SPECIATION AND RELEASE KINETICS OF THE FISSION PRODUCTS Mo, Cs, Ba and I FROM NUCLEAR FUELS IN SEVERE ACCIDENT CONDITIONS

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ABSTRACT

A speciation and release kinetics study of the gaseous fission products freed from nuclear fuels in conditions of severe accident (temperature range $1300^{\circ}\text{C} - 2500^{\circ}\text{C}$) as a function of the applied thermodynamic conditions is carried out using the experimental feedback collected at CEA within the VERCORS and VERDON programs.

The nature of the major species involved is deduced from the analysis of the specific features of the Cs-Mo-Ba-I-O chemical system in the composition range of irradiated nuclear fuels. Stable gaseous Cs dimolybdate and Ba monomolybdate species, i.e. $Cs_2Mo_2O_7(g)$ and $BaMoO_4(g)$, clearly predominates in oxidising conditions, which contributes to increase significantly the released fraction of molybdenum as compared to reducing conditions.

An analytical model accounting for thermo-chemistry in the release rate of chemically reactive elements is applied to estimate the release of Cs, I, Mo and Ba in severe accident conditions. Good agreement is achieved with on-line measurements.

1. Introduction

Thermodynamic equilibrium modelling is a powerful tool for the description of complex chemical processes and systems, like those involved in irradiated nuclear fuels. A special case of great current interest in this field is the assessment of the possible consequences of severe reactor accidents. The major concern is to determine the Source Term, i.e. the speciation, release rate and global released fraction of the fission products (FP) and other radioactive materials.

Temperature and oxygen partial pressure (i.e. oxygen potential) are the major parameters governing the speciation and release behavior of the fission products from nuclear oxide fuels. In severe accident conditions, temperature significantly exceeds the nominal one and strong variations of oxygen potential are likely to occur because of oxidation of the Zr-clad by water (giving large amount of H₂) followed by ingress of steam or air. Major variations in the speciation and release of volatile fission products are thus expected in such off-normal conditions.

A large number of research programs in the field of severe reactor accidents have been undertaken in various countries, in order to assess the Source Term. Some of them like PHEBUS-FP are devoted to integral tests while others are devoted to separate-effect experiments¹. Among the later, the VERCORS and VERDON programs are composed of a series of analytical tests conducted on samples of irradiated PWR fuels, in carefully controlled temperature and oxido-reduction conditions. From those thermal tests carried out in highly instrumented experimental facilities, extensive and invaluable data were collected and then used to improve the models for the Source Term evaluation.

Among the chemical systems arising from fission products in irradiated oxide fuels, the Cs-Mo-O system is one of the most important in assessing the consequences of off-normal operating conditions^{1,2}. In calculation of fission products (FP) releases from irradiated fuels, Cs is usually found to be freed from the reactor core as $Cs_2MoO_4(g)$, CsOH(g) and CsI(g), while Mo is freed as $Cs_2MoO_4(g)$ and $MoO_x(g)$, O< x< 3. However, discrepancies between experimental and calculated released fractions still remain³, possibly owing to limitations in the thermodynamic database (cf. §.3.1). In fact, in view of recent specific studies performed on the Mo-Cs-I-O-H system in conditions representative of severe accidents, it is highly suspected

that caesium polymolybdates should contribute significantly to the Mo and Cs releases from irradiated fuels². Thus, complementary to basic investigations on the thermodynamics of caesium monomolybdate Cs_2MoO_4 , the high temperature behaviour of higher molybdates such as $Cs_2Mo_2O_7$ has been studied in details recently².

The present paper is devoted to a speciation and release kinetics study of the gaseous fission products freed from nuclear fuel in conditions of severe accident (temperature range 1300° C – 2500° C), as a function of the applied thermodynam ic conditions. The analysis relies on the experimental feedback collected at CEA within the VERCORS and VERDON programs. At first, the nature of the major chemical species involved is deduced from the analysis of the specific features of the Cs-Mo-I Ba-O chemical system in the composition range of irradiated nuclear fuels, as reflected by the experimental released fractions curves. Then, an analytical model accounting for thermo-chemistry in the release rate of chemically reactive elements is outlined and applied to FP releases in severe accident conditions.

2. Thermochemical characteristics of the VERCORS and VERDON tests

The annealing tests performed within the VERCORS and VERDON programs aim at quantifying the release of fission products (FP) and actinides from irradiated nuclear ceramics during a severe accident. From 1989 onwards, extensive and invaluable data were collected about i) the released fraction and release rates of FPs from irradiated fuels ii) the nature and chemical composition of the vapours and aerosols emitted and iii) the degradation mechanisms of the fuel.

Full descriptions of the experimental set-ups and detailed analyses of the data collected in various temperature and redox conditions are given in reference 1 (and related references). To summarize, the fuel samples are composed of PWR sections containing three UO₂ (or MOX) pellets in their original zircaloy cladding. Before thermal testing, the samples are generally reirradiated at low linear power (in Material Testing Reactor) in order to rebuild the inventory of short-lived fission products. Then, the experimental sequence consists in applying first a temperature plateau (at 1300° C or 1500° C) under con trolled oxygen potential (steam, H_2/H_2 O mixture or H_2) in order to fully oxidize the cladding. Then a temperature ramp is applied (with occasionally intermediate temperature plateaus) under controlled oxido-reduction conditions (steam, H_2/H_2 O mixture or H_2) until the end of the sequence (i.e. collapse of the sample most often).

The present study refers specifically to the VERCORS-5, -RT3, -RT6 and VERDON-1 tests, for which a consistent set of data covering the whole range of redox conditions of interest is available. All relevant information and data for the current analysis, taken out from reference¹, are collected in Table 1. The released fraction curves for Cs. Mo. Ba and I for each test under consideration are shown in Figs 1 to 4. Figure 5 gives the thermodynamical characteristics (temperature, oxygen potential) of the tests listed in Table 1, in consistency with the thermohydraulic sequences followed during the experiments. Thus, the set of thermal tests selected in this study covers a range of redox conditions varying from highly reducing (under hydrogen atmosphere) to highly oxidising (under steam atmosphere). The oxidation step until the first temperature plateau (at 1300℃ or 1500℃) is carried out for VERDON-1. VERCORS-5 and VERCORS-RT6 in redox conditions slightly higher than that of the (MoO₂/Mo) redox buffer, which is assumed to fix the oxygen potential of the fuel in nominal operating conditions⁴ (-380 kJ/mol O₂ at ~1000℃). It is clear from Fig 5 that VERCORS-5, VERCORS-RT6 and VERDON-1 are submitted to similar conditions during the oxidation step up to the first temperature plateau (at 1300℃ or 1500℃). In contr ast, during the high temperature step, the VERCORS-5 test is carried out under highly oxidising conditions (steam), while the VERCORS-RT6 test is carried out under the same H₂O/H₂ atmosphere as that used for the oxidation step and the VERDON-1 test is carried out under highly reducing conditions similar to that of VERCORS-RT3.

By carefully examining the released fraction curves of Cs, Mo and I from fuel samples of varying burn up in varying experimental conditions (Figs 1 to 4), a number of typical features indicative of the speciation of the gaseous species involved in the process emerge clearly, as described below:

- In oxidising conditions (VERCORS-5, -RT6 and VERDON-1), caesium and molybdenum are released together from about 1200℃ onward, which suggests that Cs and Mo are very likely involved into mixed chemical species of type $Cs_xMo_yO_z$;
- Up to the oxidation plateau at 1500℃, the VERCORS -RT6 and VERDON-1 samples are submitted to similar oxidising redox conditions and they both show similar released fraction curves for Cs and Mo with a ratio %Mo/%Cs ~ 2/3;
- In highly oxidising conditions (VERCORS-5), Ba releases partially together with Cs and Mo, which would be consistent with the presence of both mixed caesium-molybdenum and barium molybdenum species. The later would thus contribute to further increase the release of molybdenum as compared to less oxidising conditions;
- The VERCORS-RT3 sample and the VERDON-1 sample (in the high temperature step only) are submitted to similar reducing conditions. Both tests clearly show that Cs, Mo and Ba are released independently in those redox conditions, which indicates that the mixed Cs-Mo species encountered in the VERCORS RT6 and VERDON-1 (at 1500℃) tests as well as the Ba-Mo species encountered in the VERCORS-5 test in oxidising conditions are not stable any more when the oxygen potential is strongly decreased;
- Above 1300°C, the released fraction curves of iodi ne and caesium evolves in parallel with time regardless the experimental conditions (burn-up, temperature, redox). The ratio of the released iodine fraction over that of caesium (%I/%Cs) is invariant and close to 1.0. This peculiar feature of the release curves of Cs and I is typical of the simultaneous release of Cs, Mo and I.

In the following section, a simplified thermodynamic speciation scheme is developed in consistency with the characteristics of the experimental FP release curves of the selected thermal tests, assuming thermodynamic equilibrium is achieved between the most likely major species, as naturally expected at high temperature conditions.

Test reference	Fuel Burn-up	Atmosphere composition			Experimental sequence
	GWj/tU	H₂O g/min	H ₂ g/min	He g/min	
VERCORS-5	38.3	1.5	0.027	0	- Ramp from 400℃ to 800℃ - 30 min. plateau at 800℃ - Ramp from 800℃ to 1000℃ - 30 min. plateau at 1000℃ - Ramp from 1000℃ to 1300℃ - 70 min. plateau at 1300℃
			Steam		- Ramp from 1300℃ to 2300℃
VERCORS- RT6	71.8	1.5	0.027	0	 Ramp from 400℃ to 1500℃ 1h plateau at 1500℃ Ramp from 1500℃ to 2200℃ 30 min. plateau at 2000℃, 2100℃ and 2200℃
VERDON-1	72.0	1.5	0.027	0	- Ramp from 400℃ to 1500℃ - 1h plateau at 1500℃
		0.018	0.020	0.24	- Ramp from 1500℃ to 2400℃
VERCORS- RT3	39.0	0.075	0.075	0.12	- Ramp from 400℃ to 1500℃ - 1h plateau at 1500℃ - Ramp from 1500℃ to 2200℃

Tab 1: Thermohydraulic experimental conditions of the thermal treatment tests under consideration in the current speciation study (Figs 1 to 5). *The data come from reference*¹.

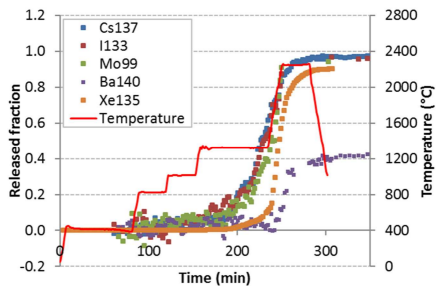


Fig 1. FP release kinetics during the VERCORS-5 test

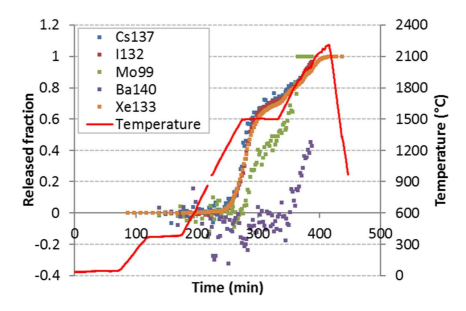


Fig 2. FP release kinetics during the VERCORS-RT6 test

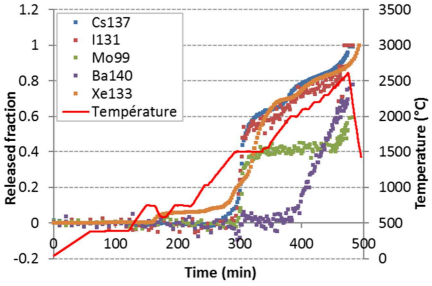


Fig 3. FP release kinetics during the VERDON 1 test

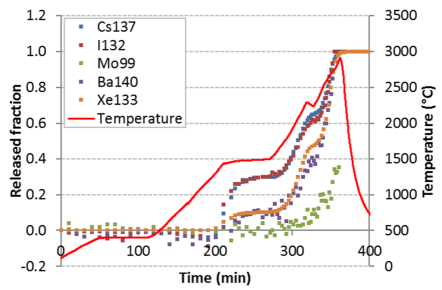


Fig 4. FP release kinetics during the VERCORS-RT3 test

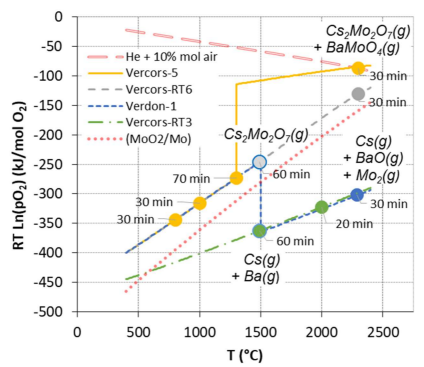


Fig 5. Thermochemical characteristics (temperature, oxygen potential) of the thermal tests listed in Table 1

The coloured discs indicate the temperature plateaus (with their duration) during the experimental sequences. All the necessary data for the thermodynamic calculations come from reference⁶.

3. Theoretical considerations

3.1. Cs-Mo-I equilibria in gaseous phase

Let us consider, in consistency with the experimental observations, a gaseous chemical system composed of the two species $Cs_xMo_yO_z(g)$ and CsI(g) released from an irradiated UO_2 fuel.

At the instant t, the released quantities of the elements Cs, Mo and I can be written:

$$Cs(t) = x * Cs_x Mo_v O_z(t) + CsI(t)$$
 (1)

$$Mo(t) = y * Cs_x Mo_y O_z(t)$$
 (2)

$$I(t) = CsI(t) (3)$$

The simultaneous releases of Cs, Mo and I are then linked by the relation:

$$Cs(t) = \frac{x}{y} * Mo(t) + I(t)$$
 (4)

with $Cs(t) \le Cs_{tot}$, $Mo(t) \le Mo_{tot}$ and $I(t) \le I_{tot}$. In terms of released fractions, equation (4) reads:

$$\%Cs(t) = \frac{x}{y} * \frac{Mo_{tot}}{Cs_{tot}} * \%Mo(t) + \frac{I_{tot}}{Cs_{tot}} * \%I(t)$$
 (5)

The released fractions of Mo and I as compared to that of Cs are therefore linked by the relation:

$$1 = \frac{x}{y} * \frac{Mo_{tot}}{Cs_{tot}} * \frac{\%Mo(t)}{\%Cs(t)} + \frac{I_{tot}}{Cs_{tot}} * \frac{\%I(t)}{\%Cs(t)}$$
 (6)

The quantities Mo_{tot} , Cs_{tot} and I_{tot} vary with the burn-up. However, in the range 0 - 90 GWj/t U, the variation of the quantity of each fission product with burn-up may be well approximated by a linear function⁵. Consequently, the ratios (Mo_{tot}/Cs_{tot}) and (I_{tot}/Cs_{tot}) may be considered constant and independent of burn-up in the range 0 - 90 GWj/t U.

Then, by assuming that no significant loss of Cs, Mo and I occurs prior to the thermal treatment test, the following equalities apply⁵ in the range 0 - 90 GWj/t U:

$$\frac{Mo_{tot}}{Cs_{tot}} = 1.371$$
 ; $\frac{I_{tot}}{Cs_{tot}} = 0.077$ (7)

Combination of the relations (6) and (7) gives:

$$1 = 1.371 * \frac{x}{y} * \frac{\% Mo(t)}{\% Cs(t)} + 0.077 * \frac{\% I(t)}{\% Cs(t)}$$
 (8)

As Cs is involved into two different chemical species including Mo and I, the released fractions of Mo, Cs and I are not independent. In fact, the released fractions of both Mo and I as compared to that of Cs are linked by the relation (8) which only depends on the stoichiometric coefficients x and y of the species $Cs_xMo_yO_z$.

The experimental feedback of the VERCORS and VERDON-1 tests shows that the ratio %I(t)/%Cs(t) is invariant and close to 1.0 in most tests, irrespective of the redox and temperature conditions. Considering $%I(t)/%Cs(t) \approx 1$ in equation (8) gives:

$$y \approx \frac{3}{2} * x * \frac{\% Mo(t)}{\% Cs(t)} \tag{9}$$

In view of recent published studies, it is essential when nuclear fuels are overheated in oxidising conditions to consider the incidence of gaseous caesium polymolybdates required in oxidising conditions to consider the incidence of gaseous caesium polymolybdates required in oxide required in oxide required in the incidence of gaseous caesium polymolybdates required in oxide required required in oxide required in oxide required re

$$y \approx 3 * \frac{\%Mo(t)}{\%Cs(t)}$$
 (10)

Applying equation (10) to the VERDON-1 or VERCORS-RT6 tests for which the ratio %Mo/%Cs is estimated to ~2/3 at 1500°C gives y = 2, which matches with the caesium dimolybdate species $Cs_2Mo_2O_7$. Note that if only caesium monomolybdate Cs_2MoO_4 is considered, then equation (10) gives Mo(t)/Mcs(t) = 1/3, which is significantly different from, and thus inconsistent with, what is observed experimentally.

It is therefore concluded in consistency with the experimental observations, that the most likely Cs species released at about 1500°C and around -250 kJ/mol O_2 is the caesium dimolybdate $Cs_2Mo_2O_7(g)$. This finding is fully consistent with recently published partitioning calculations of Cs and Mo in gaseous phase in severe accident conditions, which predict the presence in significant amount of $Cs_2Mo_2O_7(g)$ in mid-oxidising and moderately high temperature conditions².

3.2. Ba-Mo equilibria in gaseous phase

In strongly oxidising conditions such as in VERCORS-5 test, the occurrence of mixed barium-molybdenum species in addition to caesium-molybdenum species could be responsible for the rise of the Mo release as compared to less oxidising redox conditions.

Thus, by assuming that Cs and I are involved into the sole species $Cs_2Mo_2O_7(g)$ and CsI(g), and that Ba is involved into a single species of type $Ba_xMo_yO_z(g)$, the total released quantities of Cs, Mo, Ba and I at the end of the test (instant t_f) can be written as:

$$Cs(t_f) = 2 * Cs_2 Mo_2 O_7(t_f) + CsI(t_f)$$
 (11)

$$Mo(t_f) = 2 * Cs_2 Mo_2 O_7(t_f) + y * Ba_x Mo_y O_Z(t_f)$$
 (12)

$$Ba(t_f) = x * Ba_x Mo_y O_z(t_f)$$
(13)

$$I(t_f) = CsI(t_f) \tag{14}$$

 $I(t_f) = CsI(t_f)$ which gives, in terms of released fractions:

$$\%Mo(t_f) = \frac{Cs_{tot}}{Mo_{tot}} * \%Cs(t_f) + \frac{y}{x} * \frac{Ba_{tot}}{Mo_{tot}} * \%Ba(t_f) - \frac{I_{tot}}{Mo_{tot}} * \%I(t_f)$$
(15)

If no significant loss of Cs, Mo Ba and I occurs prior to the thermal treatment test, the following equalities apply⁵ in the range 0 - 90 GWj/t U:

$$\frac{Cs_{tot}}{Mo_{tot}} = 0.730$$
 ; $\frac{I_{tot}}{Mo_{tot}} = 0.0562$; $\frac{Ba_{tot}}{Mo_{tot}} = 0.592$ (16)

Combination of relations (15) and (16) gives:

$$1 = 0.730 * \frac{\%Cs(t_f)}{\%Mo(t_f)} + \frac{y}{x} * 0.592 * \frac{\%Ba(t_f)}{\%Mo(t_f)} - 0.0562 * \frac{\%I(t_f)}{\%Mo(t_f)}$$
(17)

In the VERCORS-5 test, the experimental release curves of Cs, Mo, Ba and I provide 1 %Mo(t_f) \approx %Cs(t_f) \approx %I(t_f) \approx 1 and %Ba(t_f) \approx 0.55. Inserting these data in equation (17) gives then y/x \approx 1.0, which is fully consistent with the mixed Ba-Mo species of stoichiometry $BaMoO_4$, as thermodynamically expected (Fig 5).

It is thus concluded in consistency with the experimental observations, that Ba is most likely released at high temperature and in highly oxidising conditions under the main species $BaMoO_4(g)$. Note that Ba is partially released under $BaMoO_4(g)$ during the test, the remaining part being most likely immobilized into the condensed phase $BaZrO_3(sc)$.

4. Simplified speciation model for Mo, Cs and Ba in severe accident conditions

The major gaseous species for Mo, Cs and Ba occurring in the thermal tests considered in the present study are summarised in Figs 6 and 7, at 1500 and 2200° C as a function of oxido-reduction conditions, described with the H₂ and H₂O partial pressures. The predominance diagrams are built using the FACTSAGE thermochemical software version 7.2 with the SGPS-SGTE thermodynamic database⁶. The figures clearly illustrate the high sensitivity to redox conditions of the gaseous species of Mo and Ba in the composition range of irradiated fuels in severe accident conditions.

The main species involved during the thermal sequences for each test are reported in Figs 6 and 7. The predicted speciation of Cs, Mo and Ba as well as the partitioning of Mo in the gas phase as a function of redox conditions are consistent with the corresponding experimental release curves.

The most striking feature is the predominance of gaseous polymolybdate species of Cs and Ba in oxidising conditions, which contributes to increase significantly the amount of Mo in gaseous form, as compared to reducing conditions. The main species involved are the caesium dimolybdate $Cs_2Mo_2O_7(g)$ and the barium molybdate $BaMoO_4(g)$, which are stabilised from mid-oxidising to highly oxidising conditions.

Cs-Mo-H-O, 1500 ℃ 0.333 < Mo/(Cs+Mo) < 1Vercors-RT3 0 √lo(s) $MoO_2(s)$ Cs(g)+Mo(s) log₁₀[P(H₂)] (bar) -1 Vercors-RT6 Verdon-1 Cs₂MoO₄(I) -2 Stability area of $Cs_{9}Mo_{v}O_{3v+1}($ -3 -5 -4 -3 -1 0 -2 $log_{10}[P(H_2O)]$ (bar)

Fig 6. Predominance diagram of the system Cs-Mo-O-H at 1500°C in the composition range of an irradiated fu el The red discs refer to the thermodynamic conditions of the thermal tests VERCORS-RT3, -RT6 and VERDON-1 at the oxidation step level.

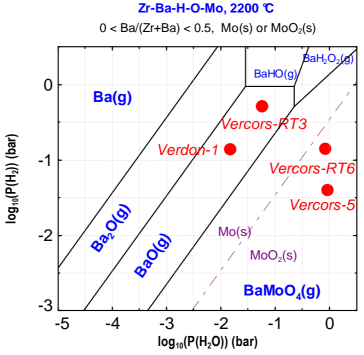


Fig 7. Predominance diagram of the gaseous species of the system Zr-Ba-H-O-Mo at 2200℃ in the composition range of an irradiated fuel

The red discs refer to the thermodynamic conditions of the thermal tests VERCORS-4, -5, -RT3, -RT6 and VERDON-1 tests, at end of the high temperature step.

5. Release kinetics

To illustrate the consistency of the proposed speciation, it is necessary first to describe the rare gas release kinetics (Xe, Kr). The simplest model that can be adopted for post-irradiation annealing experiments at high temperatures is the equivalent sphere model. It relates rare gas release to solid-state diffusion of gas atoms in an equivalent spherical fuel grain. One of the limitation of this model is related to the lack of description of trapping sites that form in irradiated fuels, such as bubbles, pores, dislocations, grain boundaries... One of the main feature of the VERCORS tests series is the dependency of fission gas release on annealing temperature (i.e. 10 to 20% at a temperature plateau of 1300-1500°C in the VERCORS-2, -4 and -5 tests, ~ 45% at a temperature plateau of 1800°C in VERCORS-1 and-2 tests, and 80 to 90% at a temperature plateau of 2200-2300°C in VERCORS-3,-4 and-5 tests). Progressive liberation of trapped gas with temperature can explain these observations and should be taken into account in the equivalent sphere model. To this end, it was assumed that the gas available for diffusion increased with temperature according to a linear regression based on the VERCORS tests results. The equivalent sphere model with a temperature-dependent source term was thus solved numerically by a finite volume scheme.

Application to Xe release and Xe release kinetics during the VERCORS-5 test is shown in Fig 8. The diffusion coefficient is obtained from the analytical solution of the Booth model considering a FGR of 10% in 20 minutes at 1300° C, a fuel grain radius of 5 µm and the activation energy proposed by Booth. The fit compares well with measurements except at the beginning of the 1300° C plateau where a burst release related to grain boundary gas is observed¹

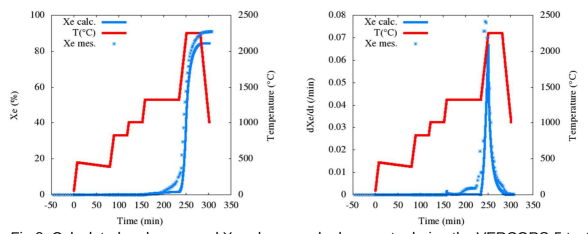


Fig 8. Calculated and measured Xe release and release rate during the VERCORS-5 test

Calculating at each time step the release rate of the bulk of fission gases (Xe, Kr), the following approach has been considered for the other (minor) gas species:

- during the 1300°C temperature plateau, all Cs, Mo and I are gaseous and released as CsI(g) and $Cs_2Mo_2O_7(g)$, see equations 1-3;
- at higher temperature and during the 2250°C final plateau, all remaining Cs, Mo and I are again gaseous and released as CsI(g), $Cs_2Mo_2O_7(g)$, and $BaMoO_4(g)$ The quantity of Ba in the gas phase is limited by the availability of Mo and thus given by equations 11-14.

At each time step, the calculated instantaneous released fraction of Xe is applied to the remaining quantities of Cs, Mo, I and Ba associated in the proposed gas species. In consistency with a previous work developed for Pellet-Clad-Interaction (PCI) analysis⁵, this approach accounts therefore for thermo-chemistry in the release rate of chemically reactive elements (Cs, I, Mo and Ba). Note that the model is based on a single diffusion equation describing the release of rare gases (Xe). The calculated releases of Cs, I, Mo and Ba during the VERCORS-5 test agree well with on-line measurements, as shown in Fig 9.

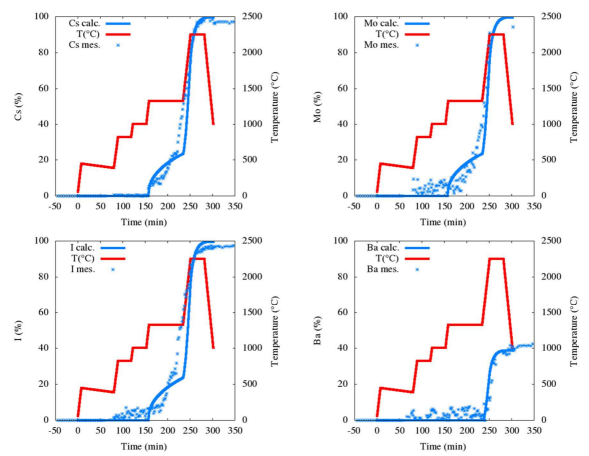


Fig 9. Calculated and measured Cs, Mo, I and Ba releases during the VERCORS-5 test

6. Conclusion

Speciation and release of fission products during some of the VERCORS and VERDON tests have been discussed. Mass balance calculations carried out from the experimental FP released fractions curves have confirmed, in agreement with recently published findings, that gaseous polymolybdate species of Cs and Ba are predominantly formed, irrespective of burnup, from mid-oxidising to highly oxidising conditions, in the range 1300℃-2500℃. Release kinetics of those volatile (minor) FP can be highly dependent on surrounding oxydo-reduction conditions. Good agreement was achieved between measured and calculated releases of Cs, I, Mo and Ba during the VERCORS-5 test using an analytical model including the proposed FP thermochemistry and describing diffusion of rare gases only, which illustrates the global consistency of the proposed approach.

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