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In situ study of the temperature activated kinetics of water sorption in an epoxy varnish

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ABSTRACT

The dielectric permittivity of a polyepoxy varnish with a glass transition temperature (T_g) of 61 °C is monitored during immersion in a 0.5 M NaCl aqueous solution, by electrochemical impedance spectroscopy, at various temperatures in the range [21; 78] °C. The sorption curves are first analysed with a Fickian approach. An Arrhenius-type dependence of the apparent coefficient of diffusion is observed, with no visible influence of T_g . Below 46 °C, the sorption curves clearly do not obey Fick's second law of diffusion. On another hand, the Kohlrausch-Williams-Watts function (KWW) satisfactorily fits all the isothermal sorption curves. The KWW time constant has the same temperature activation as the apparent coefficient of diffusion, and likewise is unaffected by T_g . The KWW stretching exponent displays a strong dependence on T_g , correlates with the Fickian behaviour observed in the rubbery state, and is thought to be governed by molecular heterogeneities in the polyepoxy varnish as a function of temperature. The originality of this work lies in the *in situ* analysis of water uptake kinetics (constant immersion), which is performed above (rubbery state) and below (glassy state) the T_g of the polyepoxy coating.

1. Introduction

When it comes to protecting metal surfaces against corrosion in aggressive environments, polymer-based coatings are the most wide-spread solution. Leaving the inclusion of inhibitive pigments aside, the "barrier effect" is the main protective mechanism of organic coatings: by limiting the diffusion of water, oxygen, and possibly ions through the coating down to the metal substrate. The ability of the coating to maintain a low ionic conductivity over time is generally accepted as the limiting factor of the corrosion initiation (high resistance between anodic and cathodic sites) [1,2] even though this assertion is still debated [3]. Regardless of the actual protective mechanism at the microscopic scale, most failure modes leading to corrosion damage at the macroscopic scale (*e.g.* blistering [4]) simply do not occur without water uptake first taking place, hence the abundant literature on the subject in the coatings community [5,6].

The diffusion of water (and other species involved in corrosion) through the coating highly depends on the polymer's physicochemical

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structure. These protective coatings often display relatively low glass transition temperatures, Tg, in the vicinity of the ambient temperature. In humid environments, or in immersion, water uptake has a plasticizing effect which leads to a T_g-depression: typically 10 °C [7] to 20 °C [8] per percent of water uptake in polyepoxies. An initially glassy polymer coating may therefore be brought to the rubbery state as a consequence of water uptake, which has great implications on its mechanical (the modulus decreases [9]) and electrical properties (ionic conductivity generally increases [10]). The glass transition may also be expected to influence the diffusion of water molecules in the polymer. A rather widespread conception is that water occupies the free volume in the polymer matrix [11,12]. This idea is debated, in particular by Merdas et al. [13], based on the observation of a unique activation energy of water diffusivity across the glass transition of several polymers, which is in contradiction with the increasing free volume in the rubbery state that suggests different activation energies on both sides of Tg [14]. On another hand, Halary [15] argues that the sub-T_g β -relaxation, mainly associated with the mobility of the hydrophilic hydroxyl groups [16],

governs the diffusion of water molecules in epoxy-based materials, rather than the mobility associated with the glass transition. Perhaps these ideas are not contradictory: as summarized by Soles and Yee [17], "the topology (nanopores), polarity, and molecular motions act in concert to control the transport" of water in amine-cured epoxies. They also emphasize the fact that the quantity of unoccupied volume (they call nanopores) accessible to the water molecules is not a rate-limiting factor for the diffusion.

Usually, studies on diffusion in polymers rely on gravimetric measurements. When performed at elevated temperatures, especially when it comes to thin coatings (typically less than 100 μ m-thick), partial desorption during the weighing step may become a problem that can be alleviated by using *in situ* apparatus. In the organic coatings community, electrochemical impedance spectroscopy (EIS) is commonly used to monitor the water uptake [18–21] but rarely at elevated temperatures [22]. This *in situ* non-destructive technique eliminates the need to periodically remove the sample from immersion.

In the present study, EIS experiments are optimized for the characterization of the water sorption kinetics (particularly at short times) in a low T_g polyepoxy varnish, that is an unfilled coating with only the required constituents to produce a continuous, macroscopic defect-free film. These measurements are performed at various immersion temperatures ranging from ambient to 78 °C, allowing to study the diffusion mechanism on both sides of T_g . The main objective of this study is to analyse the influence of the glass transition and water-induced plasticization on the sorption kinetic parameters as determined by EIS measurements. A link between the macroscopic sorption and the physical structure at the microscopic scale is sought.

2. Experimental

2.1. Polyepoxy varnish samples

A two-component high-solids polyepoxy varnish was manufactured by Peintures Maestria (Pamiers, France). It consisted of an epoxide as base and a mixture of polyamidoamines as hardener. The liquid varnish was deposited by air spraying onto S235JR steel plates (200 mm \times 100 mm \times 3 mm). Prior to spraying, the steel plates were sand-blasted to an average roughness of 8 \pm 1 μ m. After 21 days of cure at 21 °C in a climate-controlled room, the varnish was 182 \pm 15 μ m thick (measured by an ultrasonic probe). Freestanding films were also prepared by bar coating deposition onto polypropylene substrates and peeling after cure, with a final thickness of 200 \pm 20 μ m. To stabilize the physicochemical structure (elimination of residual solvent and completion of residual crosslinking) of both the supported and freestanding samples, a postcuring step was achieved in an oven at 160 °C for 30 min.

2.2. Differential scanning calorimetry (DSC)

A Q2000 calorimeter, manufactured by TA Instruments, was used to determine the glass transition temperature of the polyepoxy varnish in both dry and wet states. Small pieces of a freestanding film were cut out to a total mass of approximately 10 mg and put in a standard sealed aluminium pan. A 10 mg-sample was also harvested from a piece of film previously immersed in 0.5 M NaCl at 70 °C for 2 h, in order to saturate the water absorption by the sample and erase the physical ageing. This "wet" sample was sealed in a hermetic TzeroTM pan with 10 µL of water to limit the sample dehydration inside the pan. For each sample, three temperature ramps were applied (heating, cooling, heating) at 20 K/min under a helium flow. The temperature range for the analysis of the dry sample was [-50; 150] °C. For the wet sample, it was limited to [5; 80] °C to prevent water from freezing (below 0 °C) and vaporizing (100 °C). The glass transition temperatures were determined at the midpoint of the heat capacity jump in the cooling ramps.

2.3. Broadband dielectric spectroscopy (BDS)

A Novocontrol BDS 4000 impedance analyser was used to perform dielectric measurements (in the temperature range [-150; 150] °C and in the frequency range [5 × 10⁻²; 10⁶] Hz). A gold plated cylindrical stainless steel electrode (40 mm diameter) was placed on top of a dry polyepoxy varnish sample, in the supported film configuration (the steel substrate acted as the lower electrode). In the complex domain, the applied sinusoidal voltage (1.5 V peak) divided by the measured current yielded the complex impedance, $Z^*(\omega)$, of the sample. The material's complex dielectric permittivity, $\varepsilon^*(\omega)$, was then calculated as a function of temperature and frequency by the WinDeta software, based on Eqs. (1) and (2).

$$\varepsilon^{*}(\omega) = \varepsilon^{'}(\omega) - i \ \varepsilon^{\prime\prime}(\omega) = \frac{1}{i \ \omega C_{v} \ Z^{*}(\omega)}$$
(1)

with
$$C_v = \frac{\varepsilon_v \mathscr{A}}{\ell}$$
 (2)

where $\omega=2\,\pi\,$ f is the angular frequency of the applied voltage, $Z^*(\omega)$ the measured complex impedance of the sample, and C_v the capacitance of the equivalent vacuum-filled (ϵ_v is the vacuum permittivity) parallel plate capacitor formed by two electrodes of area $\mathscr A$ (here, the area of the 40 mm top electrode) separated by the distance ℓ (here, the coating thickness).

In the present work, only the real part of the dielectric permittivity at high frequency (94 kHz) was analysed as a function of temperature (in the range [0; 100] $^{\circ}$ C) and compared to the electrochemical impedance spectroscopy measurements.

2.4. Electrochemical impedance spectroscopy (EIS)

Impedance measurements were performed on supported polyepoxy varnish films immersed in a 0.5 M NaCl solution (conventional EIS medium and high enough electrical conductivity to have a negligible influence on the total impedance), with a REF600+ impedance analyser manufactured by Gamry Instruments. Similarly to the BDS measurements, the coated steel acted as the working electrode while a graphite rod partially immersed in the electrolyte acted as both the counter and reference electrodes (two electrodes configuration). A custom-made double-jacketed glass cell was used to regulate the solution and sample temperature by means of a water flow in the double wall. A stable cell temperature control ($\pm 0.5^{\circ}$ C over tens to hundreds of hours) was achieved by using a Corio CD-200F refrigerated/heating circulator, manufactured by Julabo. A Pt100 probe was used to accurately monitor the solution temperature without interfering in the EIS measurements. Isothermal impedance measurements were performed in the temperature range [21; 78] °C, in the frequency range [10; 10⁶] Hz, with a 500 mV^{RMS} perturbation at the open circuit potential. Loops were programmed within the Gamry software to acquire EIS spectra every 2 min right after filling the cell with the electrolytic solution. Both the reduced frequency range and short open circuit measurement time (30 s) were compromises allowing such closely spaced EIS measurements, required to capture the fast sorption processes (especially right after the immersion). The interval between measurements was increased (up to 15 min) when a slowing down of the sorption process was observed. The relatively high ac perturbation was motivated by the increased signal-tonoise ratio it provided, especially for the initial high impedance measurements (various amplitudes were tested which led to the 500 mV optimum). Moreover, no evidence of corrosion was observed after the various measurements, neither visually nor in the impedance spectra.

A unique system composed of a polyepoxy varnish sample and cell was used throughout the whole EIS study, in order to limit the sources of variability (thickness, application defects, probed area). Prior to each EIS sorption monitoring, the sample and cell assembly were put to dry overnight in an oven set to 65 $^{\circ}$ C. Immediately after taking the sample

out of the oven, a rubber film was stretched over the glass cell to limit the sorption of ambient moisture. The water circulator was then quickly connected to the cell and equilibrated to the desired temperature, while a flask containing 200 mL of 0.5 M NaCl was placed in the circulator bath. After a few minutes, once the NaCl solution reached the desired temperature, it was poured in the EIS cell, marking the start of the immersion time axis in the following figures (uncertainty on initial time: 1 s). The graphite electrode and Pt100 probe were, as quickly as possible, immersed and the EIS measurement loop started. Prior solution equilibration and fast handling allowed to limit the delay between immersion and the first EIS spectrum acquisition to an average of 120 s. A first series of isotherms between 31 and 75 °C was acquired sequentially, followed by the 21 °C and finally the 78 °C isotherms. The monotonous evolution observed between 75 and 78 °C (the most critical range considering the fast sorption kinetics at high temperatures) allowed to dismiss any significant influence of the previous temperature of exposure.

In the present work, the dielectric permittivity formalism was chosen to represent the EIS results. It was preferred to the usual capacitance or impedance formalisms because it is intrinsic (does not depend on the sample geometry) and because the rules of mixtures used to assess the fraction of absorbed water are based on the real part of the dielectric permittivity. The latter was therefore calculated from the complex impedance data by means of Eq. (3), arising from Eq. (1).

$$\varepsilon'\left(\omega\right) = \frac{-Z(\omega)''}{C_{v}(Z'(\omega)^{2} + Z''(\omega)^{2})}$$
(3)

where $\varepsilon'(\omega)$ is the real part of the complex dielectric permittivity, ω the angular frequency of the applied voltage, $Z'(\omega)$ and $Z''(\omega)$ the real and imaginary parts of the measured sample impedance, and C_v the capacitance of the equivalent vacuum-filled parallel plate capacitor (see Eq. (2)) formed by two electrodes of area \mathscr{A} (here, the area of the EIS cell = 34.2 cm^2) separated by the distance ℓ (here, the coating thickness).

3. Results and discussion

3.1. Magnitude of the water-induced plasticization process

Fig. 1 shows the first cooling DSC ramps of the polyepoxy varnish, in the dry and wet states.

The glass transition temperatures (T_g) are indicated with dotted lines in Fig. 1. Thanks to the use of a hermetic pan and the addition of a small quantity of distilled water prior to sealing the pan, the hydration level of



Fig. 1. DSC thermograms (cooling ramp, -20 K/min) of polyepoxy varnish free-standing film samples: dry (continuous line) and wet (dashed line).

the sample is thought to be similar to a complete immersion (as in EIS measurements, in particular). Therefore, the 25 °C decrease in T_g observed between the dry and wet samples represents the maximal measurable water-induced plasticization of the polyepoxy network. The magnitude of this plasticizing effect highly depends on the chemistry of the epoxy-based resin and hardener used, but comparable values of the water-induced T_g -depression have been reported in the literature: 24 °C on an epoxy-based adhesive [23], 10–20 °C on various epoxy-based materials at 2 wt% water uptake [7], 14 °C for a DGEBA/DETA system at 2.2 wt% water uptake [24], and again 20 °C for a polyepoxy matrix and some silica-filled composites [25].

During the sorption monitoring by EIS measurements, the T_g of the varnish is expected to vary between 61 $^\circ C$ at zero immersion time and 36 $^\circ C$ when the sorption reaches the saturation plateau.

3.2. In situ isothermal impedance monitoring

The sorption process is monitored by acquiring closely spaced EIS spectra as soon as the sample is immersed in the 0.5 M NaCl solution. The same protocol is repeated at various temperatures in the range [21; 78] °C. From each EIS spectrum, the real part of the dielectric permittivity at 10^5 Hz is calculated and plotted against the square root of immersion time (Fig. 2). This frequency is chosen, based on the analysis of the full spectra [26], to avoid the contribution of the α dielectric relaxation (frequency range $[10^{-2}; 10^3]$ Hz, depending on the temperature) and instrument accuracy problems in the 10^6 Hz range.

Based on the assumption that any substantial variation of the sample's permittivity during the immersion is mostly caused by the absorption of water, which has a much higher permittivity (80 at room temperature [27]) than a polymer coating (around 3 in general), the plots of the permittivity as a function of immersion time will be referred to sorption curves. This approach has been described by Bellucci and Nicodemo [28] where Eq. (4) is formulated in terms of the film capacitance, which equals the permittivity multiplied by the sample geometry. In other words, this assumption means that an increase in real permittivity (relative to the dry value) is proportional to the mass increase due to water uptake one would observe during a gravimetric measurement (as in Eq. (4)). The bulk dielectric measurement averages



Fig. 2. Isothermal real dielectric permittivity as a function of the square root of immersion time in the 0.5 M NaCl solution, as measured by EIS. The indicated temperatures correspond to the sample cell temperature, maintained during the measurement. The 21 °C experiment was monitored over 450 h to reach a saturation plateau (not entirely shown for clarity purpose).

any water concentration profile that may exist in the sample, which is equivalent to assuming a uniform concentration over the sample's thickness.

$$\frac{\Delta \varepsilon'(t)}{\Delta \varepsilon_{\infty}'} = \frac{\varepsilon'(t) - \varepsilon_{t=0}'}{\varepsilon_{\infty}' - \varepsilon_{t=0}'} \propto \frac{m_t(t)}{m_{\infty}}$$
(4)

where t is the immersion time, $\epsilon_{t=0}'$ is the permittivity at zero time (dry state), ϵ_{∞}' is the permittivity value at saturation, m_t and m_{∞} the analogous mass water uptake that would be measured during gravimetric measurements.

The sorption curves in Fig. 2 display a typical Fickian shape with an initial, apparently linear, rise in permittivity before transitioning to a saturation plateau at longer immersion times [6]. The higher the temperature is, the quicker a plateau is reached: saturation appears in less than 10 h at 78 $^{\circ}$ C, while around 300 h are needed at 21 $^{\circ}$ C. Thereafter, two methods are used to analyse these sorption curves. The first one is based on the classical assumption that the sorption process obeys Fick's second law of diffusion, and the second one makes use of the stretched exponential, in an attempt to reduce the number of initial assumptions. Both methods are described in the following.

i) Fickian analytical method

The Fickian analytical method is often encountered in the literature about water uptake in coatings. It consists in assuming a linear dependence of the initial sorption process on the square root of immersion time (up to 60% of the total absorption), based on the physical assumption that the flux of diffusing substance is proportional to the gradient in concentration. Fig. 3 is a zoom of the initial permittivity rise in Fig. 2 and features the corresponding linear regressions, performed over 50% of the total permittivity rise.

When extrapolated to t = 0, the linear regressions yield the permittivity of the polyepoxy varnish in the dry state, $\varepsilon'_{t=0}$, used as a normalization parameter in Eq. (4). J. Crank [29] formulated analytical solutions to Fick's second law of diffusion through various geometries. The relevant one for the present work is the plane sheet exposed to the diffusing solution on one side, initially derived by Daynes [30] (Eq. (5)). It should be noted that a constant (or unique) coefficient of diffusion, D_{Fick} , is assumed in Eq. (5). The slope of the linear regression allows calculating the apparent coefficient of diffusion, D_{Fick} , by means of Eq.



Fig. 3. Linear regressions (dashed lines) of the real dielectric permittivity (grey symbols) as a function of the square root of immersion time. Fits are performed over 50% of the total permittivity increase (33% at 21 °C).

(6) [28], which is the short immersion times approximation of Eq. (5) for a supported film. These equations, initially defined for gravimetric water uptake, are adapted to permittivity variations based on the assumption of Eq. (4).

$$\frac{\Delta \varepsilon'(t)}{\Delta \varepsilon'_{\infty}} = 1 - \frac{8}{\pi^2} \sum_{n=0}^{\infty} \frac{1}{(2n+1)^2} \exp\left(-D_{\text{Fick}} \frac{(2n+1)^2 \pi^2}{(2\ell)^2} t\right)$$
(5)

$$\frac{\Delta \varepsilon'(t)}{\Delta \varepsilon_{\infty}'} = \frac{2}{\ell'} \sqrt{\frac{\mathsf{D}_{\mathrm{Fick}}}{\pi}} \times \sqrt{t} \quad \text{when } \frac{\Delta \varepsilon'}{\Delta \varepsilon_{\infty}'} \lesssim 0.6$$
(6)

where t is the immersion time and ℓ the sample thickness.

The linear regressions are satisfactory (correlation factor > 0.999) for all the isotherms, except the 21 °C one, which shows a clear deviation from linearity when approaching 50% of total permittivity increase (not visible in Fig. 3). In that particular case, and even though the correlation factor is acceptable, the regression is performed up to 33% of the total permittivity increase. The permittivity increase at saturation, $\Delta \epsilon_{\infty}^{'}$, is required to determine D_{Fick} by means of Eq. (6). An average of the last values of the sorption curves in Fig. 2 is considered in this method. The fact that a constant plateau is indeed reached represents another assumption. In the present work, the saturation was arbitrarily considered reached when the overnight variation in permittivity was less than 0.1%. The D_{Fick} values are reported in the Arrhenius plot of Fig. 9a and will be discussed in section 3.3.

Knowing the permittivity both at zero time and at saturation allows normalizing (Eq. (5)) the sorption curves as in Fig. 4, in a similar way to the usual gravimetric measurements of water uptake. The normalization emphasizes the influence of temperature on the sorption kinetics, regardless of the initial and final values.

The apparent coefficient of diffusion, D_{Fick} , previously obtained from the linear regressions (Eq. (6)) can be injected in Eq. (5), to determine if the experimental sorption curves obey to it over the entire time range. A representative selection of sorption curves is reported in Fig. 5 along with their corresponding Fick's law.

Fig. 5 shows that at high temperatures (75 and 61 $^{\circ}$ C), Fick's law satisfactorily describes the experimental sorption curves. This holds true down to the 50 $^{\circ}$ C isotherm (not shown in Fig. 5 for clarity purposes). Below 46 $^{\circ}$ C, as the temperature decreases, the sorption curves gradually diverge from Fick's law. While the satisfactorily linear regressions of all



Fig. 4. Real dielectric permittivity variation, normalized to the total variation (saturation plateau), as a function of the square root of immersion time in the electrolytic solution (0.5 M NaCl).



Fig. 5. Real dielectric permittivity variation normalized to the total variation (grey symbols) and associated Fick's law based on the initial slope (black dashed lines), as a function of the square root of immersion time.

the isotherms at short immersion times could suggest a Fickian behaviour (Fig. 3), the global sorption curves below 46 °C indicate otherwise. This seemingly contradictory observation motivates the second approach, deprived of two assumptions made earlier, namely the initial linear dependence of sorption with respect to the square root of time, and the value of the saturation plateau read from experimental data.

ii) Stretched exponential approach

The second analytical method starts with the determination of the real permittivity of the polyepoxy varnish, $\varepsilon'_{t=0}$, by extrapolation of a power law of time (Eq. (7)) rather than of a linear regression with a fixed exponent of 0.5 (square root function). To be consistent with the previous linear regressions (Fig. 3), the power law fits, reported in Fig. 6, are performed over the same time ranges.

$$\hat{\epsilon}(t) = \hat{\epsilon}_{t=0} + A t^{n} \tag{7}$$



Fig. 6. Power law fits (dashed lines) of the real dielectric permittivity (grey symbols) as a function of the immersion time, over 50% of the total permittivity increase (33% at 21 °C).

where A is a constant, t the immersion time and n the power law exponent.

The fitted values of the extrapolated dry permittivities and exponent n are respectively reported as a function of temperature in Fig. 7a and Fig. 7b.

Fig. 7a indicates that $\varepsilon'_{t=0}$ is a slightly increasing function of temperature. The drying protocol comes with the expectation of a somewhat constant $\varepsilon'_{t=0}$, corresponding to the retrieval of the same initial dryness. The small variation of $\varepsilon'_{t=0}$ (overall 10%) is not inconsistent with an identical initial state from the perspective of water content. Temperature also affects properties of the polymer (thermal expansion, dipole polarizability) that can reflect on the value of $\varepsilon'_{t=0}$. This point will be more extensively discussed in section 3.4, with the analysis of broadband dielectric spectroscopy measurements in the dry state.

Fig. 7b shows that the exponent *n* increases with the temperature and spans over the range [0.43; 0.63]. A 20% deviation from the expected value of 0.5 questions the Fickian assumption made earlier. On another hand, the correlation factor of the power law fits with the exponent *n* left free are equally as satisfactory (>0.999) as those previously obtained with the linear regressions. Fit quality is therefore not decisive when it comes to determining the actual value of *n*. Nevertheless, the power law fits (as opposed to the linear regression versus the square root of time) are mainly used, here, to obtain "assumption-free" values of $\varepsilon'_{t=0}$ (Fig. 7a) for normalization purposes.

The increase in permittivity, $\Delta\epsilon'(t) = \epsilon'(t) - \epsilon'_{t=0}$, is then fitted with the Kohlrausch-Williams-Watts (KWW) function [31], also known as the stretched exponential (Eq. (8)). This parametric function was used extensively to empirically describe relaxation phenomena associated with the molecular mobility of polymers in the time domain [32]. The underlying idea is that a distribution of relaxation times yields a broadened response (mechanical, dielectric) which departs from the simple exponential decay known as a Debye relaxation.

$$\Delta \varepsilon'(t) = \Delta \varepsilon'_{\infty} \left(1 - e^{-\left(\frac{t}{\tau_{KWW}}\right)^{F_{KWW}}} \right)$$
(8)



Fig. 7. Real permittivity extrapolated at t = 0 (dry state) (a) and exponent n (b) yielded by the power law fits (Eq. (7)) of the EIS measurements (Fig. 6).

where τ_{KWW} (s) is the time constant, and $\beta_{KWW} \in]0;1]$ is the stretching exponent.

The stretching exponent of the KWW function, β_{KWW} , quantifies the wideness of the distribution, while the time constant, τ_{KWW} , is the mean relaxation time of the distribution. Today spectroscopic techniques (mechanical and dielectric) have mostly replaced static measurements in the time domain, and equivalent functions can be used in the frequency domain, such as the Havriliak-Negami equation [33]. By analogy with the analysis of polymer relaxations, using the KWW function to empirically account for a distribution of diffusion processes may make sense. For instance, Shen and Springer [34] found that a stretched exponential with a stretching exponent of 0.75 approximated well the Fickian sorption case, and Zeng and Xu [35] resorted to the stretched exponential to model water vapour sorption in porous cement-based materials.

Contrary to the first method, based on the Fickian hypothesis, the increase in permittivity at saturation, $\Delta \varepsilon_{\infty}^{'}$, can be left as a free parameter in the fitting procedure. This eliminates the assumption made earlier when reading the value on the supposed saturation plateau. The sorption curves normalized to the new sets of parameters $\{\varepsilon_{t=0}^{'}; \Delta \varepsilon_{\infty}^{'}\}_{T}$, respectively obtained from power law extrapolations at short times (Fig. 7) and KWW fits of $\Delta \varepsilon'(t)$ (not shown), are reported in Fig. 8. The corresponding KWW fits are also plotted as dotted lines.

As shown in Fig. 8, the KWW function satisfactorily fits the sorption isotherms over the entire temperature range. As opposed to Fick's laws (Fig. 5), there is no degradation of the fit quality below 46 °C. The very short immersion times (below 20–30% of the total permittivity increase) is the only region where the KWW function slightly diverges from the experimental data. However, this is not considered problematic here because the aim is to describe the overall sorption kinetics.

3.3. Analysis of the sorption kinetics parameters

The apparent coefficient of diffusion, D_{Fick} , and the KWW parameters, τ_{KWW} and β_{KWW} , are displayed in Fig. 9 as a function of reciprocal temperature to emphasize Arrhenius-type dependences, which appear linear in this representation. Diffusion processes indeed are often reported to obey an Arrhenius equation (Eq. (9)) [5,36].

$$D_{\text{Fick}}(T) = D_{\text{Fick},0} e^{-\frac{L_{d}}{k_{B}T}}$$
(9)

where D_{Fick 0} is the pre-exponential factor, E_a the activation energy and



Fig. 8. Normalized real dielectric permittivity variation (grey symbols) and associated Kohlrausch-Williams-Watts fits (black dashed lines), as a function of the square root of immersion time.



Fig. 9. Arrhenius plots of the apparent coefficient of diffusion (a), the time constant (b) and the stretching exponent (c) of the Kohlrausch-Williams-Watts function. The Arrhenius fitted curves appear as dashed lines.

 k_{B} the Boltzmann constant.

The apparent coefficient of diffusion has values in the range [9.2 10^{-8} ;2.6 10^{-6}] mm²/s between 21 and 78 °C, which is in very good agreement with a study by De'Nève and Shanahan [37] reporting values in the range [2.8 10^{-7} ;3.6 10^{-6}] mm²/s between 40 and 70 °C for a diglycidyl ether of bisphenol A/dicyandiamide network.

In first approximation, both D_{Fick} (Fig. 9a) and τ_{KWW} (Fig. 9b) display an Arrhenius-type dependence over the whole temperature range. Given the uncertainty margins, their activation energies (Table 1) can be considered equal, around 50 kJ/mol (or 0.5 eV). This suggests that both D_{Fick} and τ_{KWW} reflect the same diffusion process. A very good agreement is found with values from the literature on epoxy-based thermosets: 55 kJ/mol with an adhesive immersed in water between 30 and 45 °C [38], 54 kJ/mol with a DGEBA/TETA between 23 and 75 °C [39], 56 kJ/mol with an adhesive between 19 and 90 °C [40], 50 kJ/mol with a high-T_g glassy polyepoxy between 40 and 60 °C [41]. The excellent agreement with these previous studies, based on gravimetric measurements, shows that the temperature-dependent EIS measurements

Table 1	
Arrhenius activation energy for the sorption kinetic parameters D_{Fick} and τ	ĸww

	E _a (kJ/mol)	E _a (eV)
$D_{Fick} (mm^2/s)$ $\tau_{KWW} (s)$	$\begin{array}{c} 50\pm2\\ 54\pm2 \end{array}$	$\begin{array}{c} 0.52 \pm 0.02 \\ 0.56 \pm 0.02 \end{array}$

developed in the present work can satisfactorily be used to monitor sorption processes in polymer coatings.

Linde et al. [42] studied a series of polyepoxy materials based on a DGEBA resin with various hardeners and found that the activation energy of the water diffusivity between 20 and 200 °C lied in the range [44; 62] kJ/mol. In the same article, the authors showed that only one out of the six systems studied showed different activation energies on both sides of T_g. In the present work, the diffusion process similarly seems mostly unaffected by the glass transition of the polyepoxy network (indicated with grey vertical dotted lines in Fig. 9): the Arrhenius dependence is monotonous over the entire temperature range. This behaviour may be explained from the point of view of Merdas et al. [13] who assimilate the activation energy of diffusivity (in various polymers, including epoxy-based ones) to the formation/dissociation of hydrogen bonds between water molecules and hydrophilic (polar) sites on the polymer chain. In epoxy-based networks, the hydroxyl, amine and amide groups are likely to be the preferred sites of interaction with water molecules [17,43]. Under the assumption that the hydrogen bonding to polar sites of the polymer backbone governs the diffusion of water molecules, the impact of the main chain mobility at the glass transition on the free volume should not play a major role in the diffusion process, which would explain the monotonous temperature dependence of the coefficient of diffusion on both sides of T_g.

Fig. 9c shows that the stretching KWW exponent, β_{KWW} , has a nonmonotonous dependence on temperature: it transitions from an increasing function of temperature below 55 °C to a constant value above (unexplained deviation for 78 °C point). This more or less corresponds to the temperature above which the Fick's law starts to satisfactorily describe the sorption curves (Fig. 5), suggesting an underlying correlation of the two equations. In order to evidence this correlation, all of the Fick's laws plotted in Fig. 5 are fitted with the KWW function. The obtained stretching exponent, β_{KWW}^{Fick} = 0.79 \pm 0.01, is temperature independent and seems to be intrinsic to Fick's law (at least in the present study) and is in very good agreement with the value of 0.75 reported by Shen and Springer as a general approximation to Fick's law applied to a plane sheet [34]. A grey rectangle, labelled "Fickian zone" and corresponding to the value β_{KWW} takes in the Fickian case, is displayed in Fig. 9c. Above 55 °C, the plateau reached by the experimental β_{KWW} points matches the "Fickian zone", which confirms the visual evidence that the sorption process becomes Fickian. As 55 °C is reasonably close to the dry T_g of the polyepoxy network (Fig. 1), β_{KWW} seems to convey the influence of the molecular mobility. In the literature, it is often stated that the water sorption by polymers is Fickian in the rubbery state, and that deviations from Fick's law mainly occur in the glassy state [6]. Moreover, β_{KWW} provides a quantification of the deviation from Fickian diffusion: as β_{KWW} decreases, Fick's law less and less accurately describes the sorption curves.

While β_{KWW} is not directly associated with a distribution of relaxation times, as it is the case when the KWW function is used to analyse molecular mobility modes, the argument that some kind of distribution is present with regards to the diffusion of water molecules in the polymer network is sensible: a distribution of diffusion coefficients for instance. The temperature dependence of β_{KWW} could reflect an evolution of the diffusion medium occurring during the sorption process, as a result of the plasticization process. If the test temperature is higher than the dry $T_{g}\!,$ the polyepoxy varnish stays in the rubbery state over the whole sorption experiment. If the test temperature is lower than the dry Tg, the polymer is in the glassy state at the start of the experiment and plasticization occurs as soon as water molecules enter the network, possibly leading to the occurrence of the glass transition during the experiment (especially for temperatures between 36 and 61 °C, the wet and dry T_g, respectively). From the perspective of a water molecule diffusing in the polymer, the diffusion paths are more heterogeneous in the partially plasticized glassy polymer than in the continually rubbery one. It is striking to note that these heterogeneities reflected by low values of β_{KWW} do not influence the mean diffusional process, reflected by τ_{KWW} or D_{Fick} which both display the same activation energy. The discussion about the heterogeneity of the diffusion medium could meet with the concept of D and I areas defined by Mayne, which could be associated with low and high cross-linked regions of thermoset coatings [1]. Mainly D areas are thought to be involved in the initiation of under film corrosion, as they constitute preferential ionic pathways to the substrate [2]. There could be a link between the heterogeneity in cross-linking density and a distribution in diffusion coefficients in the glassy state, which becomes less significant in the rubbery state.

3.4. Assessment of absorbed water volume fraction

The fact that water uptake is responsible for the permittivity increase (Fig. 2) is a recurring assumption in the previous two sections. This section focuses on quantifying the water uptake by means of a rule of mixture between the real dielectric permittivity of the dry polyepoxy varnish and that of water. A simple linear 2-phase rule of mixtures [19, 44] is applied to the high-frequency EIS data (10^5 Hz) to calculate the water volume fraction at saturation, $\Phi^{v}_{H_2O}(t)$, as in Eqs. (10) and (11). This linear rule of mixture is chosen on the basis of various studies reporting its better agreement with gravimetric measurements [19,44, 45] when compared to the more widely used Brasher-Kingsbury equation [18].

$$\epsilon_{\infty}^{'}(T) = \Phi_{H_{2}O}^{v}(T)\epsilon_{H_{2}O}^{'}(T) + \Phi_{dry \ varnish}^{v}(T)\epsilon_{t=0}^{'}(T)$$
(10)

$$\Leftrightarrow \Phi_{H_2O}^{v}(T) = \frac{\varepsilon_{\infty}'(T) - \varepsilon_{t=0}'(T)}{\varepsilon_{H_2O}'(T) - \varepsilon_{t=0}'(T)}$$
(11)

where $\Phi^v_{H_2O}(T)$ and $\Phi^v_{dry\ varnish}(T)=1-\Phi^v_{H_2O}(T)$ are the volume fractions of water and dry varnish at the temperature T, $\epsilon^{'}_{H_2O}(T)$ and $\epsilon^{'}_{t=0}(T)$ their respective real permittivity and $\epsilon^{'}_{\infty}(T)$ the sample's real permittivity at saturation.

The real permittivity of water is usually considered to be a constant, close to 80 [46], but this holds at 20 °C only. It is indeed a decreasing function of the temperature, as thermal motions hinder the polarizability of a dipolar substance (except when a relaxation mode is reached). Malmberg and Maryott [27] reported accurate measurements of the real dielectric permittivity of liquid water as a function of temperature in the range [0; 100] °C and fitted the experimental data with a third degree polynomial (Eq. (12)).

$$\epsilon'_{\rm H_{2}O}(T) = 87.740 - 0.400 \ T + 9.39810^{-4} \ T^{2} - 1.41010^{-6} \ T^{3}$$
 (12)

The application of Eq. (12) shows that in the temperature range [21; 78] °C of the present study, the dielectric permittivity of liquid water is in the range [80; 61], which represents a significant 24% decrease between 21 and 78 °C that needs to be taken into account when estimating the water uptake. Eq. (11) was applied to all the isotherms in Fig. 2 with the $\varepsilon'_{t=0}(T)$ values obtained from the power law extrapolations (Fig. 6) and $\varepsilon'_{\infty}(T)$ values yielded by the KWW fits (Fig. 8). The water volume fraction absorbed at saturation by the polyepoxy network is plotted as a function of temperature in Fig. 10 (\Box symbols).

The water volume fraction at saturation increases from 1.3% at $21 \degree C$ up to 3.7% at 78 °C, which is a credible order of magnitude for a polyepoxy. However, the linear rule of mixture (Eq. (11)) does not account for any dimension changes as the temperatures increases (thermal expansion, swelling) that are to be expected, but difficult to dissociate from one another. An experimental correction of any temperature-induced permittivity contribution, except the swelling, which is expected to be in the same order of magnitude than the thickness uncertainty (typically less than 10% [23]), is proposed. It consists in comparing the temperature dependence of the permittivity at saturation (obtained by EIS) with actual measurements in the dry state, by broadband dielectric spectroscopy (BDS). Fig. 11 features BDS



Fig. 10. Water volume fraction in the polymer at saturation as a function of temperature, calculated by means of the linear rule of mixture (Eq. (11)) with: uncorrected values of both ε'_{∞} and $\varepsilon'_{t=0}$ (\Box), their correction by the BDS measurements (\bullet) and the additional correction of ε'_{∞} taking into account the plasticization effect (\blacktriangle).



Fig. 11. Real dielectric permittivity of the dry polyepoxy varnish as a function of temperature, measured by broadband dielectric spectroscopy at 94 kHz. The dashed lines correspond to linear fits and the dotted line indicates their intersection at 62 °C, which is very close to the calorimetric dry T_g .

measurements of the real permittivity of the polyepoxy varnish at high-frequency, as a function of temperature in the range [0; 105] $^\circ C.$

Two seemingly linear dependences (dashed lines) of the permittivity with respect to the temperature are observed below and above 62 °C (dotted line), with a factor of 4.6 between the two slopes. This behaviour closely resembles the shape of thermomechanical measurements in the glass transition region of polymers, which evidence thermal expansion coefficients several times higher in the rubbery state than in the glassy state. If this thermal effect on the BDS permittivity actually originates from dimensional changes, it is expected to occur during EIS measurements as well. By considering 21 °C as the reference point, this normalized increase (BDS) is subtracted to the relative increase in ε'_{∞} (T) (EIS, at saturation), as in Eq. (13). This yields the permittivity at water saturation corrected to the thermal effect also observed in the dry state,

therefore not linked to water uptake.

$$\epsilon_{\infty,corrected}'(T) = \epsilon_{\infty}'(21^{\circ}C) \times \left(1 + \left[\left(\frac{\epsilon_{\infty}'(T)}{\epsilon_{\infty}'(21^{\circ}C)}\right)_{EIS} - \left(\frac{\epsilon'(T)}{\epsilon'(21^{\circ}C)}\right)_{BDS}\right]\right)$$
(13)

The corrected permittivity values at saturation are reported in Fig. 12 (\bullet symbols), along with the uncorrected values (\square symbols).

To be consistent, the same correction has to be made to $\epsilon_{t=0}'(T)$ (not shown) before calculating the corrected water volume fraction with Eq. (11). The obtained water fraction values are displayed in Fig. 10 (\bullet symbols). Accounting for the supposedly dimensional thermal effect observed in Fig. 11 only reduces the water volume fraction by 8% at most (at 78 °C). Such a low impact can probably be explained by the fact that the subtraction in the numerator of Eq. (11), $\epsilon_{\infty}'(T) - \epsilon_{t=0}'(T)$, already cancels the majority of any contribution common to the dry and wet states.

However, if different dimensional changes occur in the wet and dry states, they do not cancel each other anymore. In particular, this would be true in the case of swelling, but also as a result of plasticization: the T_{o} -depression at saturation (61–36 °C) can be expected to accordingly shift the thermal effect observed in the dry state (Fig. 11) towards lower temperatures. This idea is the basis of the second correction of the water volume fraction reported in Fig. 10 (▲ symbols). In order to assess the effect of plasticization on the water uptake values, a correction can be generated by horizontally shifting the curve in Fig. 11 by 25 °C towards the low temperatures and applying Eq. (13) in the same way as the first correction (the $\epsilon_{t=0}^{'}(T)$ are still corrected to the unmodified BDS values of Fig. 11, because the polymer is in the dry state). This new correction, which makes physical sense in spite of the crude assumption made when simply shifting the BDS curve horizontally, has much more impact on the calculated values of the water volume fraction at saturation. A 33% decrease compared to the uncorrected values is observed at 78 °C. Perhaps more importantly, a tendency to reach a plateau is now visible in Fig. 10 (\blacktriangle symbols) when the uncorrected and first corrected water fractions seemed to keep increasing at a constant rate with increasing temperature (\Box and \bullet symbols).

The considerations about dimension changes and plasticization illustrate the complexity hidden behind simple rules of mixture like Eq. (11) or the Brasher-Kingsbury equation. They are based on debatable (sometimes clearly incorrect) assumptions. For instance, taking 80 as the



Fig. 12. Uncorrected real permittivity as extracted from the water saturation plateaus of EIS measurements (\square), its correction by the BDS measurements (\bigcirc) and an additional correction taking into account the plasticization effect in the form of a -25 °C shift of the BDS data (\blacktriangle), as a function of temperature.

permittivity of water at room temperature implies that water is in the liquid form and that it does not interact with the polymer, which is in contradiction with the plasticization phenomenon, precisely caused by water molecules bonding to the polymer chain. Plasticization, which can change the state of the polymer from glassy to rubbery during a sorption experiment, is another intrinsic limitation of the rules of mixture: they simply cannot account for this complexity. The correction attempts made earlier show that small variations of permittivity are magnified by the rule of mixture. The multiple sources of error on the permittivity values (extrapolations at zero time, value on the plateau, plasticization effect, dimension change by thermal expansion and swelling) dictate caution when using rules of mixture to obtain more than a rough esti-mate of the water uptake.

4. Conclusions

In this study, EIS was used to monitor the increase in the high-frequency real permittivity of a polyepoxy coating as a result of water uptake. These experiments were reproduced at various temperatures in the range [21; 78] °C so as to evidence the thermal dependence of the diffusion mechanisms and to assess the influence of the molecular mobility associated with the glass transition.

A traditional analysis based on the short-time approximation of the analytical solution to Fick's second law of diffusion allowed the determination of the apparent coefficient of diffusion of water in the coating. An Arrhenius-type activation (of roughly 50 kJ/mol) was found for this diffusion process, in good agreement with existing literature. Below 46 °C, Fick's second law of diffusion failed at describing the whole sorption curves. The stretched exponential (Kohlrausch-Williams-Watts function), on another hand, successfully fitted the sorption curves in the entire temperature range, providing a mean time constant and a stretching exponent both associated with the diffusion kinetics. The time constant seemingly evidenced the same Arrhenius-type activation as the apparent diffusion coefficient, which suggest that the initial sorption somewhat encompasses the whole diffusion process kinetics. The stretching exponent reflects a notion of distribution, probably related to diffusion paths. It displayed a strong dependence on the glass transition of the coating, contrary to the average time constant and the apparent coefficient of diffusion: an increasing function of the temperature leading, at the dry Tg, to a plateau value corresponding to a Fickian behaviour. This influence of the molecular mobility at Tg was explained by the water-induced modification of the diffusion medium during the sorption experiment. Above the dry Tg, plasticization did not significantly alter the diffusion medium (polymer is in the rubbery state at all times), while below the dry Tg, the plasticization of the initially glassy polymer resulted in an inhomogeneous distribution of diffusion paths responsible for the low values of the stretching exponent.

Sensible values (below 4%) for the absorbed water volume fraction were estimated by means of a simple linear rule of mixture. Corrections based on a dimensional change evidenced by dry BDS measurements were attempted, also taking into account the influence of plasticization. These corrections led to a decrease of the water uptake at saturation, and plasticization was shown to have the greater impact on the value (up to 33% decrease compared to the uncorrected value).

This study showed that temperature-controlled EIS measurements can replace the widespread gravimetric measurements when it comes to analysing the temperature dependence of the sorption kinetics in polymer coatings, which allows establishing behaviour laws and analysing the influence of the molecular mobility on diffusional processes. The partial desorption problem encountered when taking samples out of immersion to weigh them is alleviated by the *in situ* character of EIS. In particular, the very short delays between acquisitions in the order of a minute allow accurate monitoring of the fast initial sorption processes. Clearly this experimental method is suitable for the analysis of a wide range of polymer materials destined to be used in immersion (biopolymers for instance). However, when a precise determination of the water uptake is required, gravimetric measurements should be performed due to the uncertainty inherent to the rules of mixture.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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