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Thermostimulated inert gas release analysis in the diagnostics of materials l

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Abstract

The principles of thermostimulated inert gas release analysis (TIGRA), based on the measurements of inert gas migration in solids, are given. Methodological bases and the computer treatment of the TIGRA results are described. The potential and various applications of this method are demonstrated, e.g. the determination of inert gas mobility in metals, polymers, ionic crystals and ceramics, the characterization of non- crystalline solids and their changes. © 1997 Elsevier Science B.V.

Keywords: Thermostimulated; Gas release analysis; Diagnostic

During the past two decades, the family of EGA area of the sample, can be revealed by this method.
techniques has been increased by TIGRA. The TIGRA Minor effects in the solid unaccompanied by changes techniques has been increased by TIGRA. The TIGRA Minor effects in the solid unaccompanied by changes
is based on the measurement of inert gases released in sample mass or enthalpy which cannot be invesfrom solids, usually previously labelled by the inert tigated by thermogravimetry or DTA, respectively, can gas [1]. The atoms of inert gases (from helium to be investigated by TIGRA. This method makes it gas [1]. The atoms of inert gases (from helium to be investigated by TIGRA. This method makes it radon) are used as trace indicators or a microprobe of nossible to gain information about the defects and radon) are used as trace indicators or a microprobe of possible to gain information about the defects and the defect state of the solid and its changes. Radio-
irrequisities of the structure of solids, which is comthe defect state of the solid and its changes. Radio-
active nuclides of inert gases (e.g. 24 Ne, 39 Kr, 133 Xe, alementary to the information given by other methods active nuclides of inert gases (e.g. ²⁴Ne, ³⁹Kr, ¹³⁹Xe, plementary to the information given by other methods, 222Rn), can be advantageously used as the probe of $\frac{1}{2}e$ X-ray diffraction. The TIGRA can be used in 222Rn), can be advantageously used as the probe of i.e. X-ray diffraction. The TIGRA can be used in the solids due to their easy detection in trace concentrations. Noble gases of "smaller size" such as He, Ne $[2,3]$. (the atomic radius of Ne is 0.112 nm) can be applied in In this paper methodical bases for the application of the study of the single point defects, whereas the $\frac{1}{2}$ to the diagnostics of materials are given "large sized" noble gases, such as Kr, Xe, Rn (the atomic radius of Rn is 0.214 nm) can be used in the complex defects mobility assessment, in the investi-
2. Labelling of solid samples by the noble gases

1. **Introduction** 1. **Introduction gation** of the morphology formations of solids and their changes. The inert gas atoms do not react with the The Evolved Gas Analysis (EGA) is known as an solid matrix and their release is controlled primarily appropriate method for characterisation of reactions in by diffusion. The processes which influence the inert by diffusion. The processes which influence the inert solids accompanied by gas evolution.

During the past two decades, the family of EGA area of the sample can be revealed by this mathod in sample mass or enthalpy, which cannot be invesstudy of both crystalline and non-crystalline materials

the TIGRA to the diagnostics of materials are given.

¹The paper is dedicated to Dr. H.J. Matzke on the occasion of his In general, common inert (noble) gases and their

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⁶⁰th birthday, radioactive isotopes or natural radioactive inert gases

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(such as radon) can be used for the TIGRA-measurements. $\frac{1}{\infty}$ 3.1 keV 33 keV 80 keV 150 keV 30 keV in Au

Most of the solids to be investigated by TIGRA

turally do not contain inert gas and it is necessary to

bel them with a trace amount of the inert gas. The

molitions of high energy gas ions, high temperature

d high gas naturally do not contain inert gas and it is necessary to $\frac{a}{2}$ $\frac{1}{8}$ label them with a trace amount of the inert gas. The $\frac{8}{5}$ $\frac{8}{6}$ conditions of high energy gas ions, high temperature and high gas pressure are required for impelling the inert gas into solids. $\sum_{i=1}^{\infty}$ 2

There are a number of techniques which can be used for direct incorporation of inert gas atoms in solids. $\frac{0}{3}$ $\frac{5}{25}$ $\frac{0}{25}$ $\frac{1}{25}$ $\frac{30}{25}$ $\frac{25}{25}$ All of them are characterized by a definite quantity of the introduced gas which decreases during storage or Fig. 1. Differential range distribution curves normalised to the measurement. The choice of the technique for inert peak height for ¹³³Xe in A1 and Au targets. The bombarded energy gas introduction denends mainly on the character of is indicated beside each curve (\bullet -Al \times -Au). gas introduction depends mainly on the character of the solid to be labelled and the aim for which the labelled solid should be prepared, ment and the properties of the labelled matrix. A number

This is based on the diffusion of the inert gas into
for sample labelling was invented by Jech [9].
solids at elevated temperatures and pressure of the gas. The substance to be labelled and the inert gas are placed in a high-pressure vessel which is then closed
and heated for several hours at the temperature of distinctly indicate non uniform distribution The dis-
distinctly indicate non uniform distribution. The disand heated for several hours at the temperature of distinctly indicate non-uniform distribution. The dis-
approximately $0.3-0.5$ T_{melt} of the substance to be tribution of the embedded atoms is seen to consist of approximately 0.3–0.5 T_{melt} of the substance to be tribution of the embedded atoms is seen to consist of labelled, and finally quenched in liquid nitrogen. The an asymmetric neak with an exponential tail With an labelled, and finally quenched in liquid nitrogen. The analymmetric peak with an exponential tail. With an amount of inert gas introduced into the sample increase in the energy of the ions the maximum is amount of inert gas introduced into the sample
depends on time, temperature and pressure, as given
objected towards creater distances from the surface. In depends on time, temperature and pressure, as given shifted towards greater distances from the surface. In
by the diffusion equation [1].

atmosphere (e.g. argon) are automatically labelled by solid. Direction (110) exhibits the deepest penetration. the inert gas. The inert gas atoms are captured in the structure of the deposited substance. The release of the *2.4. Inert gases produced from nuclear reactions* gas can be used for diagnostics of the film during its subsequent heating.
The recoil energy of nuclear reactions producing subsequent heating.

of techniques can be used for inert gas ion bombard-*2.1. Diffusion technique* ment [6-8]. A defined ion beam can be produced using a magnetic separator. A versatile low-cost technique

Fig. 1 gives curves of the differential distribution of xenon atoms in aluminium and gold [8]. Energies from the diffusion equation [1].
The diffusion technique was used to introduce 85 Kr represent for metarials consisting of light algorithments (with The diffusion technique was used to introduce \mathbb{R}^r rower for materials consisting of light elements (with in more than 250 different solids including metals, in more than 250 different solids including metals, lower Z) such as aluminium, than for heavier elements, inorganic and organic compounds in the form of foils, $\frac{1}{2}$ such as $\frac{1}{2}$ such as $\frac{1}{2}$ and $\frac{1}{2}$ inorganic and organic compounds in the form of folls, such as gold. The distribution curves are considerably
powders and single crystals. A microdiffusion tech-
headened when a large integral flux is used powders and single crystals. A microdiffusion tech-
nique for labelling powders permitting the reduction
 $(10^{10} \text{ ions/cm}^2)$ during a long bombardment packing $(>10^{10}$ ions/cm²) during a long bombardment, peaking of the losses of residual inert gas was proposed [4,5]. at a lower penetration depth. The peak occurs at only *2.2. Physical vapour deposition* half the depth observed in the trace bombardment. The deep penetrations are typical channelling events and In this method, thin films produced in an inert gas are markedly different from the isotropic (amorphous)

inert gases can be used for the implantation of gases *2.3. Implantation of accelerated ions of inert gases* into solid samples, e.g. in alkali-metal or alkaline earth metal halides Ar, Kr, and Xe are formed by means of The amount of inert gas introduced and its concen- (n, α) and (n, p) reaction, respectively. Nuclear fission tration profile depends on the energy of ion bombard- reactions have been used for the incorporation of the

Table I

Nuclear reactions which can be used for production of inert gas $\overline{08}$ 1 atoms and their introduction into solid samples C~***** intervalse cannot be calculated their introduction into solid samples

reactions which have been used for the production of obtained either by neutron irradiation generating inert gas atoms and their introduction into solid sam- inert gas atoms in the bulk of the solid, or in natural

By varying the conditions for labelling (e.g. neutron gin, flux, type of the nuclear reaction, dose and energy in \bullet distribution with a definite concentration profile ion bombardment), it is possible to separate different beneath the surface (e.g. plane source, exponenrelease processes which could otherwise occur simul- tial, peaked with an exponential tails) obtained by taneously leading to a release behaviour which would ion bombardment, diffusion or recoil technique of be difficult to interpret. By varying the ion dose in the labelling. ion bombardment, the gas concentration can easily be varied without changing the purity of the specimen thus avoiding the disturbing impurities produced dur- *2.5. Introduction of parent nuclides* ing reactor irradiation. By varying the energy of gas ions, the position of the gas with respect to the surface Trace amount of 226 Ra can be introduced into the can be chosen thus enabling a separation of surface sample by co-precipitation during the sample preparaand bulk release processes. Moreover, the effect of tion from a solution or adsorption on the surface of the radiation damage or specific impurities on release can sample. 222 Rn is formed by spontaneous alpha decay

introducing the gas traces it is possible to obtain nuclear reaction which give rise to the radon nuclide, samples in which the noble gas is distributed uni-
has also been used for the incorporation of the inert formly throughout the volume or located beneath the gas into the solid sample. Radon atoms penetrate surface. The most common distribution profile (see several tens of nanometers, depending on the compo-Fig. 2) are: sition of the target materials.

Fig. 2. Idealized concentration profiles suited for ion bombardment, recoil and diffusion techniques of labelling.

- radionuclides of ¹³³Xe and ⁸⁵Kr. Some of the nuclear \bullet homogeneous distribution which is usually ples are listed in Table 1 in [1]. minerals containing inert gases of radiogenic ori-
	-

easily be studied in double bombardment. $\qquad \qquad$ of radium and can be introduced into the solid owing According to the method used and the conditions of to the recoil energy (80 keV per atom). The above

are situated on the structure defects, such as vacancies, vacancy clusters, dislocations, grain boundaries and pores. The mobility of the inert gases in solids is strongly dependent on the structure and its changes.
Stage IA is due to gas fortuitously located in high An important role in the gas migration is played by **Stage IA** is due to mobility sites. impurities of various kinds and by natural and artifi- mobility sites. cially induced defects. The defects of the solid can serve both as traps and as diffusion paths for the noble gas.

explain inert gas migration in solids [1]. Self-diffusion of the matrix atoms.

- assumed to occupy vacancy sites in the lattice gas atoms.
and diffuse through either the cation or anion Stage IIB vacancies. \blacksquare
- ing into adjacent interstitial sites. The pre-existing defects or in gas-filled bubbles.
- *Interstitial diffusion plus trapping.* Inert gas Fraction release F or release rate J are most atoms diffuse interstitially but may be trapped at release rate from the defect and the measured diffusion activation energy will involve both the conditions employed. defect binding energy and the energy of migration of the gas atom.
- *Mobile cluster diffusion.* Inert gas atoms associate *4.1. Normal volume diffusion mechanism* with mobile vacancy clusters and move through diffuses to the lattice position adjacent to the

Release of inert gas can be caused by a number of can be written in the form of Eq. (1) processes, such diffusion, release caused by the instability of the surface, annealing of lattice damage $\frac{6}{5}$ during heating, evaporation of surface layers, recrys-

3. State **and diffusion of noble gases in solids** tallization or phase changes, chemical reactions of labelled sample, etc. A clear separation of various The solubility of noble gas atoms in inorganic solids processes that contribute to the overall release is is small. The inert gas atoms incorporated into solids possible. These process were grouped by Matzke possible. These process were grouped by Matzke [10] into a system of stages.

- Stage I involves gas mobility at unusually low
-
- radiation damage, the most common form of which is radiation-introduced amorphous state.
- Four diffusion mechanisms have been proposed to \bullet Stage II involves gas mobility at "normal temperatures", i.e. temperature similar to those of
- *Vacancy diffusion.* The inert gas atoms are **Stage IIA** is due to unperturbed mobility of single
- Stage IIB is due to interactions of gas atoms with sublattice by jumping into appropriate adjacent radiation damage or pre-existing defects, or with
- *Interstitial diffusion*. Inert gas atoms occupy **Stage III** involves gas mobility at unusually high interstitial sites in the lattice and diffuse by jump-
temperatures and is due to strong trapping of gas at temperatures and is due to strong trapping of gas at

defects in the lattice such as vacancies. If the commonly measured in experiments where the inert
gas release is studied. The heating at various tembinding energy to the defect is sufficiently large,
peratures for a definite time, so called isothermal the rate-controlling step in diffusion will be the peratures for a definite time, so called isothermal
step-heating or isochronal heating, or the heating in conditions of increasing temperature are usually

the lattice with the cluster. Dissociation of the gas The diffusion stage of the release is usually stom from the cluster which immobilized the gas observed at temperatures when the self diffusion in atom from the cluster, which immobilized the gas observed at temperatures when the self diffusion in
atom may occur until another vacancy cluster the crystal lattice of solids takes place and when the atom, may occur until another vacancy cluster
diffuses to the lattice position adjacent to the inert gas is distributed in the whole volume of the gas atom.
gas atom. Sample. In this case the inert gas release can be described by the solution of Fick's laws by taking into account the respective initial and limit conditions and the geometrical form of the samples investigated 4. **Release of inert gas** [11]. The initial differential equation taking into account various geometrical shapes of the sample

$$
\frac{\partial C}{\partial t} = D \frac{1}{r^{\nu}} \frac{\partial}{\partial r} \left(r^{\nu} \frac{\partial C}{\partial r} \right) \tag{1}
$$

where $C(r, t)$ is the concentration of inert gas, t is time, coefficient can be described by Eq. (6) D is the inert gas diffusion coefficient, r is the radial coordinate, $\nu=0$ for a plate, cylinder or prism with impermeable sides, $\nu = 1$ for a cylinder or prism with impermeable bases (i.e., infinite cylinder), $\nu = 2$ The time dependence of the flux J of the inert gas for a sphere. Eq. (1) has to be solved at the limit

$$
C(L,t) = 0 \tag{2a}
$$

$$
\frac{\partial C(0,t)}{\partial r} = 0 \tag{2b}
$$

$$
C(r,0)=f(r) \qquad \qquad (2c) \qquad \qquad 0 \qquad \qquad T_0
$$

volume at the point where sample diffusion takes can write place, e.g., the half width of the plate or radius of the sphere or cylinder.

For homogeneous distribution of the diffusion gas in the volume of the sample $C(x, 0) = C_0$ the solution of Eq. (1) under the limit conditions (2) results in the time dependence of the density J of the flux of inert gas release, in Eq. (3)

$$
J = \frac{2(\nu+1)D}{L^2} \sum_{m=0}^{\infty} \exp\left(-\mu_{m,\nu}^2 \frac{D}{L^2} t\right)
$$
 (3)

 πm and q_m are nuclei of the Bessel functions, sample is expressed by Eq. (9)

The existence of the temperature dependence of the diffusion coefficient causes more complicated mathematical expressions for the equation of the inert gas release flux J . (9)

The temperature dependence of the diffusion coef-
ficient is usually represented as
 $\frac{1}{2}$ besting at a condition being surposed

$$
D = D_0 \exp\left(\frac{-E_d}{RT}\right) \tag{4}
$$

$$
T = T_0 + \beta t \tag{5}
$$

heating, β is heating rate (in K s⁻¹), t is heating time distribution with an exponential tail were analysed by (in s). Then the time dependence of the diffusion Kelly and Matzke [12]. The $J(T)$ curves for linearly

$$
D(t) = D_0 \exp\left[-\frac{E_d}{R}(T_0 + \beta t)\right]
$$
 (6)

for a sphere. Eq. (1) has to be solved at the limit release on constant heating rate of the sample can be conditions obtained by replacement in Eq. (3) of *D* by $D(t)$ (see Eq. (6)) and *D*, by the integral time τ_i which in given by

$$
\tau_i = \int\limits_0^t D(\tau) \, \mathrm{d}\tau = \int\limits_{T_0}^T D(\theta) \, \mathrm{d}\theta \tag{7}
$$

where L is the characteristic size of the elementary Under conditions of a linear temperature increase we

$$
e\tau_i = \frac{D_0 E_d}{\beta R} \left[-\frac{\exp(\tau)}{\tau} + E_i(\tau) \right]_{\tau = -E_d/RT_0}^{\tau = -[E_d/R(T_0 + \beta t)]}
$$
\n(8)

$$
E_i(z) = \int_{-\infty}^{z} \frac{\exp(\tau)}{\tau} d\tau
$$

Finally, in the case of a uniformly labelled sphere Where $\mu_{m,\nu=0} = (2m+1)\pi/2$, $\mu_{m,\nu=1} = q_m$, $\mu_{m,\nu=2} =$ with radius r_0 , the flux J of the inert gas from the

$$
J = \frac{2C_0SD_0}{r_0} \exp\left(-\frac{E_d}{RT}\right) \sum_{n=1}^{\infty} \exp\left(-\frac{n^2\pi^2\tau_i}{r_0^2}\right)
$$
\n(9)

heating rate condition being supposed.

As it follows from Eq. (9) with conditions of linearly increasing temperature, the time dependence of the inert gas flux J can be represented by an asymwhere D_0 and E_d are the pre-exponential factor and metric peak-like curve (see Fig. 3). The peak max-
activation energy of inert gas diffusion, respectively, R imm temperature rises with the increase of the imum temperature rises with the increase of the is the molar gas constant, and T is temperature (in K). diffusion activation energy, radius of the grain and In conditions of constant heating rate the tempera-
the sample heating rate. The decrease of the preture increase according to the relationship exponential factor should also lead to an increase in the temperature of the peak maximum.

Cases of plane source and exponentially decreasing where T_0 is the temperature of the onset of sample profile, linearly decreasing gas distribution, and peak

lated inert gas release during constant rate heating of samples of retarded gas release. An important role in the gas various shape, supposing various diffusion stages: curve (1) migration is played by impurities of various kind
spherical sample, diffusion stage of gas release; curve (2) plate, and by natural and artificially induced defe diffusion stage of gas release; curve (3) kinetic stage of gas release.

 (T_m) of which are governed mainly by the value of imperfection existing in a material [14]. The sample is the activation energy, E_d . The shapes of peaks depend believed to contain the inert gas in two energetically the activation energy, E_d . The shapes of peaks depend believed to contain the inert gas in two energetically markedly on the type of diffusion and on the heating distinguishable sites; in the continuous phase of the

$$
\frac{E_{\rm d}}{T_m} = (A \pm 5) + 4.6 \frac{\log_{10} T_m}{R_m^2 \beta} \tag{10}
$$

heating rate β in K min β . Constant A depends on the cally, a trapping effect has been introduced in Fick's distribution profile and the diffusion type [1]. R_m is the law by adding reaction between mobile gas atom median range of ions in units of atomic spacings a , stable distribution of traps [15,16]. The process, called which for single jump mechanism equals 1 and for a "gas diffusion with reversible trapping", can be sphere is taken as radius r_0 . The uncertainty ± 5 arises described by following differential equations: from the assumption of D_0 values. The range $D_0 = 3 \times 10^{-1} \pm 1$ cm² s⁻¹ is obeyed in metal diffu- σC_1 σC_2 sion and also applies well to self-diffusion in the ionic crystals as alkali halides.

The widths of the peaks depends mainly on the character of the gas distribution. The narrowest one corresponds to a single jump mechanism, while a \ddot{a} wider peak was obtained with a plane source, the

 $\left(\begin{array}{ccc} \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \end{array} \right)$ range R_m [13]. The expression for the width of a gas temperature, $T_{1/2}$ can be written is general form as follows

$$
\frac{\Delta T_{1/2}}{T_m} = B - C \frac{\log_{10} T_m D_0}{R_m^2 \beta \cdot 0.3}
$$
(11)

 B and C are constants which depend on the geometry of samples.

4.2. Retarded gas release due to a trapping

Theoretical considerations suppose an ideal volume $\overline{\smash{\big)}\ 320}$ $\overline{\smash{\big)}\ 570}$ diffusion mechanism of the inert gas release. A large number of gas release experiments follows TEMPERATURE T [K] this ideal diffusion pattern. However, the larger part Fig. 3. Computed temperature dependences of the thermostimu- of the experimental work showed deviations, e.g. and by natural and artificially induced defects: vacan-The values of E_d =20 kcal mol⁻¹, $In(K_0R/E_d = 23$ are assumed. cies, vacancy clusters, dislocations, grain boundaries and pores.

In the retarded gas release mechanism gas atoms are considered that migrate in the solid matrix via random increasing temperature exhibit peaks, the maxima walks that are interrupted by trapping into various markedly on the type of diffusion and on the heating distinguishable sites: in the continuous phase of the rate β and the median or 50% range R_m . solid (i.e. "normal") and in the defect (i.e. "trapped") The expression for E_d/T_m is ones. Isolated point defects with an unlimited capacity are randomly distributed in the bulk of the solid. In the course of their random motion, inert gas atoms are trapped by the defects and are excluded for a definite where activation energy E_d is given in cal mole⁻¹ and time interval from the diffusion process. Mathemati-
heating rate β in K min⁻¹. Constant A depends on the cally a transing effect has been introduced in Fick' law by adding reaction between mobile gas atoms and described by following differential equations:

$$
\frac{\partial C_1}{\partial t} = D \frac{\partial^2 c_1}{\partial x^2} - k_1 N_2 c_1 + k_2 N_1 c_2
$$

=
$$
D \frac{\partial^2 c}{\partial x^2} - k_1^* c_1 + k_2^* c_2
$$
 (12a)

$$
\frac{\partial C_2}{\partial t} = k_1 N_2 c_1 - k_2 N_1 c_2 = k_1^* c_1 - k_2^* c_2 \qquad (12b)
$$

sion agent in the diffusion channels and traps, respec- experiments. Temperature dependences of the effectively, N_1 and N_2 are the number of states 1 and 2, k_i is tive diffusion coefficient of inert gases in highly the rate constant for the passage of inert gas atoms disordered media obtained by mathematical modelfrom one energy state to another, D is the undisturbed ling are given in [17]. diffusion coefficient, and

$$
k_1^* = k_1 N_2
$$
 and $k_2^* = k_2 N_1$

The first-order chemical reaction kinetics is used for describing the trapping of gas atoms by the matrix and Matzke [18] was the first to suggest that inert gas describing the trapping of gas atoms by the matrix and describing the trapping of gas atoms by the matrix and atoms could possibly associate with mobile defect subsequent release of the gas.

$$
\text{defect} + \text{gas atoms} \underset{k_2}{\rightleftarrows} \text{defect}
$$
\n
$$
= \text{atoms complex (immobile)} \tag{13}
$$

hand side at high temperatures, favouring movement relates to gas-defect cluster diffusion. When a gas of the gas to its mobile state and vice versa. The atom, diffusing in a solid, is exchanged between two equilibrium constant of the trapping reaction is diffusion streams. so-called dissociative diffusion

$$
K_H = \frac{n_2}{n_1} = \frac{k_1 N_2}{k_2 N_1} = \frac{k_1^*}{k_2^*} = \frac{c_2}{c_1} = K \frac{N_2}{N_1}
$$
 (14)

sample has been considered in the mathematical mod-
mobile cluster, is given by the following differential elling experiments [1]. equations:

An important special case of the model is given when the trapping reaction has reached its thermal $\frac{C_1}{C_1}$ equilibrium. Differential Eqn. 12 reduces to the simple Fick's type, with $D_{app'}$ which is smaller than D for undisturbed diffusion. If the local equilibrium is $\ddot{\theta}$ reached during the experiment $(k_1^*C_1 = k_2^*C_2)$, the observed diffusion coefficient for the mobile gas, D_{app} where C_1 and C_2 are the concentrations of the inert gas
by the following expressions:

$$
\frac{\partial c}{\partial t} = \frac{D}{(1 + K_H)} \frac{\partial^2 c}{\partial x^2} = D_{\text{app}} \frac{\partial^2 c}{\partial x^2}
$$
 (15a)

$$
D_{\rm app} = \frac{D}{1 + K\phi_2/\phi_1} = \frac{D}{1 + K_H} \tag{15b}
$$

These results are derived by considering the equilibrium distribution of gas atoms between normal sites and traps. The expressions for the diffusion coefficient If the local equilibrium is reached during the experiare correct if this equilibrium is established rapidly ment $(k_1^*C_1 = k_2^*C_2)$, the observed diffusion coeffi-

where C_1 and C_2 are the concentrations of the diffu- the solid, and this condition is satisfied in many

4.3. A mobile cluster diffusion mechanism of inert

clusters and move with clusters through the lattice. The trapping reaction is The gas atoms may periodically dissociate from the gas atoms may periodically dissociate from the k_1 clusters, and become trapped in the lattice. These defect + gas atoms \rightleftarrows defect diffusion models consider the transport of an atom from a solid with two diffusion coefficient: D_1 and D_2 $(D_2 \gg D_1)$. The diffusion coefficient D_1 relates to the A quasi-chemical equilibrium is shifted to the left- simple inert gas diffusion. The diffusion coefficient $D₂$ diffusion streams, so-called dissociative diffusion takes place [19].

The system of the differential equations describing one-dimension diffusion in a planar sample with a where $K=k_1/k_2$ **thickness l, when a reversible quasi-chemical reaction** The inert gas flow from the spherical or plate-like of the first order takes place between gas atoms and

$$
\frac{\partial C_1}{\partial t} = D_1 \frac{\partial^2 c_1}{\partial x^2} - k_1^* c_1 + k_2^* c_2 \tag{16a}
$$

$$
\frac{\partial C_2}{\partial t} = D_2 \frac{\partial^2 c_2}{\partial x^2} + k_1^* c_1 - k_2^* c_2 \tag{16b}
$$

atoms in dissolved form and in the complex form with mobile defects.

In the absence of local equilibrium the system of the differential Eqn. 16 should be solved using the general where, where initial and boundary conditions. The unidirectional flux J_x is given by the linear combination of two $Fick's$ -law contributions:

$$
J_x(T) = -D_1(T) \frac{dc_1}{dx} - D_2(T) \frac{dc_2}{dx}
$$
 (17)

compared with the rate of diffusion of the gas out of cient for the mobile gas, D_{apo} , can be written:

$$
D_{\rm app} = \frac{D_1 + K_H D_2}{1 + K_H} = \frac{D_{10} \exp(-E_1/RT + H_{H0} D_{20} \exp[-(E_2 - E_{KH})/RT]}{1 + K_{H0} \exp(E_{KH}/RT)}
$$
(18)

xponential factors and activation energies, and K_H is present in the sample at time t. the equilibrium constant of the inert gas exchange The inert gas flux J from the sample heated at a between the two channels (see Eq. (14)). constant rate is given by Eq. (20) .

Like the trapped interstitial model, the mobile cluster diffusion qualitatively explains most of the features in inert gas diffusion experiments.

4.4. "Single-jump" diffusion mechanism: the In Fig. 3 the temperature dependences of the ther-

observed after long diffusion times, i.e. when the 1 and 2) and the single jump diffusion mechanism majority (>80%) of inert gas has been released. How- (curve 3, Fig. 3) were considered. The main influenever, in many cases the kinetic stage of inert gas cing factors are the geometrical shape of the grains and release can be considered as the beginning of the the diffusion mechanism of the inert gas. Supposing TIGRA-experiment $[11]$. This is for e.g. the case of the same parameters of the inert gas diffusion, the size inert gas desorption from solid surfaces, the inert gas of the samples and the volume diffusion mechanism of diffusion in highly defected solid media (e.g. meteor- inert gas, curve 1 in Fig. 3 results for spherical samites and natural minerals) the inert gas diffusion from pies, with the maximum at a lower temperature than in the surface layers of solids labelled by ionic bombard- the case of plates or samples of other shapes (Fig. 3, ment or nuclear reactions, the case of solids with large curve 2). When "single-jump" diffusion is considsurface area (e.g. zeolites) and when intense solid state ered, the inert gas release curves are more symmetric processes take place during the heating of the sample and the peak is sharper (Fig. 3, curve 3) than in the (e.g. annealing of defects, phase transitions, solid state curves of the volume diffusion mechanism, and are reactions, etc.). independent of the shape of the sample.

In the kinetic stage, the inert gas release is almost independent on the initial concentration profile of the inert gas in the solid sample, the geometrical shape and size of the sample investigated; usually it takes 5. Analysis of the TIGRA curves place at temperatures considerably lower than the

This behaviour can be explained by the "singlejump" diffusion model considering that for the release *between the inert gas and the defect sites in* of inert gas atoms a small number (less than 100) of *the solid* gas-atom jumps in the lattice is sufficient. Considering the kinetic stage of inert gas release, the classical In the critical review [11] a solid sample labelled by diffusion equation (see Eq. (1)) can be substituted by an inert gas which is trapped in various defect sites, the equation of the first order chemical reaction in was supposed. It was considered that N_i is the number Eq. (19) of sites of *i*th type and that the sample contains *m* types

$$
\frac{dC}{dt} = -K_d C \tag{19}
$$

where $D_{1,0}$, $D_{2,0}$, E_1 and E_2 are the respective pre- jump, C is the number of the inert gas atoms which are

$$
J = C_0 K_0 \exp\left(-\frac{E_d}{RT}\right) \exp\left\{-K_0 t \exp\left(-\frac{E_d}{RT}\right)\right\}
$$
\n(20)

desorption of gas as first order kinetics reaction mostimulated inert gas release during constant heating of spherical samples or plates are demonstrated. In The kinetic stage of the inert gas release is usually calculations, the volume diffusion mechanism (curves

volume diffusion of inert gases in the respective solids. *5.1. Thermostimulated inert gas release as*

of defects [20].

For characterization of the solids labelled by inert gas before the sample heating, the energy spectrum where $K_d = \pi^2 D/d^2$, d being the length of diffusion $N(E)$ of the defect sites, and the energy spectrum $n(E)$

of the inert gas in the solid were used. In the case of can be written in the form total occupation of the defect sites by the inert gas, the spectra $N(E)$ and $n(E)$ were considered as identical.

The inert gas release can be described by a firstorder reaction mechanism, determined by the activa-
where the integral time $\tau_i = (k_0, E_i/\beta R)$ tion energy, E_i , then the time dependence of the inert $\{-\left[\exp(\xi)/\xi\right] + E_i(\xi)\}\$ (see Eq. (8)). gas flow $J(t)$ from the sample heated at constant In order to analyse the temperature dependences of $\frac{1}{2}$ in order to analyse the temperature dependences of $\frac{1}{2}$ in order to analyse the temperature dependences o

$$
J = \frac{\mathrm{d}\theta_i^*}{\mathrm{d}t} = K_{0,i} \exp\left(-\frac{E_i}{RT}\right)\theta_i^* \tag{21}
$$

where θ_i^* is the dimensionless expression for the number of inert gas atoms located on *i*th type sites, normalized on the total number of the defects N where $N = \sum_{i=1}^{m} N_i$.
The total inert gas flow *J* from sample is

$$
J = \frac{d\theta}{dt} = \sum_{i=1}^{m} K_{0,i} \exp\left(-\frac{E_i}{RT}\right) \theta_i^*
$$
 (22)

occupation of the sites by the inert gas.

the potential sites of the *i*th type normalized to the total spectrum of the activation energy of the inert gas
release was investigated. The principles are pointed number of defects of the *i*th type was introduced [20]. release was introduced α . The principles are pointed. The principles are pointed. The principles are pointed. The pointed are pointed. The pointed are pointed. T Eq. (22) can be written using the parameter ϕ_i as follows 1. when the spectrum consists of a single activation

$$
\frac{\mathrm{d}\theta}{\mathrm{d}t} = \sum_{i=1}^{m} K_{0,i} \exp\left(-\frac{E_i}{RT}\right) \phi_i \theta_i^* \tag{23}
$$

activation energies of inert gas release, for inert gas linear decreasing when the energy values are difflow J flow \int field to distinguish, and has a minimum when the

$$
J = \frac{d\theta}{dt} = -k_{\text{ef}}\theta = \int_{0}^{\theta} k_{0}(\theta^{*}) \exp\left(-\frac{E(\theta^{*})}{RT}\right) d\theta_{i}^{*}
$$
\n(24)

$$
J = \frac{d\theta}{dt} = -\int_{0}^{1} k_0(\phi) \exp\left(-\frac{E(\theta^*)}{RT}\right) \theta(\phi) d\phi
$$
\n(25)

rate are taken into account. The solution of Eq. (23) values of E_2 .

$$
J(t) = \sum_{i=1}^{m} \theta_{0,i} k_{0,i} \exp\left(-\frac{E_i}{RT}\right) \exp\left(\tau_i\right) \qquad (26)
$$

 $E_{\rm ef}$, the values of the effective diffusion constant were first determined at every point on the experiment curve $J(t)$. The effective diffusion constant k_{ef} can be expressed

$$
k_{\rm ef} = \frac{J(T)}{\theta T} = \frac{J(T)/F}{1 - 1/F \int_{T_0}^{T_F} J(T)} \tag{27}
$$

where $F = \int_{T_0}^{T_f} J(T)$ is the area of the peak of the TIGRA curve and T_0 and T_F are the onset and final temperature of the heating respectively. The calcu*d Ink_{eff}* values were plotted against *1/T* at every point on this curve; and the activation energy E_{ef} and where $\theta = \sum_{i=1}^{m} \theta_i^* = 1/N \sum_{i=1}^{m} N_i$ is the complete $k_{0,\text{eff}}$ were evaluated. Subsequently, the temperature dependences of the plots E_{eff} against T and $k_{0,\text{eff}}$ against The term of $\phi_i = N_i/N$ equalling the proportion of T were constructed. On the basis of these plots, the proportion of the activation energy of the inert gas

- energy, the plot of $E_{\rm ef}$ against T is linear and *parallel to the x-axis; and*
- 2. when the spectrum consists of several values of In the case of the continuous spectrum of the activation energy, the plot of E_{ef} against T is nonenergy values are clearly distinguished.

The application of this method in the analysis of the model TIGRA curves is demonstrated in Figs. 4 and 5 and Fig. 6. It was assumed that the energy spectrum consists of two discrete lined E_1 and E_2 , where and after substituting $d\theta^* = \theta d\phi$ we obtain $E_1 < E_2$.
In Fig. 4 schemes of the energy spectra $n(E)$ of the

defect sites in the solids considered during the mathematical modelling are given. The spectrum is characterised by two activation energy values E_1 and $E₂$, assuming the same occupation of the respective defect sites by the inert gas($\theta_1 = \theta_2$). The spectra The experimental conditions of the constant heating shown are assumed to have the same E_1 but different

Fig. 4. Schematic diagrams of the discrete energy spectra $n(E)$ of the solid state defect sites taking part in the inert gas diffusion. Two 5.2 . *Influence of the labelling conditions on the* different values F , and different values E_1 and E_2 of the effective activation energy of inert gas diffusion and the equal occupation of the defect sites by the inert gas $\theta_1 = \theta_2$ are supposed:(a) $E_1 = 10.0$ kcal mol⁻¹, inert gas $\theta_1 = \theta_2$ are supposed:(a) $E_1 = 10.0$ kcal mol⁻¹, In the case of the diffusion technique of labelling $E_2 = 10.1$ kcal mol⁻¹(b) $E_1 = 10.0$ kcal mol⁻¹, $E_2 = 10.4$ kcal mol⁻¹, the change of the TIGPA c and (c) $E_1 = 10.0$ kcal mol⁻¹, $E_2 = 11.0$ kcal mol⁻¹.

In Fig. 5 typical curves of the temperature depen- of samples. dences of the inert gas release from solids are demon-
A solid sample containing several defect sites charstrated. It is obvious from Fig. 5 that the dependences acterized by the discrete-energy Spectrum $N(E)$ was are single peaked curves, irrespective of whether one considered in [21]. When such a solid is maintained or two activation energy values are considered in the under high inert-gas pressure, the defect sites will discrete spectrum. A higher symmetry was observed become occupied by the gas. Supposing that the defect in the case where the energy spectrum has two lines. sites are equally accessible for the inert gas, the inert However, when the method of the analysis of tem- gas will first be trapped on the sites of maximum perature dependences, *lnko* versus T and Eef versus T, energy. The distribution of the gas between the defect was used, the influence of both values of activation sites is either given by the Boltzman distribution energy in the spectrum was found (see Fig. 6). In function (in the case of partially occupied defect sites), Fig. 6, the influence of energy lines in the spectrum or by the Fermi-Dirac distribution (in the case of the differing by only 1% can be seen. A difference in the complete occupation of the defect sites). When the energy lines of 4% is more obvious, pressure of the inert gas is increased, the energy

thermostimulated inert gas release (TIGRA curves) for the selected cases of the discrete energy spectra of the single energy value E_1 = **0,2 10.0** and 10.5 kcal mol⁻¹, respectively: curve 3 corresponds to the spectrum characterized by two discrete energy values $E_1=$ 10.0 kcal mol⁻¹ and E_2 =10.5 kcal mol⁻¹. In the modelling, the $\frac{1}{10.0}$ $\frac{1}{10.4}$ $\frac{1}{10.8}$ heating rate 2.0 Ks⁻¹ and the pre-exponential factor 10^{13} s⁻¹ were considered.

the shape of the TIGRA curves depends considerably on the conditions of the sample labelling, e.g. the inert gas pressure and the duration of the diffusion labelling

Fig. 6. Tempereature dependences of the effective activation where E_{min} and E_{max} are the limits of the energy energy E_{ef} of the inert gas diffusion calculated at every point of spectrum $N(E)$ of the defect sites.
the model TIGRA curves corresponding to the discrete energetic spectrum characterized by Fig.4 (curves 1-3 correspond to the

the TIGRA measurements, the experiments must be At low pressure the relative occupation (θ_1 and θ_2) of performed under the conditions of complete occupa-
the energy sites E_1 and E_2 is $\theta_1 < \theta_2$, at medium tion of the defect sites by the inert gas. Supposing that pressures $\theta_1 \cong \theta_2$ and only at sufficiently high presthe inert gas adsorption at every site is controlled by sures does θ_1 become higher than θ_2 . In practice this the Laangmuir law, the amount of the inert gas located means that as the inert-gas pressure is increased, the at the defect sites of ith type with energy E is given by defect sites of potential energy E_2 are occupied first.

$$
\theta_i(E) = \frac{n_i}{n_{i,\infty}} = \frac{K_i(E)p}{1 + K_i(E)p}
$$
\n(28)

defect of the inert gas. K_i is the equilibrium constant much lower amount of inert gas than do the defects of with respect to the defect site, and p is the partial potential energy E_2 , the occupation of the defect sites pressure of the inert gas. The relationship $n_i = K_i p$ of energy E_1 takes place. results from the Henry law. At low-pressure values the It is obvious that the inert-gas spectrum $\theta(E)$ dependence of θ on p is linear, independent of p. reflected by the TIGRA curves may differ substan-

sample is **If this fact is neglected, substantial errors can be made**

$$
n = \int_{E_{\min}}^{E_{\max}} n(E) \, \mathrm{d}E \tag{29}
$$

Fig. 7. Schematic diagrams of the discrete-energy spectra of the 2 defect sites $\phi(E)$ and the degree of occupation of the defect sites by inert gas introduced at: (a) labelling pressure $p=0.5$ relative units; and (b) labelling pressure $p = 10$ relative units. The effective $\overline{140}$ 150 160 170 180 T(K) activation energies E and corresponding degrees of occupation of the defect sites used are: $E_1=10$ kcalmol $\frac{1}{2}$, $\phi=0.7$ and

The schemes of the discrete-energy spectra $\phi(E)$ of spectral characterized by T_{B} . Centres T_{B} correspond to the spectra of the spectra of their occupation the defect site and of the spectra of their occupation $\theta(E)$ under different inert gas function $\phi(E)$ and $\theta(E)$ for a single solid sample may differ substantially.

spectrum $n(E)$ of the inert gas located at the defects is Fig. 7 illustrates schematically two discrete-energy similar to the energy spectrum $N(E)$ of the defect sites. spectra characterized by the energies E_1 and E_2 When the function $\phi(E)$ is to be determined from $(E_1 < E_2)$ and the degrees of occupation ϕ_1 and ϕ_2 . equation: The redistribution of the inert-gas atoms between the potential-energy sites E_1 and E_2 depends on the temperature dependence of the equilibrium constant K. The defect sites of potential energy E_1 usually (at the where $n_{i,\infty}$ is the maximum capacity of the *i*th type low pressures used for sample labelling) contain a

The total amount, n_i of inert gas located in the solid tially as the partial pressure of the inert gas is changed. in the interpretation of the TIGRA curves.

> Fig. 8 shows the TIGRA curves computed for linearly rising temperature in the cases of the discrete defect-site spectra shown in Fig. 7. At low partial

Fig. 8. The model TIGRA curves computed for the case of the defect sites of two discrete energy lines $(E_1 = 10 \text{ kcal mol}^{-1}, \phi_1 =$ Fig. 9. Model TIGRA curves for the linear heating rate $\beta = 2 \text{ Ks}^{-1}$
0.7 K -- 0.01; and $E_2 = 11 \text{ kcal mol}^{-1}$ $\phi_1 = 0.3 \text{ K}^{-1}$ and different and the par 0.7 K= 0.01; and E_2 =11 kcal mol⁻¹, ϕ_2 =0.3, K=1) and different and the partial outgassing of samples at 140 K. The defect sites
partial pressures during sample labelling. The pre-exponential characterized in Fig. partial pressures during sample labelling. The pre-exponential characterized in Fig.7 were assumed. Curves 1–5 correspond to prediction rate $\ell = 10^{13} e^{-1}$ and the linear besting rate $\ell = 2 \kappa e^{-1}$ were assumed uration factor $K_0 = 10^{13} \text{ s}^{-1}$ and the linear heating rate $\beta = 2 \text{ Ks}^{-1}$ were preliminary outgassing durations of β , and β respectively. supposed. Curves $1-4$ correspond to relative partial pressures of (in relative units): 1, 10, 100 and saturation gas pressure respectively.

expected, in the energy of the defect sites determining The preliminary thermal treatment caused the partial

labelling is increased, the height of the $(J_{\text{max}})_2$ peak ment makes it possible to influence the shape of the rises, in correspondence with the Langmuir isothermal TIGRA curves. Therefore, the determination of the curve. In Fig. 8, curve 3, it is shown that at a relative energy spectrum of the inert gas labelling can be partial pressure of $p = 100$ during sample labelling, simplified. the maximum height of $(J_{\text{max}})_2$ is attained whereas the Fig. 9 shows the model TIGRA curves computed

becomes most significant in the TIGRA curve. The 140 K assuming various time intervals for the sample importance of the sample labelling conditions on the outgassing. It can be seen from Fig. 9 that the height TIGRA model curves has been shown here. The of the first peak diminished with increasing duration of results have been used for the assessment of the the thermal treatment. When the treatment is suffifunctions $\theta(T)$ and $\phi(T)$ from the experimental TIGRA ciently long, the first peak disappears completely and curves, the second peak diminishes substantially.

5.3. Influence of preliminary sample thermal treatment on the TIGRA curves 6. **Measurement of inert gas release**

The amount of inert gas located at the defect sites of The apparatus for TIGRA consists of several comsolids also depends on the preliminary thermal treat- ponents designed to ensure the detection of inert gas, ment of the solid samples. The solid sample charac- and to provide sample heating and temperature conterized by the discrete-energy spectrum shown in trol. In addition, the instrument supplies the carrier gas

Fig. 7 can be obtained by the diffusion technique of labelling at partial pressures which ensured the compressures the one-peak TIGRA curve result, as plete occupation of the defect sites by the inert gas. the maximum J_{max} . J_{max} outgassing of the labelled sample. The choice of a When the partial pressure of the inert gas during suitable temperature and duration of the thermal treat-

height of (J_{max}) increases. For the samples whose energy spectra are given in At the high pressures of inert gas used, (T_{max}) Fig. 7, and for preliminary sample outgassing at

with flow stabilisation, and measures the gas flow and the sample can be evaluated. Other diffusion paracomplementary parameters, meters can be determined and the fitness of adequate

another suitable gas) carries the inert gas released by diffusion models has been organized on the basis of the sample in the vessel (situated in a furnace) into a both analytical and numerical solutions of diffusion detector for the inert gas. For example, to measure the cases. In the bank of experimental data the standard α -activity of Rn, a scintillation counter, ionisation interface for the input of experimental data is used. α -activity of Rn, a scintillation counter, ionisation chamber or semiconductors can be used. On the other The set of the programmes involves four parts hand, all β -activity measurements involving Kr, Xe which give following possibilities: and Ar are made by Geiger-Muller tubes. Gamma active gases can be measured by a γ -spectrometer. The stable nuclides of inert gases are measured by a mass cal diffusion model. spectrometer. To ensure optimum conditions for a direct comparison of TIGRA data with results fusion model. obtained by other thermoanalytical methods, devices were constructed to provide simultaneous measurement of additional parameters [1].

results. The TIGRA of composites, mixtures, natural order rocks and other materials as well as multistage diffusion processes accompanied by solid-state reactions make mathematical description rather complicated, which can be solved by means of computers.

A set of computer programmes called DIGS (diffu-
sion of inert gas in solids) for the statistical treatment
the true curve $y(t)$ is the response of the apparatus to and interpretation of the results of various methods the short-time pulse. used for the investigation of transport properties of solids was designed [22-25]. Following parameters *7.2. Reconstruction of "non-distorted" TIGRA* were considered: *curves*

- the gas concentration distribution along the width

Fig. 10 shows the results of the reconstruction of

Fig. 10 shows the results of the reconstruction of
-
-
-

can be taken into account and diffusion coefficients can be seen from curve 1, Fig. 10 which shows depending on time, concentration and coordinates of apparatus response to the "one second pulse", the

During an analysis, the carrier gas (air, nitrogen or diffusion models can be verified. A bank of general

- preliminary treatment of experimental results,
- evaluation of the results on the basis of the classi-
- choice of a more suitable phenomenological dif-
- choice of a suitable physical diffusion model.

7.1. Testing the TIGRA apparatus

7. **Evaluation of TIGRA experimental data** The TIGRA apparatus should be tested in order to estimate the distortion effect of the apparatus on the The computer software for mathematical modelling experimental curve. The testing consists of measuring of thermostimulated inert gas release kinetics, and for the apparatus response to a short impulse of the gas. evaluation of experimental data of TIGRA methods The resulting apparatus response $\phi(t)$ is usually norrepresents an important part of modern thermal ana- malized with respect to the total amount of the inert lysis. The computer is used in the control of TIGRA gas; true form of the experimental curve can be equipment operation and in the evaluation of the obtained by solving the Fredholm equation of the first

$$
f(t) = \int_{0}^{t} \psi(t-\tau)^{*} y(\tau) d\tau
$$
 (30)

the true curve, $\psi(t)$ is the response of the apparatus to

of the sample,
the quantity of the gas in the solid,
 100 cm likelled by gadax by a diffusion to history • the quantity of the gas in the solid, $100 \mu m$, labelled by radon by a diffusion technique the quantity of the inert gas in gaseous phase, $100 \mu m$, labelled by radon by a diffusion technique the quantity of the inert gas in gaseous phase,
 $[26]$. The TIGRA curve was measured during heating
 $[26]$. The TIGRA curve was measured during heating
 $[26]$. The TIGRA curve was measured during heating • the gas fluxes at the output of the sample. $\left\{ \begin{array}{c} \text{[Fe]} \\ \text{[Fe]} \end{array} \right\}$ and $\left\{ \begin{array}{c} \text{[Fe]} \\ \text{[Fe]} \end{array} \right\}$ and $\left\{ \begin{array}{c} \text{[Fe]} \\ \text{[Fe]} \end{array} \right\}$ and $\left\{ \begin{array}{c} \text{[Fe]} \\ \text{[Fe]} \end{array} \right\}$ Various geometrical shapes (sphere, cylinder, plate) over the temperature range from -196 to 400° C. As

Fig. 10. The reconstruction of the experimentally obtained TIGRA The experimental curves of the thermally stimucurve of a polypropylene foil in the temperature range from -196 to $+400^{\circ}$ C (heating rate 4 min⁻¹) which was labelled with 222 Rn: lated diffusion in normal scale and in the functional curve (1)-the apparatus response to the "one second pulse", curve (2) - the experimentally obtained TIGRA curve; curve (3)-the reconstructed TIGRA curve. **I, 0.**

apparatus may cause a significant distortion of the $\begin{bmatrix} rel \\ units \end{bmatrix}$ apparatus may cause a significant distortion of the $\begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$ $\begin{bmatrix} 3 & 3 \\ 4 & 4 \end{bmatrix}$ high value of the time constant 7" i of the count-rate *///! --------L4* meter. The experimentally obtained TIGRA curve (Fig. 10, curve 2) exhibits a symmetrical form which $\qquad \qquad \text{0.5}$ could be ascribed to the inert gas release obeying first $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ order release kinetics with a spectrum of activation $\sqrt{2\pi r}$ energies.

However, as it follows from the reconstructed curve (Fig. 10, curve 3) the true form of the TIGRA curve is characterized by two effects, the first corresponding to the gas release controlled by first order kinetics with a $\begin{bmatrix} 0.0 | \mathbf{r} \end{bmatrix}$ the basis of the reconstructed (true) TIGRA curve it can be stated that in the polypropylene foil sample, Fig. 11. Temperature dependences of TIGRA influenced by which has been prepared under a pressure of 100 the inert gas trapping in the lattice of the solid. Differential atmospheres, radon atoms are situated in both amor-
 $K_0C^n \exp(-E_d/R(T_0+\beta t))$ where *n* is the formal order of
 $K_0C^n \exp(-E_d/R(T_0+\beta t))$ where *n* is the formal order of phous and crystalline phases. These phases are characterized by different diffusion resistances, 84 and
126 kJ mol⁻¹, respectively. From the amount of inert expectively $n = 0.5 \times 0.8 \times 10^{-15}$ are considered, curves 1-8, and 14.6, and 14.6, and 15, and 15, and 15, and 1 gas release in the respective release processes it scale (b) Functional scale.

Fret.
 Can Example 1 can be found.
 Can be found.

" : 7.3. Determination of the inert gas diffusion parameters

After the reconstruction of "true", non-distorted experimental curves the diffusion parameters can determined. The task of the treatment of experimental $\begin{array}{c|c}\n\hline\n0.5 & \begin{array}{\hline}\n\end{array}\n\end{array}$ diffusion data can be substantially simplified when using functional scales which give the possibility of linearizing non-linear dependences between the parameters investigated. There exists a possibility to construct special diffusion scale sheets by means of which the linearisation of the non-linear dependences can be made directly and simply. For every technique used in $\frac{1}{10}$ 10 $\frac{1}{20}$ TIGRA a functional scale was designed [11]. The **lime** [min] diffusion parameters and errors of their determination can be evaluated by the least squares method.

reaction. The following order n of reaction mutual interactions respectively: $n= 0.5, 0.8, 1.0, 1.5, 2.0, 3.0, 4.0, 5.0$ (a) Normal

]O00/T

Fig. II. (Continued).

scale are represented in Fig. 11 (a) and (b), respec- **8. Examples of TIGRA application** tively. In this way the initial peak-shape temperature dependence of the inert gas release is transformed to a A number of authors in the past years used the linear dependence. There are several advantages of Evolved Gas Analysis measuring the release of inert this simple method of treatment of diffusion experi- gases from solid samples. Although the inert gas mental data. The linearized dependences of inert release measurements carried out by various authors gas release enabled us to calculate easily the differed, a high sensitivity of the method to fine diffusion parameters using all points of the TIGRA changes in the solid materials, phase and structure curve. The suitable diffusion model to be used for the changes was observed. Results of additional measureconstruction of the inert gas release curve can be ments, such as electrical conductivity, DTA, TG, estimated according to the correlation coefficient dilatometry, X-ray diffraction patterns, etc, were used value. The most suitable diffusion model is the one in the interpretation of the EGA results. where the corresponding correlation coefficient is In this part of the paper we shall demonstrate the close to unity, high sensitivity of the EGA to various processes taking

do not correspond to the model of classical diffusion, It should be mentioned that methods of computer another more adequate diffusion model has to be treatment of the experimental data developed in the chosen for the determination of the diffusion coeffi- recent years (and described in this paper) were not cient. The basic expressions necessary for the corn- used in the demonstrated cases. Nevertheless, the puter treatment are stored in a special bank of demonstration of the results should promote interest phenomenological diffusion models. The choice of in the TIGRA applications. an adequate diffusion model is made by a step-wise trial of various models, starting from a simple to *8.1. Diagnostics of the defect state* more complicated ones. In the bank of phenomenological models analytical and numerical solutions of Inert gas diffusion parameters evaluated from diffusion equations are stored, taking into account TIGRA measurements reflect the mobility of noble irreversible and reversible chemical reactions of the gases in the solids, which can be used in the determi-Ist and 2nd order between the gas and traps in the nation of the defects in the solids. Inert gas atoms solid, various cases of parallel diffusion, diffusion incorporated into solids are situated on the natural and/ taking into account the exchange of diffusing gas or artificial defects produced, for example, by ion between the diffusion channels, etc. In solutions to bombardment, neutron irradiation or mechanical the diffusion equations various concentration profiles treatment. The release of the inert gases on sample of the gas in the solid, various concentration and time beating reflects the thermally stimulated processes, i.e. dependences of the diffusion parameters, etc. are diffusion, annealing of defects, etc. It was shown that considered. The choice of an adequate phenomeno- the mobility of inert gas atoms in the ionic crystals logical model of diffusion is made by means of differs invarious crystallographic directions, owing to standard methods of mathematical statistics. A diffu-
the channelling effect. The mobility of the inert gases sion model is considered to be adequate when the (the activation energy) can be used as a parameter difference between the experimental curve and the characterising the defect state of an ionic crystal theoretically constructed curve corresponds to the lattice and may cause the formation of the metamict normal distribution, and no drift of residues is phase. observed. Kornelsen and Sinha [27] presented the results

curves did not give us a possibility for determining argon, krypton and xenon ions in a tungsten single the diffusion mechanism. In these cases dependences crystal. The temperature dependences of the Ne, Ar, of the effective diffusion parameters on temperature, Kr, and Xe release from W single crystal introduced inert gas partial pressure, size of the sample and other in W single crystal by ion implantation at the energy parameters should be investigated. of 600 eV and the dose of 5×10^{12} ion cm⁻², in

Evolved Gas Analysis measuring the release of inert

In the case that the experimental data treated so far place during thermal treatment of different materials.

In some cases the analysis of the experimental indicating differences in the mobility of neon,

tungsten single crystal. The inert gases were incorporated by ion tungsten crystals. Fig. 13 shows variations of Xe

Fig. 12. The results of inert gas release from other $(8\times10^{10} \text{ ions sm}^{-2})$ diffusion release along and permaterials, such as Si, Ge, GaAs and SiC are given in pendicular to the c -axis dominated with activation [28]. energy values of 243 and 255.4 kJ mole^{-1}, respec-

Fig. 13. Xenon release from the three different faces of single tungsten crystals after 400 eV Xe bombardment.

8.2. Crystal orientation effects.

Several authors found that different crystal phases gave different gas release rates and apparent diffusion coefficients [1]. The crystal orientation effect is $\frac{\lambda}{\lambda}$ is the λ λ λ λ λ λ emphasized for the ion bombardment and fission λ *4 8 12 16 20 24* recoil techniques of labelling, i.e., where channelling of the inert gas atoms is possible. Kornelsen and Sinha *T (IO0°K)* [29] observed an effect of crystallographic direction of Fig. 12. Temperature dependences of Ne, Ar and Xe release from ion bombardment on the inert gas release from single bombardment at the energy of 600 V and the dose of release spectra of tungsten crystals for identical 5×10^{12} ions cm⁻² in the crystallographic direction <100>. 400 eV-Xe bombardment of three different faces $\langle 110 \rangle$, $\langle 100 \rangle$ and $\langle 211 \rangle$. Similarly Matzke [10] obtained different values of diffusion activation energy from release curves of $SiO₂$ crystals bomthe crystallographic direction (100) are given in barded with 40keV xenon. At low dose

tively. A value 301 kJ mole⁻¹ was found for Xe- $n(E)$ bombarded fused silica. Diffusion in the amorphous phase was markedly slower than the diffusion in the crystalline phase, o \overline{a} **b** \overline{a} **b** \overline{a} **b** \overline{a} **b** \overline{a}

8.3. Morphology changes in bulk ceramics and thin $\left\{\n \begin{array}{ccc}\n \text{if } & \text{if } \\
 \text{if } & \text{if } \\$

aging, drying, calcination and firing of feed or intermediate geleous materials can be successfully investigated by TIGRA. The method was used for quality \overline{a} **ao a**² **a**² **a**² **a**² **E**, [kc**al**/mol] testing of intermediate geleous products of urania, J , J , titonia, $\frac{1}{2}$ is estimated in $\frac{1}{2}$. E'. Units titania, silica, zirconia, etc. [30-33]. Thin ceramic films can also be investigated. TIGRA is sensitive to the morphology changes caused by deviations of technological conditions; therefore it has been recommended as the control method for quality testing intermediate products of sol-gel processed ceramics both in bulk and thin films. The morphology changes $\overline{500}$ $\overline{500}$ $\overline{700}$ $\overline{600}$ $\overline{100}$ 1300 $\overline{100}$ $\overline{100}$ of TaSi₂ thin layers (transition from a very disordered state to a crystalline one) were revealed by the mea-
surement of argon release which was included in the $\frac{1}{\alpha}$, $\frac{1}{\alpha}$, determined from the TIGRA experiments in the diagnostics structure defects of the thin film prepared by sputter- thermostimulated radon release from futile powder. (Heating in air ing $[34]$ (see Fig. 14). at a heating rate of 5 K min⁻¹).

TIGRA was used in the characterization of thermal behaviour of titania (rutile) The experimental results obtained by Beckman [17], made it possible to deter- mine the activation energies of the inert gas migration

250 nm thick TaSi₂ thin film deposited by sputtering in argon and the rutile diffusion of the temperature dependence of radon release antimosphere (argon atoms were captured during the sputtering imum of the temperature atmosphere (argon atoms were captured during the sputtering process in the thin film defects). The rate at the temperature of 1450 K corresponds to the

Fig. 15. Upper part: The spectra of activation energy values (parts of rutile sample. Lower part: The temperature dependence of the

controlled by the mobility of the structure defects of different types. The experimental results were treated by means of the DIGS software using the "singlejump diffusion" model. The inert gas (radon) was introduced into the surface near layers of rutile in the low temperature glow discharge according to Jech [9]. The release of radon at the temperature from 300 to 900K (see Fig. 15-the temperature interval denoted α) corresponds to the continuous activation energy spectrum (spectrum a), where the E_{min} =19 kcal mol⁻¹, E_{max} =63 kcal mol⁻¹. The radon release rate at the temperatures above 1250 K is controlled by the diffusion characterized by a narrow spectrum of the activation energies: $E_{\text{min}} =$
45 kcal mol⁻¹, $E_{\text{mean}} =$ 57 kcal mol⁻¹, $E_{\text{min}} =$ 64 $\frac{1}{200}$ $\frac{1}{400}$ $\frac{1}{600}$ $\frac{1}{600}$ 1000 1200 1400 $\frac{1}{1400}$ $\$ kcal mol⁻¹ (the energy spectrum b). The activation Fig. 14. Temperature dependence of argon release rate from energy values of this process correlate well that of self-
250 an this T-Si, this film dependent to contrain in second diffusion of Ti cation in the rutile lattice

inert gas release denoted as **stage IIb**, by Matzke [8] I. Bergstrom, F. Brown, J.A. Davies, J.S. Geiger, R.L. Craham
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9. Conclusion

The thermostimulated inert gas release analysis, [17] I.N. Beckman, in V. Balek and J. Tolgyessy (Eds.), Emanation TIGRA, can be recommended as a sensitive tool for Thermal Analysis, Suppl, to Russian Translation, Mir, the diagnostics of materials, making it possible to Moscow, Russia (1986) p. 309.

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