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# ANALYSES OF RADIOLYTIC GASES RESULTING FROM GAMMA IRRADIATION OF ASSE ROCKSALT PERFORMED AT SACLAY.

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#### **ABSTRACT**

The radiolytic generation of gases is an issue which has been considered when studying the safety of radioactive waste repositories in rocksalt. During these 5 years of research over 350 samples of ground salt have been irradiated and analyzed in order to elucidate which are the mechanisms in the radiolytic generation of gases in salt.

We have studied the effect of dose and dose rate at different temperature in the amount and nature of generated gases. The irradiated samples were of different composition and granulometry, in this way different backfill characteristics could be simulated. the samples were encapsulated in glass vials prior to irradiation. The gases contained in the vials were analyzed after irradiation. Samples from the same batch were encapsulated in vials inside which different atmospheres were created. In this way the relationship between original oxidizing, reducing, or inert atmosphere, and the obtained gases could be studied.

The way in which the experiments were performed and the obtained results are here described.

#### 1 INTRODUCTION

ANDRA, the French National Radioactive Waste Management Agency, joined the HAW project in 1988 in the framework of European Community agreements FI1W 0199 and F12W 0002. Andra's involvement relates to calculations, laboratory testing and *in situ* measurements. The objective of laboratory work summarized in this report was parametric study, in a laboratory,

of gas generation by radiolysis of Asse salt. From 1988 to 1993, the test program involved over 350 samples. The first phase of the test protocol developed included preparation of samples and their irradiation at the Saclay Research Center, primarily with spent fuel from the Osiris test reactor. The second phase consisted of analyzing the generated gases by various methods suited to the context of the study: chromatography, mass spectrometry, infrared spectrometry. Test results presented and discussed in this report reveal the preponderance of certain irradiation parameters in gas generation, such as integrated dose, the type of initial gas, salt granulometry and the content of impurities in the salt.

#### 1.1. Purpose of Study

The purpose of the study is qualitative and quantitative assessment of gaseous releases in salt caused by radiation, primarily gamma radiation, emitted by waste canisters. Salt is considered as both the host rock and the principal component of the engineered barriers. The radiation emitted by the waste can induce radiolytic reactions, especially of water or fluids coming from the geologic formation (gases or liquids). These reactions, combined with a temperature rise in the near field, may produce toxic, corrosive or even explosive substances.

Gamma irradiation of salt samples therefore had to be performed under a variety of conditions by varying the essential parameters of the phenomenon studied in realistic areas in terms of disposal and test duration: dose rate, temperature and duration of irradiation, salt granulometry and composition of the atmosphere under which the salt is irradiated. The combination of issues raised also prompted us to look into the material's behavior and damage to the crystalline lattice caused by  $\gamma$  radiation. An understanding of the chemical and mineralogical composition of salt is indispensable in understanding and observing the formation of gaseous compounds.

#### 1.2. Partners

ANDRA cooperated with different CEA teams at the Saclay and Fontenay aux Roses sites in performing these tasks:

- DRN/DRE/SRO for salt irradiation;
- DCC/DSD/SCS to analyze radiolytic gases and process results; and
- DCC/DPE/SPEA for mass spectrometry analyses.

While salt domes are plentiful in Germany, they are rarer in France, where saliferous formations are usually layered. CEA's DCC/DSD/SCS group at Fontenay aux Roses has also carried out work focused specifically on the phenomenology of radiolysis of a French salt from the Mines de Potasse d'Alsace [Akram, 1993].

#### 1.3. Objectives and Planning

The general purpose can be split into some specific experimental objectives which were worked out into a test plan established together with the GSF. Table 1 gives an overview of the parameters which relationships were studied in this program. Carrying out the program meant sample collection and packaging, irradiation of the samples and post irradiation analysis. The irradiation experiments and the methodology of irradiation are described by Marchand (1995, artc. nr. this volume), and will not be repeated here.

The original program as stablished in Table 1 was supplemented with additional studies to better identify the various reactive mechanisms involved, summarized in Table 2.

Previous to carrying out the test plan, during the first half of 1988, two series of irradiations were performed to select sample packaging, sampling procedures and analytical methods suited to the different radiolytic gases. These tests were performed on a 2 kg batch of crushed salt (batch 1) supplied in April 1988 by GSF.

Table 1 : Asse salt radiolysis : initial programme of irradiations

Parameter	Number of samples (except blanks)	Dose rate (Gy/h)	Total dose (Gy)	Duration (hours)	Temperature (°C)
Salt cores	9	1. E+4	1. E+6	100	ambient
Crushed salt	15	1. E+4	1. E+6	100	ambient
Grainsize	6 6 6 6	1. E+4 1. E+4 1. E+4 1. E+4 1. E+4	1. E+3 1. E+4 1. E+5 1. E+6 1. E+7	0.1 1 10 100 1000	ambient
Dose rate	3 3 3 3	1. E+2 1. E+3 1. E+4 1. E+5	1. E+6 1. E+6 1. E+6 1. E+6	10000 1000 100 10	ambient
Total dose	3 3 3 3 3 3	1. E+3 1. E+3 1. E+3 1. E+3 1. E+3 1. E+3	1. E+2 1. E+3 1. E+4 1. E+5 1. E+6 1. E+7	0.1 1 10 100 1000 10000	ambient
	3 3 3 3 3	1. E+4 1. E+4 1. E+4 1. E+4 1. E+4	1. E+3 1. E+4 1. E+5 1. E+6 1. E+7	0.1 1 10 100 1000	
Temperature	3 3 3 3 3	1. E+4 1. E+4 1. E+4 1. E+4 1. E+4	1. E+6 1. E+6 1. E+6 1. E+6 1. E+6	100* 100* 100* 100* 100*	50 100 150 200 250
Pure minerals	21	1. E+4	1. E+6	100	ambient
Containers	15	1. E+3	1. E+6	1000	200
Filling atmos. (N <sub>2</sub> , O <sub>2</sub> , Ar, He)	12	1. E+4	1. E+6	100	ambient

Table 2 : Asse salt radiolysis : complementary programme of irradiations

Parameter	Number of samples (except blanks)	Dose rate (Gy/h)	Total dose (Gy)	Duration (hours)	Temperature (°C)
Dose rate and total dose	3 3	5. E+4 4. E+4	1. E+6 4.9 E+6	20 120	120 80
Crushed salt (with 60 Co)	3 3 3	1. E+4 4. E+4 3.3 E+4	1. E+6 2.9 E+6 1.3 E+8	100	ambient 80 65
Massive samples	3	1. E+4	1. E+6	100	ambient
Dose rate, dose and temperature	3 3 3 3 3 3 3	1. E+5 1. E+4 1. E+4 1. E+3 1. E+5 1. E+4 1. E+4 1. E+4	1. E+6 1. E+4 1. E+5 1. E+6 1. E+6 1. E+4 1. E+5 1. E+6	10 1 10 1000 10 1 1 10 1000	150 150 150 150 200 200 200 200
Filling atmos.	3 3 3 3 3 3	1. E+5 1. E+4 1. E+4 1. E+4 1. E+3 1. E+5	1. E+6 1. E+4 1. E+5 1. E+6 1. E+6 1. E+6	10 1 10 100 1000 10	
with increasing O <sub>2</sub> content	3 3 3 3 3	1. E+4 1. E+4 1. E+4 1. E+3 1. E+5	1. E+4 1. E+5 1. E+6 1. E+6 1. E+6	1 10 100 1000 10	ambient

# 1.4. General Methodology

On the occasion of this research program, we developed an experimental approach which enabled the influence of radiation on gas generation in rock salt to be assessed. This methodology has already been applied to salt samples coming from other sites than the Asse mine, e.g. salts including variable quantities of marl, anhydrite, sylvinite and organic matter. Based on a consistent set of test results, we proposed interpretations in terms of mechanisms,

and initiated predictive modelling of the evolution of the system in a disposal context. This methodology comprises the following points:

- design of the test plan to optimize the parametric area to be explored;
- development of a special irradiator using irradiated or spent fuel supplying a large range of dose rates and integrated doses and providing constant dose rates over several months; optimization of dosimetry on a continuous basis;
- development of procedures for preparation and packaging of samples for irradiation and analysis;
- use of different characterization methods for solid samples, before and after irradiation, some of them being completely new (nuclear microprobe);
- development of various special accurate analytical methods for identification of the gases, particularly chlorides (GPC after chemical reaction) and sulfured products (GPC coupled with mass spectrometry), adaptation of the FTIR to the determination of complex gas mixtures, measurement of certain acidic gases by ion chromatography after trapping, in addition to the conventional methods (GPC and direct mass spectrometry); and
- development of desorption procedures followed by on-line analyses.

#### 2. PRE-IRRADIATION METHODOLOGY.

Salt characterization, especially identification of organic compounds was mainly carried out by microscopic examination. The approach consisted of investigating the medium's characteristics at the moment of geologic formation of the rock, when minerals precipitated. Microscopic inclusions trapped in intracrystalline cavities are representative of original brines. Different techniques, described hereunder, had to be used for qualitative analysis of the medium's different compounds.

## 2.1. Methodology of the Analysis of Fluids.

*Microscopic examination* is based on criteria relating to the location, size, form and content of inclusions at ambient temperature. Two types of inclusions may tell the history of the rock: inclusions trapped during the salt deposition, or primary inclusions, and inclusions trapped later, or secondary inclusions.

Microthermometric examination consists of measuring phase change temperatures in an inclusion under a microscope at varying temperature. This gives information on density, overall salinity and the presence of dissolved gases such as CO<sub>2</sub>, CH<sub>4</sub> and of other hydrocarbons.

Microthermometry can be combined with other analytical techniques, such as Raman mol microprobe with laser excitation, Fourier Transform InfraRed microspectroscopy (FITR), Electron Spectroscopy (ESCA) and nuclear microprobe.

This latter technique developed in the framework of this program is described below.

# 2.1.1. The Nuclear Microprobe ( A New application of the Instrument)

The characterization of the light elements (carbon and oxygen) of the salt was carried out with the nuclear microprobe at CEA's Bruyères le Châtel site. The test device for focusing and analysis is coupled with a 4 MeV Van De Graaff single stage accelerator. Using various spectroscopic modes (elastic diffusion spectrometry, nuclear reactivity analysis, X and  $\gamma$  ray emissions induced by charged particles) information can be gained on three-dimensional elemental and sometimes isotopic distributions of most of the elements in the periodic table. The nuclear microprobe make it possible to do non-destructive elemental characterization of salt samples and to analyze the light element composition of the fluid inclusions.

#### Experimental Procedure

Nuclear microprobe analysis of salt samples was done by detecting protons and gamma photons emitted during nuclear reactions induced by deuterons. The resulting reactions induced by the deuterons were used to analyze, respectively:

- carbon:

 $^{12}C(d,p)^{13}C;$ 

oxygen:

 $^{16}O(d,p)^{17}O$  and  $^{16}O(d,p\gamma)^{17}O$ , E $\gamma = 871$  keV;

- sodium:

 $^{23}$ Na(d,p $\gamma$ ) $^{24}$ Na, E $\gamma$  = 472 and 1369 keV.

Salt behavior during the irradiations was also observed by analyzing emitted X photons.

The carbon and oxygen content was determined by reference to graphite and silica standards. The salt's contribution to the charged particle spectra was deduced by using  $(d,\alpha)$  reactions induced in <sup>23</sup>Na and <sup>35</sup>Cl. Gamma photon spectra were interpreted taking into account the interferences due to  $(n, n'\gamma)$  and  $(n,\alpha\gamma)$  reactions induced in the germanium of the gamma detector (energies  $E\gamma = 868$  and 875 keV respectively).

#### Experimental configuration

The following experimental configuration was used:

- deuteron energy: 1.4 MeV;
- microbeam diameter of 25 to 50  $\mu$ m adjusted to the size of the inclusions;
- current density of 8 to 26 pA/ $\mu$ m<sup>2</sup> inclusive; and
- charged particles and  $\gamma$  and X photons detected by surface barrier detectors (detection angle in relation to incident beam = 120°), Ge HP (angle = 135°) and SiLi (angle = 112°) respectively.

Approximately 2 mm thick sections of salt were polished with alumina powder lubricated with ethylene glycol. A 1/1 solution of dichloromethane/ethanol was used to clean the surfaces. The inclusions analyzed are at a depth of 0 to 15  $\mu$ m beneath the surface. The polished surfaces were plated with an approximately 30 nm thick gold film to drain off charges and reduce sample damage linked to the temperature increase and ionic erosion at the point of impact.

Verification of salt and inclusion behavior when exposed to an ion beam

The development of F centers and sodium colloids when exposed to an ion beam depends both on the total dose and on the dose rate. For ions of a given type and energy, the density of the current determines the heat applied to the sample during irradiation. Because alkaline halides are particularly sensitive to erosion by electronic sputtering, the use of a high density current that increases the yield of primary defects will also contribute to the diffusion of interstitial halogens towards the surface and vaporization of metal sodium in the surface. The use of a sufficiently low density of current therefore helps to limit the development and diffusion of defects.

The metal-plating film cools the outermost surface and prevents its erosion. In addition this film stabilizes fluid inclusions that have a tendency to migrate along the thermal gradient and, in our case, in the direction of the surface level.

An analysis of signals detected from charged particles and  $\gamma$  and X photons shows 1) no measurable variation in salt composition for current densities of 12 to 15 pA/ $\mu$ m<sup>2</sup> inclusive, and 2) fluid inclusions could be analyzed with a current density of as much as 26 pA/ $\mu$ m<sup>2</sup>.

## 2.2. Methods of Organic Matter Characterization .

The important role played by organic matter during radiolytic reactions prompted us to characterize organic compounds present in the material [GAUDIN and GAUDEZ, 1993].

Some organic compounds present in the material were identified and estimated by chromatographic analysis coupled with mass spectrometric detection. The organic matter was extracted from the salt with dichloromethane.

Silylation of a fraction of the sample by the BSA helped to identify non-soluble products, such as alcohols and acids. These compounds, which are principally linear or branched hydrocarbons and organic acids, are present in very low concentrations (<10<sup>-7</sup> g/g) in the Asse salt. Examination of the unidentified insoluble fraction in the dichloromethane suggests the

presence of heavier hydrocarbons. The behavior of these products under  $\gamma$  irradiation may lead to the formation of toxic compounds by reaction with the principal constituent (NaCl), giving rise to the formation of chlorinated compounds. The research begun as part of this program together with conclusions pertaining to unhomogeneous saline materials (MDPA) revealed the importance of organic matter during irradiation and will require appropriate analytical development during later work.

## 2.3. Starting Material (Sample Supply)

The test plan required different types of sample composition and granulometries which were supplied by the GSF as the project progressed and are listed below. The methodology used to produce the samples of natural rocksalt of homogenous composition is extensively described in Gies (1995), art. Nr.. This volume. The samples were prepared from ground Asse rocksalt samples, or constituted by container wall material pieces. The following samples were used:

- 2 kg of crushed salt for preliminary tests (batch 1);
- 50 kg of crushed salt (batch 1988) for the irradiation program;
- five samples of granulometries ranging from 0.125 to 8.0 mm;
- 2 batches of granulometries in the ranges of  $(0.1 < \phi < 0.25 \text{ mm})$  and  $(1.0 < \phi < 2.0 \text{ mm})$  to study the influence of grain size as a function of integrated dose;
- core drilled samples measuring 20, 30 and 50 mm in diameter and 50 mm high;
- the principal mineralogical compounds of the salt in the form of pure products:
  - \* halite (NaCl),
  - \* sylvite (KCl),
  - \* anhydrite (CaSO<sub>4</sub>),
  - \* gypsum (CaSO<sub>4</sub>, 2H<sub>2</sub>O),
  - \* kieserite (MgSO $_4$ , H $_2$ O), and
  - \* polyhalite ( $K_2Ca_2Mg(SO_4)_4$ ,  $2H_2O$ );
  - \*carnallite (MgCl<sub>2</sub>KCl, 6H<sub>2</sub>O) could not be procured in sufficient quantity; and

samples of steel to observe the effect of the simultaneous action of radiation and salt on steel corrosion. These experiments are useful in relation to the construction of steel containers or overpacks for radioactive waste repositories in salt formations.

#### 2.4. Method of Sample Packaging

Before being irradiated, the salt samples were packaged in a special vessel able to retain the gases produced for later analysis. This vial was designed to be gastight up to few bars and resistant to temperature, gamma rays and corrosion by salt or brine. Attenuation of gamma-rays should also be as low as possible.

The design of the irradiation vial was determined after preliminary tests involving crushed salt irradiated under an atmosphere of synthetic air.

The vial is designed to enclose 200 g of crushed salt and fit a certain geometry, with free volume equal to the apparent volume of the salt. A ratio of A/v = 1/3 was maintained throughout the program, except for the 50 mm diameter core drillings, where A/v is equal to 1/4, because otherwise the sample would heat up during vial sealing. A is the free volume left by the salt and v is the real volume of the salt, calculated from its mass and density.

The vial selected is manufactured from a tube of Pyrex glass measuring 44 mm in diameter and 330 mm long (Fig. 1). Its sealing concept prevents the introduction of water vapor or combustion gas during sealingwhich is done by welding the glass.

At one end he vial has a 12 mm and 6 mm diameter neck pipes, one for salt filling and the other for pumping and re-filling the selected atmosphere.

The other end of the vial is closed with a "pig tail" inside a 12 mm diameter neck pipe; a 19/9 female connector is welded to this end after irradiation. A magnetic Teflon-coated bar ( $\phi$  = 7 mm, L = 60 mm) is introduced for later break off of the pig tail. Then a glass/metal valve with grease-free connectors is added, so that the vial can be adjusted to fit different analytical apparatus and, samples of a given quantity of gas can be taken without any pollution.

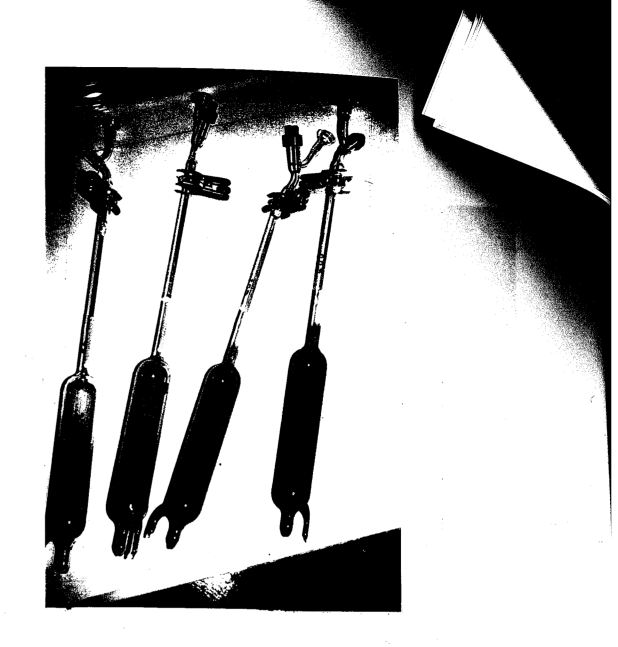
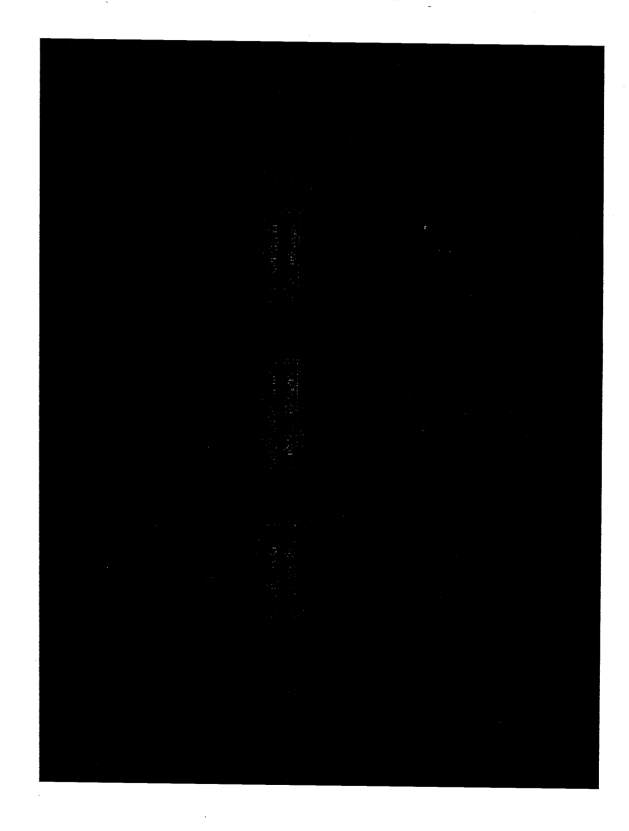


Figure 1: Vials after irradiation, fitted for gas analysis

Each vial has and engraved number and is heated to 500°C before use.

Most irradiations were performed on 200 g samples of crushed salt. Three irradiations were performed simultaneously on the same sample as well as on a blank of selected atmosphere to determine the order of magnitude of the deviation in the measurements. However, after a program delay incurred when the irradiator was off-line (November 1989 to July 1990), irradiations were first performed in groups of eight on two levels in the irradiator, then vial diameter was reduced to 40 mm while maintaining the A/v ratio; the irradiated salt mass was 150 g. The good test reproducibility and reliability of test methods also allowed the reduction in gas volume to be analyzed.

Fig 2 : Coloration of salt samples irradiated at different doses



The samples must be prepared carefully to control initial conditions and ensure good test reproducibility. The set of four vials is filled with salt, then the air in the vials is expelled for one minute and replaced with an atmosphere of known composition at less than atmospheric pressure. This atmosphere could be N50 reconstituted air, inert gas, oxygen or media in a 50/50 or 20/80 N<sub>2</sub>/O<sub>2</sub> ratio supplied by Air Liquide/Alphagaz. This operation is repeated three times before the vials are sealed. The final packaging pressure is 930 mb.

After irradiation, the vial receive the valves and connectors described above. A vacuum pumping of these parts removes any traces of humidity or pollution before releasing the gases by breaking the pig tail with the magnetic bar.

#### 3. POST-IRRADIATION PROCEDURES.

This section describes the procedures selected to analyze the different products of irradiation namely: to study of radiation damage in the halite crystals of the salt rock, and to study the gases released by radiolysis of the salt rock.

Gamma rays induce radiolytic reactions in constituents of a saline medium, particularly water, fluid inclusions and the salt itself. When  $\gamma$  rays enter a solid, Compton electrons can be produced. As these electrons move, they produce ionization along their trajectory. In alkali halides, such as sodium chloride, excited Cl<sup>-</sup> ions can be ejected from their normal position in the lattice leaving behind them an electron. The void left behind by the chlorine with the trapped electron constitute the F centers which agglomeration produces Na colloids regions in the crystal. The chlorine itself occupies an interstitial position giving rise to H-center development. The F centers are responsible for the brownish yellow coloration observed at low doses. The colloidal particles of metallic sodium are responsible for the blue coloration observed in the samples of salt irradiated at high doses (Fig. 2).

Different techniques, described below, to analyze gases were developed gradually as the experimental program advanced.

Basically, these techniques are, on one hand, mass spectrometry to analyze hydrogen and constituents of the atmosphere ( $N_2$ ,  $O_2$ , possible traces of argon) and of NO, principally in the blanks of synthetic air, and, on the other hand, gas chromatography for systematically analyzed compounds such as  $CO_2$ ,  $N_2O$ , CO and  $CH_4$ .

Based on results of the foregoing, different supplemental analyses to detect SO<sub>2</sub>, H<sub>2</sub>S, HCl, etc., that may be present after irradiations at a high total dose.

#### 3.1. Method of Colloidal Sodium Content Determination.

Analysis of the amount of sodium colloid contained in the samples irradiated in this program took place by dissolving them in water and measuring the produced hydrogen. This indirect determination method is based on the assumption that every two Na atoms from the colloids will decompose two water molecules and produce an hydrogen molecule and sodium hydroxide.

Irradiated salt is dissolved in deionized water to quantify the formation of defects from irradiation. The following reaction occurs :

$$2 \text{ Na}^+\text{e}^- + 2 \text{ H}_2\text{O} \ge \text{H}_2 + 2 \text{ Na}^+ + 2 \text{ OH}^-$$

The hydrogen content measured by mass spectrometry enables observation of sodium colloid formation. Four grams of irradiated salt are poured in a flask which is then vacuum emptied. Then 20 ml of deionized water are introduced through a septum. Bubbles observed when water is added correspond to the formation of hydrogen. Dissolution lasts approximately one hour, then the flask is cooled with liquid nitrogen to trap the water before analysis by mass spectrometry.

# 3.2. Direct Mass Spectrometry Methodology

The spectrometer used to analyze light gases (atomic weight< 200) is an apparatus with single focusing magnetic sectors manufactured by the CEA. Separation is carried out by a magnetic prism producing an homogeneous field perpendicular to the ion beam. The resolution of this type of apparatus under normal conditions is 250. The pipe of introduction is made of stainless steel and completely tight (vacuum < 10<sup>-5</sup> torrs). The mixture to be studied is released into a flask calibrated to a working pressure of approx. 1 torr, measured with a membrane micromanometer. The interface between the flask and the spectrometer source is a molecular leak of 30 lusecs (millitorr-liter per second). The fragmentation of each compound and the sensitivity of each gas must be known before proceeding to an examination of the spectrum (signal/pressure ratio of the source). Standard gaseous mixtures are used for calibration.

The results of mixture compositions are estimated in relation to nitrogen content, which is assumed to be constant. Gamma irradiations at total doses higher than  $10^7$  Gy have revealed variation in the  $N_2/O_2$  ratios and the formation of nitrogen oxides. A future experiment might introduce a noble gas into the composition of the irradiation atmosphere, such as neon, which is very easily detectable by mass spectrometry and unaffected by radiolysis. Deeper studies of the absorption of nitrogenated compounds by salt could be perform by marking the air with  $^{15}N$ . The hydrogen detection limit of 3 ppm in volume made it possible to observe this gas even after irradiations at very low integrated doses ( $10^3$  Gy). NO content values did not give rise to any special calibration.

#### 3.3. Gaseous Phase Chromatography (GPC) Methodology

Two types of chromatographic separation analysis with an identical "on-column" injection system were used. Because the quantities of the different compounds to be detected could be very low, we established a fixed gas injection volume of 2 ml (injection loop) when performing the first tests. The adaptation of a six-way valve, when used in different positions, allows the column to be flushed, the sampling system and sampling loop to be degassed, and the sampled fraction to be carried away for chromatographic separation with the carrier gas. Under

these conditions, the injected volume remains constant, both for the calibrating standard and for the sample. Sample or standard pressure at injection is measured with a manometer.

The mixtures of calibrating standards in nitrogen make it possible to obtain the respective calibration curves of the analyzed compounds by dilution in a known volume of nitrogen using volumetric flasks. In these calibration curves the peak area obtained at the characteristic retention time of the analyzed compound is plotted as a function of its concentration. The partial pressures are taken into account when interpreting the results.

The methods and conditions used to analyze the different compounds were:

- Chromatography with catharometric detection for analysis of CO<sub>2</sub> and N<sub>2</sub>O: Separation is done on a steel column measuring 1/8" in diameter and 3 m long packed with a material made of 80-100 mesh porous polymer beads (Porapack Q, Chrompack); helium is the carrier gas (16 ml/min) and a constant temperature of 30°C is maintained during analysis. Catharometric detection is carried out with a 600 ml cell containing four WX (tungsten-rhenium) filaments fed with a 200 mA current and heated to 50°C. This equipment enables also measurement of CO (> 40 ppm in volume).
- Flame ionization detection after methanation: Because chromatographic separation on a molecular sieve followed by catharometric detection to analyze traces of CO is not sensitive enough, flame ionization detection is used. Hydrogen (15 ml/min) is the carrier gas for separation. The 2 m long, 1/8" diameter steel column is packed with a 60-80 mesh molecular screen and heated to 30°C. Hydrogen reduction of CO into CH<sub>4</sub> occurs in a platinum sponge furnace heated to 350°C.
- Chromatography with flame ionization detection for CH<sub>4</sub> and light hydrocarbons with a molecular weight of less than 60: Separation take place in a 3 m long, 1/8" diameter steel column packed with 50-80 mesh porous polymer (Porapack T, Chrompack), whose polarity is greater than that of Porapack Q, with helium as

the carrier gas. The flame ionization detector (q = 250°C) uses a hydrogen flame burning in air. Separation of the other hydrocarbons is achieved by increasing the temperature of the sample from 30 to 120°C with a rate of 10°C/min.

Flame photometry detection (FPD) of SO<sub>2</sub> and H<sub>2</sub>S was planned after chromatographic separation at a temperature of 80°C in a 1/8" glass column packed with 80-100 mesh Tenax (preheated during 2 hours at 100°C), with the detector temperature raised to 140°C. Feasibility tests served to define detection limits of 5 ppm for H<sub>2</sub>S and 0.2 ppm for SO<sub>2</sub> in the case of a calibrating mixture in nitrogen. The advantages of flame photometry are the simultaneous detection of sulfured compounds and the time saved over the combined chromatography/mass spectrometry technique. Detection can be improved by injecting a larger volume (2 ml) with a valve that is inert to sulfured products.

Gaseous chlorine detection by GPC with flame ionization detection: For simultaneous analysis of chlorine and hypochlorite contained in the gaseous phase, the property of phenols to react with these species by an electrophilic substitution mechanism is used. All or part of the available mixture in the vial is complexed with a 2.6 dimethylphenol (DMP) solution in a 0.05 M sulfuric medium. The chloro-4-DMP complex is extracted in an n-hexane medium and can be concentrated to improve sensitivity (this was done for low doses of radiation). The presence of an internal standard, such as chloro-4 dimethylphenol-3.5, makes it possible to observe the progress of the operations. After injection of the sample at a temperature of 230°C, chromatographic separation is done in helium. The column used is a CP-sil-5-CB semi-capillary column (Chrompack) of apolar melted silica measuring 25 m long with a 0.53 mm inner diameter and a phase thickness of 5 μm. Detection is done by flame ionization with a furnace temperature programmed from 80 to 200°C. Calibration curves based on a calibrating mixture of 1000 ppm of Cl<sub>2</sub> in nitrogen were plotted.

# 3.4. Gaseous Phase Chromatography Coupled with Mass Spectrometry Methodology

The coupling of a fragmentation technique such as chromatography with an analytical technique such as mass spectrometry makes it possible to achieve good detection sensitivities for compounds in trace amounts. Analysis may be qualitative or quantitative. In the first case, the spectrometer is used to scan the magnetic field to achieve a mass spectrum or currentogram; in the second case, it is used for fragmentography. In the second case, the apparatus is set, in succession, for each ion characteristic of the compound to be analyzed or that has the least amount of interference with another compound by varying the source's acceleration voltage with the computer control system.

This technique was used to identify SO<sub>2</sub> and H<sub>2</sub>S after irradiation. In this case, the operating conditions were as follows:

- glass chromatographic column (1/4" dia. and 2 m long) filled with 80-100 mesh Tenax;
- two constant temperature chromatographic separations: 25°C for H<sub>2</sub>S and 50°C for SO<sub>2</sub>;
- a VG 7035 magnetic spectrometer with a resolution of 2000;
- quantitative analysis based on fragmentation by observing characteristic ions:

ions 64 and 66 for  $SO_2$   $^{32}S^{16}O^{16}O$  and  $^{34}S^{16}O^{16}O$  ions 34 and 33 for  $H_2S$   $^{1}H^{1}H^{32}S$  and  $^{1}H^{33}S^*$  for ion 34,  $^{1}H^{32}S$  and  $^{33}S^*$  for ion 33

(\* being very low)

The product is identified in this manner by three criteria: its chromatographic retention time, the two characteristic ions and the concentration ratio of these two ions (for example, for  $SO_2$ , the 66/64 = 4% ratio corresponds to the natural isotopic ratio of  $^{34}S/^{32}S$ ).

The quantitative range of analysis is 1 to 50 ppm. Calibrations are performed using standard mixtures in nitrogen supplied by Air Liquid/Alphagaz (500 ppm content), by dilution in vials with unlubricated spigots and with a LB1 "thermogreen" septum (Supelco). These dilutions are not stable for more than one week.

Chromatographic separation for analysis of  $SO_2$  made identification of  $Cl_2$  possible. The chlorine present in the currentogram upstream from the  $SO_2$  disturbs measurement of the latter, mainly at high doses. Direct quantitative analysis of gaseous chlorine is not feasible with this technique, since this element is rapidly trapped on the metal sections of the apparatus.

# 3.5. Ion Chromatography Methodology

From the irradiated vial, two samples of gas are simultaneously taken into two 20 ml calibrated flasks. Then, all of the chloride ions as well as the other anionic species are trapped in 5 ml of an NaOH 10<sup>4</sup>M solution. To obtain all of the chloride ions, the transformation of the ClO ions into Cl is accelerated by the addition of 100 ml of  $H_2O_2$ . Analysis is done after separation in an HR anionic column (Waters) by conductimetric detection and determination of the areas of characteristic peaks. Calibration based on sampling of gaseous compounds in the nitrogen is done under identical conditions. By proceeding in this manner, the "total chlorides" concentration of the gaseous mixture analyzed is derived; the concentration of hydrochloric gas is the difference:

[total chloride] - [chloride + hypochlorite]

# 3.6. Fourier Transform InfraRed Spectrometry Methodology

Analysis by infrared spectrometry of the asymmetric gaseous compounds is based on the excitation of different types of molecular movements: rotation and vibration. The energies corresponding to these movements lead to absorptions of electromagnetic radiation characteristic of the molecules observed. The rotational absorption spectra are located in the outer infrared band (0.1 to 200 cm<sup>-1</sup>). The vibrational absorption spectra, often associated with vibration-rotation movements, in the case of gases, is located in the spectral range of medium infrared (200 to 4000 cm<sup>-1</sup>), This was therefore the analytical range chosen to perform analysis of the gaseous compounds developed or liberated during our irradiation experiments (CO<sub>2</sub>, N<sub>2</sub>O, CO, NO, NO<sub>2</sub>, HCl, CH<sub>4</sub>, hydrocarbons, H<sub>2</sub>O).

The improvements introduced by us on the existing standard method consisted of adaptation of the sampling and measuring cell, and modification of data processing in order to allow quantitative analysis to take place. It was necessary to constitute a library of standards of each compound studied as a function of temperature.

The apparatus used is an IFS 48 from Bruker equipped with a mercury, cadmium, tellurium (MCT) detector cooled with liquid nitrogen for use in the medium infrared (600-4000 cm<sup>-1</sup>) range. IR radiation is emitted by a source of silicon carbide cooled by water. In order to study the detailed structure of the gas molecules, we work with a resolution of 0.5 cm<sup>-1</sup>.

The IR beam crosses the measuring cell, where the gaseous sample is enclosed, before being detected. A special cell for the analysis of corrosive gases (Fig. 3) has been developed in the framework of this program in cooperation with the Bruker/Spectrospin company. This cell is small (100 ml), but the length of the optical path is increased by using mirrors. This path is adjustable from 25 cm to 1 meter by acting upon the concavity of one of the reflex mirrors with a micrometric screw. The number of reflections thus ranges from 4 to 16. The choice of materials is important; Pyrex was chosen for the body of the cell, and the windows are in BaF<sub>2</sub>. This material offers the advantage of being less soluble than KBr and more resistant to light and temperature than standard materials (such as AgBr). The cut-off of the spectrum is around 800 cm<sup>-1</sup>. The mirrors of the cell are gold-plated and their supports are of Teflon. The cell can be heated to 200°C.

The analysis is made more complex by the presence of molecules whose spectrum characteristic rays can superpose. Because the range of concentrations of certain major compounds such as CO<sub>2</sub> and N<sub>2</sub>O can vary from a few ppm to several % in volume, the choice of specific rays that are unsaturated and without overlap is restricted for the analysis of the other constituents. However, quantitative analysis was possible by combining a certain number of standards which spectrum rays are on either side of the sample spectrum, with a sufficient number of characteristic rays, up to ten.

The processing software (GAZ) [GAUDEZ & al 1993] developed over the course of this study is based on the principle of multipeak analysis. It permits determination of the

content of each of the compounds in the mixture, and takes into account the possible non-linearity of detection.

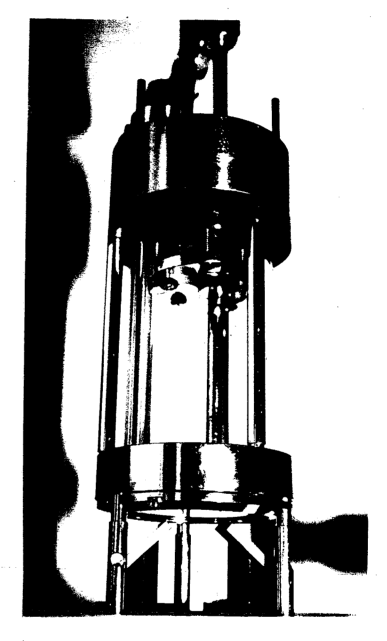


Figure 3 : Cell developed for FITR spectroscopy

# 3.7. Detection Limits for Different Gases

The adaptations to the apparatus and the analytical procedures described previously allowed to reach the detection limits listed in Tables 3 and 4 and expressed in ppm (mg of gas formed by kg of salt  $(mg/kg_{salt})$ ).

Table 3: Detection Limits (mg/kg<sub>sall</sub>) of Mass Spectrometry and Chromatography

CO <sub>2</sub>	N <sub>2</sub> O	СО	CH <sub>4</sub>	H <sub>2</sub>	SO <sub>2</sub>	H <sub>2</sub> S	Cl-	Cl <sub>2</sub>
10-2		5 . 10-4						2 . 10-1

Table 4: Detection Limits (mg/kg<sub>salt</sub>) of FTIR/Gas at Ambient Temperature

CO <sub>2</sub>	N <sub>2</sub> O	NO	СО	NO <sub>2</sub>	HNO <sub>2</sub>	HNO <sub>3</sub>	CH <sub>4</sub>	C <sub>2</sub> H <sub>6</sub>	C <sub>2</sub> H <sub>4</sub>	HCl	SO <sub>2</sub>
				5.10-3							

# 4. RESULTS OF PRE-IRRADIATION DETERMINATION

# 4.1. Mineralogical and Chemical Composition

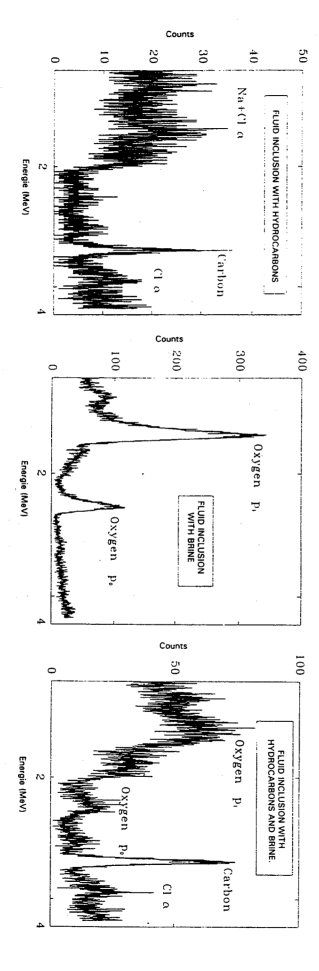
The granulometry of the salt delivered in August 1988 is very heterogeneous. Grain sizes vary from a few millimeters to the size of very fine dust. Moreover, salt blocks were observed, leading one to assume substantial humidity. The mineralogical compositions of the batches supplied by our colleagues of GSF/IfT and their water contents are given in Tables 5 and 6.

Table 5: Mineralogical composition of Asse Salt (Batch I)

Mineral Phase	Halite	Polyhalite	Anhydrite	Others	H <sub>2</sub> O
Mass % (mineralogical analysis)	95.2	0.8	3.8	< 0.3	0.066

Table 6: Mineralogical composition of Asse Salt (August 1988)

Mineral Phase	Halite	Polyhalite	Anhydrite	Kieserite	H <sub>2</sub> O	Carbonates
Mass % (mineralogical	91.39	3.53	5.08			
analysis)						
Mass % (chemical analysis)	92.84	3.82	3.06	0.06	0.24	0.10



CEA performed additional chemical analyses on the final batch with the following

results:

$\operatorname{assm} \% \left[ c.0 \pm 0.8 c \right]$	Cl- analyzed by potentiometry	-
$885 \text{m} \% [20.0 \pm 89.1]$	$\mathrm{SO}_4^{2\text{-}}$ analyzed by ion chromatography	-
sssm %1.0 >	$\mathrm{CO}_3^2$ analyzed with exclusion chromatography	-

Total organic carbon (TOC) of this batch is  $[8 \pm 3]$  ppm

Cation distribution is given in Table 7.

Table 7: Cation composition

200.0>	>0.005	₱90.0	62.0	ç6.0	10	730	tlas to marg / gm
эŦ	İS	ΙA	gМ	Ca	K	ьИ	Mineral

# 4.2. Composition of Fluid Inclusions.

The charged particle spectra obtained with the Nuclear Microprobe?? make it possible to identify three types of fluid inclusions : fluid inclusions with hydrocarbons, with brine and fluid inclusions with both hydrocarbons and brine (Fig. 4).

Inclusions' carbon and oxygen content have been calculated for each spectrum by subtracting the contribution of the salt, deduced from a particle signals characteristic of chlorine that are emitted at a higher energy than the carbon protons. Corrections in the measurements were made to take into account interferences existing on the carbon and oxygen proton signals. These interferences were attributed to charged particles (alphas) produced by sodium and chlorine. They were estimated from an halite piece devoid of fluid inclusions.

Secondly, and despite rather poor counting statistics, oxygen contents were estimated from the p<sub>o</sub> signal (see Fig. 4) because of intense interferences with salt signals in the p<sub>1</sub> proton regions. From the non-offset position, in relation to information from the surface, of the carbon

and oxygen peaks and their width at mid-height, the following information can be deduced:

- in the biphased inclusion, an approximately 2  $\mu$ m thick, low-density, carbon-rich layer covers the surface of the inclusion; it could be liquid methane;
- inclusions with hydrocarbons and biphased inclusions with hydrocarbons and brine are small in size: 30  $\mu$ m wide and a maximum of approximately 5  $\mu$ m thick; the brine inclusion is 5  $\mu$ m wide and 7  $\mu$ m thick.

Measured carbon and oxygen contents are, respectively, 0.015 mol cm<sup>-3</sup> of carbon in the inclusion with hydrocarbons, 0.01 mol cm<sup>-3</sup> of oxygen in the biphased inclusion and 0.08 mol cm<sup>-3</sup> of oxygen in the inclusion with brine only. For the latter inclusion, the measured oxygen content based on the gamma photon signal is 0.04 mol cm<sup>-3</sup>. Analytical accuracy is 50% due to corrections for salt and germanium contributions. Measured carbon and oxygen contents for pure constituents correspond to the contents in pure water and methane.

The overlap of the different techniques enabled us to specify the composition of the fluid inclusions, esp. presence of carbon and heterogeneity of chemical composition from one fluid inclusion to another in the same sample; the inclusions may be rich in carbon and/or oxygen.

#### 5. RESULTS OF POST-IRRADIATION ANALYSIS

The results from nearly 350 experiments were provided as work progressed and in the framework of meetings of the CEC/B1 (Asse) project committee. Most of the irradiations planned at  $10^{-3}$  Gy/hr for 0.1, 1 and 10 hours were not carried out because the quantities of the gaseous compounds,  $CO_2$  and  $N_2O$ , which were expected to develop, laid at the level of the detection limits.

### 5.1. The Determined Amount of Colloids.

Samples irradiated in a synthetic air atmosphere with a dose rate of 10<sup>4</sup> Gy/hr at 50°C, were dissolved in H<sub>2</sub>O and the amount of hydrogen obtained was measured. The Hydrogen results show that the amount of Na colloids developed by irradiation increases as a function of the integrated dose. The photograph in Fig. 2 shows the blue coloration of the salt caused by the development of colloidal sodium particles. The amount of hydrogen which can be obtained by dissolving the samples containing colloids can be expressed as:

$$hydrogen = K (Integrated Dose)^x$$

with x = 0.64 from the plot of Fig. 5b in the irradiation conditions (dose rate and temperature).

The amount of colloids developed in samples irradiated at temperatures of 50 to 250°C, with a dose rate of 10<sup>4</sup> Gy hr<sup>-1</sup> and an integrated dose of 10<sup>6</sup> Gy reaches a maximum between 100 to 150°C. It is linked to the mobility of the F centers and to the facility of diffusion of sodium atoms based on colloidal particles of metallic sodium, both of which depend on temperature.

The results obtained for samples of salt with different grain sizes, irradiated with a dose rate of 10<sup>4</sup> Gy/hr and an integrated dose of 10<sup>6</sup> Gy at 50°C in a synthetic air medium shows that the formation of colloids is greater when the salt is crushed more finely.

The production of sodium colloids increases slightly when the dose rate increases between 10<sup>3</sup> and 10<sup>5</sup> Gy/hr.

#### 5.2. Results of Analysis of Gases Released During Radiolysis

The gaseous compounds present in the vial after irradiation can be formed or released either from the atmosphere surrounding the salt (whether oxidizing, reducing or inert) or from the salt sample itself including crystalline matrix (mainly halite), secondary minerals, adsorbed gases and fluid inclusions.

Figures 5 and 6 as well as the attached result curves (Fig. 7 to 26) make it possible to observe the different interpretations presented hereunder.

In the Fig. 7 to 26, the contents of the different gaseous compounds are expressed in  $\mu$ mole per kilogram of salt in order to better point up the importance of hydrogen in the mechanisms in play.

## 5.2.1. Evolution of the Filling Atmosphere.

For irradiations done in synthetic air atmosphere, the  $O_2/N_2$  ratio, initially at 0.25, is monitored as a function of the total dose. For an integrated dose of  $10^6$  Gy, variations of the dose rate from  $10^2$  and  $10^5$  Gy·hr<sup>-1</sup> does not affect this ratio.

This ratio, which is practically constant as a function of the integrated dose up to  $10^6$  Gy, decreases rapidly beyond  $10^7$  Gy. This is explained by the consumption of the oxygen of the filling atmosphere related to the formation of  $CO_2$ . High temperatures enhance this phenomenon.

The drop of the  $O_2/N_2$  ratio above total doses of  $10^7$  Gy has been observed for two different granulometries as plotted in Fig. 10 and  $11(.1 < \phi < .25 \text{ mm})$ .

Irradiations of batches of salt of different granulometries did not reveal a variation in the  $O_2/N_2$  ratio. However, during irradiation at  $10^4$  Gy/hr for a dose of  $10^6$  Gy, for the batch of salt whose particle diameter is from 0.125 to 0.25 mm, the formation of  $CO_2$  is abnormally high. The explanation proposed by our German colleagues was a possible pollution of the batch during its storage after crushing in the mining equipment.

Irradiations performed at  $10^{+6}$  Gy in the presence of mixtures of oxygen and nitrogen with  $O_2/N_2$  ratios of 50/50 or 20/80 (Fig. 17 and 18) hardly change the original ratios. However, a 10% decrease in the nitrogen content of one vial, initially filled with an  $O_2/N_2$  ratio of 20/80 have been observed.

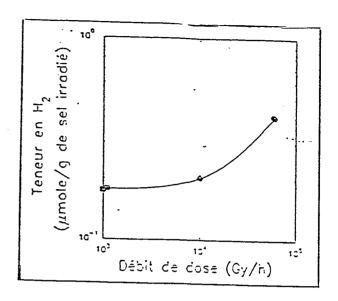


Figure 5a Production of colloidal sodium vs dose rate (106 Gy in air @ 50°c)

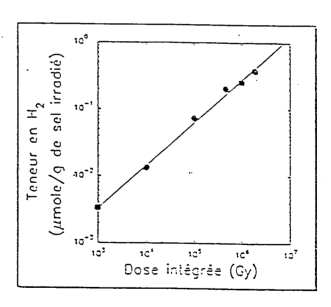


Figure 5b Production of defects in crystals vs integrated dose (10<sup>4</sup> Gy/h in air @ 50°c)

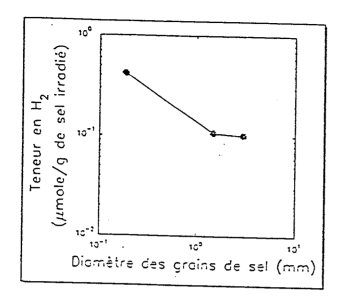


Figure 6a Formation of colloidal sodium vs grain size (10<sup>4</sup> Gy/h and 10<sup>6</sup> Gy in air @ 50°c)

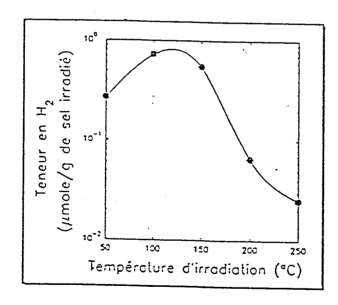


Figure 6b Formation of colloidal sodium vs temperature ( $10^4$  Gy/h and  $10^6$  Gy in air)

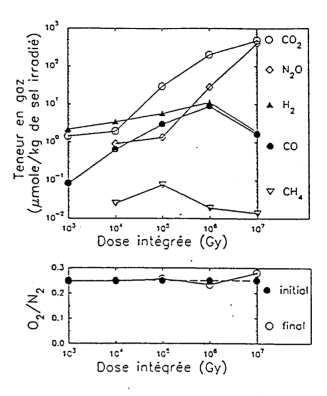


Figure 7 Gas production vs integrated dose (10<sup>4</sup> Gy/h in air @ 50°c)

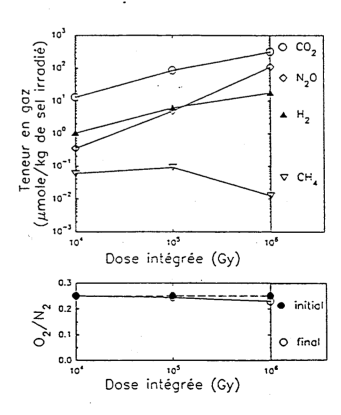


Figure 8 Gas production vs integrated dose (10 $^4$  Gy/h in air @ 150 $^\circ$ c)

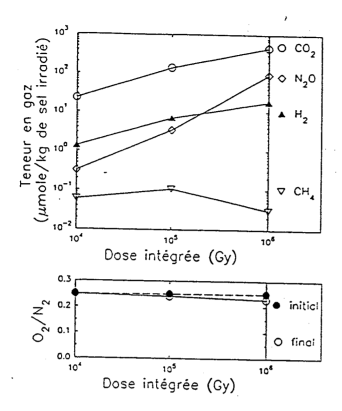


Figure 9 Gas production vs integrated dose (10<sup>4</sup> Gy/h in air @ 200°c)

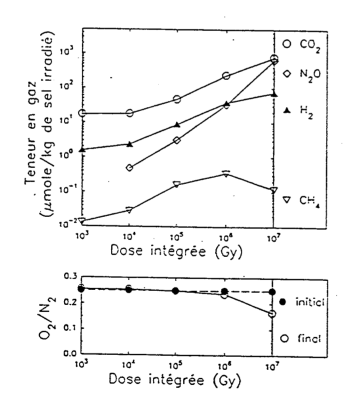


Figure 10 Gas production vs integrated dose : Grain size between 0.1 mm and 0.25 mm (10<sup>4</sup> Gy/h in air @ 50°c)

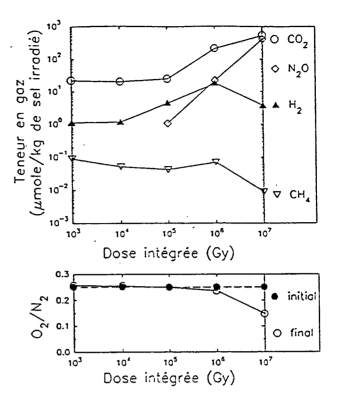


Figure 11 Gas production vs integrated dose : Grain size between 1 mm and 2 mm  $(10^4 \text{ Gy/h in air } @ 50^{\circ}\text{c})$ 

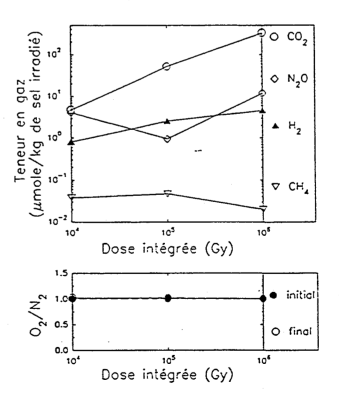


Figure 12 Gas production vs integrated dose : Initial filling gas 50% N<sub>2</sub> + 50% O<sub>2</sub> ( $10^4$  Gy/h @  $50^{\circ}$ c)

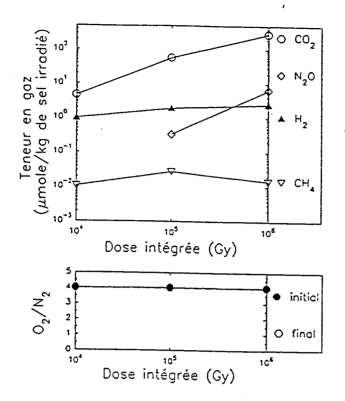


Figure 13 Gas production vs integrated dose : Initial filling gas 20%  $N_2$  + 80%  $O_2$  (10<sup>4</sup> Gy/h @ 50°c)

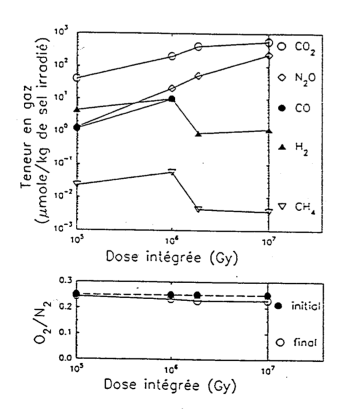


Figure 14 Gas production vs integrated dose (10<sup>3</sup> Gy/h in air @ 50°c)

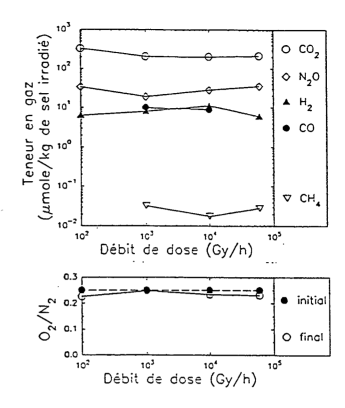


Figure 15 Gas production vs dose rate (106 Gy in air @ 50°c)

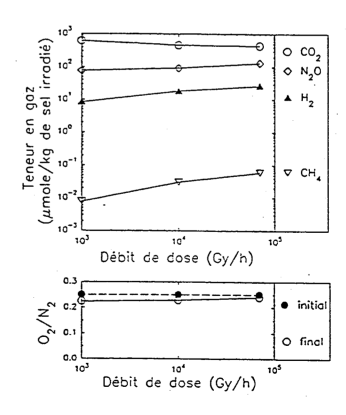


Figure 16 Gas production vs dose rate (106 Gy in air @ 200°c)

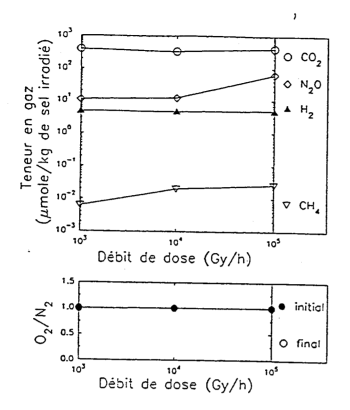


Figure 17 Gas production vs dose rate : Initial filling gas 50% O<sub>2</sub> + 50% N<sub>2</sub> ( $10^6$  Gy @  $50^{\circ}$ c)

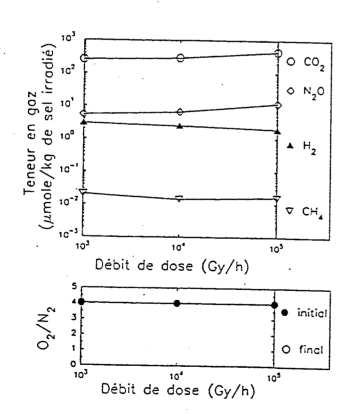


Figure 18 Gas production vs dose rate : Initial filling gas 20%  $N_2$  + 80%  $O_2$   $(10^6~\mbox{Gy}~\mbox{@}~50\mbox{^{\circ}}\mbox{c})$ 

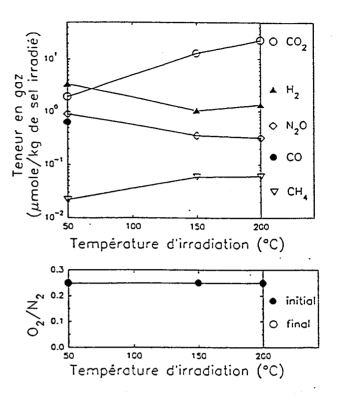


Figure 19 Gas production vs temperature (10<sup>4</sup> Gy/h - 10<sup>4</sup> Gy in air)

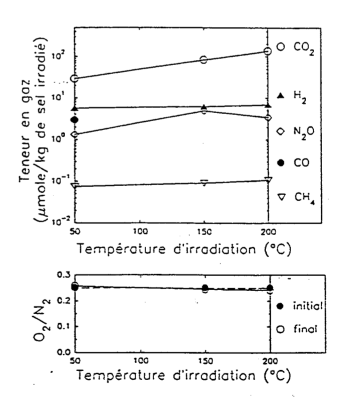


Figure 20 Gas production vs temperature (10<sup>4</sup> Gy/h - 10<sup>5</sup> Gy in air)

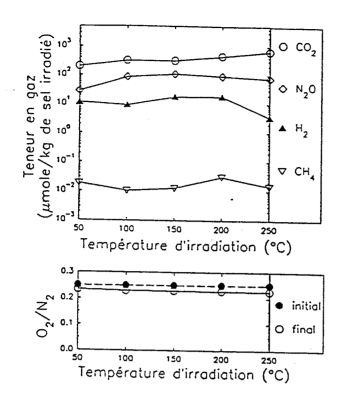


Figure 21 Gas production vs temperature (10<sup>4</sup> Gy/h - 10<sup>6</sup> Gy in air)

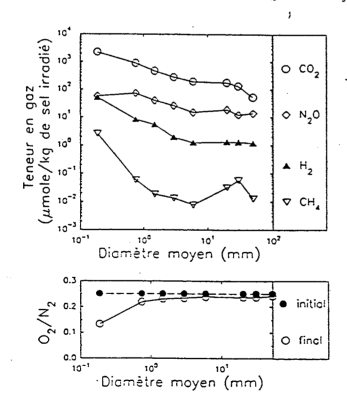


Figure 22 Gas production vs grainsize : Crushed salt and core samples  $(10^4 \text{ Gy/h in air } @ 50^{\circ}\text{C})$ 

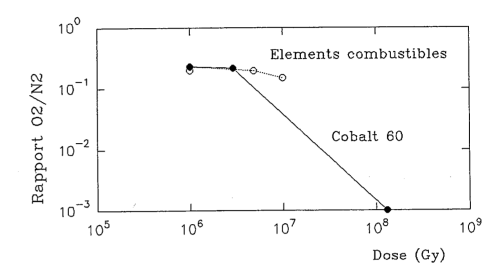


Figure 23 Evolution of O<sub>2</sub>/N<sub>2</sub> ratio vs dose

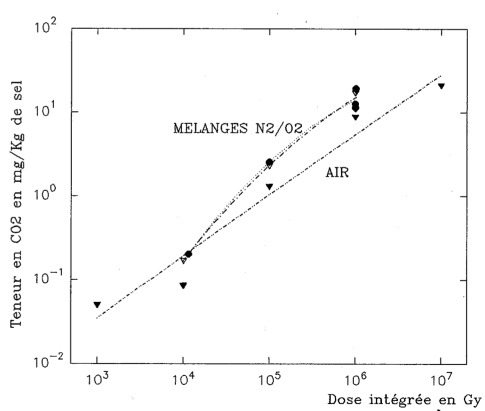


Figure 24 Evolution of CO<sub>2</sub> content vs O<sub>2</sub>/N<sub>2</sub> ratio in the initial filling gas

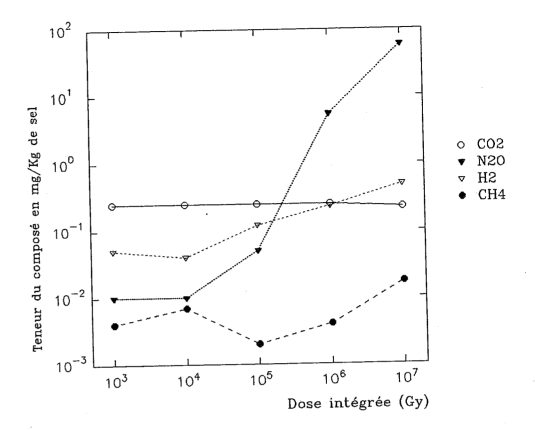


Figure 25 Evolution of irradiated blanks vs dose (10<sup>4</sup> Gy/h)

Blank = ampoule filled with air

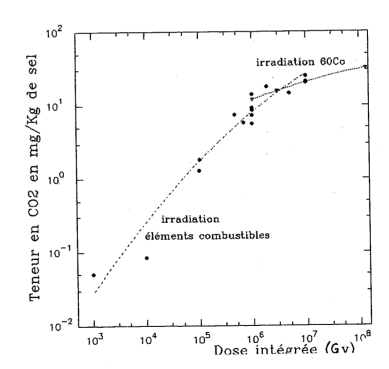


Figure 26 Evolution of CO<sub>2</sub> vs dose including irradiations with <sup>60</sup>Co

# 5.2.2. Type of Compounds Formed as a Function of the Filling Athmosphere.

The principal gaseous compounds released during  $\gamma$  irradiation observed as a function of the various parameters are:  $H_2$ ,  $CO_2$ ,  $N_2O$ ,  $CH_4$  and the initially present gas compounds. Others, such as CO,  $Cl_2$ ,  $Cl_1$ , NO,  $SO_2$ ,  $H_2S$  and the light hydrocarbons were identified during certain irradiations, mainly at high doses (>  $10^7$  Gy).

It is convenient for further study to distinguish between gases developed in an oxidizing atmosphere and those developed in an inert atmosphere. These two types of environments and of course the mixtures of  $O_2$  and  $N_2$  in a 50/50 and 20/80 ratio were studied during this program. An irradiation in vacuum with a dose rate of  $10^4$  Gy/hr and a dose of  $10^6$  Gy rounded off the study.

The composition of the gaseous phase in the repository will evolve over the course of time, with consumption of ambient oxygen; it would be interesting to pursue this study with mixtures whose  $O_2/N_2$  ratio diminishes to observe the kinetics of formation of the different compounds indicated.

Compounds formed in an oxidizing medium: these are mainly  $N_2O$  and  $CO_2$ , which are the ultimate phases of the reactions. NO and CO were analyzed, these are no doubt intermediate compounds, principally detected during incomplete oxidation reactions.

Compounds formed in an inert medium or in vacuum: these are mainly H<sub>2</sub> and hydrocarbons.

#### 6. INTERPRETATION

# 6.1. The origin of N2O.

 $N_2O$  is due to radiolysis of the air, the quantities detected in the inert medium are close to the detection limits.

The sequence of reactions that bring the nitrogen oxides into play culminates, when the quantity of oxygen is sufficient, in the formation of  $N_2O$ . In the irradiation blanks, the presence of NO is explained by the oxidation reaction of  $N_2 + O_2 - > 2$  NO. In the samples, the combination of  $N_2$  with  $O_2$  as 1/2  $O_2 + N_2 --> N_2O$  may be enhanced by the presence of salt.

The N<sub>2</sub>O contents obtained during the desorption tests at 300°C of samples irradiated in synthetic air, when subject to heating for more than one day, are not higher than those obtained immediately after irradiation, confirming that this compound is an end product of the reaction.

The generation of  $N_2O$  decreases slightly as the samples grain size increases, as plotted Fig. 22. For equal volume and composition of the atmosphere, the gas included in the crushed salt is more affected by radiation when the grain size is smaller.

The quantities of  $N_2O$  increase with integrated dose as plotted Fig. 7. This has been confirmed by the comparison of the results obtained with real samples and those obtained with those of "air blanks" simultaneously irradiated under the same conditions (Fig. 25). In the whole experiment, the observation of  $N_2O$  in the blanks was a way to control that the irradiation was correct.

The dose rate ( $10^2$  to  $10^5$  Gy/hr) does not influence the development of  $N_2O$  as it is plotted Fig. 15 for a constant total dose of  $10^6$  Gy.

The heated irradiations show that the production of  $N_2O$  seems to reach a maximum between 150 and 200°C for total doses of  $10^5$  and  $10^6$  Gy (Fig. 20 and 21). However, this was not confirmed at  $10^4$  Gy.

After irradiations carried out in an inert atmosphere, the  $N_2O$  contents are below the detection limit. The formation of  $N_2O$  depends on the presence of oxygen, as inferred from the results of an irradiation in synthetic air and reaching a total dose of 1.3  $10^8$  Gy ( $^{60}$ Co source), where the formation of other oxygenated compounds was related to reduction of  $N_2O$  yields.

The presence of HNO<sub>2</sub> (sometimes detected by FTIR) in the samples of irradiated air is

It is not know whether salt has any influence on the radiolysis of air. Our results indicate the amount of air radilysis on the presence of salt, which is relevant for a repository.

### 6.2. The Origin of $CO_2$ .

The  $CO_2$  of the salt samples can be present as adsorbed or as gas in fluid inclusions which can be mono or polyphasic. Polyphasic fluid inclusions use to contain as well hydrocarbons.

Analyses by ESCA and FTIR showed that the carbon present in the salt is organic in origin. The content of total organic carbon (TOC) is 8 ppm.

A thermal desorption of a non-irradiated sample of crushed salt (batch 1988) packaged in synthetic air has been carried out up to  $300^{\circ}$ C for 24 hours. The analysis of the  $CO_2$  produced give a value of 35 mg of  $CO_2$  per kg of salt.

The generation of CO<sub>2</sub> takes place essentially in an oxidizing medium, the quantities formed in an inert medium are close to the detection limits.

The carbon dioxide development increases with total dose up to 10<sup>6</sup> Gy, then seems to reach a plateau at high doses (Fig. 7).

The dose rate has little influence on the release of CO<sub>2</sub> (Fig. 15 to 18).

The irradiations carried out to study the dependence of gas production on the temperature took place at various dose rates (10<sup>3</sup>, 10<sup>4</sup> and 10<sup>5</sup> Gy/hr) and total doses (10<sup>4</sup>, 10<sup>5</sup> and 10<sup>6</sup> Gy). The yields of CO<sub>2</sub> obtained from them show that at low doses, CO<sub>2</sub> development is enhanced by temperature (Fig. 15 and 16, and 19 to 21). For an irradiation at a total dose of 10<sup>4</sup> Gy, an increase of temperature 150°C increases the CO<sub>2</sub> yields in a factor of 10; at 10<sup>5</sup> Gy the same temperature rise encrease the CO<sub>2</sub> yields in a factor of 5. After irradiation at a

total dose of 106 Gy, heat seems having nearly no influence on CO2 production.

At low dose rates, the  ${\rm CO}_2$  generation is intensified by the quicker decomposition of hydrocarbons.

The influence of the salt grain size, as well as salt core drillings diameter, on the compounds produced has been studied at a dose of  $10^6$  Gy and at ambient temperature (see Fig. 22). The results confirm that  $CO_2$  generation increases as the granulometry decreases. The greater the specific surface area of the material, the easier the interaction of  $\gamma$  radiation will be with the organic compounds contained in the fluid inclusions.

At a high doses the oxidation reactions slow down because the oxygen originally present in the vial has been consumed already. This is inferred from an irradiation performed up to 1.3 10<sup>8</sup> Gy (using Co<sup>60</sup> sources) and at 65°C (Fig. 26). The quantity obtained stabilized around 20 mg of CO<sub>2</sub> per kg of salt above 10<sup>7</sup> Gy.

The formation of carbon dioxide is closely related to that of the other compounds, such as CO,  $CH_4$  and the existing hydrocarbons.

# 6.3. The Origin of $H_2$ , CO, $CH_4$

In the studied gaseous phase,  $H_2$ , CO and  $CH_4$  are less abundant than the rest of the gases. However, they provide valuable information on the reaction and on the equilibrium of the gaseous phase.

The *hydrogen* originates primarily by the action of gamma rays on the  $H_2O$  molecules and on the organic compounds present in the fluid inclusions.

Hydrogen yields reaches a maximum around  $10^6$  Gy then decreases at higher doses ( $10^7$  Gy), for irradiations in synthetic air. However, a greatest content has been observed at  $1.3 \ 10^8$  Gy. This may indicates that the reactions in play are interrupted by lack of  $O_2$  and

formation of oxygenated radical-like compounds allowing linkages with H<sub>2</sub>.

In an inert atmosphere, or in vacuum, where hydrogen cannot react with the oxygenated radicals which are also being developed by radiolysis, the yields of  $H_2$  are higher than in other environments where these reactions are possible.

Some irradiations have been carried out with samples of steel introduced in the vials together with the salt. One of the products of steel corrosion is hydrogen, which develops due to the reaction of the brines contained in salt with the metal. Contrary to the expectations, the amount of  $H_2$  produced within these steel containing vials is not significantly higher than that of other vials. Probably integrated doses of  $10^6$  Gy, with dose rate of  $10^3$  Gy/hr, and at temperatures of  $200^{\circ}$ C are not damaging enough to change the  $H_2$  production.

The <u>carbon monoxide</u> production is a function of the integrated dose. In the presence of oxygen the amount of CO produced reaches a maximum at  $10^6$  Gy, and then decreases (Fig. 7). It is therefore inferred that CO is produced as a the results of the oxidation of the organic matter contained in the salt samples, but that provided the amount of oxygen is sufficient,  $CO_2$  is produced to the expense of CO.

Hydrocarbons detected in our experiments have their origin in the organic mater contained in the salt samples. They use to be either absorbed at the surface of the grain or ocluded in the fluid inclussions. The organic mater undergoes radiolytic fragmentation during gamma irradiation. Heavy molecules can become light hydrocarbons in an iner environment. The hydrogen being produced by the brine hydrolysis can help in this fragmentation, and moreover there is an additional thermal fragmentation of heavy hydrocarbons which has to be taken into account. As a result of these processes in an inert environment mainly light hydrocarbons are found. The main product is methane, although  $C_2H_4$ ,  $C_2H_6$ ,  $C_2H_2$  and  $C_3H_8$  are also found in trace amounts. The amounts of light hydrocarbons other than methane are close to the detection limits of our methods.

In oxidizing environments, with increasing time, and dose,  $CH_4$  becomes CO and finally  $CO_2$ . Production of  $CH_4$  at a dose rate of  $10^4$  Gy/h and at  $50^{\circ}C$  in a synthetic air

environment has been observed to reach a maximum at 10<sup>5</sup> Gy, and then decrease in coincidence with the start up of the oxidation of the organic material. However, for irradiations up to very high doses (130 MGy) CH<sub>4</sub> content can increase if the amount of exygen in the environment is depleted: CH<sub>4</sub> would be constantly produced by fragmentation but not be able to oxidize and become CO<sub>2</sub>.

The characterization of the organic matter present in the salt inclusions would allow prediction of the formation of compounds from degradation during the radiolysis reactions.

# 7. CONCLUSIONS

The methodology developped and refined in the course of this research was adequated to produce the sought answers and can now be used to study the gas by radiolysis in other possible candidate host rocks for radioactive wast. It is important to remark that the detection limits were so low that they allow us to draw the following conclusions.

Two big groups of gas types can be distinguished amongst those generated during irradiation of salt; those gases obtained by radiaolysis of brine and salt, and those which are obtained by fragmentation and eventual oxidation of organic mater. We have found that small grain sizes enhance both types of gas generation.

The gases produced by radiolytic decomposition of brine and /or of the salt itself are mainly H<sub>2</sub> and Cl<sub>2</sub>, and minor amounts of other chlorinated compounds. Their production mainly depends on total dose, temperature and dose rate. Besides this direct production of H<sub>2</sub>, hydrogen could be produced in repositories in the eventuality of their accidental flooding. The water would then react with the sodium colloids produced in the rocks salt by radiation damage, and produce hydrogen, amongst other substances. The amounts of hydrogen produced would be proportional to that of sodium metal colloids.

The gases produced from, first fragmentation, and then eventual oxidation of originally heavy molecules of hydrocarbons, are mainly methane and carbon dioxide. Methane can

survive in higher or lower amounts depending on the oxidizing characteristics of the environment. In oxydating environments it will oxidize up to carbon dioxide, but if the environment is depleted in oxygen, the production of methane would be enhanced, at least relatively to that of carbon dioxide. Intermediate products can as well develop, e.g. carbon monoxide.

The amount of produced gases is limited by the volume of rocksalt affected by the radiation and the temperature encrease on one hand, and on the other hand by the amount of source material for gas production contained in this volume. The most important sources for radilytic gas production are already present fluids and the organic material. Moreover, gases such as hydrogen and methane will oxidize to produce wtaer and carbon dioxide in so far as oxygen is available.

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