ precipitates obtained at different acidities are compared in Fig. 4a. While there is no obvious difference between 0.5, 1.0 and 2.0 M H $^+$, the lower acidity studied (0.1 M) clearly resulted in much larger diffraction peaks. Similarly, the effect of the $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratio was investigated in 0.5 M H $^+$ (Fig. 4b). Here, the studtite synthesized with a molar ratio of 10 has sharper peaks than the other samples.

The difference in PXRD line widths reflects a difference in crystallite size. Crystallite size and unit cell volume were determined by Le Bail refinement from the PXRD patterns (Fig. 5). The data confirm that the crystallite size increases as the

Paper Dalton Transactions

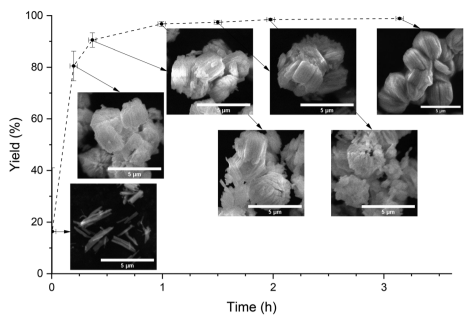


Fig. 3 SEM micrographs of uranium(vi) peroxide obtained during precipitation for $C(H^+)_{ini} = 0.5$ M, $C(H_2O_2)_{ini}/C(U)_{ini} = 20$ and $C(U)_{ini} = 0.07$ M. The dashed line is a quide to the eye and does not represent any mathematical fit.

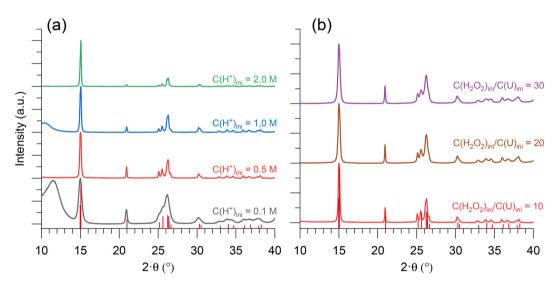


Fig. 4 PXRD patterns of studtite precipitates obtained for various H^+ concentrations (0.1 M in black, 0.5 M in red, 1.0 M in blue, and 2.0 M in green), with $C(H_2O_2)_{ini}/C(U)_{ini}=10$ and $C(U)_{ini}=0.07$ M (a) and for different initial $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratios (10 in red, 20 in brown and 30 in purple) with $C(H^+)_{ini}=0.5$ M and $C(U)_{ini}=0.07$ M (b); PXRD lines in red correspond to uranium(v) peroxide from Burns *et al.*³⁸

initial acidity increases. This crystallite size also decreases as the $C(H_2O_2)_{\rm ini}/C(U)_{\rm ini}$ molar ratio increases.

Except for the uranium(v1) peroxide synthesized at 0.1 M $\mathrm{H^+}$, no changes in the volume of the unit cell (681 ų) are observed. The volume of 684 ų observed for the uranium(v1) peroxide synthesized in 0.1 M $\mathrm{H^+}$ can be explained by nanocrystalline particles. This lattice parameter expansion for nanocrystallites is commonly reported^{39,40} and can be attributed to (i) the accumulation of sorbed species on the nanoparticle surfaces, which has a tensile effect on the crystal lattice⁴¹ and/or

(ii) the accumulation of defects such as vacancies at the boundaries of nanocrystals, which locally increases the lattice parameters. 42

Le Bail refinements were also performed on the other precipitates (Table ESI $2\dagger$). From these data, the crystallite size and lattice parameters of the studtite crystals can then be expressed as a function of the SI_{ini} value (Fig. 6). First of all, regarding the lattice parameters, b and c do not seem to vary with SI_{ini} . In contrast, a and b increase with SI_{ini} above b. b. for conditions where spherical agglomerates are obtained. A

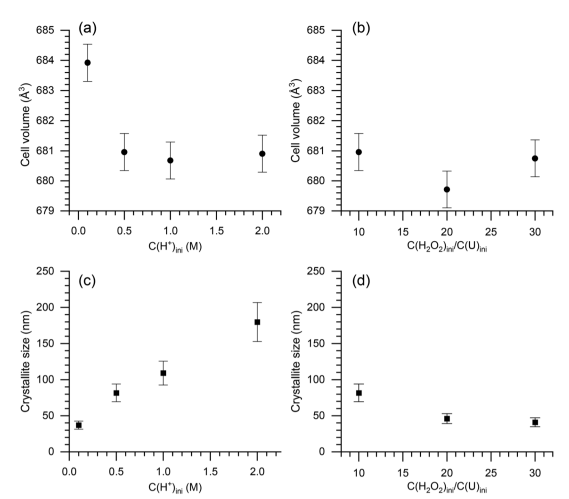


Fig. 5 Unit cell volume of uranium(vi) peroxide for different initial H^+ concentrations (a) and for different initial $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratios (b); crystallite size of uranium(vi) peroxide for different initial H^+ concentrations (c) and for different initial $C(H_2O_2)_{ini}/C(U)_{ini}$ molar ratios (d).

plateau seems to be reached for $SI_{ini} > 2.7$, *i.e.* under conditions where nanoparticles are formed. The cell volume logically follows this tendency. The evolution of these parameters is probably related to the three morphologies obtained. For nanoparticles, the effect of the particle surface is isotropic and greater than for the other two morphologies. In contrast, for the needle-like morphologies, the effect of the surface on the lattice parameter is more likely to be anisotropic. This could then explain the variation of a and β as a function of particle morphology.

Finally, there is a good correlation between the SI_{ini} value and crystallite size. Indeed, as the SI_{ini} increases, the crystallite size decreases from about 150–200 nm to about 10–50 nm. For $SI_{ini} < 2.7$, which corresponds to the formation of studtite needle-like crystals, the crystallite sizes are about 150 nm, which is smaller than the size of the crystals observed in the SEM images for these SI_{ini} values (Fig. 2). This implies that the needle crystals are composed of multiple crystallites. In contrast, it is likely that the crystallite sizes obtained for $SI_{ini} < 2.7$ correspond to the nanoparticles observed in the SEM images.

4. Discussion

In the context of the high importance of powder morphology for fuel fabrication, the saturation index seems to be the key parameter controlling the morphology and the microstructure of the precipitate. The higher the SI_{ini}, the smaller the crystallite and the crystal size of the agglomerates. These observations suggest that neither the initial concentration of nitric acid nor that of peroxide ions, considered individually and for the same initial SI, influences the morphology of the precipitate. Instead, the morphology appears to be primarily influenced by the value of the saturation index in the solution. Indeed, it appears to be identical for a given saturation index value (images within frames of matching colors in Fig. 2), even though the initial acidity and peroxide concentration are different. The relative lack of effect of the initial acidity and C(H₂O₂)_{ini}/C(U)_{ini} ratio on the particle characteristics compared to the saturation index is somewhat surprising, since the kinetics of precipitation would be expected to differ under different initial chemical conditions. Therefore, the nucleation processes should be different and thus the final powder mor-

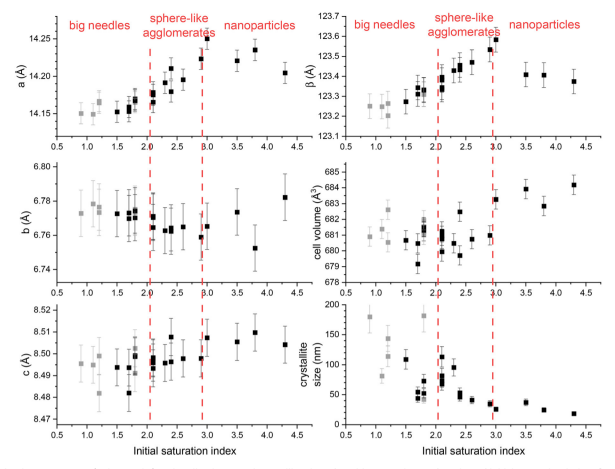


Fig. 6 Lattice parameters $(a, b, c \text{ and } \beta)$, unit cell volume and crystallite size of studtite crystals as a function of initial saturation index. Grey points are U(vi) peroxide samples from precipitation that have not reached equilibrium. Refinement results are available in Table ESI 2.†

phology should be distinct. Two additional precipitation experiments with kinetic monitoring were performed with the same initial saturation index (*i.e.* 1.7) but different initial reactant concentrations (*i.e.* $C(U)_{ini} = 0.07$ M, $C(H^+)_{ini} = 1.0$ M and 2.0 M, and $C(H_2O_2)_{ini}/C(U)_{ini} = 14.7$ and 59.4) to confirm the predominant influence of the initial saturation on the final morphology (Fig. 7).

First, the morphologies at the end of the precipitation are indeed identical (Fig. 7). However, the evolution of the yield does not follow the same curve, although the plateau reached by the two precipitation processes is almost the same. For the precipitation in 1.0 M H⁺ (Fig. 7, red squares), a first plateau seems to be formed at the beginning of the experiment, unlike under other conditions where precipitation occurs faster. This difference is probably explained by a different evolution of the saturation index during precipitation. The precipitation in 2.0 M H⁺, with more peroxide and no plateau at the beginning, exhibits a higher saturation index that is maintained in the reacting system during the precipitation compared to the first experiment. This agreement of the morphologies but not of the kinetics for the same SI_{ini} could be explained by the fact that the evolution of these two precipitation processes is not sufficiently different to form different morphologies or that

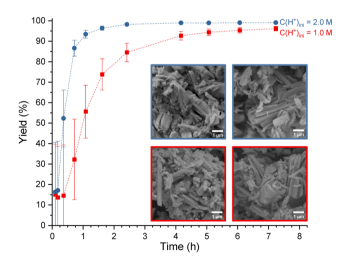


Fig. 7 Uranium(vi) precipitation yield evolution and morphologies for different initial reactant concentrations $(C(U)_{ini}=0.07\ M,\ C(H^+)_{ini}=1.0\ M$, and $C(H_2O_2)_{ini}/C(U)_{ini}=14.7$ in red squares and $C(U)_{ini}=0.07\ M$, $C(H^+)_{ini}=2.0\ M$, and $C(H_2O_2)_{ini}/C(U)_{ini}=59.4$ in blue circles) but the same initial supersaturation index (1.7). The dashed lines are a guide to the eye and do not represent real values. The scale bars on SEM images represent 1 um.

 Table 1
 Specific surface areas (SSA) of powders obtained for different

Dalton Transactions

IS _{ini}	3.5	2.4	2.1	1.8
$C(H^+)_{ini}(M)$	0.1	0.5	1.0	2.0
$C(H_2O_2)_{ini}/C(U)_{ini}$	10	20	40	68
$C(H_2O_2)_{ini}/C(U)_{ini}$ SSA $(m^2 g^{-1})$	21 ± 1	7 ± 1	5 ± 1	5 ± 1

the difference between the particles is not visible at the SEM scale. Nevertheless, it seems that in very acidic media, particle morphologies are controlled only by supersaturation. A wide variety of particle morphologies from nanoparticles to needles or spherical agglomerates can then be synthesized on demand.

Finally, given that thermal treatment preserves the morphology of U(vi) peroxides, ²¹ oxides with corresponding morphologies can be produced. However, it remains difficult to determine whether a specific morphology is optimal for nuclear fuel fabrication. As mentioned above, the two key parameters to consider are powder flowability and sinterability.

In terms of flowability, and based on the correlations established by Cayla Arianer *et al.*, 25 the most influential morphological factor appears to be the particle size distribution for particle size below 75 μ m. While spherical morphologies (such as those observed for SI_{ini} values between 2.4 and 2.9) are generally expected to improve flowability, this advantage is offset by the strong cohesion induced by small particle sizes.

From a reactivity standpoint, which directly influences sinterability, the nanometric powder obtained at $\mathrm{SI}_{\mathrm{ini}}$ values above 2.9 appears to be the most favorable. Specifically, the specific surface areas (SSA) of four representative morphologies from this study (Table 1) were determined *via* the BET method, revealing that the nanometric powder synthesized at $\mathrm{SI}_{\mathrm{ini}}=3.5$ exhibits the highest SSA. In general, a higher specific surface area translates into lower required pressing pressure and sintering temperature, which ultimately leads to a higher final pellet density.⁴³

5. Conclusion

The effect of initial chemical conditions on the precipitation of studtite in acidic media has been studied. Initial acidity and peroxide concentration (expressed as the molar ratio of initial peroxide concentration to uranium concentration at constant initial C(U)) both affect the kinetics of precipitation. Increasing acidity and decreasing peroxide concentration both slow down the precipitation of studtite.

Secondly, a correlation was found between the initial saturation index and the final morphology and crystallite size of the powder. The higher the $\mathrm{SI}_{\mathrm{ini}}$ value, the smaller the particles and the smaller the crystallite size. Conversely, low $\mathrm{SI}_{\mathrm{ini}}$ values promote the formation of larger crystals with larger crystallite sizes. An intermediate morphology consisting of spherical needle clusters is also observed for 2.4 < $\mathrm{SI}_{\mathrm{ini}}$ < 2.9. This corre-

lation is explained by the fact that a high saturation index favors the nucleation phenomenon over crystal growth, which increases the formation of small particles. At low values of $SI_{\rm ini}$, larger crystals are formed because the crystal growth phenomenon is promoted. Under the conditions studied, the saturation conditions seem to be the only parameter controlling the morphology. In fact, for the same $SI_{\rm ini}$ value, but different initial concentrations of the reactants, the final morphologies were found to be similar. Although the obtained morphologies cannot be considered as optimal for fuel fabrication at this stage, these results show that the morphology of $U(v_1)$ peroxide powder precipitated from an acidic uranyl solution can be controlled by adding an appropriate amount of peroxide ions, corresponding to the correct initial saturation index.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

The authors are grateful to Stéphanie Szenknect for her help with Phreeqc simulations and Renaud Podor and Joseph Lautru for SEM micrographs. The authors would like to thank the CEA for funding the PhD work of Lucas Muller.

References

- 1 K. Walenta, On Studtite and Its Composition, *Am. Mineral.*, 1974, **59**(1–2), 166–171.
- 2 D. Lunt, P. Boshoff, M. Boylett and Z. El-Ansary, Uranium Extraction: The Key Process Drivers, *J. South. Afr. Inst. Min. Metall.*, 2007, **107**(7), 419–426.
- 3 J. Dugua, Uranium Peroxide in the Form of Spherical Particles Having Good Flowability and Processes for Preparing Same, US4530823A, 1985.
- 4 F. E. Caropreso and D. F. Kreuz, *Production of Uranium Peroxide*, US4024215A, 1977.
- 5 B. Morel, D. Amaraggi, M. Arab, R. Thomas, M. Rivenet and F. Abraham, *Process for Converting UO3 or U308 to Hydrated UO4*, FR2969659B1, 2013.
- 6 H. Lagrave, C. Beretti and P. Bros, Treatment of Uranium and Plutonium Solutions Generated in Atalante by R and D Activities, 2008.
- 7 B. Pastina and J. A. LaVerne, Hydrogen Peroxide Production in the Radiolysis of Water with Heavy Ions, J. Phys. Chem. A, 1999, 103(11), 1592–1597.

Paper

8 B. D. Hanson, B. McNamara, E. C. Buck, J. I. Friese, E. Jenson, K. Krupka and B. W. Arey, Corrosion of Commercial Spent Nuclear Fuel. 1. Formation of Studtite and Metastudtite, *Radiochim. Acta*, 2005, **93**(3), 159–168.

- 9 B. McNamara, B. D. Hanson, E. C. Buck and C. Soderquist, Corrosion of Commercial Spent Nuclear Fuel. 2. Radiochemical Analyses of Metastudtite and Leachates, *Radiochim. Acta*, 2005, **93**(3), 169–175.
- 10 C. Jégou, R. Caraballo, J. De Bonfils, V. Broudic, S. Peuget, T. Vercouter and D. Roudil, Oxidizing Dissolution of Spent MOX47 Fuel Subjected to Water Radiolysis: Solution Chemistry and Surface Characterization by Raman Spectroscopy, J. Nucl. Mater., 2010, 399, (1), 68–80.
- 11 A. Perrot, A. Canizares, S. Miro, L. Claparede, R. Podor, T. Sauvage, S. Peuget, C. Jegou and N. Dacheux, *In Situ* Raman Monitoring of Studtite Formation under Alpha Radiolysis in ¹⁸O-Labeled Water, *J. Nucl. Mater.*, 2024, 600, 155267.
- 12 M. L. Schlegel and C. Jegou, Uraninite Alteration by H₂O₂ Solutions and Formation of Secondary Phases: An, in Situ MicroRaman Spectroscopy and Synchrotron X-Ray Diffraction Study, J. Nucl. Mater., 2022, 572, 154056.
- 13 J. S. Hopkins, J. A. Golding and G. M. Ritcey, The Critical Factors Influencing Uranium Precipitation by Hydrogen Peroxide: The Use of Experimental Design Techniques, *Hydrometallurgy*, 1987, 17(3), 315–334.
- 14 M. Shabbir and K. E. Tame, Hydrogen Peroxide Precipitation of Uranium; Report of investigations RI7931, U.S, Department of the Interior, Bureau of Mines, 1974.
- 15 R. Gupta, V. M. Pandey, S. R. Pranesh and A. B. Chakravarty, Study of an Improved Technique for Precipitation of Uranium from Eluted Solution, *Hydrometallurgy*, 2004, 71(3-4), 429-434.
- 16 A. E. Cahill and L. E. Burkhart, Continuous Precipitation of Uranium with Hydrogen Peroxide, *Metall. Trans. B*, 1990, 21(5), 819–826.
- 17 L. Mojica-Rodríguez, M. Bertrand, J. Gaillard, H. Muhr, E. Plasari, F. Auger and E. Brackx, Study on Uranium Peroxide Precipitation in a Continuous Fluidized-Bed Reactor with Mechanical Stirring, *Nucl. Eng. Des.*, 2015, 293, 150–158.
- 18 S. Planteur, M. Bertrand, E. Plasari, B. Courtaud and J.-P. Gaillard, Thermodynamic and Crystal Growth Kinetic Study of Uranium Peroxide, *CrystEngComm*, 2013, 15(12), 2305.
- 19 S. Planteur, M. Bertrand, E. Plasari, B. Courtaud and J. P. Gaillard, Crystal Growth Kinetics of the Uranium Peroxide, *Procedia Chem.*, 2012, 7, 725–730.
- 20 K.-W. Kim, J.-T. Hyun, K.-Y. Lee, E.-H. Lee, K.-W. Lee, K.-C. Song and J.-K. Moon, Effects of the Different Conditions of Uranyl and Hydrogen Peroxide Solutions on the Behavior of the Uranium Peroxide Precipitation, *J. Hazard. Mater.*, 2011, 193, 52–58.
- 21 I. J. Schwerdt, A. Olsen, R. Lusk, S. Heffernan, M. Klosterman, B. Collins, S. Martinson, T. Kirkham and L. W. McDonald, Nuclear Forensics Investigation of

- Morphological Signatures in the Thermal Decomposition of Uranyl Peroxide, *Talanta*, 2018, **176**, 284–292.
- 22 A. S. Kornilov, E. V. Piterkina, K. O. Shcherbakova, A. O. Makarov and O. S. Dmitrieva, Specific Features of Peroxide Precipitation of Uranium from Acid Water– Ethanol Solutions, *Radiochemistry*, 2020, 62(2), 173–176.
- 23 N. Hibert, B. Arab-Chapelet, M. Rivenet, L. Venault, C. Tamain and O. Tougait, *Dalton Trans.*, 2022, 51(34), 12928–12942.
- 24 A. L. Tamasi, L. J. Cash, W. Tyler Mullen, A. R. Ross, C. E. Ruggiero, B. L. Scott, G. L. Wagner, J. R. Walensky, S. A. Zerkle and M. P. Wilkerson, Comparison of Morphologies of a Uranyl Peroxide Precursor and Calcination Products, *J. Radioanal. Nucl. Chem.*, 2016, 309(2), 827–832.
- 25 L. Cayla Arianer, M. Leturia, A.-C. Robisson, C. Ablitzer, E. Daouk and K. Saleh, Assessment of the Influence of Model Powder Characteristics on Their Flowability Using Correlation Matrices, *Powder Technol.*, 2025, 453, 120636.
- 26 Powder Technology Handbook, ed. H. Masuda, K. Higashitani and H. Yoshida, CRC Press, Boca Raton, 3rd edn, 2006.
- 27 D. Sicsic, F. Balbaud-Célérier and B. Tribollet, Mechanism of Nitric Acid Reduction and Kinetic Modelling, Eur. J. Inorg. Chem., 2014, 6174–6184.
- 28 R. E. Elson, Lawrence Radiation Laboratory, in *The Homogenous Decomposition of Hydrogen Peroxide by Plutonium(IV)*, UCRL; 6536; Lawrence Radiation Laboratory, Livermore, CA, 1961.
- 29 N. Dacheux and J. Aupiais, Determination of Uranium, Thorium, Plutonium, Americium, and Curium Ultratraces by Photon Electron Rejecting α Liquid Scintillation, *Anal. Chem.*, 1997, **6**(13), 2275–2282.
- 30 A. Le Bail and D. Louër, Smoothing and Validity of Crystallite-Size Distributions from X-Ray Line-Profile Analysis, *J. Appl. Crystallogr.*, 1978, 11(1), 50–55.
- 31 C. Frontera and J. Rodríguez-Carvajal, FullProf as a New Tool for Flipping Ratio Analysis, *Physica B: Condens. Matter*, 2003, 335(1), 219–222.
- 32 P. Thompson, D. E. Cox and J. B. Hastings, Rietveld Refinement of Debye–Scherrer Synchrotron X-Ray Data from Al₂O₃, *J. Appl. Crystallogr.*, 1987, **20**(2), 79–83.
- 33 OECD, Second update on the Chemical Thermodynamics of Uranium, Neptunium, Plutonium, Americium And Technetium, Organisation for Economic Co-operation and Development, Paris, 2021, vol. 14.
- 34 K.-A. H. Kubatko, K. B. Helean, A. Navrotsky and P. C. Burns, Stability of Peroxide-Containing Uranyl Minerals, *Science*, 2003, **302**(5648), 1191–1193.
- 35 B. Madé, W. Bower, S. Brassinnes, E. Colàs, L. Duro, P. Blanc, A. Lassin, L. Harvey and J. D. Begg, Recent Developments in ThermoChimie A Thermodynamic Database Used in Radioactive Waste Management, Appl. Geochem., 2025, 180, 106273.
- 36 S. Shaddel, S. Ucar, J.-P. Andreassen and S. W. Østerhus, Engineering of Struvite Crystals by Regulating Supersaturation – Correlation with Phosphorus Recovery,

Crystal Morphology and Process Efficiency, *J. Environ. Chem. Eng.*, 2019, 7(1), 102918.

Dalton Transactions

- 37 L. Wang, J.-P. Andreassen and S. Ucar, Precipitation of Silver Particles with Controlled Morphologies from Aqueous Solutions, *CrystEngComm*, 2020, 22(3), 478–486.
- 38 P. C. Burns and K.-A. Hughes, Studtite, [(UO₂)(O₂)(H₂O)₂] (H₂O)₂: The First Structure of a Peroxide Mineral., *Am. Mineral.*, 2003, **88**(7), 1165–1168.
- 39 K. Lu and Y. H. Zhao, Experimental Evidences of Lattice Distortion in Nanocrystalline Materials, *Nanostruct. Mater.*, 1999, 12(1-4), 559–562.
- 40 X. D. Liu, H. Y. Zhang, K. Lu and Z. Q. Hu, The Lattice Expansion in Nanometre-Sized Ni Polycrystals, *J. Phys.: Condens. Matter*, 1994, **6**(34), L497.
- 41 T. V. Plakhova, A. Y. Romanchuk, D. V. Likhosherstova, A. E. Baranchikov, P. V. Dorovatovskii, R. D. Svetogorov, T. B. Shatalova, T. B. Egorova, A. L. Trigub, K. O. Kvashnina, V. K. Ivanov and S. N. Kalmykov, Size Effects in Nanocrystalline Thoria, *J. Phys. Chem. C*, 2019, 123(37), 23167–23176.
- 42 W. Qin, T. Nagase, Y. Umakoshi and J. A. Szpunar, Relationship between Microstrain and Lattice Parameter Change in Nanocrystalline Materials, *Philos. Mag. Lett.*, 2008, **88**(3), 169–179.
- 43 I. R. Gibson, S. Ke, S. M. Best and W. Bonfield, Effect of Powder Characteristics on the Sinterability of Hydroxyapatite Powders, *J. Mater. Sci.:Mater. Med.*, 2001, 12(2), 163–171.