UCRL-51818

DEVELOPMENT AND EVALUATION OF RADON SEALANTS FOR URANIUM MINES

H.G. Hammon

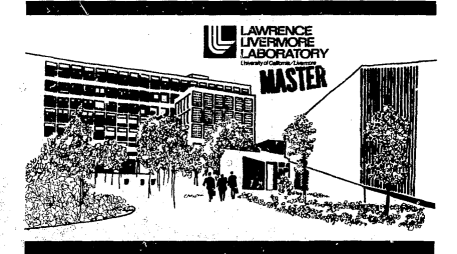
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May 21, 1975

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Foreword

This report was prepared by Lawrence Livermore Laboratory, Livermore, California under USBM Contract Number H0232047. The contract was initiated under the Metal and Nonmetal Health and Safety Research Program, It was administered under the technical direction of SMRC with Mr. John C. Franklin acting as the Technical Project Officer. Mr. Frank Pavlich was the contract administrator for the Bureau of Mines.

This report is a summary of the work recently completed as part of this contract during the period October, 1972, to February, 1975. This report was submitted by the authors on May 29, 1975.

Work was performed under the auspices of the U.S. Energy Research & Development Administration. Reference to a company or product name does not imply approval or recommendation of the product by the University of California or the U.S. Energy Research & Development Administration to the exclusion of others that may be suitable.

The authors acknowledge gratefully the assistance of many others in this work, especially 'Floyd Momyer for direct radon permeation measurements, James S. Johnson for toxicity evaluations, and Norriss Hetherington for statistical and mathematical analysis.

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Introduction

Uranium miners have been shown to develop lung cancer in direct proportion to the time of exposure in the mine and the concentration of radioactivity (cumulative working level).1 Radon gas is continually evolved from the uranium ore, but because of its short half-life, 3.82 days, it is soon decomposed into particulate daughter products, which, along with the dust in the mine, become deposited in the miners' lungs. At present, massive ventilation is the only effective method of reducing the radioactivity concentration in the mines. However, the Bureau of Mines is evaluating the possibility of using coatings on the mine walls that will have sufficiently low permeability to radon that the gas will decompose into the particulate daughter products before it can permeate the coating. Permanently trapped in the coating, the daughter products would be prevented from dispersing in the mine atmosphere

Lawrence Livermore Law ratory was asked to aid in this investigation. The direct determination of radon permeability is more difficult than that of the other noble gases because of the short half-life of the gas and radioactive contamination of the apparatus. We felt that it should be possible to estimate radon

permeation coefficients from other noble gas permeation coefficients. It was noted that the log of the diffusion constants for helium, neon, argon, and krypton, as reported by Meares²⁻²³ plotted to a straight line against the square of the gas molecular diameter. The log of the permeation coefficients also plotted to a straight line against the square of the gas molecular diameter. Thus, by determining the permeation coefficients of such gases us argon and krypton through various films and coatings, it appeared likely that radon permeation coefficients could be estimated. In addition, we felt it should be possible to find some relationship between polymer structure and the noble gas permeation coefficient that would aid in the selection of coatings materials as radon barriers.

As the program progressed, we were asked to evaluate selected coatings for their possible toxicity during application or a mine fire. As the coatings were of industrial origin, it was possible that they might contain volatile materials that would pose a toxicological hazard to personnel while applying coatings to the mine walls. Also, a mine fire might be made more hazardous by dense smoke from the coating or toxic materials emitted during its pyrolysis.

Summary and Conclusions

The purpose of this study was to evaluate selected coatings as radon barriers in uranium mines; first on their effectiveness in reducing radon permeation, and second on the basis of their potential toxicity during application or in a possible mine fire. Hopefully, this information would be helpful in developing coatings more effective than those commercially available.

The program was organized in three tasks: (1) Film Selection and Preparation; (2) Measurement of Permeability; and (3) Evaluation of Toxicity of Coatings.

Films for the permeation study were selected from commercially available polymer systems of known chemical composition. Coating systems were selected on the basis of their chemical composition using our own permeability data or published data to aid in selections; other coatings previously evaluated at Bureau of Mines were also studied. In agreement with the Bureau of Mines seven coatings were selected for full evaluation, including smoke chamber and toxicity studies. Two more were added later for smoke chamber study only. Permeation coefficients of the films and coatings to noble gases were determined. When the log of the permeation coefficients for argon, krypton and xenon is plotted against the square of the molecular diameter a linear relationship is obtained. Thus, a straight-line extrapolation of this line permits the estimation of the permeatility coefficient for radon. The latter may be undesirable as surface coatings because they yield large amounts of hydrogen

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chloride (HCV) when exposed to heat or fire; however, they produce negligible amounts of smoke. They might be useful as penetrating compositions to bond loose uranium ore. The one-component polyester develops only light smoke, but would be hazardous during application because it releases relatively large amounts of styrene vapors, and has a low flush point.

The furan coating has very low permeability to noble gases, and develops negligible amounts of smoke. However, it has an objectionable odor which would be difficult to mask, and its black color might be undesirable.

Three epoxy coatings were evaluated. Two of these were unpigmented; they developed dense smoke when exposed to heat or fire. The pigmented epoxy coating produced only a light smoke, and its vapors contained no toxic components.

Toxic components evolved from the uncured polymer systems were determined by gas chromotography/mass spectrography (GC'MS). The only materials of toxicological significance that were found were vinylidene chloride (a suspected carcinogen) in the vinylidene chloride copolymer, vinyl chloride (a carcinogen) in one of the unpigmented epoxy coatings, and epichlorohydrin (a sensitizer), in the same epoxy coating.

Pyrolysis studies were made on the cured coatings. Other than carbon monoxide and HC® no significant quantities of other toxic species were found. Carbon monoxide was not generated from any of the coatings in amounts that would be large compared with those that would be evolved from other materials usually involved in a mine fire, such as wood, diesel fuel, and waste materials.

In ranking the coatings studied, we should note that they represent only a few of such commercial coatings available; other manufacturers may make similar coatings. Also, the ranking gives relatively little weight to permeability, as all are adequate to stop radon in the normal thickness of application, providing a continuous coating is effected. Finally,

coatings 8 and 9 were evaluated only in the smoke chamber, but produced more smoke than most of the other coatings under these conditions. Thus, we rank the coatings in decreasing order of overall effectiveness as follows:

Name and Type

- Hydr Cpoxy 300, pigmented water-dispersed epoxy
- Resitron II, furan (catalyzed furfuryl alcohol polymer)
- Essex Polyester, pigmented one-component styrenated polyester
- Aerospray 70, plasticized polyvinyl acetate latex
- Saran XD-7151, vinylidene chloride copolymer
- EpiRez WD-510/EpiCure
 872, unnigmented waterdispersed epoxy
- 7. WSU-118, modified epoxy
- Promulsion 200, unidentified composition
- 9. Hydro Seal, acrylic emulsion

Comments

Bad odor

Flammable; contains styrene

Possible smoke problem Liberates hydrogen chloride in possible fire Possible smoke problem

Possible smoke problem
Possible smoke problem
Possible smoke problem

We have calculated that virtually all roatings with permeation constants lower than $10^{+10}~{\rm cm}^2_{\rm STP}$, cm/s·cm Hg and thickness between 5 and 10 mil will provide nearly 100% effectiveness. The selection of a suitable coating then ought to be based on other factors such as ${\rm cost/m^2}$, vapor toxicity during application, ability to bind to the uranium ore in continuous films free of pin-holes etc. Obviously, any openings in the mine shafts and tunnels which are not covered with a film or any discontinuities of the film will act like the spillway on a water dam, i.e., the radion contained behind the barrier will flow out into the tunnel.

Experimental Work

The study of radon scalants for uranium mines was organized as three tasks: (1) Coating Selection and Preparation; (2) Measurement of Permeability and Diffusion Constants; and (3) Evaluation of Toxicity of Coatings. Although there was overlapping of these tasks, this was also the chronological order in which the study was made.

TASK 1: COATING SELECTION AND PREPARATION

Many of the coatings evaluated at LLL had previously been studied at Bureau of Mines, SMRC. Additionally, we surveyed the chemical industry and obtained specimens of coatings materials for evaluation. As much as possible, the composition of the coatings was obtained from the manufacturer. Permeability studies were also made on commercial films of known composition, and these results were used as a basis for selection of coatings.

Most of the coatings were prepared by drawdown from solution or latex using a doctor blade on a substrate of known permeability. This substrate was usually a polyethylene ionomer (Surlyn A). This material had high permeability and wetted better than conventional polyethylene. The coatings were allowed to dry thoroughly at room temperature and were conditioned for several days at about 50% relative humidity. The thickness of unsupported coatings or coatings on the Surlyn A substrate was measured with an electronic micrometer and averaged values were rounded off to the nearest 0.025 mm (0.0001 in.).

Coatings for evaluation in the LLL/NBS Smoke Chamber were applied, usually by spraying, to the surface of 73 × 73 × 25 mm cement-asbestos board specimens. In some cases, similar size specimens were cut from low-grade uranium ore and were coated for smoke-chamber study. Results with coatings on uranium ore specimens were essentially the same as with cement-asbestos board specimens. It was costly to cut the ore specimens and their uranium content required special monitoring procedures. Therefore, mort determinations were made on cement-asbestos board

specimens. Coatings on the cement-asbestos hoard or uranium ore specimens were measured indirectly by bonding a thin steel steeh to the surface of a similar specimen, coating it at the same time and in the same nanner as the speciments to be evaluated in the smoke chamber, and measuring the thickness of the cured coating with a magnetic coating thickness gage.

TASK 2: MEASUREMENT OF PERMEABILITY AND DIFFUSION CONSTANTS

Dynamic Method Using a Mass Spectrometer as a Gas Detector

In this dynamic technique a small mass spectrometer (a converted CEC21-614 Residual Gas Analyzer) serves as gas detector. The specificity, sensitivity and precision are superior to most other detectors. For thin films (up to 15 mils) this method gives results rapidly, and interfering membrane outgassing and desorption gases can be distinguished from the permeant. The presence of pinholes in, or leaks around. the membrane is readily recognized. A schematic of the measurement system is shown in Fig. 1. The specially designed permeation cell is connected to the ion source via an isolation valve. In the open position the flow impedance of the valve port is negligible relative to the flow impedance of the membrane. By using two Viton O-rings and continuously pumping on the volume between the O-rings, air leaks are virtually eliminated.

The O-rings are vacuum baked before assembling the cell to remove dissolved gases and water vapor. This cell can be heated to about 150°C and the membrane thickness can be varied from 1 mil to about 65 mils. In operation, the permeation cell is evacuated on both sides of the membrane and a spectrum of the residual gases is obtained. If any of these gases are identical to the permeant to be used (the test gas) evacuation continues until the background is negligible. If this requires too much time, the outgassing and/or desorption rates are measured as function of time to be used as correction factors to the permeation rate. The isolation valve to the spectrometer is

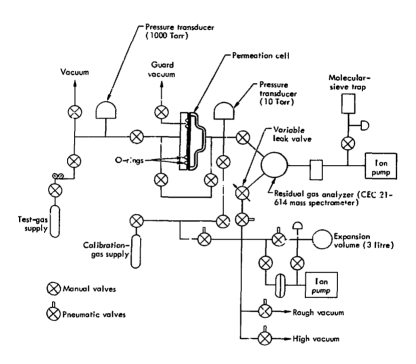


Fig. 1. Diagram of the mass spectrometer and associated apparatus that was used for measurement of permeability constants.

then opened and the test gas is applied at a known pressure (P_i) on the in-going side c f the membrane. A suitable ion peak (m/c = 4) for helium: m/c = 40 for Ar; m/c = 83, 84 or 86 for Kr f is continuously monitored until the recorder trace or digital output reaches a steady state value.

The system is calibrated by measuring the mass spectrometer's electrometer response to a known, constant flow of the test gas which furnishes the sensitivity (S). The steady-state permeation rate ($\hat{\mathbf{Q}}$) is obtained from the relation

where

$$\dot{Q}$$
 = permeation rate (cm $\frac{3}{STP}/s$)
S = sensitivity (cm $\frac{3}{STP}/s \cdot mV$)

V = electrometer signal at steady state (mV).

From this, the permeation coefficient is calculated using the following equation:

$$P = \dot{Q} \cdot d/A\Delta_{D}$$
.

where

d = membrane thickness (cm)

A = membrane area (cm2)

 $\Delta p = p_1 \sim p_2 \simeq p_1 = \text{test gas pressure (cm Hy)}$

P = permeation coefficient (cm³_{STP} · cm/s·cm² ·cm Hg)).

From a single experiment we can also calculate the disffusion constant (D) by the method of Pasternak⁴ if the complete permeation curve from zero transmission to the steady state value is obtained. Since

$$P(T) = D(T) s(T)$$
.

the solubility (s) can be calculated with P and D determined. All three terms are temperature dependent. Experiments at different temperatures permit the measurement of the temperature coefficients.

2. Use of Dow Cell for Permeability Measurements

Determinations of permeability coefficients using the mass spectrometer often required long times, expecially with low permeability films and coatings and with the higher molecular weight gases such as kyrpton and xenon. Therefore, we made many of our determinations with a Dow film permeability cell, as described in ASTM D-1434-66. A computer program was written for a PDP-8 to calculate permeation coefficients from the experimental data: the program is shown in Appendix B.

3. Direct Determinations of Radon Permeability

Radon permeation measurements are more difficult to make than those of other noble gases for a number of reasons. For most polar materials, the radon permeation rate will be slow due to the larger gas molecule and its slower diffusion rate. It has a relatively short half life, thus deposits its daughter products in the measuring apparatus. We did not want to so contaminate the mass spectrometer of the Dow cell. Therefore, direct determination of radon permeation through polyethylene and through polycarhonate films was made using radiochemical techniques with apparatus made for the purpose from 80-mm ovrex glass tubing. Radon, derived from a radium solution, was admitted to one side of the apparatus, and allowed to permeate the film which separated the apparatus into two chambers. The gas which permeated the film was determined on the other side by alpha counting. Complete details of the determination of the radon permeability coefficient of polycarbonate film are described in Appendix C by Dr. Floyd Momver, who was in charge of this phase of the work.

RESULTS AND DISCUSSION OF PERMEABILITY STUDY

The permeation coefficients of some commercial films are listed in Table 1 and the permeation coefficients of some supported coatings are listed in Table 2. A complete listing of the permeation coefficients of the supported coatings will be found in Table A-1 in Appendix A. Some of these data are plotted in Figs. 2 and 3 as the logs of the permeation coefficients vs the square of the diameter of the permeant pas. In general, good straight line fits were obtained for argon, krypton, and zenon permeation coefficients, although the helium and neon permeation coefficients did not

usually fit well with the other data. Direct measurements of the radon permeation coefficients were made only through polyethytene and polycarbonate films. As may be seen from Fig. 2, these values agreed well with values that would be predicted by extrapolation of the argon, krypton, and xenon values.

The permeation coefficients for noble gases through polycthylene and Surlyn A increase with increasing gas molecular weight and diameter. Similar effects have been reported for neon and argon permeation through ethylene-propylene copolymers, styrene-butadiene copolymers, and a cis-polybutadiene⁵, for neon, argon, ktypton, and xenon through natural rubber⁶, and for neon, argon, ktypton, and xenon through nitrough silicone rubber⁷. In polymers of higher cohesive energy

Table 1. Permeation coefficients of commercial films.

Film	Gas	Permeation coeffice	cients
		cm3rp · cm	nm²
		sec · cm² · cm Hg	Pa · s
Polyethylene	He	6.7 × 10 ⁻¹⁰	50
	Ar	3.4 × 10 ⁻¹⁰	25
	Kr	4.5 × 10 ⁻¹⁰	34
	Xe	8 × 10 ⁻¹⁰	60
	Rn	1.5 × 10 ⁻⁹	113
Surlyn A	Ne	$2.0 \times 10^{-1.0}$	15
(polyethylene ionomer)	Ar	1.3×10^{-10}	9.8
	Kr	1.5 × 10 ⁻¹⁰	11
	Xe	1.9 × 10 ⁻¹⁰	14
Polycart onate	He	1.5 × 10 ⁻⁹	113
	Ar	5.5 X 10 ^{-1.1}	4.1
	Kr	5 × 10 ^{-1 1}	3.8
	Xe	2 × 10 ⁻¹¹	1.5
	Rn	1.8 × 10 ^{-1 l}	1.4
Połyvinylchloride,	He	4.0×10^{-10}	30
unplasticized	Ar	7.0×10^{-12}	0.53
	Kг	3.5 X 10 ⁻¹²	0.26
	Xe	4 × 10 ⁻¹³	0.03
Phenoxy	Ne	2.5 × 10 ⁻¹¹	1.9
	Ar	2.6×10^{-12}	0.20
	Kr	$1.1 \times 10^{-1.2}$	80.0
	Xe	2.1×10^{-13}	0.016

Table 2. Permeation coefficients of supported coatings.

Permention coefficients,							
Ar	Kr	Xe					
3.9 × 10 ⁻¹¹ (2.9)	N.D.	N.D.					
$6.0 \times 10^{-12} (0.45)$	N.D.	N.D.					
4.4 × 10 ⁻¹³ (0.033)	$2.2 \times 10^{-13} (0.017)$	8.9 X 10 ⁻¹⁴ (0.0067)					
7.3 X 10 ⁻¹² (0.55)	6.8 × 10 ⁻¹² (0.51)	5.0 × 10 ^{-1 2} (0.38)					
1.1 × 10 ⁻¹ 1 (0.83)	7.7 × 10 ⁻¹² (0.58)	3.5 × 10 ^{-1 2} (0.26)					
3.4 × 10 ⁻¹¹ (1.8)	1.1 × 10 ⁻¹¹ (0.83)	1.9 X 10 ^{-1 2} (0.14)					
1.7 × 10 ⁻¹⁰ (13)	N.D.	N.D.					
	Ar 3.9 × 10 ⁻¹¹ (2.9) 6.0 × 10 ⁻¹² (0.45) 4.4 × 10 ⁻¹³ (0.033) 7.3 × 10 ⁻¹² (0.55) 1.1 × 10 ⁻¹¹ (0.83) 2.4 × 10 ⁻¹¹ (1.8)	$\begin{array}{c c} \frac{cm_{STP}^3 \cdot cm}{s \cdot cm^2 \cdot cm Hg} & \frac{nr}{Pa} \\ \hline Ar & Kr \\ \hline 3.9 \times 10^{-11} (2.9) & N.D. \\ 6.0 \times 10^{-12} (0.45) & N.D. \\ 4.4 \times 10^{-13} (0.033) & 2.2 \times 10^{-13} (0.017) \\ 7.3 \times 10^{-12} (0.55) & 6.8 \times 10^{-12} (0.51) \\ \hline 1.1 \times 10^{-11} (0.83) & 7.7 \times 10^{-12} (0.58) \\ 2.4 \times 10^{-11} (1.8) & 1.1 \times 10^{-11} (0.83) \\ \hline \end{array}$					

density*, however, the permeation coefficients of the noble gases decrease with increasing gas molecular weight and diameter, as shown by our data and the data of Meares²⁻¹³.

It has been shown⁸ that P = DS, where P is the permeation coefficient, D is the diffusion coefficient, and S is the solubility coefficient, provided D is constant. Up to pressures of 1 or 2 atmospheres, the solubility of the noble gases in polymers obeys Hen.y's $l_0...$, i.e. C = SD, where

- C = concentration of gas dissolved in unit volume of polymer (cm³_{STP}/cm³)
- p = partial pressure of noble gas in contact with the polymer (1 atm)
- the polymer (1 atm) $s = solubility coefficient \left(\frac{cm_{STP}^3}{cm^3 \cdot atm}\right).$

For gases that dissolve only sparingly, the solubility coefficient (S) is a constant. Once dissolved the permated molecules diffuse to the other side of the membrane. The driving force is the concentration

difference that exists between the incoming and outgoing membrane surfaces. The flux of permeant (J) per unit membrane surface area is then given by

$$T = D (C_{in} - C_{out})/d$$
or
$$J = SD (p_{in} - p_{out}/d,$$

where

d = thickness of the membrane (cm)

D = diffusion coefficient
$$\left(\frac{\text{cm}^2}{\text{s}}\right)$$

 $f = \text{flux of permeant } \left(\frac{\text{cm}^3\text{STP}}{\text{s} \cdot \text{cm}^2}\right)$

The diffusion coefficient (D) is essentially a constant for sparingly soluble gases because the membrane structure is not perturbed by the dissolved molecules. Up to and near atmospheric pressures, the diffusion of noble gases in polymers is independent of concentration. This is mainly because of the very low concentrations encountered at all normal pressures, but it also reflects the comparative lack of interaction between these gases and the polymer.

At pressures of 1 or 2 atmospheres, the solubility of the gases in the polymers obeys Henry's law. In

^{*}The cohesive energy density (CED) is approximately equal to the heat of vaporization or sublimation at constant volume and can be estimated from thermodynamic data.

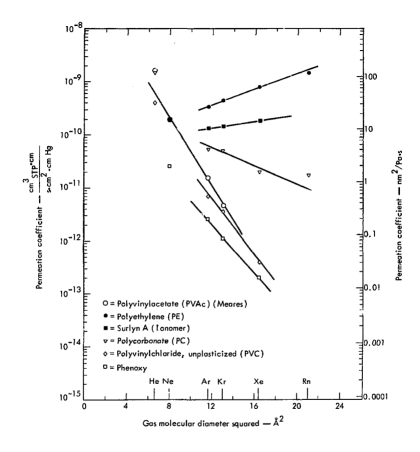


Fig. 2. Permeation coefficients of films vs the square of the gas molecular diameter.

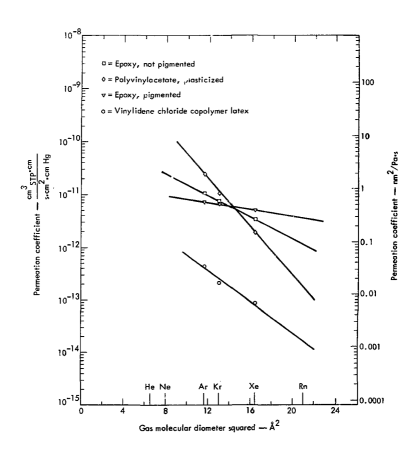


Fig. 3. Permeation coefficients of coatings vs the square of the gas molecular diameter.

the generally accepted picture of the activated diffusion process, larger holes need to be formed in the polymer for the diffusion of larger molecules. These larger holes require a larger energy for their formation, thus the activation energy will be larger for the diffusion of larger molecules and the diffusivity will be smaller. This has been found to be true. However, the solubility increases with increasing gas molecular weight and has been shown9,10 to be directly related to the critical temperatures of the gas. Solubility of simple gases in hydrocarbon polymers has been found to be directly proportional to the volume fraction of amorphous polymer;11 however, introduction of polar groups decreases solubility. Thus, in looking at the permeability coefficients of the noble gases in a particular polymer film, if the solubility coefficient increases more (in going from one noble gas to the next in order of increasing molecular weight) than the diffusion coefficient decreases, the net effect on the permeation coefficient is an increase.

Modification of Coatings

Effects of Pigments in Coatings

Many of the coatings evaluated in this study were pigmented as obtained from the manufacturer. In some cases, it was possible to compare pigmented with non-pigmented coatings, but, as in the case of the epoxy coatings reported (compare the Hydr Epoxy

coatings with the EpiRez/EpiCure coatings) there appeared to be little difference between them. It would be reasonable to expect, however, that some pigments would reduce the permeability of the coating by an overlapping or leafing structure in the dried coating. Two pigments were selected with this effect in mind: a leafing aluminum pigment, and powdered mica. In a Saran latex (Dow XD-4624), the addition of the leafing aluminum pigment reduced the permeation coefficient by about a factor of 2, but the mica modified coating had a permeation coefficient higher than the unmodified coating by about a factor of 10. The pigments were also studied in an ethylene/vinvl chloride copolymer, Monflex 4514. The permeation coefficient to argon of the mica modified Monflex 4514 was found to be 1.2 x 10-11, about half the value found for the latex compound with aluminum pigment, but still twice as high as the unpigmented latex. Because the permeability coefficients appeared to be affected more by the choice of the base polymer than by pigment compounding, this work was not carried further. Data are listed in Table 3.

Effects of Crosslinking

Crosslinking of polymer film has been shown to reduce permeation, apparently by reduction in the diffusion constant. 12-15 No direct attempt was made in this program to reduce permeation by crosslinking

Table 3. Effects of pigments in coatings.

Coating (type)	Pigment	Permeation coefficier	it of coating
		cm3 · cm	nm ² Pa·s
		s · cm² · cm rig	Pa·s
Saran XD-4624 (Vinylidene chloride copolymer)	None	4.4×10^{-13}	0.033
Saran XD-4624 (Vinylidene chloride copolymer)	Leafing Juminum	1.9×10^{-13}	0.014
Saran XD-4624 (Vinylidene chloride copolymer)	Ground mica	4.2×10^{-12}	0.32
Monflex 4514 (ethylene/vinyl chloride copolymer)	None	6.0×10^{-12}	0.45
Monflex 4514 (ethylene/vinyl chloride copolymer)	Leafing aluminum	2.5×10^{-11}	1.9
Monflex 4514 (ethylene/vinyl chloride copolymer)	Ground mica	1.2×10^{-11}	0.9

because of the difficulty of introducing crosslinking agents that would be effective in room temperature cures. However, we did observe a definite lowering, by about an order of magnitude, in the permeation coefficient of a HydrEpoxy coating that was retested after standing for several months. We attribute this to the increase in crosslinking taking place in the room temperature curing epoxy. A polyvinyl acetate coating prepared at the same time as the epoxy coating showed no change on remeasurement after standing.

Thickening Agents

Some of the coatings were low in viscosity and tended to sink into porous substrates such as the uranium ore. If this happened, a coherent film was not formed and the coating was not a good barrier. Thickening agents such as polyvinyl alcohol solution or carboxymethyl cellulose were recommended by the manufacturer. Used at the recommended levels, these thickening agents did not change the permeation coefficients. However, the thickened coating made a coherent film when applied to uranium ore, and thus should be a better barrier.

Correlation of Permeability with Polymer Structure

In selection of coatings as radon barriers, it would be desirable to be able to predict the radon permeation coefficient based on the structure of the base polymer used in the coating. For this reason, we made some saudies of possible relationships. Many chemical, morphological, and structural properties of the polymer can affect the permeability of a polymer.14,16-18 These include chemical properties such as cohesive energy density (CED) and hydrogen bonding, morphological properties such as crystallinity and orientation, and the effect of structural groups in the polymer chain. It was quickly apparent that CED alone would not correlate with permeability, but we did find a reasonable correlation of the product of CED and polymer density with argon permeability. as shown in Fig. 4. This correlation did not appear to hold with the higher molecular weight noble gases. We then tried linear regression analysis with a number

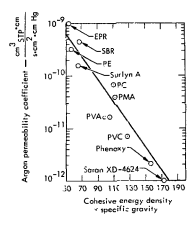


Fig. 4. Argon permeability coefficient vs cohesive energy density times specific gravity.

of polymer properties including CED, density, refractive index, gas molecular diameter (d), d^4 , etc. The results were not particularly promising. We observed, however, that there was a correlation of the argon permeability of a number of polymers with the oxygen Permachor as developed by Salame¹⁹ (see Fig. 5). This relationship was later extended to the other noble gases. Although Salame developed the Permachor equation empirically, he showed that there was a definite relationship to the classical relationship between permeability, diffusion, solubility, and temperature:

$$P = DS = (D_0 S_0)e^{-1}[(E_d + \Delta H)/RT]_{(see Ref. 8)}$$

where

P = perireability coefficient

D= diffusion coefficient

S = solubility coefficient

E, a energy of diffusion

 $\Delta H = heat$ of solution of the gas in the polymer.

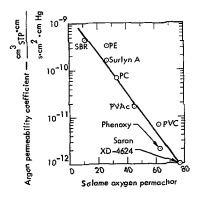


Fig. 5. Argon permeability coefficient vs Salame oxygen Pennachor.

 \mathbf{D}_0 and \mathbf{S}_0 are pre-exponential factors from the Arrhenius equations \mathbf{R} is the gas constant and \mathbf{T} is absolute temperature.

Salame's Permachor equation for the oxygen Permeability of polymers at 25°C is: $P \approx (6.1 \times 10^{-9})e^{-0.115 \text{ I}}$, where I is the oxygen Permachor. The Permachor equation can also be written

$$P = e^{-0.115} I e^{-(11200/RT)}$$

which can be written in the form

$$P = e^{A+BI}$$
 or $\ln P = A + BI$,

where Λ can be related to $(E_d + \Delta H)/RT$, and the coefficient B is determined by fitting experimental data.

We have studied the permeability of the noble gases, especially argon, krypton, and xenon, and have found that a relationship exists between the factors A and B and the square of the gas molecular diameters.

Apparently it is possible to extrapolate to radon permeability through the determination of these relationships.

Because the data were limited to Ar, Kr, and Xe, a linear relationship was used. A least-squares fit was made of the oxygen Permachor to the permeability data we had determined for phenoxy, polyvinyl chloride, polyvarbonate, polyvethy ene ionomer (Surlyn A), and Meares^{2,3} data for polyvinyl acetate. The coefficients A and B of the gases argon, krypton, and xenon thus obtained appeared to be linearly related to the d² of the gases. The least squares fit of this relationship resulted in the tinal equation

$$\ln (10^2 \text{P}) = (4.2354 + 0.4165 \text{ d}^2) + (0.088 - 0.0155 \text{ d}^2) \text{I},$$

where P = permeability coefficient in nm²/Pa*s $<math>d^2 = gas diameter squared, in Å^2$ I = Salame's oxygen Permachor.*

Calculated values, using this equation, are shown in Table 4 along with our experimentally determined values for a number of polymers and coatings. The fit is quite good in most cases, but is rather poor for polyethylene. Because the Permachor is related to the structure of the polymer, this equation establishes a definite relationship to polymer structure and can be used to estimate the permeability of radon as well as the other noble gases through polymers.

There are relatively few permeability data in the literature for the noble gases as the major interest is in nitrogen. Oxygen, carbon dioxide, and water as they may affect packaging of foods, etc. However, Burgess et al. 20 have measured the permeability of polymethyl acrylate to argon and krypton and reported values equivalent to 1.3 and 0.75 nm²/Pa-s at 20°C. The oxygen Permachor of polymethyl acrylate calculated from Salame's segmental values is 51. Using this value in the above equation, we calculate values for argon and krypton of 0.8 and 0.48 nm²/Pa-s, which agree to better than a factor of 2 with the experimentally determined values.

^{*}Salame used the symbol π (pi) for Permachor. We use I to avoid confusion with the conventional use of π .

Table 4. Permeability coefficients of polymers calculated from Permachor.²

Material	Permachor			Perme	ability, n	m²/Pa·s	s
	(1)		Ne	Ar	Kr	Xe	Rn
Phenoxy film	63	Experimental	1.9	0.20	0.08	0.016	_
		Calculated	2.0	0.27	0.12	0.02	0.0015
Polyvinyl chloride film,	62		_	0.53	0.26	0.03	_
unplasticized			2.1	0.29	0.14	0.02	0.0019
Polycarbonate film	33		_	4.1	3.8	1.5	1.4
			5.9	4.2	3.7	2.7	1.7
Surlyn A film	26		15	9.8	11	14	_
			7.6	7.8	8.1	8.5	9.1
Polyvinyl acetate (data of Meares)	45		15	1.1	0,36	_	_
			3.8	1.4	0.94	0.36	0.10
Epoxy, pigmented coating ^b	50		-	0.55	0.51	0.38	-
			3.2	88.0	0.53	0.16	0.03
Epoxy, unpigmented coating ^c	55		3.4	0.83	0.58	0.26	-
			2.7	0.56	0.30	0.07	0.001
Polyvinyl acetate, d plasticized	43		-	1.8	0.83	0.14	_
coating			4.1	1.7	1.2	0.50	0.16
Polyethylene film	26		_	25	34	60	113
			7	7.8	8.1	8.5	2.1
d^2 , A^2			8.0	11.6	13	16.4	20.9

 $^{^{4}}$ kn(10²P) = (4.2354+0.4166 d²) + (0.0880 - 0.0155 d²) 1

^bPermachor estimated and arbitrarily lowered for pigment content.

^cPermachor estimated.

^dPermachor arbitrarily lowered for plasticizer content.

Permeability of Uranium Ore

The permeability was measured in a sample of uranium ore supplied by USBM. The ore sample was retrieved from the Dakota Mine located 29 miles northwest of Grants, New Mexico. The exact location within the mine was 657 ft from the portal. The change in sample permeability with confining pressure up to 800 psi was also measured. This pressure corresponds to approximately 800 ft of overburden.

Sample preparation was fairly standard. Several cored specimens were machined to the desired dimensions and attached to end adaptors for tubing connections. Confining oil pressure was exerted on a sample in both the radial and axial directions. Oil pressure was transmitted without penetration through a flexible plastic jacket cast around each sample. Steady state, axial flow of dry N₂ gas through the sample provided the conditions required for permeability determination.

The results of the measurements are summarized in Table 5.

Based on these measurements, ore permeability in the unconfined state ranges from 0.25 to 0.46 darcys, decreasing at most 25% at 800 psi confining pressure. Since the samples were oven dried at 50 to 60°C prior to the measurements, the above values represent the upper limit of permeability, and would be greatly reduced with the presence of interstitial water. The permeability of rock is normally reported in darcys, but this can be converted into the units we have been using by introducing the appropriate gas viscosity value and the appropriate pressure correction. For argon gas, the viscosity is 222 x 10⁻⁶ poise (222 x 10⁻⁷ Pa·s). To convert darcys to [(cm³-cm)/(s·cm²-cm Hg)], divide darcys by about 1.7. To convert darcys to (nm²/Pa·s), multiply darcys by 4.4 x 10¹⁰. Thus, 0.25 darcy equals 0.15 [(cm³-cm)/(s·cm²-cm Hg)] or 1.1 x 10¹⁰ (nm²/Pa·s).

TASK 3: EVALUATION OF TOXICITY OF COATINGS

Coatings suitable for use in a mine should be non-toxic storing application, and the combustion and pyrolysis products should be of low toxicity. The volatile components of the coatings materials were identified qualitatively and semiqualitatively using gas chromatography and mass spectrography (GC/MS). The combustion and pyrolysis products were evaluated in two separate studies: (1) on cement-asbesto: board specimens in the LLL-NBS Smoke Chamber, and (2) small scale pyrolysis studies of unsupported films of the coatings. Finally, toxicological evaluations were made of the data from these studies from published toxicological information: no animal toxicity studies were made.

Table 5. Permeability of uranium ore samples at 0 to 800 psi confining pressure.

Confining pressure, psig		Permeability, darcys	
······································	Sample I	Sample 2	Sample 3
0	0.25	0.29	0.46
100	0.25	0.38	0.46
200	0.25	0.27	0.46
400	0.24	0.25	0.46
600	0.24	0.23	0.46
800	0.24	0.21	0.45

Smoke Chamber Studies of Coatings

Preparation of Specimens

The coatings were tested on cement-asbestos board substrates about 73 × 73 × 25 mm; the area exposed to heat and flame is about 65 X 65 mm. Thermocouples were imbedded at the center surface and at the lower surface, 6 mm from the lower edge. The lower thermocouple was impinged by the flame in the early tests (HydrEpoxy 300, Saran XD-7151, Essex polyester, and Ventron's Resitron II), but due to a slight modification of the smoke chamber, the thermocouple was slightly out of the flame area in the later tests. Some tests were also made on similar sized test specimens cut from uranium ore. There was no significant difference in the test results, and because use of uranium ore in the smoke chamber area required close monitoring by Hazard Control personnel (concern about release of alpha radiation), most tests were made on cement-asbestos board substrates only.

The coatings were prepared on the substrate usually by spraying, but sometimes by doctoring the coating material. It was usually necessary to use multiple coating to reach the desired coating thickness of 0.010 in. (0.25 mm). Although most coatings were tested at this thickness, the Resitron II coatings were 0.015 and 0.020 in. thick (0.38 and 0.51 mm), and the Hydro Seal coatings were only 0.007 and 0.008 in. thick (0.18 to 0.20 mm) and the Hydro Seal did not spread uniformly (did not wet) the previously dried coating.

Smoke Chamber Tests

This test studies the time dependence of toxic gas evolution under conditions designed to simulate a "fire" exposure. We used the LLL version of the NBS-developed Smoke Density Chamber. The procedure is given in NBS Technical Note 708 and was also desribed in the literature.²¹

In this test, the specimen is exposed to a radiant heat flux of 2.5 W/cm², (a) without the presence of an igniting flame (NF in tables and figures), and (b) in the presence of a small igniting flame (F in tables and figures). The radiant heat flux of 2.5 W/cm² is equivalent to that emitted by a black body at a temperature of 527°C (980°F). This is considered to be a moderately severe fire exposure. The chamber is continually

monitored for smoke density. CO. and CO₂ content. The LLL chamber is also fitted with a continuous sampling device to measure HC°C this was used in the case of the Saran XD-7151 coatings tests. The results, however, were lower than expected, based on the results of our small scale pyrolysis tests. We now believe these HC°C values from the smoke chamber tests to be invalid; they are low due to absorption on the chamber walls and sampling tubes, as well as incomplete recovery of HC°C from the chamber gas.

The data from the smoke chainber tests are listed in detail in Table A-2 and Figs. A-1 to A-15 in Appendix A. The results may be summarized as follows:

- HydrEpoxy 300, a water-based, pigmented epoxy coating. Under non-flaming conditions, the coating slowly produced a light smoke over a 30-min period, without any detectable production of CO or CO₂. Under flaming conditions, a light smoke was produced with an accompanying evolution of both CO (350 ppm) and CO₂ (1.4%). This amount of CO₂ is due to the pilot flame used under flame conditions. The quantities of these gases are similar to those found for other epoxy coatings previously tested in the same manner.
- 2. Saran XD-7151, a vinylidene chloride copolymer latex. Very little if any smoke was evolved from these coatings under either flaming or non-flaming conditions. No CO₂ was detected under non-flaming exposure. Under the flaming condition, the concentration of CO₂ reached about 1.4% in about 30 min, about the same as that for uncoated samples. We ascribe this to the combustion of the pilot flame. The Saran coating yielded 50 ppm CO in 25 min under a non-flaming condition and 300 to 500 ppm in 25 min under the flaming exposure. As indicated above, the HCV values are low.
- 3. Essex Polyester, a one-component (internally catalyzed, initiated by atmospheric moisture) styrenated polyester. This material slowly evolved a light smoke under the non-flaming exposure and a slightly denser smoke under the flaming exposure. Essentially no CO or

- CO_2 was evolved under non-flaming exposure. In the flaming case a slow evolution of CO was observed with a maximum of 70 ppin in 30 min. Likewise a small amount of CO_2 was slowly evolved up to a maximum of 1.5% at 30 min; most of this can be ascribed to the pilot flame.
- 4. Resitro., II, a catalyzed furfuryl alcohol polymer coating. This coating gave off no smoke under the non-flanning condition and a very light smoke under the flaming mode. Essentially no CO or CO₂ was evolved under non-flaming exposure. In the flaming mode, a slow evolution of CO was observed with a maximum of 100 ppm in 30 min. The CO₂ was evolved slowly up to a maximum of 1.5% at 30 min, most of which can be ascribed to the pilot flame.
- 5. WSU-118, a low viscosity modified epoxy developed for Bureau of Mines by Washington State University. The coating slowly evolved a dense smoke under non-flaming exposure; under flaming exposure a dense smoke was evolved moderately rapidly. No CO was detected under non-flaming exposure; under flaming exposure, a maximum of 300 ppm was evolved.
- 6. Aerospray 70, a polyvinyl acetate latex containing dibutyl phthalate plasticizer. Under both non-flaming and flaming exposure, the coating evolved a dense smoke moderately rapidly. No CO was detected under non-flaming exposure; under flaming exposure a maximum of 200 ppm was evolved.
- 7. EpiRez WD-510/EpiCure 872, a water-dispersed polyamine-cured epoxy coating. This was also evaluated cured with a blend of EpiCure 872 and 879; this cures harder and faster. Dense smoke evolved under both flaming and non-flaming conditions; evolution was slower under the non-flaming condition. No CO was detected under non-flaming condition; flaming exposure produced a maximum of 300 ppm.
- Promulsion 200, an unidentified pigmented emulsion. This slowly produced a moderately

- dense smoke under non-flaming exposure; under flaming exposure it produced a somewhat denser smoke more rapidly. In the latter case transient ignition and flame extinction occurred throughout the test. No CO was detected under the non-flaming condition and only a minimal amount, 25 ppm, was detected under flaming exposure.
- Hydro Seal, a water-based acrylic sealer, unpigmented. This slowly produced a moderate smoke in the non-flaming mode; in the flaming condition no ignition occurred, although a slightly denser smoke was produced, and a bit more rapidly. No CO was detected under either condition.

Smoke density curves, the specific optical density versus time, for several of the coatings are plotted in Fig. 6. Resitron II evolved a very light smoke and its specific optical density did not reach the valvo of 16, which is considered severe visual obscuration for a firefighter wearing a mask or other protective equipment. HydrEpoxy 300 produced a light smoke, with little difference on the transite substrate under flaming (F) or non-flaming (NF) conditions. Essex Polyester, also a light smoker, produced a slightly denser smoke more rapidly in the flaming mode. The EpiRez WD-510/EpiCure 872 coating evolved a dense smoke rapidly, especially in the flaming mode.

Flash Points of Coatings

Coatings materials for use in mines must have relatively high flash points to avoid danger of fire and explosion during application. The seven coatings selected for detail: A evaluation were tested for flash point by the closed cup method (ASTM D-56). These coatings were: Acme's HydrEpoxy 300, Dow's Saran XD-7151, Ventron's Resistron II, Essex's styrenated polyester, Bureau of Mine's WSU-118, Cyanamid's Aerospray 70, and Celanese's EpiRez WD-510/EpiCure 872. Except for Essex's styrenated polyester, none of the coatings systems had a measurable flash point under the conditions of the test, i.e. less than 79°C (175°F).

The Essex Polyester had a closed cup flash point (ASTM D-56) of 20°C (68°F), and an open cup flash

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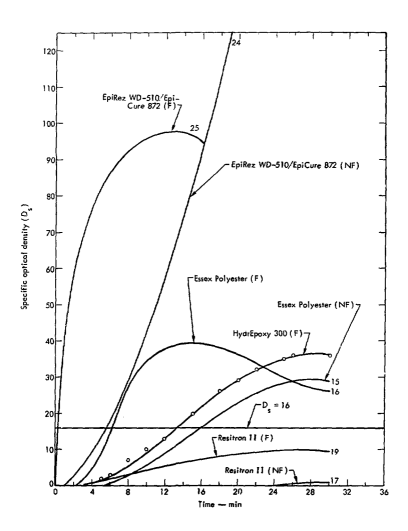


Fig 6. Smoke density curves for some representative coatings.

point (ASTM D-1310) of 39°C (102°F). This latter value is somewhat higher than the literature value²² for styrene of 31°C, but may be due to the fact that the material tends to "skin over" rapidly, so must be stirred occasionally during the test.

Laboratory Pyrolysis Test Under Non-Flaming Conditions

In this section the pyrolysis data for seven selected polymer systems are summarized. It should be noted that our objectives were to determine in a semi-qualitative way the extent of toxic gas evolution and to compare the results for the different polymer systems.

The gas concentration data were obtained from small-scale laboratory experiments under non-flaming conditions. Samples, weighing between 25 and 75 mg, were isothermally heated in closed quartz tubes with a free volume of about 0.13 litre. The samples were pyrolyzed in normal laboratory air at temperatures ranging from 200 to 600°C. Reaction times varied between 0.25 and 4 h. The longer reaction times were used at temperatures below 300°C to achieve a measurable degree of degradation. For each polymer system at least nine separate pyrolysis tests were made. In each test the experimental variables (time, temperature, and initial weight and/or film surface area of the sample) were varied to sample the infinite population of measurements.

At the completion of each pyrolysis experiment the gases were analyzed by mass spectroscopy (MS). For the Saran polymer two identical pyrolysis tests were done: one to obtain the gas analysis by MS: the second for the analysis of HCl. Hyd-ochloric acid was determined by dissolving the gaseous and absorbed HCl with dilute base followed by titration of the chloride ion. Selected gas mixtures, which were previously analyzed by MS, were also analyzed by gas chromatography (GC) to detect low level pyrolysis products. Standard gas mixtures containing CH₄, $\rm C_2H_6$, and CO₂ were used to calibrate the chromatograph. The CO concentration is accurately determined

by MS. Worst condition detection limit is about 0.3%. With a thermal conductivity detector the limit of detection for these gases was about 75 ppm. The results of the GC work (1) verified the presence and approximate concentration of the gases as measured by MS and, (2) showed that no other co-aponents were present at concentrations above 75 ppm.

Results and Discussion

This discussion will be mainly concerned with three topics: (1) HC@ production from Saran; /2) Comparison of polymer stability and gas evolution data; (3) Comparison of CO evolution data from laboratory and smoke chamber tests.

HCQ Production from Saran — It has previously been pointed out that the analytical system for determining HCQ during smoke chamber tests is unreliable. In these tests most of the HCQ appears to have been lost by adsorption on smoke particlen or by dissolution in water vapor. Thus, the HCQ quad from the small scale laboratory tests (Table 1) are the only reliable data that can be used to predict the evolution of this gas. These data indicate that the HCQ evolution is essentially time-dependent after 15 min and temperature-independent between 300 and 800°C. The best value appears to be the average of the 11 runs made under these conditions which is 0.26 \pm 0.07 cm $_{\rm STP}^3$ HCQ (gas) per mg of sample weight.

Comparison of Polymer Stability and Gas Evolution
Data — In Fig. 7 the percent weight loss is plotted as
a function of temperature for each polymer system.
The use of somewhat longer reaction times at temperatures below 300°C relative to the reaction times used
at the higher temperature distorts the ph. a somewhat.
In effect the low temperature degradation is weighted
more than the degradation at the higher temperature.
However, at each temperature the data are comparable. Figure 7 clearly shows that Ventron's Resitron
II, Essex Polyester, and HydrEpoxy 300 have significantly better thermal hability than the other polymer
systems.

The evolved CO and CO₂ gas volumes as function of pyrolysis temperature are presented in Figs. A-16

^{*}The tubes were filled with air at atmospheric pressure; the relative humidity was about 40-50%.

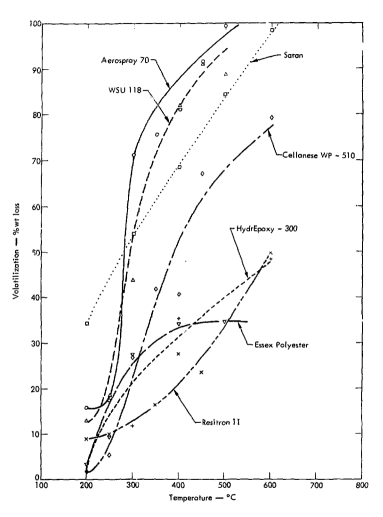


Fig. 7. Weight loss vs temperature.

and A-17, respectively. A generalized CO evolution is shown in Fig. 8. The CO evolution for most coatings falls within the band. The Essex Polvester and Resitron show much different behavior. The gas volumes have been normalized to the initial sample weight (Wo). Under isothermal conditions, and in the presence of oxygen, the effect of increasing the reaction time is to decrease the CO production rate. In the case of Saran, where the largest number of data points were obtained, it appears that the CO evolution rate decreases as t1/2. This is consistent with a rate controlled by a diffusion process. Our samples were films of nonuniform thickness, but generally of thickness larger than 10 mils. It is not unreasonable to postulate that the diffusion of oxygen into the film is the rate-controlling step, and that the reaction products (CO and CO2) must diffuse out of the film. Figures 7 and A-16 also indicate that the gas evolution has an exponential temperature dependence. These facts lead us to postulate the following reaction rate expression for relatively thick Saran films:

$$\frac{dC_{CO}}{dt} = Ke^{-\frac{1}{2}a/RT}\rho \frac{A}{L}t^{-\frac{1}{2}},$$

where K = Constant

E = The apparent activation energy

T = Temperature in K

A = Surface area of the film

L = Film thickness

R = Gas constant

 $\rho = \text{Film density}$.

Similar rate expressions can be fitted to the other polymer systems.

Of principal interest in this study is the evolution of the highly toxic gas, carbon monoxide. Other potentially harmful gases, such as hydrocarbons and hydrogen, were also produced but at significantly

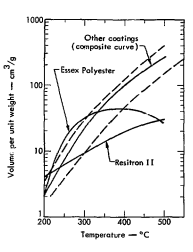


Fig. 8. Carbon monoxide evolution.

lower levels. Based on the slopes of the CO-evolution curves (Fig. A-16) in the 300 to 500°C temperature interval the seven polymers are ranked as follows:

Best: Ventron's Resitron

Essex Polvester

HydrEpoxy 300

Saran (considering the HCl data in addition

to the CO data)

Aerospray 70

WSU-118

EpiRez WD-510/EpiCure 872

These results are in general agreement with our conclusions obtained from the thermal stability data (Fig. 7).

For the degradation below 300°C this ranking may not be quite correct. Materials with low activation energies, such as Ventron's Resitron II and Essex Polyester, produce relatively more CO at these low temperatures than the other polymers.

^{*}Data obtained from runs above 500°C have been omitted from these graphs because in nearly all cases the oxygen depletion was near 100%. Under this condition the combustion mechanism must be expected to differ from the condition with high oxygen concentration.

Comparison of CO Gas Evolution Data from Laboratory and Smoke Chamber Tests — Smoke chamber tests were conducted under both flaming and non-flaming conditions. In the latter situation only the thermal flux from the heater produces the polymer degradation. This situation is comparable to the small-scale laboratory tests. In contrast to the laboratory tests, no CO was detected in the smoke chamber studies (non-flaming mode), except for Saran (~50 ppm).

In an attempt to predict the CO concentration for m the laboratory tests we make the following assumptions:

- a) The average temperature of the polymer film was either 300°C or 250°C for 0.5 h.
- b) The film was uniformly heated (no flame).
- c) The film thickness was uniform in both test situations and equal to 0.025 cm (10 mil).

These assumptions are expected to yield results that are upper limits for the predicted CO concentrations.*

The formula used to make these predictions is:

$${\rm CO}_{SC}(ppm) = 10^6 \ X \frac{V_L^{CO}}{V_{SC}} \, X \, \frac{A_{SC}}{A_L} \, X \, \frac{t_{SC}}{t_L} \, , \label{eq:cosc}$$

where

CO_{SC} = predicted CO concentration in the smoke chamber (SC)

V^{CO}_L = volume of CO (cm³_{STP}) measured in laboratory test (L) at 300°C or 250°C

A = film surface area

t = reaction time.

The laboratory data was taken from Tables A-2 through A-8: $A_{SC}=42~{\rm cm}^2\cdot V_{SC}\sim 5 \times 10^5~{\rm cm}^3$, $t_{SC}=0.5$ b. The predicted CO concentrations, based on the 250°C and 300°C laboratory results are summarized in Table 6.

The predicted CO concentrations are below the minimum detectable level of the CO analyzer on the smoke chamber (25 ppm).

Comparisons between laboratory tests and the flaming smoke chamber tests are probably not valid. The degradation mechanisms are expected to differ considerably as is evident from the observed CO concentrations (~300 ppm).

GC/MS Analysis of Volatiles from Polymers

This section summarizes the analysis of the head-space volatiles from the seven selected polymer systems. In the case of two-component systems, each component was analyzed separately. Analysis was done using a computer automated gas cirromatography/mass spectrometer (GC/MS) system. Initially, we attempted to identify the volatiles using only gas chromatography. This was not successful mainly because the elution times were not sufficiently reproducible to be matched against listed or suspected compounds.

We then used the computer-automated GC/MS system to identify most of the volatiles. The experimental conditions were as follows:

Columns: 1. 3 m X 3 mm o.d. glass, packed with Poropak Q, 80 to 100 mesh.

> 2. 2.8 m X 3 mm o.d. glass, packed with 10% OV-3 on 80 to 100 mesh Chromosorb WAD.

Carrier gas: Helium: flowrate 8 to 12 cm³/min. Temperature program: 65 to 220°C at 10 or 15°C/

min.

Split ratio: 70% to MS and 30% to the flame ionization detector (FID).

Injector temperature: 200°C. Separator temperature: 150°C.

Mass range: 12 to 250 automatic mass units (amu).

Integration time: 15 or 20 ms/amu.

Samples of selected liquid polymers and curing agents were stored in glass bottles and sealed with septum caps. The volatiles from the polymers were allowed to equilibrate with the normal air in the containers for at least several hours. The headspace was sampled through the septum cap with a 5 cm³ gas-tight syringe. The syringe and needle were heated to about 80°C and flushed three times with the headspace gas before withdrawing a sample. This procedure was adopted to minimize loss of material through adsorption on the internal

^{*}For example, thermocouple measurements indicate that the film surface reaches about 300°C after 0.5 h. For most of the time the film surface is at a lower temperature where the degradation is proceeding at a slower rate.

Table 6. Predicted maximum CO concentrations in the smoke chamber under non-flaming conditions.

Polymer	mer Reaction temp					
	300°C	250°C				
	CO (ppm)	CCO ppm				
Saran	11°	7				
HydrEpoxy-300	7	3				
Essex Polyester	32	5				
Resitron	9	12				
EpiCure	12	4				
Aerospray 70	11	4				
WSU-118	19	4				

^aThis value is predicted from the 2-h experiment. From the 0.5-h experiment we would predict 37 ppm; i.e., a value reasonably in agreement with the measured value.

syringe surfaces. The gas sample was then injected into the gas chromatograph. As each component was detected by the FID, the mass spectrum was obtained under computer control by automatically initiating the mass can near the top of the GC peak. Background spectra were manually taken between GC peaks. All samples were first analyzed on the Porapak Q column, which is suitable for the separation of relatively low molecular weight compounds (up to C-6 hydrocarbons). Subæquently we also used the OV-3 column to separate and detect compounds with molecular weights up to about 200 amu.

After we had tentatively identified some of the vapors by their characteristic mass spectrum we prepared calibration mixtures containing 1000 ppm (by volume) of these compounds in methanol. A 0.5-µ sample of this mixture was then injected to calibrate the FID response and to confirm the mass spectrometric identification. From the recorded GC peak area we calculated calibration factors that took into account the change in GC peak shape with increasing elution time. The calibration factors for both columns were averaged to estimate the concentrations of all identified compounds in the 5 cm³ headspace samples. The results are manarized in Table A-10. A typical mass spectrum, which was obtained during analysis of the headspace sample from Saran XD-7151 Latex, is

shown in Fig. 9. This compound was readily identified as vinylidene chloride.

Although we were informed that WSU-118H contained DMP-30 [tris(dimethylaminomethyl)phenol] and DETA (diethylenetriamine), these materials were not identified in the vapors.

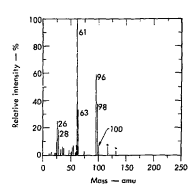


Fig. 9. Mass spectrum of vinylidene chloride from Saran latex (* = background peaks).

Toxicological Evaluation of Coatings

To evaluate the toxicity of the materials found by GC/MS in the study of the seven selected coatings, a table comparing the concentration found to the threshold limit value (TLV)* has been compiled (Table A-11 in Appendix A). The sample analyzed by GC/MS was collected from above the coating (or its components) in a closed container. Extrapolation of concentrations to a real mine situation is difficult. However, if one assumes these values are the maximum concentrations that can result from these coatings in a stagnant air situation, proper ventilation can be used to reduce them to a lower value where necessary. Most of the materials found in the coatings are innocuous with normal ventilation. However, because of the nature of one or more of the components of their curing vapors, the Saran XD-7151 latex and the WSU-118 coating must be handled carefully. The Saran coating liberates vinvlidene chloride (suspect carcinogen) and the WSU-118 liberates vinyl chloride (carcinogen) and epichlorohydrin (sensitizer).

The smoke and pyrofysis vapors appear to contain little of toxicological significance other than CO and HCC: the latter is present only on pyrofysis of the Saran latex coating. The amount of CO generated would probably be small compared to that evolved from the other fire components, i.e., burning wood, diesel oil, or other waste materials. Possible "worst condition" calculations have been made as follows:

Calculation conditions:

Mine ventilation rate = 100 ft/min.

Mine dimensions = 8 ft/8 ft

Film area liberating $CO = 32 \text{ ft}^2$.

Volume of air moved through CO source = 6400 ft³/min

CO liberation rate = $400 \times 550 \text{ ppm/ft}^2/\text{ft}^3$ in 30 min or $7320 \text{ ppm/ft}^2/\text{ft}^3$ in 1 min.

Calculation: $(32 \text{ ft}^2)(7320 \text{ ppm/ft}^2/\text{ft}^3)$ $(1/6400 \text{ ft}^3) = 36.6 \text{ ppm CO average}$ concentration/min.

This calculation was made for the Saran coating under flaming conditions and under very poor ventilation conditions. The Saran coating gave the highest CO concentration in the smoke chamber. Under these conditions the calculated CO concentration is clust to the TLV for CO which is 50 ppm. Under more normal ventilation conditions, the CO concentration would be much lower.

HCl concentration in mine shaft:

Using the data in Table A-4 and assuming: (1) a 10-mil (0.254 mm) coating of the Saran XD-7151, (2) an 8×8 ft mine cross section. (3) a minimum air flow of 250 ft/min, and (4) that the coating teaches 400° C with adequate oxygen, we calculate that the pyrolysis of about 1/3 11° per min of the coating would exceed the 10 ppm level. Even 5 ppm of HCl is highly irritating and may be the maximum allowable working level.

The second second second second second second

CO concentration in mine shaft:

^{*}Threshold limit values refer to airborne concentrations of substances and represent conditions under which it is believed that nearly all workers may be repeatedly exposed day after day without adverse effect. Because of wide variation in individual susceptibility, however, a small percentage of workers may experience disconfort from some substances at concentrations at or Low the threshold limit, a smaller percentage may be affected more seriously by aggravation of pre-existing conditions or by development of an occupational illness.

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Appendix A: Tables and Figures

cmc, HydrEpoxy 300 Mod 1 cme, HydrEpoxy 300 Mod 2 cme, HydrEpoxy 300 Mod 2 cme, HydrEpoxy 300 Mod 2 cme, HydrEpoxy 156/300° cme, HydrEpoxy 156/300	r, Name Type						
			cm ³ _{STP} · cm s · cm ² · cm Hg	nm² Pa•s			
Acme, HydrEpoxy 300	Pigmented, water-dispersed Epoxy	Ar	6.7 X 10 ^{-1 1}	5.0			
Acme, HydrEpoxy 300 Mod !	Pigmented, water-dispersed Epoxy	Ar	1.4×10^{-11}	1.1			
Acme, HydrEpoxy 300 Mod 2	Pigmented, water-dispersed Epoxy	Αr	7.3×10^{-12}	0.55			
Acme, HydrEpoxy 300 Mod 2	Pigmented, water-dispersed Epoxy	K,	6.8 X 10 ⁻¹²	0.51			
Acme, HydrEpoxy 300 Mod 2	Pigmented, water-dispersed Epoxy	Χe	5.0 × 10 ⁻¹²	0.38			
		(R _n) ^b	(4 × 10 ⁻¹²)	(0.3)			
Acme, HydrEpoxy 156/300 ^c	Pigmented, water-dispersed Epoxy	٨r	1.7 × 10 ⁻¹¹	1.3			
Acme, HydrEpoxy 156/300	Pigmented, water-dispersed Epoxy	Κr	1.4 × 10 ⁻¹¹	1.1			
Acme, HydrEpoxy 156/300	Pigmented, water-dispersed Epoxy	Хe	1.4 × 10 ^{-1.1}	1.1			
		(Rn)	(1.3×10^{-11})	(1)			
Am. Cyanamid, Aerospray 52	Alkyd resin dispersion	Ar	1.1 × 10 ^{-1 1}	0.83			
Am. Cyanamid, Aerospray 70	Plasticized polyvinyl acetate emulsion	Ar	2.4 × 10 ⁻¹¹	1.8			
Am. Cyanamid, Aerospray 70	Plasticized polyvinyl acetate emulsion	Κr	1.1×10^{-11}	0.83			
Am. Cyanamid, Aerospray 70	Plasticized polyvinyl acetate emulsion	Χe	1.9 × 10 ⁻¹²	0.14			
		(R _{n)}	(4×10^{-13})	(0.03)			
Celanese, EpiRez WD-\$10/EpiCure 879/EpiCure 872	Non-pigmented, water-dispersed epoxy	Ar	8.7 × 10 ^{-1 2}	0.65			
Celanese, EpiRez WE-3520/EpiCure BCT-60-8533	Non-pigmental, water-dispersed epoxy	Ne	4.5 × 10 ⁻¹¹	3.4			
Celanese, EpiRez WE-3520/EpiCure BCT-60-8533	Non-pigmented, water-dispersed epoxy	Ar	1.1 × 10 ⁻¹	0.83			
Celanese, EpiRez WE-3520/EpiCure BCT-60-8533	Non-pigmented, water-dispersed epoxy	Κr	7.7 × 10 ⁻¹²	0.58			
Celanese, EpiRcz WE-3520/EpiCure BCT-60-8533	Non-pigmented, water-dispersed epoxy	Хe	3.5×10^{-12}	0.26			
	•	(Rn)	1.1×10^{-12}	0.083			
Dow, Saran XD-4624	Vinylidene chloride copulymer latex	Ar	4.4 × 10 ⁻¹³	0.033			
Dow, Saran XD-4624	Vinylidene chloride copolymer latex	Kr	2.2×10^{-13}	0.017			
Dow, Saran XD-4624	Vinylidene chloride copolymer latex	Xe	8.9 × 10 ⁻¹⁴	0.0067			
	•	(Rn)	1.8×10^{-14}	0.0014			

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Table A-1. Permeation coefficients of supported coatings (continued).

Maker, Name	Туре	Gas	Permention coefficient of coating		
			em ³ _{STP} · cm s · cm ² · cm Hg	nm² Pa - s	
Dow, Saran XD-7151	Vinylidene chloride copolymer latex	Ar	2.0 × 10 ⁻¹³	0.015	
Dow, Saran XD-7828	Vinylidene chloride copolymer latex	Ar	2.7×10^{-13}	0.020	
Dowell, M159/M171	Vinylidene chloride copolymer latex plus thickener	Ar	3.9×10^{-13}	0.029	
Dowell, M175/M171	Vinylidene chloride copolymer latex plus thickener	Ar	7.6 × 10 ⁻¹³	0.057	
Essex, moisture curing unsaturated polyester	Pigmented styrenated polyester	Ar	1.7 × 10 ⁻¹⁰	13.	
Goodrich, Geon 660X1	Vinylidene chloride copolymer latex	Ar	4.9×10^{-12}	0.37	
Goodrich, Geon 652	Vinylidene chloride copolymer latex	Ar	$1.7 \times 10^{-1.2}$	0.13	
Monsanto, Monflex 4500	Vinyl chloride copolymer latex	Ar	8.0 × 10 ⁻¹²	0.60	
Monsanto, Monslex 4514	Vinyl chloride copolymer latex	Ar	6.0×10^{-12}	0.45	
Monsanto/Rohm and Haas, Monflex 4500/Rhoplex HA-20	Vinyl chloride copolymer latex/vinylidene chloride- acrylic copolymer latex	Ar	4.2 × 10 ⁻¹²	0.32	
Pan. Am. R&D, GPC-500	Acrylic solution	Ar	8.5 × 10 ⁻¹¹	6.4	
Preserv-O-Paint, CP-5005	Epoxyized polyester urethane	Ar	2.9 × 10 ⁻¹¹	2.2	
Rohm and Haus, Acryloid C-10 LV	Polymethyl acrylate solution	Ar	3.9 × 10 ⁻¹¹	2.9	
Quaker, Quaker Coat		Ar	3.5 × 10.11	2.6	
Union Carhide, 9484-153-100	Acrylonitrile copolymer latex	Aı	1.0 × 10 ⁻¹⁰	7.5	
Ventron, Resitron II	Furan polymer	Ar	1.5×10^{-13}	0.011	

⁴ All coatings were applied on polycthylene ionomer, Surlyn A, about 0.05 mm, except Resitron II which was an unsupported film. Coefficients have been calculated for coating alone.

b Radon coefficients are estimated values obtained by extrapolating values of Ar, Kr, and Xe coefficients vs gas diameter squared.

^c HydrEpoxy 300 (0.0025 mm) on top of HydrEpoxy 156 (0.0018 mm) on Surlyn A.

^{*}NF = non flaming; F = flaming; NR = not reached.

CO in ppm; CO2 in %. CO2 meter not functioning after GH-19. Lowest detectable funts "O2 25 ppm, CO2 04%

Obscuration time = time to reach specific optical density of 16 (severe visual obscuration for life lighter wearing mask)

Table A-3. Pyrolysis study of HydrEpoxy 300.

	Sample wt, nig	Residue wt, mg	Wt loss, mg	Loss wt, %	Surface area, mm ²	Pyro temp, °C					Gas evolved. cm _{STP}				dved/sample st. cm _{STP} g		
_			_					со	CO ₂	CH ₄	C2H6	co	co,	CH4	C2116		
	25.244	23.909	1.335	5.3	166	200	16.0	0.18	0.38	~0	∿ 0	7	15	~0	\sim o	4.3	
	25.476	23.742	1.734	6.8	133	200	29.5	0.50	0.53	~0	~ 0	20	21	\sim 0	~ 0	7.8	
	25.348	22.337	3.011	11.9	139	300	2.0	0.42	1.08	~ 0	~0	17	43	~0	~0	10.4	
	24.633	20.150	4.483	18.2	133	300	16.0	1.13	2.28	0.02	∿0	44	90	4	∨ 0	37.2	
	25.400	16.360	9.040	35.6	132	400	2.0	2.29	6.24	0.02	∿ 0	90	246	4	~0	41.1	
	25.354	18.234	7.120	28.	95	400	0.5	0.96	1.86	0.04	0.01	38	7,3	2	1	18.4	
	25.074	12.880	12.194	48.6	99	600	0.5	~0	11.6	~0	~0	~0	402	~0	∿0	53.6	
	25.110	11.965	13.145	52.4	96	800	0.5	~0	12.7	~0	~0	~0	506	~ 0	~0	57.9	

Table A-4. Pyrolysis study of Saran XD-7151 film (95% Saran plus 5% polyvinyl alcohol solution).

	Sample wt, mg	Residue wt, mg	Wt loss,	Loss	Surface area, mm ²	Pyro temp, °C	Pyro time, h	G	as evolve	ed. cm ³ STP		Gas evo	lved/sam	ple st, cm	STP/g	O ₂ loss,
	* -							co	CO ₂	HCk	CH ₄	0	CO2	HCf	CH ₄	
:	50.448	48.380	2.068	4.1	106	150	3.0	~0	0.07	0.57	~0	~0	1	11	~0	5.6
4	49.232	32.270	16.962	34.5	104	200	2.0	0.13	0.11	7.14	~0	3	2	15	~0	~0
:	25.501	11.883	13.618	53.4	55	300	0.25	0.24	0.24	5.26	0.01	10	10	21	1	0.2
:	25.074	11.205	13.869	55.3	56	300	0.5	0.35	0.50	5.05	0.04	14	20	202	2	1.5
:	24.924	12,175	12.749	51.2	57	300	1.0	0.44	0.47	5.43	~0	18	19	215	~0	1.5
:	25.143	11.538	13.605	54.1	62	300	2.0	0.32	0.50	5.85	~0	13	20	233	~0	3.1
:	24.550	9.550	15.000	61.1	56	400	0.25	0.76	0.80	7.45	~0	31	33	304	~0	0.8
:	24.410	10.195	14.215	58.2	47	400	0.5	0,95	1.43	8.28	~0	39	58	330	~0	8.6
:	25.352	9.450	14.902	58.9	56	400	1.0	1.09	1.66	6.57	0.03	43	66	259	1	8.6
:	25.290	8.120	17.170	67.9	55	400	2.0	1.42	3.08	5.45	~0	56	122	215	~0	16.9
:	25.053	7.385	17.668	70.5	48	400	4.0	1.92	3.19	7.97	G.04	77	127	318	2	20.9
:	24.3	8.819	15.481	63.7	74	400	0.25	1.55	3.65		80.0	64	150		3	20.5
	74.9	23.474	51.426	68.7	148	400	0.75	7.35	7.95		0.59	98	106		8	45.9
	24.4	.343	24.06	98.6	86	600	0.5	9.55	6.80		0.04	391	279		2	46.9
:	23.153	.336	22.817	98.5	83	600	0.5	8.55	6.15		~0	382	266		~0	64.5
	49.69	~0	49.69	100		600	0.5	18,85	8.70	(10.9)	0.06	379	175	219	1	76.0
	23.1	~o	23.1	100	82	800	0.75	0.62	14.6		~0	27	632		~0	39.5
	22.2	~ 0	22.2	100	82	800	0.25	3,14	10.9	(6.7)	~0	141	491	302	~0	46.3
	75.1	13.77	61.32	81.7		800	0.25	13,14	13.4	(17.35)	~0	178	178	231	~0	81.0

Table A-5. Pyrolysis study of Essex Polyester.

Sample wt, mg	Residue wt, mg	Wt loss, mg	Loss wt, %	Surface area, mm ²	Pyro temp, °C	Pyro time, h	Gas evolved, cm ³ _{STP}				Gas evolved/sample st, cm ³ _{STP} /g				O2 loss
							co	co,	CH ₄	H ₂	co	CO2	CH4	H ₂	
49.793	48.069	1.724	3.5	117	200	4.0	~0	1.05	~0	~0	~0	21	~0	~0	10.5
25.631	23.109	2.522	9.8	42	250	4.5	0.24	0.65	~0	0.04	9	25	~0	2	2.9
75.200	62.600	12.600	16.8	105	300	1.0	1.42	3,49	0.03	0.08	19	46	1	1	20,4
50.654	36.710	13.944	27.5	104	300	2.0	1.57	4.81	0.04	0.09	31	95	ı	2	23.5
25.583	17.641	7.942	31.0	56	350	2.6	1.09	3.10	~0	0.07	43	121	~0	3	13,6
49.292	32.340	16.952	34.3	104	400	1.0	2.13	10.58	0.04	0.20	43	215	1	4	45.3
49.322	32.631	16.691	33.8	108	450	1.5	~0	15.52	0.08	0.25	_	315	2	5	60.2
25.678	16.792	8.886	34.6	52	500	0.5	0.71	7.95	0.13	0.17	28	310	5	7	32.4
25.070	16.402	8.668	34.6	56	500	1.5	0.52	8.33	0.01	~0	21	332	1	~0	34.8
74,828	48.971	25.857	34.6	105	600	0.5	7.06	19.39	0.72	1.63	94	259	to	22	98.3

Table A-6. Pyrolysis study of Ventron's Resitron II.

	imple t, mg	Residue wt, mg	Wt loss,	Loss wt,%	Surface area, mm ²	Pyro temp, °C	Pyro time, h	C	ias evolve	d, cm ³		Gas evo	lved/sam	ple st, cm	STP/g	O ₂ loss,
								co	CO ₂	CH ₄	C ₂ H ₆	co	CO ₂	CH ₄	C ₂ H ₆	
40	0.500	45.364	4.136	8.4	127	150	4.0	~0	0.20	∿ 0	~0	~0	4	~0	~0	1.4
75	5.222	72.712	2.510	3.3	165	150	6.0	0.32	0.30	\sim 0	~0	4	4	~0	~ 0	2.5
50	0.416	45.892	4.524	9.0	138	200	2.0	0.18	0.43	~0	~0	4	9	~0	~0	4.6
7-	4.557 ·	67.284	7.273	9.8	150	250	1.0	0.42	1.74	~0	~0	6	23	~0	~0	5.4
50	0.359	44.900	5.459	10.8	127	300	1.0	0.27	1.38	0.01	~0	5	27	~0	~0	4.3
2:	5.506	21.300	4.206	16.5	56	350	1.0	0.38	0.83	0.05	~0	15	32	2	~0	2.6
	5.314	18.312	7.002	27,7	25	400	0.5	0.45	1.23	0.07	~0	18	49	3	~0	6.4
7:	5.178	57.403	17.775	23.6	138	450	0.25	1.56	3.87	0.44	0.08	21	51	6	ı	14.2
2	5.165	12.650	12.515	49.7	27	600	0.5	0.85	14.61	~0	~0	34	581	~0	~0	74.2

Table A-7. Pyrolysis study of Aerospray 70.

	Sample wt. mg	Residue wt. mg	Wt loss.	Loss wt, %	Surface area, mm ²	Pyro temp. °C	Pyro time, h	C	ias evolve	d. cin <mark>3</mark>		Gas evo	lved/sam	ple st. em	STP g	O ₂ loss.
								(0	CO ₂	CH ₄	H ₂	co	CO ₂	CH ⁴	H ₂	
	75.602	63.423	12.179	16.1	195	200	4.0	0 14	0.52	\sim 0	0.06	2	7	\sim 0	1	2.7
	75.043	62.042	13.001	17.3	163	250	3.0	.50	1.25	0.04	~0	7	17	1	~0	10.8
	51.001	14.709	36.392	71.4	144	300	3.0	1.12	2.58	.18	~0	22	50	4	~0	20.0
	50.326	12.197	38.129	75.8	98	350	2.0	1.97	4.09	0.10	0.07	39	80	2	1	28.3
	50.550	9.554	40.996	81.1	154	400	3.0	4.52	13.01	0.28	0.16	90	257	6	3	88.5
ž.	74.905	6.953	67.952	90.7	175	450	1.75	8.36	17.39	0.84	0.35	1!2	230	10	5	99.1
33	25.673	0.155	25.518	99.4	55	500	1.0	7.01	14.30	0.74	0.36	273	557	29	14	71.8
	25.452	0.086	25.366	99.7	42	550	1.0	6.53	13.52	0.61	0.30	256	531	24	12	83.9
	25.494	0.412	25.082	98.4	46	600	0.5	4.80	20.96	0.09	0.10	188	822	4	4	88.0

Lable A-S. Pyrolysis study of EpiRez WD-510/EpiCure 872

Sample wt. mg	Residue w.t. mg	Withous, mg	-	Surface area, mm ²	Pyro ichip. C	Pyrec tame, h	J	Gas evolved, cm _{S1P}	L. cm31p		Gas evolved sample st. cm _{S1p} # = 0, hoss.	dues per	le st. cm	SIP #	0, 1055
		1					8	6,1	≣"	= 1	8 1	ę'	E	<i>≡</i> ′	
74,584	13,091	1.473	2.0	ž	700	0.4	0.18	0.30	90:0	٥	т	7	-	20	8:
75.405	71.345	4.06	5.4	ôh	250	3.0	0.27	0.79	90.0	2	4	10	-	Ş	8.3
49,791	36.452	13,339	26.8	ī	300	3.0	0.71	1.43	90'0	٥٧	7	61	C1	2	14.9
51.707	29.963	21.744	77	7	350	2.0	31:18	3.79	0.24	0.06	52	06	9	-	50.6
50.534	30.0	20.534	40.6	27	007	2.5	5.78	8.94	0.24	0.04	11	177	\$	-	73.9
24.715	8.1	16.615	67.3	5	450	5.	8.54	9.17	0.31	0.03	346	371	13	-	61.2
25.314	0.4	21.314	84.2	7.2	200	1.0	10.22	10.04	0.47	0.22	104	397	19	6	72.7
75.326	35.4	39.926	53.0	55	550	1.0	17.36	7.73	4.53	15.0	131	103	09	7	99.1
25.700	5.3	20.40	79.4	20	009	0.5	5.27	19.63	90.0	0.22	205	764	c)	6	4,66

Table A.9. Pyrolysis study of WSU 118 (5.3 parts R, 1.0 parts H).

Sample wt, mg	Residue wt. mg	Wt loss.	Loss wt, 7	Surface area, mm²	Pyro temp. °t	Pyro time. h	ĕ	Gas evolved, cm _{STP}	I. ст ³ т		Gas evol	ved/sam	ple st, cm	STP/B	Gas evolved/sample st, cm ³ _{TP} /g O ₂ loss,
							e	ုင်	ੂ	£	e	8	₽	±21	
75.412	65.578	9.894	13.1	63	200	4	0.17	0.26	20	20	2	ю	20	Ş	6.9
75.340	61.744	13.596	18.05	95	250	~:	0.28	0.39	c'	2	4	S	20	5	7.7
50.379	28.262	22.117	43.9	92	300	٣	0.97	2.12	90.0	20	61	43	-	Ş	37.6
50.632	{	;	1	30	350	2	17.7	6.22	0.13	5	54	123	æ	5	57.7
49.492	8.989	40.503	81.8	48	400	2.5	6.23	11.28	0.17	0.07	126	228	3	-	80.7
25.668	2.079	23.589	616	22	420	1.5	5.74	10.21	0.13	0.08	224	398	\$	٣	71.9
25.012	2.686	22.326	80.3	23	200	1.25	7.33	12.69	0.23	0.13	293	507	6	-	83.9
74.927	10.170	64.757	86.4	\$\$	880	-	15.44	8.71	3.46	0.64	206	116	46	æ	98.6
25.805	2.056	23.749	92.0	45	009	0.5	7.76	15.76	0.11	0.33	301	119	4	13	99.4

Table A-10. TLV of organic materials released during curing of several radon barrier coatings.

Coating	Compound	Volume found, ppm	TLV, ppm	Personal guess, ppm
EpiRez WD-510	Acetone	25-49	1000	
	MEK	10-24	200	
	Batanol	10-24	100	
	Toluene	10-24	100	
	Xylene	0-9	100	
	2-Ethoxy ethylacetate	10-24	100	
	2-Methyl butanol	100-199		100
EpiCure 872	Ethylene	10-24	1000	
•	Propane	0-9	1000	
	Vinyl Acetate	10-24	10	
	Toluene	10-24	100	
	Xylene	10-24	100	
	Ethyl benzene	200-399	100	
Resitron II	Methanol	0-9	200	
(Resin)	2-Methyl furan	0-9		10
(Vinyl acetate	0.9	10	
	MEK	0-9	200	
	Acetone	25-49	1000	
	Furfural	25-49	5	
	Furfuryl alcohol	200-399	5	
Aerospray 70	Acetaldehyde	0.9	100	
	Ethanol	0-9	1000	
	Vinyl acetate	200-399	10	
	Acetone	400-699	1000	
HydrEpoxy 300	Ethylene	25-49	1000	
(A component)	Acetone	10-24	1000	
(// component)	Vinyl acetate	25-49	10	
HydrEpoxy 300	Acetone	0-9	1000	
(B component)	Vinyl acetate	0-9	10	
Saran XD-7151	2-Methyl propene	0-9		200
Latex	Butanol	0-9	100	
	2-Methyl-2-propanol	25-49	100	
	Butyl acrylate	50-99		10-100
	Vinylidene chloride	100-199		1-10
	Acetone	200-399	1000	
	Acrylonitrile	20-399	20	

Table A-10. TLV of organic materials released during curing of several radon barrier coatings (continued).

Coating	Compound	Volume found, ppm	TLV, ppm	Personal guess, ppm
Essex Polyester	MEK	25-49	200	
(fresh ^a)	Styrene	200-399	100	
•	Toluene	400-699	100	
	Acetone	10,000 +	1000	
Essex Polyester	Ethylene chloride	10-24	50	
(after 48 hb)	MEK	25-49	200	
	Styrene	700-999	100	
	Acetone	> 20,000	1000	
Essex Polyester	Toluene	10-24	100	
(after curing	Propyl heptanol	200-399		10-100
total gas)	Ethyl benzene	400-699	100	
	Xylene	700-999	100	
	Benzene	1000-2499	25	
	Isopropyl benzene	1000-2499	50	
	Acetone	2500-10,000	0001	
Essex Polyester	Benzene	10-24	25	
(vapor over cured	Benzaldelhyde	50- 9 9		10-100
material)	Acetone	200-399	1000	
WSU-118	Vinyl chloride	25-49	1	
(R component ^a)	Ethylene chloride	50.99	50	
	Epichlorohydrin	50.99	5	
	Acetone	100-199	1000	
WSU-118	Trimethylamine	2 00 -399		10
(H component ^a)	Toluene	400-699	100	
	Dimethylamine	1000-2499	10	
	A ^c			?
	$\mathbf{B}^{\mathbf{c}}$?

^a Poropak Q column only: To check styrene content in fresh material.

b Stabilized styrene: different elution time.

c Unresolved compound: excludes, DMP-30, DETA.

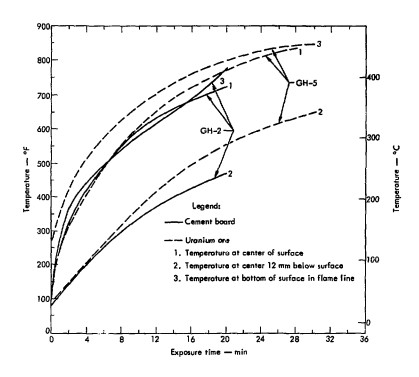


Fig. A-1. Surface and body temperatues of uncoated cement-board and uranium-ore squares exposed to 2.5 W/ cm² radiant heat with pilot flame.

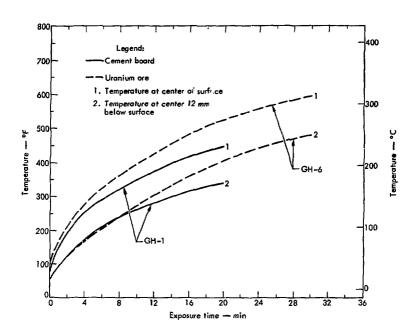


Fig. A-2. Surface and body temperatures of uncoated cement-board and uranium-ore squares exposed to 2.5 W/cm² radiant heat, without pilot flame.

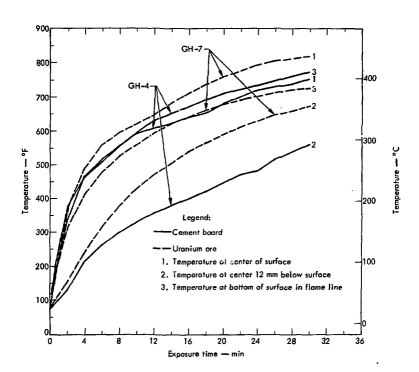


Fig. A-3. Surface and body temperature of cement-board and uranium-ore squares whose front surface had been coated with 10 mils of HydrEpoxy 300. The squares were exposed to 2.5 W/cm² radiant heat with pilot flame.

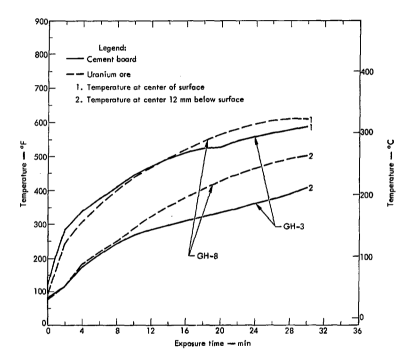


Fig. A-4. Surface and body temperatures of cement-board and uranium-ore squares whose front surface had been coated with 10 mils of HydrEpoxy 300. The squares were exposed to 2.5 W/cm² radiant heat, without pilot flame.

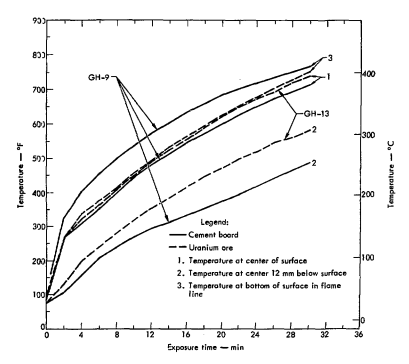


Fig. A-5. Surface and body temperature of coment-board and uranium-ore squares whose front surface was coated with 10 mils of Saran XD-7151. The squares were exposed to 2.5 W/cm² radiant heat with pilot flame.

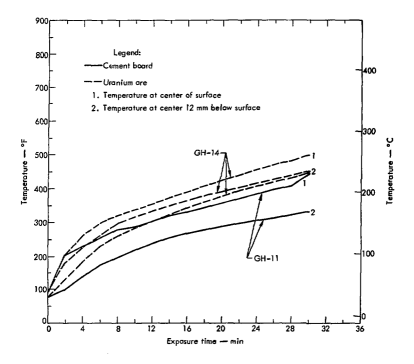


Fig. A-6. Surface and body temperature of cement-board and uranium-ore squares whose front surface was coated with Saran XD-7151. The squares were exposed to 2.5 W/cm² radiant heat, without pilot flame.

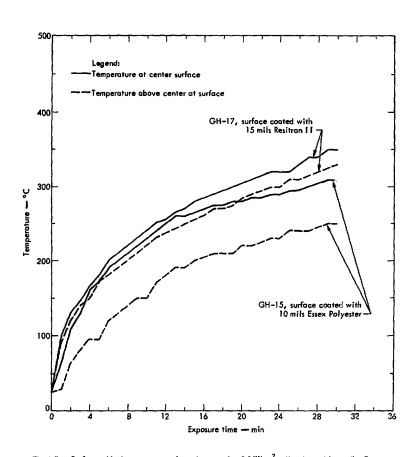


Fig. A-7. Surface and body temperature of sample exposed to 2.5 W/cm² radiant heat without pilot flame.

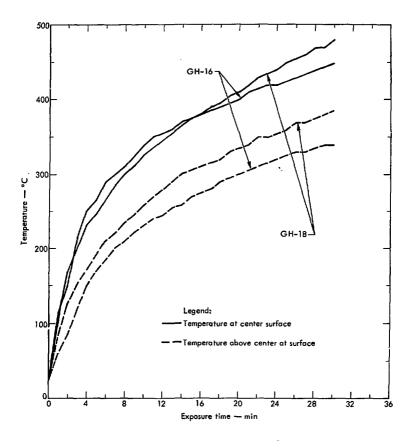


Fig. A-8. Surface and body temperature of sample exposed to 2.5 W/cm² radiant heat with pilot flame.

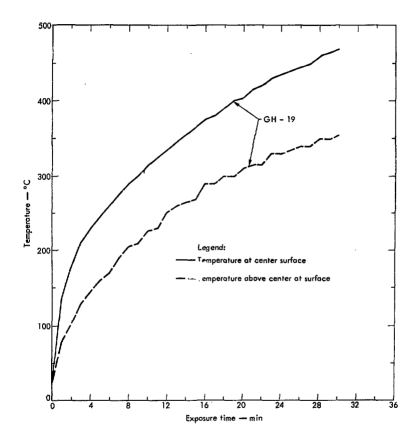


Fig. A.9. Surface and body temperature of sample GH-19, surface coated with 20 mils Resitron II and exposed to 2.5 W/cm² radiant heat with pilot flame.

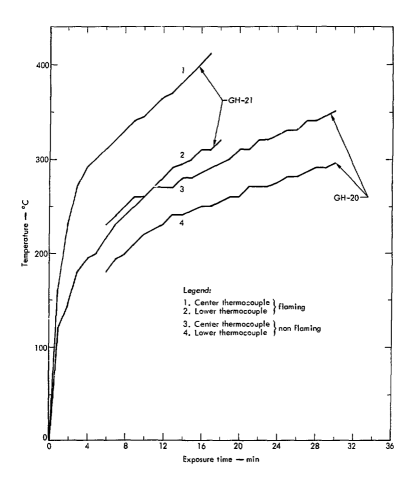


Fig. A-10. Surface and body temperature of a sample whose surface was coated with WSU-118 and exposed to 2.5 W/cm² radiant heat with and without flame.

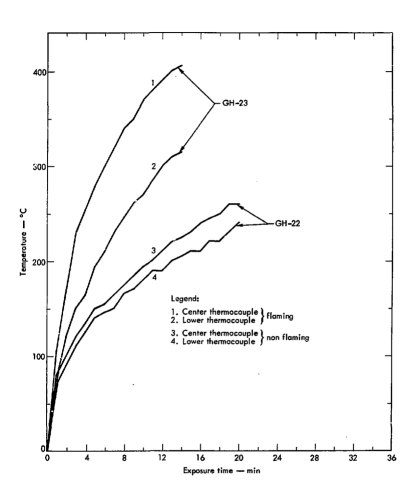


Fig. A-11. Temperatures recorded at the center and lower thermocouple of a sample coated with Aerospray 70 and exposed to 2.5 W/cm² of radiant heat with and without flame.

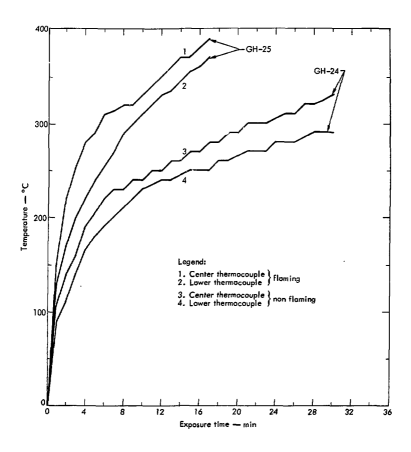


Fig. A-12. Temperatures recorded at the center and lower thermocouple of a sample coated with EpiRez WD-510/ EpiCure 872 and exposed to 2.5 W/cm² of radiant heat with and without flame.

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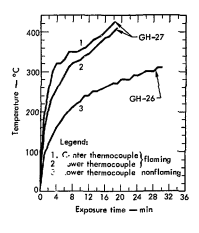


Fig. A-13. Temperatures recorded at the center and lower thermocouple of a sample coated with EpiRez WD-510/EpiCure 872 and 879. The sample was exposed to 2.5 W/cm² of radiant heat with and without flame.

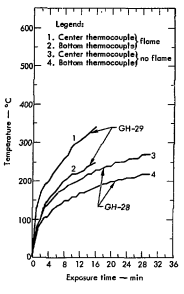


Fig. A-14. Temperatures at the center and bottom of samples coated with Promulsion 200.

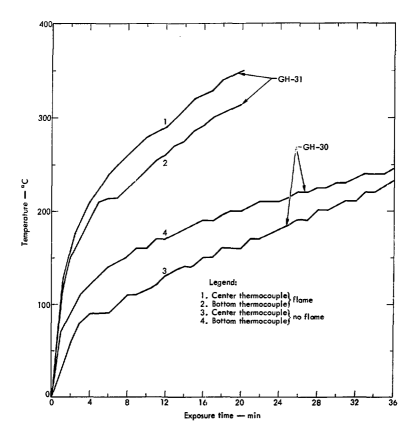


Fig. A-15. Temperatures at the center and bottom of samples coated with Hydro Seal.

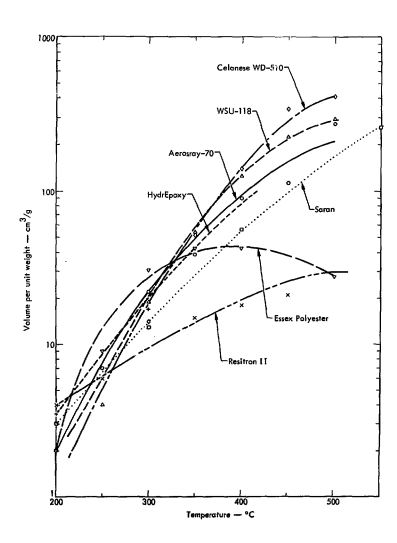


Fig. A-16. Carbon monoxide evolution.

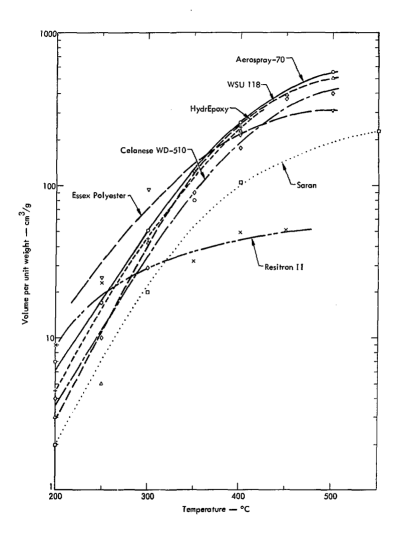


Fig. A-17. Carbon dioxide evolution.

Appendix B:

Computer Program (PDP-8) for Calculation of Permeability Coefficients from Dow Cell Data

```
.R PEAN
700.00
*!; A
C POLYMERIC MATERIALS FOCAL - 7/15/73
01.01 T "PERMEAPILITY PROCESM.".!
01.02 C FILE PERM.50 1/9/74
01.03 T "ENIER UNITS PER ASIM-L-1434-66 METHOD M.".!!
01.05 C PROGRAM TO COMPUTE WAS THANSMISSION THROUGH THIN FILMS.
01.07 CHANGE DATE 8-17-73
01.10 $ A=1.872
61.20 S AT=63.62
01.30 A "H (MM) = ",H
01.40 S HR=65.5
01.50 S HL=65.5
01.60 A "P (MMHG) = ",P
Ø1.70 5 T=296
01.80 A "UF (MM13) = ".VF
01.90 A "RTC (MMHG/IN:-1) = ",RC
01.95 A "RTP (IN/H) = ", HP
01.96 S T1=0
02.10 \text{ A "SLP (IN(MMHG)/IN(H))} = \text{".SP}
02.20 A "T1 (THICKNESS OF CARRIER - MIL) = ",T1
02.30 A "T3 (1HICKNESS OF SAMPLE - MIL) = ",T3
02.40 I (T1) 2.5,3.1,2.5
02.50 A "PC (PERM. OF CAREIER) = ".FC
03.10 S FT=HL-H;5 DH=(RC+hF)/SF
03.20 S GH=(4.09*10*12/((F-PT)*AT))*DH
03.30 S GH=GR*(((2*A*H)-(A*(HL+HF))-VF)/(6.23*10*7)*1)
03.40 S FM=GR*T3*(3.87/10:14)
03.50 I (T1) 3.6,4.1,3.6
03.60 S 12=(T3-T1)/10:3;5 14=T1/10:3;5 15=T3/10:3
03.70 S PL=(T2*PM*PC)/((15*PC)=(T4*PM))
04.10 T !."CTR = ". %6.06.GR.!
04.20 T "PERM OF SAMPLE = ", %, PM, %6.06, !
04.30 IF (T1) 4.4,4.9,4.4
04.40 T "PERM OF COATING = ", %, PL, %6.06, !
04.90 T !!:S CT=10:C 10.1
```

```
10.10 A "CHANGE - ", CH
10.20 I (CT) 10.3,10.4
10.30 I (CH) 10.4,19.1
10.40 I (CH) 10.5.3.1
10.50 S CT=0
10.80 I (CH-OH) 11.2,14.3
11.20 I (CH-OF) 11.4,14.6
11.40 I (CH-0VF) 11.5,14.8
11.50 I (CH-0k1C) 11.6,14.9
11.60 [ (CH-OHTP) 11.7,15.1
11.70 I (CH-05LF) 11.8,15.2
11.80 [ (CH-071) 11.9,15.3
11.90 I (CH-0T3) 12.1,15.4
12.10 I (CH-0FC) 12.2 15.5
12.20 T "? ? ?". 1; 6 10.1
14.30 D 1.3; G 10.1
14.60 D 1.6;G 10.1
14.80 D 1.8;6 10.1
14.90 D 1.936 10.1
15.10 D 1.95;G 10.1
15.20 D 2.1;6 10.1
15.30 D 2.2;D 2.5;G 10.1
15.40 D 2.3; G 10.1
15.50 D 2.5; G 10.1
19.10 0
*701.00
*701.00
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Appendix C:*

Direct Determination of Radon Permeability Through Polycarbonate Film

In connection with the program to find sealants for uranium mine tunnels, an experiment to determine the permeation of radon-222 through polycarbonate (PC) film was performed. The cell used in the experiment is shown in Fig. C-1.

The two halves of the cell were constructed of 80-mm Pyrex glass tubing. This tubing has an inner diameter of 74 mm and the exposed area of the membrane mounted between the two halves was thus 43 cm². The volume of the upper half of the celt was about 200 cm³ and the lower half about 20 cm³. The flanges of the celt were greased with Apiezon N and a square portion of the membrane about 10 cm on a side mounted between them. The two halves of the cell were wired together and then simultaneously evacuated to check that a vacuum-tight seal had been obtained. Once this was confirmed, air was simultaneously admitted to the two halves of the cell to a pressure of about 70 cm Hg (93 kPa).

A flask of about 400 cm³ volume containing about 100 cm³ of a solution of about 10⁹ DPM of radon-226 had been previously evacuated and allowed to stand for a month until the 3.8-day radon-222 had grown into equilibrium. The flask was attached to the vacuum manifold and filled with air to a pressure of about 70 cm Hg (93 kPa). Subsequent to the experiment, samples of the gas were taken and the atoms of radon-222 per unit volume determined by alpha-counting.

The experiment was initiated by attaching the radon-226 flask to the lower cell and opening stopcocks to admit the radon-222 to the lower side of the membrane. Initially both halves of the cell contained a total pressure of about 70 cm Hg, and once radon-222 air diffusion from the flask to the lower cell was complete, the "pressure" of radon-222 on the lower side of the cell would be constant.

Referring to Fig. C-1, helium from the upper left port was periodically flushed through the upper coll and into the U-trap packed with molecular sieve SA. The MSSA trap was cooled in liquid nitrogen during these sampling operations. Pumping on the trap exit at upper right maintained pressure at 96 ± 1.3 kPa. While there is no stable isotopic carrier for radon-222, ethane has similar adsorption characteristics on MSSA. Approximately $30~{\rm cm}_{\rm STP}^2$ (0°C, $101~{\rm kPa}$) of ethane were added to each collected radon-222 fraction and the mixture desorbed at 350° C into a liquid nitrogen cooled trap. Counting tubes were calibrated on the alpha plateau using the radon-222 at equilibrium with a known amount of radon-226. By expanding a known amount of the radon-222 ethane mixture into one of these tubes and counting radon-222, one determines the radon-222 per unit volume of mixture. From the known total of the mixture one then calculates total radon-222 in the sample. Measurements were made at least in duplicate. The percent errors quoted in the table of results are standard deviations based on the differences annong replicates.

The experiment extended over a period of 32 days during which 14 collections of radon-222 from the upper cell were made. The results are given in Table C-1. Columns 1 and 2 list sample number and sampling time. Column 3 gives the number of atoms in the sample at sampling time per cm² of exposed membrane. As noted above, this area was 43 cm². Column 4 gives a cumulative total obtained by sunnining the atoms per cm² in the latest sample and those in all previous samples decay-corrected to the latest time of sampling. These results are thus the total which would have been observed in the sample if no prior sample had been taken. It will be seen later that this is one of the easier quantities to calculate theoretically. The half-life of radon-222 was taken as 3.82 days. Columns 5, 6, and 7 will be elucidated later.

^{*}The work reported in this section was done and reported by Floyd Momyer.

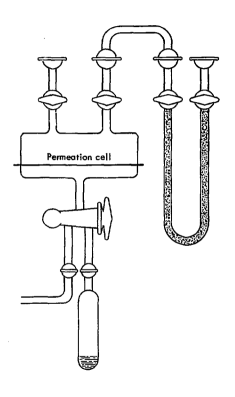


Fig. C-1. Schematic of radon-222 permeation cell and experimental setup.

Table C-1. Results for Radon-222 Permeation Through Polycarbonate. a, b

Sample	Time, h	Atoms/sample ^c	Σ Atoms ^c	C ₁ (Q)	C ₁ (S)	C ₁ (R)
1	2.42	6.25 × 10 ² 4.9%	6.25 × 10 ²			
2	69.43	4.99 × 10 ⁶ 1.5%	4.99 × 10 ⁶	3.23 × 10 ¹¹	3.23×10^{11}	(4.32×10^{10})
3	74.18	1.73 × 10 ⁶ 1.1%	6.54 × 10 ⁶	3.19 × 10 ¹¹	3.07 × 10 ¹¹	(1.60 × 10 ¹¹)
4	96.82	$1.07 \times 10^{7}_{2.9\%}$	1.62 × 10 ⁷	2.93 × 10 ¹¹	2.82×10^{11}	(1.73 × 10 ¹¹)
5	141.53	$3.14 \times 10^{7}_{0.4\%}$	4.30×10^{7}	2.76 × 10 ¹¹	2.70 × 10 ¹¹	(2.09 × 10 ¹¹)
6	165.87	$2.29 \times 10^{7}_{0.4\%}$	5.86 × 10 ⁷	2.71 × 10 ¹¹	2.64×10^{11}	(2.39×10^{11})
7	237.80	$7.86 \times 10^{7}_{2.0\%}$	1.13 × 10 ⁸	3.01 × 10 ¹¹	3.16 × 10 ¹¹	2.98 × 10 ¹¹
8	288.53	$7.21 \times 10^7_{-0.8\%}$	1.49×10^{8}	3.26 × 10 ¹¹	(3.58×10^{11})	(3.52×10^{11})
9	333.98	5.56 × 10 ⁷ 0.4%	1.61×10^{8}	3.15×10^{11}	2.97×10^{11}	2.95 × 10 ¹¹
10	405.85	8.53 × 10 ⁷ 1.2%	1.79×10^{8}	3.14 × 10 ¹¹	3.13 × 10 ¹¹	3.12×10^{11}
11	477.23	$9.10 \times 10^{7}_{1.8\%}$	1.95 × 10 ⁸	3.23×10^{11}	3.34 × 10 ¹¹	3.34 × 10 ¹¹
12	573.77	$1.06 \times 10^8_{-0.8\%}$	2.00×10^{8}	3.18×10^{11}	3.13×10^{11}	3.13 × 10 ¹¹
13	764.77	1.50×10^{8} 1.7%	1.98 × 10 ⁸	3.05×10^{11}	3.01×10^{11}	3.01×10^{11}
14	766.27	$2.64 \times 10^6_{2.6\%}$	1.98 × 10 ⁸	3.05 × 10 ¹¹	(3.58×10^{11})	(3.58 × 10 ¹¹)
				3.07 × 10 ¹¹ 5.8%	3.02 × 10 ¹¹ 7.3%	3.09 × 10 ¹¹ 4

b D=5.28 × 10⁻¹¹ cm²/s in calculations.

The resulting best value for C₁ is 3.06 × 10¹¹ atoms radon-222 per cm³ of polycarbonate at the coating - radon-222 interface.

At sample time per cm².

If the concentration in the membrane is given by a function C (x,t), then for a stable species the boundary value problem we are interested in is:

$$\frac{\partial C}{\partial t} = D \frac{\partial^2 C}{\partial x^2} \tag{1}$$

$$C(x,0) = 0, 0 < x < \ell \tag{2}$$

$$C(0,t) = C_1, \text{ all } t \tag{3}$$

$$C(\ell,t) = 0, \text{ all } t \tag{4}$$

In the above, x = distance into the film, $\ell = total$ thickness of the film, t = time. D = the diffusion constant ofthe species in the membrane, C(x,t) = atoms per unit volume of membrane at given x and t, and C_1 is the (constant) concentration in the membrane at the x = 0 interface.

If one assumes that C_1 is proportional to pressure (Henry's law), then $C_1 = KP$, where C_1 is the concentration

cm³_{STP} 222Rn rembrane . P is the pressure of radon-222 on the upstream side in kPa, and K is the solubility (Henry's

law constant) in units of $\frac{\text{cm}_{STP}^3 \text{ of } ^{222}\text{Rn/cm}^3 \text{ membrane}}{\text{kPa of } ^{222}\text{Pn}}.$

If we call this solution $C^{\theta}(x,t)$, then:

$$C^{0}(x,t) = C_{1}\left(1 - \frac{x}{\ell}\right) - \frac{2C_{1}}{\pi} \sum_{n=1}^{\infty} \frac{1}{n} \sin \frac{n\pi x}{\ell} = \frac{Dn^{2}\pi^{2}t}{\ell^{2}}.$$
 (5)

However, for a radioactive species the decay law says $dN/dt = -\lambda N$, where N is the number of atoms present at a given time, and λ is the disintegration constant of the diffusing species. Thus, if C(x,t) is the number of atoms per unit volume present in the membrane, the differential Eq (1) must be modified to

$$\frac{\partial C}{\partial t} = D \frac{\partial^2 C}{\partial v^2} - \lambda C . \tag{1a}$$

The other conditions remain the same. Following Carslaw and Jaeger

$$C(x;t) \approx \lambda \int_0^t e^{-\lambda t'} C^0(t') dt' + C^0 e^{-\lambda t},$$

where $C^0(x,t)$ is the function given above, and t' is a variable of integration.

The result of this integration is

$$C = C_1 \left(1 - \frac{x}{\ell} \right) - \frac{2C_1 \lambda \ell^2}{D\pi^3} \sum_{n=1}^{\infty} \frac{1}{n} \sin \frac{n\pi x}{\ell} \left\{ \frac{1 + \frac{D\pi^2 n^2}{\lambda \ell^2} e^{-\left(\frac{D\pi^2 n^2}{\ell^2} + \lambda\right) t}}{n^2 + \frac{\lambda \ell^2}{D\pi^2}} \right\} . \tag{6}$$

^{*}H.S. Carslaw and J.C. Jaeger, Conduction of Heat in Solids (Oxford Press, 1959) p. 33; also Danckwertz, Trans. Faraday Soc. 47 (1951) 1014-23.

Also if q is the rate of passage of atoms across the plane at x, then

$$q = -D \frac{\partial C}{\partial x}$$

and

$$q = \frac{DC_1}{\ell} + \frac{2C_1\lambda\ell}{\pi^2} \sum_{n=1}^{\infty} \cos\frac{n\pi x}{\ell} \left\{ \frac{1 + \frac{D\pi^2n^2}{\lambda\ell^2} e^{-\left(\frac{D\pi^2n^2}{\ell^2} + \lambda\right)t}}{n^2 + \frac{\lambda\ell^2}{D\pi^2}} \right\} . \tag{7}$$

The above equation is, of course, generally evaluated at $x = \ell$ where $\cos \frac{n\pi x}{\ell} = (-1)^n$. It should be noted at this point that for $\lambda = 0$ the late time rate of permention at $x = \ell$ is simply

$$(q)_{x=\hat{\chi}} = \frac{DC_1}{\hat{\chi}}$$
.

However, for a finite disintegration constant, at late times

$$(q)_{x=\ell} = \frac{DC_1}{\ell} \cdot \frac{\sqrt{\frac{\lambda}{D}} \cdot \ell}{\sinh \frac{1}{2} \sqrt{\frac{\lambda}{D}} \cdot \ell} . \tag{8}$$

Evaluation of the series in the second member on the right shows this member to be always negative. In fact, depending upon values of D, λ , and ℓ the first and second members may be comparable in absolute value. One may think of this as meaning that, on the average, the time an atom spends in the membrane after leaving the constant source of supply and before emerging on the other side may be significant compared to its radioactive lifetime. Unfortunately, this "holdup" time does not appear to be a very simple function of the parameters involved. However, the result is that the effect of a membrane in reducing downstream concentration of a radioactive species is always enhanced by radioactive decay — and this effect may be quite significant. The late-time rate of permeation is still a simple lipear function of C_1 ; however, it is no longer inversely proportional to thickness but decreases more strongly with increase in thickness. The product of solubility and diffusion constant no longer determines late-time rate of permeation. Unless films having extremely low solubility are found, the diffusion constant alone is the critical parameter.

To proceed with the solution of the boundary value problem, observations were actually made on the amount of radon-222 existing on the downstream side at various times. If the amount having crossed the plane x and existing at a time T is Q, then

$$Q = \int_0^T q(t)e^{-\lambda(T-t)} dt = e^{-\lambda T} \int_0^T q(t)e^{\lambda t} dt.$$

The result of this integration is

$$Q = \frac{DC_1}{\lambda \ell} (1 - e^{-\lambda T}) + \frac{2C_1 \ell}{\pi^2} \sum_{n=1}^{\infty} \cos \frac{n\pi x}{\ell} \left\{ \frac{1 - e^{-\left(\frac{D\pi^2 n^2}{\ell^2} + \lambda\right)T}}{n^2 + \frac{\lambda \ell^2}{D\pi^2}} \right\} . \tag{9}$$

Again, at late time and for $x = \ell$ the total radon-222 existing on the far side of the film is

$$(Q)_{x=\ell} = \frac{DC_1}{\lambda \ell} \cdot \frac{\sqrt{\frac{\lambda}{D}} \cdot \ell}{\sinh \left\{ \sqrt{\frac{\lambda}{D}} \cdot \ell \right\}}$$
 (10)

Thus, not only the rate of permeation eventually becomes constant, but the total radon-222 existing on the downstream side of the membrane is constant. It will also be noted that $\{\mathbf{q}\}_{n=0}^{\infty} \neq \lambda(\mathbf{Q})_{n=0}^{\infty}$ at late times, as it should when the rate of entrance into the downstream region finally equals the rate of decay of those atoms already downstream.

Before attempting interpretation of the results, several comments on departures from the theory on which the interpretation will be based should be made. While in principle these departures are all correctable, it was decided to design the experiment on the basis of convenience and maximum utilization of existing equipment with the aim of obtaining results uncertain by less than the factor of two that we understood was acceptable.

The first violation of the boundary conditions used in the solution of the differential equation for the permeation process is that the concentration of radon-222 on the upper side of the membrane is not always zero. Intuitively, one would expect that after a long interval between samplings the observed rate of permeation would be depressed and that a period of increased permeation would follow sampling. It will be seen that the next-to-last sampling interval was nearly 200 h and the last 1.5 h. While the last sample does indeed show an increased rate of permeation beyond analytical error, the effect is in any case less than 20%.

The temperature was not rigorously controlled in the box containing the equipment. However, periodic logging of the box comperature indicated that it remained within the range $(21 \pm 2)^{\circ}$ C.

At the initiation of the experiment, the pressure at the lower side of the membrane was not constant until interdiffusion of air and radon-222 was complete in the lower cell. Since this process should take hours and the experiment ran hundreds of hours this was considered acceptable.

Similarly, the reduction of radon-222 pressure in the radon-226 flask by an initial sampling operation was considered acceptable on the time scale of the experiment since equilibrium would be reestablished to within a fraction of a percent within a week.

One worries about the possibility of pinholes in the membrane, also the 10⁹ DPM solution of radon-226 was at least connected to the vacuum system to fill it with air prior to the experiment. Although care was taken in this operation, the possibility of contamination in the vacuum system could not be ruled out. The first sample at 2.42 h was taken as a check on these possibilities. Theory says their should be no detectable radon-222 in this sample. Whether the value reported is real is not unambiguous since the observations in the counting tubes were a few tenths of a count per minute over about one count per minute background. In any case, the observed amount is 4 orders of magnitude less than that in any subsequent sample. Thus leaks or contamination appear to have been at acceptable levels.

There are a number of ways in which one can handle the results of this experiment.

Method I

In the first place, as was done in Table C-1, one can sum all samples suitably decay-corrected to obtain the total permeation at the time of a given sample. This has two advantages. Inspection of the equation for $(Q)_{\chi=Q}$

indicates that at late times Q becomes an approximately linear function of the parameter $\tau = 1 - e^{-\lambda T}$. If τ^0 is the intercept of this line with the τ axis, then

$$D = \frac{2\ell^2 \lambda}{\pi^2 \tau^0} \sum_{n=1}^{\infty} \frac{(-1)^n}{n^2 + \frac{\lambda \ell^2}{Dn^2}} . \tag{11}$$

Such a plot was indeed made on data from the last eight samples to obtain a least-squares fit for the late Q = 0 intercept, and a value of D was calculated from this intercept.

Once the value of D is known it will be seen that the data from the experiment can be used to derive a value of C_1 from total permeation at each sampling. Since C_1 is assumed to be a constant, the above value obtained graphically for D was taken as the starting point for an iterative routine on D that resulted in the least sum of squares about the mean for C_1 for all but the initial sample. This process resulted in a value for D about 10% lower than the original estimate. Figure C-2 presents the data as a function of τ . The theoretical curves (solid lines) are from the result for D of the iterative process.

The second advantage of this mode of presentation is that the results should be a smooth curve plotted against time which can be compared with a theoretical curve. Figure C-3 is a plot of the experimental data compared to a theoretical curve derived from final values of the parameters D and \mathbf{C}_1 .

It might be noted that for D = 5.28×10^{-11} cm²/s, the average for C₁ in 12 samples is 3.06×10^{11} 5.8%. If D is increased 10%, C₁ from these same samples is 2.39×10^{11} 11.3%, and individual values show a distinct upward trend with increasing sampling time. Decreasing D by 10% results in C₁ equal to 4.15×10^{11} 14.3% and a downward trend with increasing time.

The disadvantage of this procedure is that an error in any one of the previous samples will be propagated until radioactive decay has reduced it to insignificance.

Method 2

The data can also be used to calculate a value of C_1 from the number of atoms in each sample. Thus, if s refers to the number of the sample,

$$Q_s = Q_{T_s} - Q_{T_{s-1}} e^{-\lambda (T_s - T_{s-1})}$$
.

Regarding the data in this manner suggests that sample 8 is too high by about 15%, probably due to an error in the amount of ethane carrier added to the radon-222. Results from the other data would allow correction for this, and such calculations were performed. Since they made less than 1% difference in the final conclusions, the data are presented as obtained. Such an error is considered insignificant compared to other uncertainties in the experiment.

Method 3

A third method of treating the data is suggested by noting that once the rate of permeation (q) is constant, the radon-222 in each sample satisfies the equation

$$q = \frac{\lambda Q_s}{1 - e^{-\lambda(T_s - T_{s-1})}} . (12)$$

Thus, once the value of D has been determined, a value of C₁ can again be calculated from the result of each sampling.

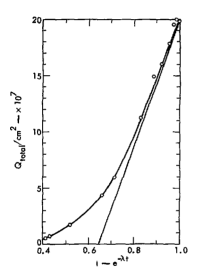


Fig. C-2. Total permeation vs $1 - e^{-\lambda t}$

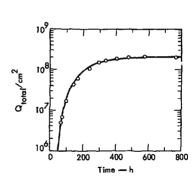


Fig. C-3. Total permeation vs time.

It might be noted that once steady-state has been achieved, methods two and three are equivalent in principle.

For purposes of data reduction, a program was written for the CDC 7600 which would calculate q/C_1 and Q/C_1 at $x=\ell$ as a function of λ , D, ℓ , and t. The disintegration constant, λ , was taken throughout as 7.55 \times 10⁻³/h. At early times all of these series, particularly that for q, are not very well behaved. Thus, calculations on q at early times are suspect both for calculational reasons and the fact that q is actually increasing during the early sampling intervals. All values of rate of permeation calculated will be too small, and yield a low value for C_1 . Although fewer terms would suffice in most instances, calculations were rather arbitrarily based on the first 5000 terms of each of the series since this required only about a second of computer time.

Referring back to Table C-1, column 5 lists derived values of $C_1(Q)$ based on method 1 above, or total permeation to sample time. Column 6 lists values of $C_1(s)$ based on method 2 or the permeation in each sample. Column 7 lists values of C_1 derived from method 3, or rate of permeation during each interval. It will be noted that the values in column 7 are indeed less than those in column 6 at early times, and later approach them almost exactly. Parenthesized values were not used in calculating quoted averages.

The samples taken at the end of the experiment indicated 1.09×10^{10} 1.5% atoms of radon-222 per cm³ of flask volume or a pressure of radon-222 = 4.44×10^{-5} Pa at 21° C. For $D = 5.28 \times 10^{-11}$ cm²/s, the value, C_1 , of the concentration in the membrane at x = 0 is 3.06×10^{11} atoms $\frac{222}{100}$ Rn/cm³ PC or 1.138×10^{-8} cm³ $\frac{37}{100}$ $\frac{222}{100}$ Rn per cm³ of PC. Assuming Henry's law,

$$C_1 = \left(0.257 \frac{\text{cm}_{\text{STP}}^3 ^{222} \text{Rn/cm}^3 \text{ PC}}{\text{kPa}^{222} \text{Rn}}\right) \times P \text{ (kPa) of }^{222} \text{Rn}.$$

Alternatively,

$$\frac{\text{atoms}^{222}\text{Rn}}{\text{cm}^3\text{ PC}} = 28.0 \times \frac{\text{atoms}^{222}\text{Rn}}{\text{cm}^3\text{ void}} \quad .$$

To illustrate the effect of radioactive decay, one may use Eq (8) to calculate late time permeation as a function of 2. He may also calculate permeation for a stable species with the same values of D and C_1 . Figure C-4 is a plot of the results. The ordinate is atoms/cm²/h (into the tunnel) per atoms/per cm³ void (in the mine wall). For radon-222 this ratio has the same numerical value if one expresses radon-222 in disintegrations per minute (DPM). The thickness of the film in this experiment (0.137 mm) is marked on the plot.

Given a thickness of 0.137 mm for the membrane, Fig. C-5 is a plot of total atoms of radon-222 downstream per atom/cm³ upstream, and rate of permeation (per hour) of radon-222 per atom per cm³ upstream.

A previous experiment extending over 5 h failed because the diffusion constant was almost two orders of magnitude less than expected. In an attempt to gain information for planning the current experiment, the alpha-activity remaining in the foil was determined with a survey meter at various times for two weeks after removal from the radon-222 source. From the observed "half-life", a value of $4.6 \times 10^{-11}~\rm cm^2/s$ was calculated for D. These were crude measurements and this is quite good agreement with the current best value $5.28 \times 10^{-11}~\rm cm^2/s$. Even with more sophisticated counting techniques, the determination of total radon-222 in the film (and thus solubility) is complicated by the fact that the film corresponded to 2 to 3 ranges for the alpha-particles involved. This makes absolute counting efficiency very difficult to determine.

However, these observations suggest an attractive technique for determining diffusion constant and solubility of radon-222 in membranes. For films "thin" compared to alpha-particle ranges, the counting efficiency is known. One is speaking generally of films about 1 mil in thickness. One could expose a membrane (or a number of membranes) to a constant known pressure of radon-222 until saturation is reached. The membrane(s) could then be removed from the source and total radon-222 determined as a function of time by alpha-counting. For the class

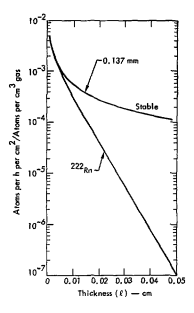


Fig. C-4. Steady-state permeation vs thickness (ℓ).

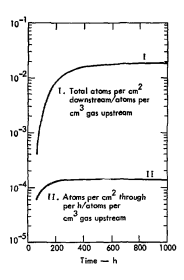


Fig. C-5. Total permeation and rate of permeation vs time.

of substances with low diffusion constants, theory would then allow determination of both diffusion constant and solubility.

We have the following boundary value problem:

$$\frac{\partial C}{\partial t} = \frac{D\partial^2 C}{\partial x^2} - \lambda C$$

$$C(x,0) = C_1 \cdot 0 < x < 0$$

$$C(0,t) = 0$$
, all t
 $C(0,t) = 0$, all t.

A substitution, $C = \mu e^{-\lambda t}$, results in

$$\frac{\partial \mu}{\partial t} = \frac{D \partial^2 \mu}{\partial x^2} . \tag{1a}$$

Other boundary conditions on μ remain the same.

Ultimately, the solution for C is

$$C = \frac{4 C_1}{\pi} e^{-\lambda t} \sum_{n=1}^{\infty} \frac{\sin \left\{ \frac{(2n-1) \pi x}{\ell} \right\}}{(2n-1)} e^{-\frac{D(2n-1)^2 \pi^2 t}{\ell^2}}.$$
 (13)

Also, Q, the amount in the membrane at a given time is

$$Q = \int_0^{\varrho} C dx = \frac{8 \, \ell \, C_1 \, e^{-\lambda t}}{\pi^2} \sum_{n=1}^{\infty} \frac{e^{-\frac{D(2n-1)^2 \pi^2 t}{\varrho^2}}}{(2n-1)^2} \, . \tag{14}$$

This series converges quite rapidly so that one approaches

$$Q = \frac{8\theta C_1}{\pi^2} e^{-\left(\frac{\mathbf{D}\pi^2}{\varrho^2} + \lambda\right) t} . \tag{15}$$

Q (t) can be determined directly by alpha-counting the membrane at a series of times. From its dependence on time, the argument of t in the exponential can be determined. As ℓ and λ are known, D can thus be calculated. Also, once the argument of t is known a value of C_1 can be calculated from each alpha-count. For values of D and ℓ leading to a time argument corresponding to a "half-life" of less than a few hours, the presence of the daughters polonium-218 and polonium-214 of radon-222, introduces some complication in the calculations as their alpha-particles will also be counted. However, this can be handled in a straightforward manner.