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Alkylating Potential of Styrene Oxide: Reactions and Factors Involved in the Alkylation Process

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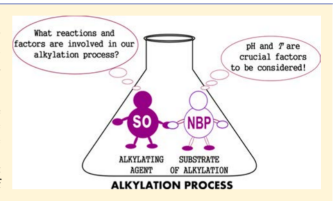
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ABSTRACT: The chemical reactivity of styrene-7,8-oxide (SO), an alkylating agent with high affinity for the guanine—N7 position and a probable carcinogen for humans, with 4-(p-nitrobenzyl)pyridine (NBP), a trap for alkylating agents with nucleophilic characteristics similar to those of DNA bases, was investigated kinetically in water/dioxane media. UV—vis spectrophotometry and ultrafast liquid chromatography were used to monitor the reactions involved. It was found that in the alkylation process four reactions occur simultaneously: (a) the formation of a β-NBP—SO adduct through an S_N2 mechanism; (b) the acid-catalyzed formation of the stable α-NBP—SO adduct through an S_N2′ mechanism; (c) the base-catalyzed hydrolysis of the β-adduct, and (d) the acid-catalyzed hydrolysis of SO. At 37.5



°C and pH = 7.0 (in 7:3 water/dioxane medium), the values of the respective reaction rate constants were as follows: $k_{\text{alk}\beta} = (2.1 \pm 0.3) \times 10^{-4} \,\text{M}^{-1} \,\text{s}^{-1}$, $k_{\text{alk}\alpha} = (1.0 \pm 0.1) \times 10^{-4} \,\text{M}^{-1} \,\text{s}^{-1}$, $k_{\text{hyd}}^{\text{AD}} = (3.06 \pm 0.09) \times 10^{-6} \,\text{s}^{-1}$, and $k_{\text{hyd}} = (4.2 \pm 0.9) \times 10^{-6} \,\text{s}^{-1}$. These values show that, in order to determine the alkylating potential of SO, none of the four reactions involved can be neglected. Temperature and pH were found to exert a strong influence on the values of some parameters that may be useful to investigate possible chemicobiological correlations (e.g., in the pH 5.81–7.69 range, the fraction of total adducts formed increased from 24% to 90% of the initial SO, whereas the adduct lifetime of the unstable β -adduct, which gives an idea of the permanence of the adduct over time, decreased from 32358 to 13313 min). A consequence of these results is that the conclusions drawn in studies addressing alkylation reactions at temperatures and/or pH far from those of biological conditions should be considered with some reserve.

INTRODUCTION

30 The reactivity of styrene-7,8-oxide (SO) has been the subject of 31 intense research from 1900 to the present, mainly due to its 32 industrial applications and biological activity. This oxirane is 33 widely used as a reactive plasticizer or diluent for epoxy resins, 34 in the production of phenethyl alcohol and styrene glycol and 35 its derivatives, and as a precursor for cosmetics, surface 36 coatings, and agricultural and biological chemicals. ^{1,2} It is also 37 the main *in vivo* mutagenic metabolite of styrene, a compound 38 extensively used in the production of plastic, rubber, fiberglass, 39 pipes, and food containers. ³ Thus, humans and other living 40 organisms are exposed to SO, whose carcinogenic, neurotoxic, 41 and mutagenic effects are well-known. ⁴

SO is a potent alkylating agent with the capacity to form adducts *in vivo* and *in vitro* with DNA bases, mainly at the N7-44 position of guanine. It has been classified as probable carcinogen for humans (2A IARC).

46 In contrast to other epoxides that only react with 47 nucleophiles through their less hindered carbon, SO can 48 achieve this through the two electrophilic carbons of the 49 oxirane ring, the primary (β) and the secondary (α) , 5–10 to 50 form the corresponding adducts.

Regarding the chemical reactivity and biological effectiveness 51 of SO with different nucleophiles, including DNA bases, there 52 are abundant references in the literature. ^{6,9,11–16} In spite of this, 53 many authors overlook some reactions that may be involved in 54 the alkylation mechanism and, hence, in the alkylation efficacy 55 of SO, such as (a) the potential formation of more than one 56 adduct between SO and the alkylation substrate ^{5–10,17–20} and 57 (b) the solvolysis reactions of the SO and of the adducts 58 formed. ¹⁹ Accordingly, in the present study the chemical 59 reactivity and alkylating potential of SO were investigated in 60 depth. The molecule 4-(*p*-nitrobenzyl)pyridine (NBP), which 61 has several advantages as an alkylation substrate, ²¹ was used as a 62 nucleophilicity model for DNA.

■ EXPERIMENTAL PROCEDURES

Styrene oxide (99%), 4-(p-nitrobenzyl)pyridine (98%), and triethyl- 65 amine (Et₃N 99%) were purchased from Sigma-Aldrich (Steinheim, 66 Germany). 1,4-Dioxane was from Panreac (Barcelona, Spain).

Caution: Styrene oxide is hazardous and should be handled 68 carefully. 69

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Scheme 1. Alkylation Mechanism of NBP by SO

To obtain deionize water, a Milli Q-Gradient (Millipore) system was used. Absorbance measurements were made with a Shimadzu UV-72 2401-PC spectrophotometer with a thermoelectric six-cell holder 3 temperature control system ($\pm 0.1~^{\circ}$ C). For pH measurements, a Metrohm 827 pH-meter with a glass electrode was used. The reaction temperature was kept constant ($\pm 0.05~^{\circ}$ C) with a Lauda Ecoline RE120 thermostat. All kinetic runs were performed in triplicate.

The alkylation reaction of NBP by SO was carried out (i) in an 78 excess of NBP to inhibit the formation of bis-alkylation adducts (1:2 79 NBP:SO) and to work in pseudo-first-order conditions and (ii) in water/dioxane medium, to render the NBP soluble. Alkylation mixtures were prepared by adding 1.0 mL of 0.005–0.03 M epoxide stock solution (SO in dioxane) to 100 mL of NBP solution (0.01–0.02 83 M) in water/dioxane medium. Acetate, phosphate, and borate buffers were used when needed to adjust the pH of the alkylation mixture.

Two different complementary techniques were used to investigate the formation and characterization of the SO-NBP adducts kinetically:

Vis Spectrophotometry. At different times, aliquots (2.4 mL) of the alkylation mixture (SO + NBP in a water/dioxane medium) were

the alkylation mixture (SO + NBP in a water/dioxane medium) were sollected and placed in a cuvette containing 0.6 mL of Et_3N to induce the deprotonation of the uncolored adducts $(AD_{un\alpha}$ and $AD_{un\beta})$, yielding the SO-NBP colored adducts $(AD_{\alpha}$ and AD_{β} ; Scheme 1), whose absorbance was measured at λ = 560 nm, where only both adducts absorb. Detailed experimental conditions are given in the figure and table legends. Rate constants were obtained by nonlinear pregression analysis of the absorbance/time data.

HPLC. A gradient-controlled Shimadzu ultrafast liquid chromatog-97 raphy system equipped with a diode array detector (UFLC-DAD) was 98 used to monitor the formation of each adduct. Chromatographic 99 separation was achieved with a Mediterranea Sea C18 column (25 cm $_{100} \times 1$ cm, 5 μm). Mobile phase A was acetate buffer (pH = 4.75 and 0.1 $_{101} M$), and mobile phase B was acetonitrile. 10% B was held for 5 min, $_{102}$ after which a gradient from 10% B up to 16% B was run over 8 min. $_{103} Then$, a gradient of up to 80% B was run over 1 min. 80% B was held $_{104} for 14$ min. The flow rate was set at 1.0 mL min⁻¹, and 200 μL of the $_{105} alkylation$ mixture was injected.

The hydrolysis reaction of SO was monitored by UV spectrophotometry. Aliquots of $50-200~\mu L$ of the stock solution of the epoxide in dioxane (0.085 M) were added to a thermostated cuvette containing 3.0 mL of the reaction mixture (HClO₄ + water + dioxane) at a constant pH. To convert the pH-meter output obtained in water/ dioxane mixtures into hydrogen ion concentrations, it is necessary to use a correction factor. However, with the solutions studied in this work (7:3 water:dioxane and [HClO₄] = 0.1–0.02 M), this correction can be neglected. However,

Absorbance at λ = 226 nm, where the difference between the signals 116 of SO and its hydrolysis product (styrene glycol, SG)¹⁶ is maximum, 117 was measured until no change was observed.

■ RESULTS AND DISCUSSION

Reaction Mechanism. In order to determine the reactions, 119 besides SO hydrolysis, involved in the alkylation mechanism, 120 UFLC-DAD analysis was performed. The results obtained 121 revealed the formation of two SO–NBP adducts with different 122 retention times (see Scheme 1). This result is in agreement 123 with their distinctive fragmentation patterns, obtained by 124 analysis of the alkylation mixtures by mass spectrometry in an 125 excess of NBP ($m/z=335,\ 317,\ 304$ for α-adduct, and m/z=126 335, 121 for β-adduct), as well as with previous results obtained 127 with other nucleophiles. 6,13,19,20,25

Assignment of the UFLC-DAD peaks was performed by 129 comparison of their retention times, UV—vis spectra, and 130 kinetic profiles with those of the β -adduct formed between 131 NBP and p-nitrostyrene oxide, a substituted styrene oxide 132 whose alkylating potential has been studied by us previously. ²⁶ 133 The adducts identified were α -NBP—SO (AD_{un α}), $t_{\rm R}=11.0$ 134 min, and β -NBP—SO (AD_{un β}), $t_{\rm R}=8.5$ min (see Scheme 1). 135 These two types of adduct also result in alkylation at the N-7 136 position in the reaction between SO and guanosine, 137 deoxyguanosine, and the single- or double-stranded DNA *in* 138 *vitro* under physiological conditions. ^{5,7,8,18}

Figure 1 shows the profile of a typical kinetic run with 140 ft UFLC-DAD, carried out by monitoring the area of the $AD_{un\alpha}$ 141

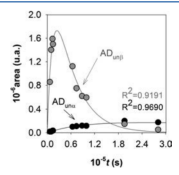


Figure 1. Kinetic profiles obtained by monitoring the variation in the UFLC-DAD peak area, AD_{un β} (gray circles) and AD_{un α} (black circles), over time in 7:3 (v:v) water/dioxane media. T=37.5 °C; pH = 7.50; [SO]_o = 1.7×10^{-4} M; [NBP]_o = 2.0×10^{-2} M; λ = 254 nm.

and $AD_{un\beta}$ peaks over time. As can be observed, while the 142 $AD_{un\alpha}$ signal remains fairly stable along the reaction time, 143 $AD_{un\beta}$ undergoes hydrolysis.

It should be pointed out that depending on the reaction rates 145 involved in the NBP alkylation process, two different 146

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Scheme 2. Reactions Involved in NBP Alkylation by SO in Aquo-Organic Media

Styrene glycol (SG)

$$k_{alk\beta}$$

NBP

HO

N

 $k_{alk\beta}$

NBP

HO

 $k_{alk\alpha}$

NBP

OH

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

 $k_{alk\alpha}$

NBP

147 approaches can be used: (i) considering only the alkylation 148 reaction; (ii) taking into account all the reactions occurring 149 simultaneously. Since abundant instances of the use of a poor 150 mechanistic framework to determine the alkylating potential of 151 SO can be found in the literature, $^{5-10,17-20}$ in our case the 152 second approach was applied (Scheme 2) and included the 153 following: (a) the formation of the α -NBP-SO and β -NBP-154 SO adducts; (b) hydrolysis of the β -adduct (the α -adduct 155 remained stable along the reaction); and (c) SO hydrolysis.

Equations 1–3 can be readily deduced from the mechanism depicted in Scheme 2. In these equations, $k_{\rm alk}$ is the global salkylation rate constant, defined as the sum of the second-order rate constants for alkylation through the α and β carbons, $k_{\rm alk}\alpha$ and and $k_{\rm alk}\beta$, respectively. Since NBP is present in a large excess, its concentration can be assumed to be constant, i.e. [NBP] = 162 [NBP]₀.

$$r = \frac{d[AD_{un\alpha}]}{dt} = k_{alk\alpha}[NBP][SO]$$
 (1)

$$r = \frac{\mathrm{d}[\mathrm{AD}_{\mathrm{un}\beta}]}{\mathrm{d}t} = k_{\mathrm{alk}\beta}[\mathrm{NBP}][\mathrm{SO}] - k_{\mathrm{hyd}}^{\mathrm{AD}}[\mathrm{AD}_{\mathrm{un}\beta}]$$
(2)

$$r = \frac{-d[SO]}{dt} = [SO](k_{alk}[NBP] + k_{hyd})$$
 (3)

Rate constants were determined using the NBP test²¹ (see 167 Scheme 1) by monitoring the variation in absorbance at $\lambda =$ 168 560 nm (see above).

Absorbance at time t ($A_{\rm AD}$) can be expressed as the sum of 170 the absorbances of the colored β - and α -adducts:

$$A_{\rm AD} = A_{\rm AD\beta} + A_{\rm AD\alpha} \tag{4}$$

The absorbance of each adduct at time t was obtained from 173 eqs. 1 and 2 by applying the Lambert—Beer law to the 174 respective integrated rate equations in terms of concentration:

$$A_{\text{AD}\alpha} = \frac{\varepsilon_{\alpha} l k_{\text{alk}\alpha} [\text{NBP}] [\text{SO}]_{\text{o}}}{k_{\text{alk}} [\text{NBP}] + k_{\text{hyd}}} (1 - e^{-(k_{\text{alk}} [\text{NBP}] + k_{\text{hyd}})t})$$
(5)

$$A_{\text{AD}\beta} = \frac{\varepsilon_{\beta} l k_{\text{alk}\beta} [\text{NBP}] [\text{SO}]_{\text{o}}}{(k_{\text{alk}} [\text{NBP}] + k_{\text{hyd}}) - k_{\text{hyd}}^{\text{AD}}} (e^{-(k_{\text{hyd}}^{\text{AD}})t} - e^{-(k_{\text{alk}} [\text{NBP}] + k_{\text{hyd}})t})$$

$$(6) _{176}$$

Equation 4 can thus be written as follows:

$$A_{\rm AD} = \frac{a}{b-c} (e^{-ct} - e^{-bt}) + \frac{d}{b} (1 - e^{-bt})$$
(7) ₁₇₈

with a, b, c, and d being parameters obtained by the nonlinear 179 fitting of the $A_{\rm AD}$ /t data and defined as

$$\begin{split} a &= k_{\rm alk\beta} [{\rm NBP}] [{\rm SO}]_{\rm o} \varepsilon_{\beta} l; \quad b = k_{\rm alk} [{\rm NBP}] + k_{\rm hyd}; \\ c &= k_{\rm hyd}^{\rm AD}; \quad d = k_{\rm alk\alpha} [{\rm NBP}] [{\rm SO}]_{\rm o} \varepsilon_{\alpha} l \end{split} \tag{8}$$

The goodness of the data fit to eq 7, where the parameters a 182 and d include a first-order reaction dependence on [SO] and 183 [NBP], while the parameter b includes a first-order dependence 184 on [NBP], reveals that the formation of the α - and β -adducts 185 occurs through S_N2' and S_N2 mechanisms, as expected. This, 186 together with the preference of SO for the N7-position of 187 guanine over other nucleophile positions of the DNA 188 suggests that NBP is a suitable model to 189 investigate the reactivity of SO with DNA. 21

The good fitting of the experimental $A_{\rm AD}/t$ data obtained 191 under neutral conditions (Figure 2) to eq 7 supports the 192 f2 proposed mechanism.

Determination of the Kinetic Parameters Involved in 194 **the NBP Alkylation Process.** As regards the global alkylation 195 rate constant, $k_{\rm alk}$ (eq 3), the value of this was obtained from 196 the slope of the $b/[{\rm NBP}]$ plot ($b=k_{\rm alk}[{\rm NBP}]+k_{\rm hyd}$), $k_{\rm alk}=197$ (3.1 \pm 0.4) \times 10⁻⁴ M⁻¹ s⁻¹ (37.5 °C, pH 7.0). Equation 9 was 198 used to determine $k_{\rm alk\beta}=(2.1\pm0.3)\times10^{-4}$ M⁻¹ s⁻¹, and 199 hence $k_{\rm alk\alpha}=(1.0\pm0.1)\cdot10^{-4}$ M⁻¹ s⁻¹ (37.5 °C, pH 7.0), 200 assuming the ratio $k_{\rm alk\alpha}/k_{\rm alk\beta}=33/67$ (obtained previously for 201 the reaction of SO with deoxyguanosine in water/ethanol, 202 50:50 (v:v), at 37.0 °C) ¹⁹ for the alkylation of NBP by SO at 203 37.5 °C. This value was accepted, since it is known that in 204 protic solvents the $k_{\rm alk\alpha}/k_{\rm alk\beta}$ ratio is solvent-independent 11 and 205 is similar for the alkylation of substrates analogous to 206

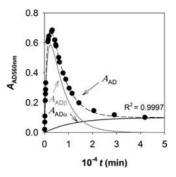


Figure 2. Experimental $A_{\rm AD}/t$ kinetic profiles in the formation of the SO–NBP adducts in 7:3 water/dioxane media (black circles), nonlinear fitting to eq 7 (dashed line), and the corresponding profiles calculated from eqs 5 and 6 for the α-NBP–SO (black line) and β-NBP–SO (gray line) adducts, respectively. $\lambda = 560$ nm; T = 37.5 °C; pH = 7.0; [NBP] $_{\rm O} = 1.99 \times 10^{-2}$ M; [SO] $_{\rm O} = 7.69 \times 10^{-5}$ M.

 $_{207}$ guanosine $^{18-20}$ by p-substituted styrene oxides at the same $_{208}$ temperature.

$$k_{\text{alk}\beta} = \frac{k_{\text{alk}}}{(k_{\text{alk}\alpha}/k_{\text{alk}\beta}) + 1} \tag{9}$$

With the values of the alkylation rate constants and those of the parameters a and d, the values of ε_{β} and ε_{α} (λ = 560 nm) were obtained (eqs 10 and 11): ε_{β} = (8 ± 1) × 10³ M⁻¹ cm⁻¹ and ε_{α} = (40 ± 8) × 10³ M⁻¹ cm⁻¹, respectively.

$$\varepsilon_{\beta} = \frac{a}{k_{\text{alk}\beta} l[\text{SO}]_{\text{o}}[\text{NBP}]} \tag{10}$$

214

2.15

$$\varepsilon_{\alpha} = \frac{d}{k_{\text{alk}\alpha} l[\text{SO}]_{\text{o}}[\text{NBP}]}$$
(11)

Regarding the SO hydrolysis reaction, its rate constant was obtained: (i) indirectly, by using the values of parameters b and $k_{\rm alk}$ (eq 12), and (ii) directly, by spectrophotometric measurements (see Experimental Procedures).

$$k_{\text{hvd}} = b - k_{\text{alk}}[\text{NBP}] \tag{12}$$

Since the absorbance, A, of the hydrolysis reaction mixture at time t is the sum of the resulting styrene glycol $(SG)^{16}$ and 223 epoxide contributions, eq 13 can be written.

$$A = l[SO]_{o}(\varepsilon_{SG} + (\varepsilon_{SO} - \varepsilon_{SG})e^{-k_{hyd}t})$$
(13)

Excellent fitting of the experimental data to eq 13 was 226 obtained (Figure 3a). In addition, the SO hydrolysis rate 227 constants determined from eq 13 revealed a high degree of 228 convergence with respect to those calculated from the b 229 parameter (Figure 3b). This supports the methodology used to 230 determine the reaction rate constants involved in the alkylation 231 process.

The values of the β -NBP-SO adduct hydrolysis rate constant, $k_{\rm hyd}^{\rm AD}$ (see Table 1), are given directly by parameter c 4 (eq 8).

The values of the fitting parameters a, b, c, and d, as well as those of $k_{\text{alk}\beta}$, $k_{\text{alk}\alpha}$, k_{hyd} , and $k_{\text{hyd}}^{\text{AD}}$ were used to determine some associated kinetic parameters that may be useful to investigate possible correlations between chemical reactivity and biological activity. Three such useful parameters are the fraction of the alkylating agent that eventually forms the α - and β -adducts, f_{α} and f_{β} , respectively (eqs 14 and 15), and the adduct lifetime, AL. The sum $f = f_{\alpha} + f_{\beta}$ ranges from 0 to 1, with 0 meaning that

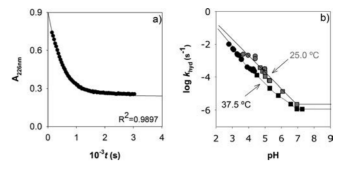


Figure 3. (a) Kinetic profile of a typical kinetic run for SO hydrolysis (experimental data (circles) and fitting to eq 13 (continuous line)). (b) Variation in the SO hydrolysis rate constant with the acidity of the medium in 7:3 (v:v) water/dioxane media obtained directly (circles) and indirectly (squares) at 25.0 °C (gray circles and squares) and 37.5 °C (black circles and squares), respectively.

Table 1. Influence of Temperature on the Rate Constants of the Reactions Involved in the Alkylation of NBP by SO and Values of Some of the Associated Kinetic Parameters^a

T (°C)	25.0	27.5	30.0	32.5	35.0	37.5			
$10^4 k_{\rm alk\beta} \ ({ m M}^{-1} \ { m s}^{-1})$	0.7	0.9	1.1	1.3	1.5	2.1			
$10^4 k_{\rm alk} ({ m M}^{-1} { m s}^{-1})$	0.2	0.3	0.5	0.4	0.7	1.1			
$10^4 k_{\rm alk} \ ({ m M}^{-1} \ { m s}^{-1})$	0.9	1.2	1.6	1.7	2.2	3.2			
$10^6 k_{\rm hyd} \ ({\rm s}^{-1})$	1.0	0.8	0.8	0.8	2.8	4.2			
$10^6 k_{ m hyd}^{ m AD}~({ m s}^{-1})$	0.5	0.6	1.0	1.5	1.9	3.1			
f_{eta}	0.50	0.57	0.54	0.62	0.42	0.40			
f_{lpha}	0.13	0.18	0.24	0.19	0.19	0.20			
f	0.63	0.75	0.79	0.81	0.61	0.60			
AL_{eta} (min)	17377	15209	8419	6810	3622	2185			
$^{a}7:3$ (v:v) water/dioxane media; pH = 7.0.									

all the initial epoxide is hydrolyzed and with 1 meaning that the 243 epoxide reacts exclusively with the NBP, yielding alkylation 244 adducts. These parameters have been correlated previously 245 with the mutagenic potency of different compounds. 21,26,28–31 246

The adduct lifetime gives an idea of the permanence of the 247 adduct over time. Thus, it is only meaningful for the unstable β - 248 adduct. AL $_{\beta}$ is defined as the area under the curve $[AD_{\beta}]/t$ per 249 unit of alkylating agent concentration (eq 16). 21,26,28–31 250 Consequently, it can be calculated from the area under the 251 curve $A_{AD\beta}/t$ (i.e., Figure 2), as shown in eq 16. Calculation of 252 this parameter is based on the rate constants of the three 253 reactions involved, $k_{\rm hyd}$, $k_{\rm alk}$, and $k_{\rm hyd}^{\rm AD}$. As may be seen from eq 254 16, AL $_{\beta}$ is also related to the alkylating efficacy, f_{β} (i.e., the 255 higher the f_{β} value and the lower the adduct hydrolysis rate 256 constant, the higher the AL $_{\beta}$).

$$f_{\alpha} = \frac{k_{\text{alk}\alpha}[\text{NBP}]}{(k_{\text{alk}}[\text{NBP}] + k_{\text{hyd}})} = \frac{k_{\text{alk}\alpha}[\text{NBP}]}{b}$$
(14) ₂₅₈

$$f_{\beta} = \frac{k_{\text{alk}\beta}[\text{NBP}]}{(k_{\text{alk}}[\text{NBP}] + k_{\text{hyd}})} = \frac{k_{\text{alk}\beta}[\text{NBP}]}{b}$$
(15) ₂₅₉

$$AL_{\beta} = \frac{\int_{t=0}^{t=\infty} [AD]_{\beta} dt}{[SO]_{o}} = \frac{\int_{t=0}^{t=\infty} A_{AD\beta} dt}{\varepsilon_{\beta} l [SO]_{o}}$$
$$= \frac{k_{\text{alk}\beta} [NBP]}{(k_{\text{alk}} [NBP] + k_{\text{hyd}}) k_{\text{hyd}}^{AD}} = \frac{f_{\beta}}{k_{\text{hyd}}^{AD}} = \frac{f_{\beta}}{c}$$
(16) 260

Influence of Temperature on the Kinetic Parameters lively Involved in the Alkylation Process. Table 1 shows the values of the above parameters determined in the 25.0–37.5 °C range.

As can be observed, whereas the values of f_{α} and f_{β} do not 266 show any clear trend with temperature, AL_{β} is strongly 267 influenced by it, being higher at lower temperatures. This 268 means that the stability of the β -adduct is higher at lower 269 temperatures; that is, a greater accumulation of the β -adduct 270 occurs with time.

It can also be seen that the $k_{\rm alk\beta}/k_{\rm alk\alpha}$ ratio is inversely related to temperature, showing an increase from 67/33 to 78/22 with a decrease in temperature from 37.5 to 25.0 °C. This trend, which seems to be common for the reaction of SO with nucleophiles similar to NBP in protic solvents, highlights the relevant role played by temperature on the alkylating capacity of SO. This result also suggests that any conclusions drawn about the chemicobiological correlations established from alkylation reactions with styrene oxide at temperatures far from those of biological conditions should be taken with some reserve.

The activation parameters calculated with the Arrhenius and 283 Eyring–Wynne–Jones equations, $E_{a\beta}=(64\pm4)~{\rm kJ~mol^{-1}};~E_{a\alpha}=284=(99\pm5)~{\rm kJ~mol^{-1}};~\Delta G^\#_{\beta(25.0~^{\circ}{\rm C})}=(96\pm8)~{\rm kJ~mol^{-1}};~{\rm and}~285~\Delta G^\#_{\alpha(25.0~^{\circ}{\rm C})}=(99\pm19)~{\rm kJ~mol^{-1}},~{\rm revealed~that~the~formation}$ 286 of the activated intermediate state of the α -adduct is enthalpy-287 controlled whereas that of the β -adduct is not.

Influence of the Acidity of the Medium on the Kinetic 289 Parameters Involved in the Alkylation Process. A series of 290 experiments was performed in media of different acidities in the 5.0 < pH < 8.0 range, because experimental limitations did not 292 allow the reaction rate constants from eqs 7 and 8 to be determined outside this range. (Since at pH < 5.0 the values of the hydrolysis rate constant are equal to or higher than those of the pseudo-first-order alkylation rate constant, calculation of the individual values is not possible. At pH > 7.7 some unidentified byproducts appear.) For the reaction at the β carbon of SO, it was observed that the alkylation rate constant was not subject to any influence of pH. The β -NBP-SO adduct was significantly hydrolyzed in neutral aquo-organic media (Table 2), whereas the hydrolysis of the β -adduct is basic-302 catalyzed and can be considered negligible in acid aquo-organic 303 media (Table 2), as occurs with other oxiranes. ^{26,30} In contrast, 304 alkylation through the α carbon is acid-catalyzed (Table 2),

Table 2. Influence of pH on the Rate Constants of the Reactions Involved in the Alkylation of NBP by SO and Associated Kinetic Parameters^a

pН	5.29	5.81	6.57	6.95	7.27	7.69
${10^5 k_{\rm alk\beta} \atop { m s}^{-1}} \ ({ m M}^{-1}$	7.30	7.15	7.09	6.98	7.29	7.20
$10^{5}k_{\text{alk}\alpha} \left(\text{M}^{-1}\right)$		4.27	1.94	1.81	0.71	0.58
${10^5 k_{\rm alk} \choose { m s}^{-1}} ({ m M}^{-1}$		11.42	9.03	8.79	8.00	7.78
$10^6 k_{ m hyd} \ ({ m s}^{-1})$	20.89	7.08	2.03	1.05	0.89	0.19
$10^7 k_{\rm hyd}^{\rm AD}~({\rm s}^{-1})$		0.79	2.59	4.81	8.44	10.43
f_{eta}	0.07	0.15	0.37	0.50	0.59	0.83
f_{α}		0.09	0.10	0.13	0.06	0.07
f		0.24	0.47	0.63	0.65	0.90
AL (min)		32358	23874	17377	11635	13313

^a7:3 water/dioxane media; T = 25.0 °C.

with the protonated transition state probably being favored by 305 the positive charge stabilization on that atom, 9,32 such that the 306 α -NBP—SO adduct formed remains stable along the reaction 307 (Figure 2). It should be noted that although consideration of 308 the stabilities of the adducts formed by SO is important, in the 309 literature they are usually overlooked. 7,27,33

The hydrolysis of styrene oxide is also influenced by the 311 acidity of the medium. In the pH 2–8 range, the rate constant 312 k_{hyd} can be expressed as the sum of the spontaneous and 313 catalytic hydrolysis rate constants (eq 17).

$$k_{\text{hyd}} = k_{\text{hyd}}^{\,\text{o}} + k_{\text{H}^{+}}[\text{H}^{+}]$$
 (17) ₃₁₅

In 7:3 water/dioxane medium the values of $k_{\rm hyd}$ at 37.5 and 316 25.0 °C are given by the expressions $k_{\rm hyd37.5} = (2.5 \pm 0.8) \times$ 317 $10^{-6} + (18 \pm 1)$ [H⁺] (s⁻¹) and $k_{\rm hyd25.0} = (1.0 \pm 0.1) \times 10^{-6} +$ 318 (6.5 \pm 0.5) [H⁺] (s⁻¹) (Figure 3b).

The reaction order with respect to $[H^+]$, determined in pH < 320 5 acidic media at 25.0 and 37.5 °C, was found to be one.

The importance of the concurrent SO hydrolysis in the study 322 of NBP alkylation by the epoxide is such that at pH < 5 we 323 observed that the value of the above parameter $b = k_{\rm alk} [{\rm NBP}] + 324$ $k_{\rm hyd}$ gave the value of $k_{\rm hyd}$ directly because it was significantly 325 higher than that of $k_{\rm alk} [{\rm NBP}]$. In spite of this, in the literature it 326 is unusual to find studies that take SO hydrolysis into account 327 (see above).

As can be observed (Table 2), all the associated kinetic 329 parameters are clearly influenced by the acidity of the medium, 330 except f_{α} . This means that a small fraction of the initial SO, 331 around 9–13%, is converted into α -adduct regardless of the pH 332 of the medium. Additionally, the fraction of β -adduct formed is 333 substantially influenced by pH and is higher than that of the α - 334 adduct. Thus, the fraction of total adducts formed increases 335 from 24% to 90% of the initial SO in the pH 5.81-7.69 range. 336 This means that in acid media SO hydrolysis is predominant, 337 whereas in neutral-basic media adduct formation predom- 338 inates, particularly that of the β -adduct (the $k_{\text{alk}\beta}/k_{\text{alk}\alpha}$ ratio 339 increases from 63/37 to 93/7 due to an increase in the pH of 340 the medium from 5.8 to 7.7). This is important because it could 341 be related to the alkylating activity of styrene oxide in the 342 neutral biological medium. The results highlight the importance 343 of choosing the appropriate kinetic approach and of considering 344 all the reactions involved in the alkylation process. Neglecting 345 them would lead to mistaken conclusions about the SO 346 alkylating potential. Despite this, scrutiny of the literature $_{347}$ shows how frequently this is the case; $_{5-10,17-20}^{5-10,17-20}$ that is, when $_{348}$ using the NBP test or another spectroscopic method, the 349 increase in absorbance at a fixed time is considered to be a 350 measure of the reactivity or alkylating potential of SO³⁴⁻³⁷ (this 351 alkylating potential has sometimes even been used to establish 352 correlations between chemical and biological activity). 34,36

Another interesting result is that although the f_{β} value 354 declines with the acidity of the medium, β -adduct stability 355 clearly increases with it.

A consequence of the above is the strong influence of the 357 acidity of the medium on the reactivity of styrene oxide as an 358 alkylating agent, which underscores the importance of this 359 analysis in the search for possible chemicobiological correla- 360 tions.

Influence of the Composition of the Reaction $_{362}$ Medium on the Alkylation Process. Figure 4a shows the $_{363}$ f4 strong dependence of $k_{\rm alk\beta}$ and $k_{\rm alk\alpha}$ on the dielectric constant, $_{364}$ D, of the reaction medium in different water/dioxane mixtures, $_{365}$

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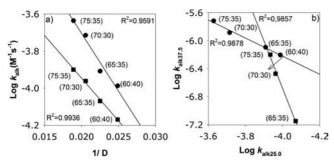


Figure 4. (a) Variation in the alkylation rate constants (37.5 °C) with the dielectric constant of the reaction medium, and (b) isokinetic correlations in the formation of both the α - and β -adducts (black squares and black circles, respectively) in media of different composition.

366 with the alkylation rate increasing upon increasing *D*. This 367 behavior, consistent with a reaction between polar molecules in 368 which the activated complex is more polar than the reagents, ³⁸ 369 supports the proposed mechanism.

As is known, ^{30⁴} the existence of an isokinetic relationship can ³⁷¹ serve to defend the argument that the reactions of a series share ³⁷² a common mechanism. The members may differ in the ³⁷³ identities of a functional group, the length of a side chain, the ³⁷⁴ composition of the solvent, and so on. In this work we ³⁷⁵ considered the solvent composition, that is, the percentage of ³⁷⁶ dioxane in the reaction medium. A mathematical formulation of ³⁷⁷ the isokinetic effect is the linear relationship between two series ³⁷⁸ of log $k_{\rm alk}$ values measured at two temperatures T1 and T2: log ³⁷⁹ $k_{\rm alk}(T2) = a + b \log k_{\rm alk}(T1)$.

It should be noted that the meaning of the isokinetic 381 relationship is the existence of a *compensation effect* between the 382 values of the enthalpy, $\Delta H^\#$, and entropy of activation, $\Delta S^\#$, 383 such that the Gibbs free energy of activation, $\Delta G^\#$, is 384 approximately constant. The results shown in Figure 4b 385 support the idea of common mechanisms of formation of 386 both the α - and β -SO-NBP adducts, respectively bimolecular 387 nucleophilic $S_N 2'$ and $S_N 2$ substitutions, across the whole 388 composition range of the solvent mixture.

389 CONCLUSIONS

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- (i) NBP alkylation reactions through the α and β carbons of SO occur via S_N2' and S_N2 mechanisms, respectively.
- (ii) Neglecting some of the reactions involved in the alkylation process would lead to mistaken conclusions about the alkylating potential of SO.
- (iii) Temperature and pH are crucial factors because: (a) The lifetime of the unstable β-adduct is strongly influenced by temperature, and hence, the accumulation of this adduct is greater at lower temperatures; (b) The hydrolysis of the β-NBP-SO adduct is basic-catalyzed. The formation of the α-NBP-SO adduct is acid-catalyzed, with the α-NBP-SO adduct remaining stable along the reaction. The hydrolysis of SO is acid-catalyzed.
- (iv) Any conclusions drawn in studies addressing alkylation reactions at temperature and/or pH far from those found in biological conditions should be considered with some reserve when searching for chemicobiological correlations.
- (v) The existence of an isokinetic relationship between alkylation rate constants in different water/dioxane ratios reveals a compensation effect that supports the idea of

common mechanisms of formation of both adducts 411 across the whole composition range of the solvent 412 mixture.

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ABBREVIATIONS

fellowship.

 $A_{\rm AD}$, sum of the absorbances of the colored β - and α -adducts at 438 time t; ${\rm AD}_{\alpha}$, colored α -NBP—SO adduct; ${\rm AD}_{\beta}$, colored β - 439 NBP—SO adduct; ${\rm AD}_{{\rm un}\alpha}$, uncolored α -NBP—SO adduct; 440 ${\rm AD}_{{\rm un}\beta}$, uncolored β -NBP—SO adduct; AL, adduct lifetime; D, 441 dielectric constant; f_{α} and f_{β} , fraction of the alkylating agent that 442 eventually forms the α - and β -adducts, respectively; NBP, 4-(p- 443 nitrobenzyl)pyridine; SG, styrene glycol; SO, styrene-7,8-oxide; 444 UFLC-DAD, ultrafast liquid chromatography system equipped 445 with a diode array detector

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