# Spin probe interaction and mobility in confined *cyclohexane*: Effects of pore size and pore surface composition of *silica gel* matrices

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#### ABSTRACT

The spectral behavior of the spin probe *TEMPO* in the bulk *cyclohexane* (*CHX*) and its confined states in a series of *silica gels* (*SG*) using ESR is reported. The spectral parameter  $A_{zz'}$  and the characteristic ESR temperature  $T_{50G}$  of slow to fast regime transition of *TEMPO* change dramatically in the virgin *SG-SIL*'s compared to the bulk *CHX* sligtly depending on the pore size. These observations result from the interrelation between the mutual interaction of the *TEMPO* with the *SG*'s and the altered phase behavior of the *CHX* from DSC. This is supported by the modified (silanized) *SG matrices*.

#### 1. Introduction

The topic of bulk vs. confined organic *materials* is usually studied by a variety of classic experimental techniques. Thermodynamic, structural and dynamic behaviors of various *organic media* are investigated by means of calorimetry, diffraction and scattering, resonance as well as relaxation spectroscopies determining characteristic material properties, such as enthalpy, static and dynamic density fluctuations, electric and magnetic dipoles, etc - see general and special reviews [1–8]. Significant changes in various physical properties are often observed which are related to phase transitions and eventually, to a formation of new phase(s) induced in confined organic *medium* (*filler*) by spatial restriction or/and by wall surface of *confining* inorganic *matrix* (*confiner*) [1–4] as well as to complex dynamic properties [5–8]. The overall confinement effect is considered to be a result of the complex mutual interplay of the two main factors: (i) geometric restricting effects of the pores on the *medium* in a given *matrix* and (ii) the mutual interaction effects of the *medium* with the pore surface wall of the *matrix* [1-8].

On the other hand, non-standard experimental techniques using various molecular and even atomic-sized extrinsic probes, such as stable free radicals or positronium, respectively, via electron spin resonance (ESR) [9–18] or positron annihilation lifetime spectroscopy (PALS) [19–22] are utilized to characterize the bulk state [9–13] and especially, the confined organic media [14-18,19-22] in essentially smaller extent. One of the promising microscopic probes are the so-called spin probes of nitroxide type, e.g., 2,2,6,6-tetramethyl-piperidinyl-1-oxy (TEMPO) via ESR [14,15,23]. Advantages of nitroxide probes are their simple ESR spectra and their sensitivity to interaction with surroundings and to their reorientation dynamics in a given medium or/and matrix. However, in comparison to the standard techniques, one additional parameter comes into the characterization problem. Although the spin probe concentration in a given organic medium is very small, typically of  $1-5x10^{-4}$  M, and the related *medium* perturbation by *extrinsic* probe seems to be apparently negligible, it is necessary to address the

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special aspect of potential interparticle interaction between this incorporated *probe* and the confined organic *filler* versus that between *the probe* and the pore wall of *the* confining inorganic *confiner*.

Recently, a potential role of this interaction aspect together with its impact into reorientation dynamics was suggested for apolar linear hydrocarbon, namely, n-hexadecane (n-HXD) in a series of polar unmodified silica gel (SG) matrices as probed by polar spin probe TEMPO [14]. The spectral behavior of TEMPO was charaterized via the extrema line separation of the triplet signal,  $2A_{zz}$  [9,11,13], as a function of temperature *T* from 100 K up to 360 K. The most pronounced features in the quasi-sigmoidal  $2A_{zz}$  vs. T plot are i) the relatively high  $2A_{zz}$  value in the immobilized state around 80 Gauss and ii) a transition from slow to fast motion regime at the convential characteristic ESR temperature,  $T_{50G}$ , i.e., the temperature at which  $2A_{zz^{\prime}} = 50$  Gauss corresponding to the correlation time of the *spin probe* of a few nanoseconds [11,13]. It was found that the TEMPO dynamics is very strongly slowed down with respect to that in the bulk medium because of the dramatic increase of the  $T_{50G}$ (bulk) value by about 100 K. Moreover, in a set of three unmodified SG-SIL matrices with the mean pore size ranging from 300 Å through 100 Å down to 60 Å, the  $T_{50G}$  value increases moderately within ca. 15 K. These findings were interpreted as a result of the preferential interaction of the polar TEMPO molecule with the polar silanol groups at the pore surface of the unmodified SG-SIL matrices. This explanation is supported by (i) the empirical fact that the high  $2A_{zz^{t}}$  value of the TEMPO in the fast regime in the confined state of all the unmodified SG matrices reaches about 40 Gauss which is comparable to the reported one of 39 Gauss for the similar nitroxide-type spin probe di-terc-butyl nitroxide (DTBN) adsorbed in bare silica gel at RT [24] and by (ii) theoretical results from quantum-mechanical calculations of the Azz' parameter for the isolated and the attached (bound) nitroxide radical to ...H-O group [25]. Consequently, the following natural question arises: Whether and under what conditions ESR is able to reflect *correctly* the altered structural-dynamic state of the confined organics of a given apolar structural type in various kinds of SiO<sub>2</sub>-based matrices differing in their surface structure, i.e., in unmodified vs. modified ones in comparison with the reference bulk state of the organics ?

Cyclohexane (CHX) was often used as a typical representative of apolar crystallizing organic media with the globular molecules in the confinement studies by standard thermodynamic, structural and dynamic techniques, such as DSC [26-33], x- and n-DIFF [34,35] and NMR [36-39]. Thus, Jackson and McKenna were the first to demonstrate systematically using DSC a strong effect of the spatial restriction on the main solid-to-liquid phase transition at  $T_{\rm m}$  of various apolar organics, including CHX, in a series of irregular control porous glasses (CPG) matrices modified by silalyzation to make their pore surface more hydrophobic with consequently better wetting to apolar media [26]. Later, Malhotra et al. showed using DSC that also the solid-to-solid (plastic crystal) phase transition temperature, T<sub>ss</sub> of CHX caused by small-scale reorientation of the quasi-spherical molecule is altered by confinement in a series of irregular modified silica gel (SG) matrices after the dehydratation and dehydroxylation treatment [27,28]. Next, the latter authors found out that both the freezing  $T_{\rm f}$  and melting  $T_{\rm m}$  temperatures of CHX in the three modified *silica gels* exhibit the non-monotonous course as a function the alkyl-type modifier substituent's length in the surface wall structure [29]. Finally, CHX was also investigated in various regular matrices, such as silicon [31] and unmodified (virgin) MCM-41 and SBA-15 [32]. Dosseh et al. [33] used the three regular unmodified SBA-15 matrices and reported the apparently very strong difference in the reduction of the melting temperature in comparison with the irregular modified CPG [26] and SG [27] ones ascribed to the difference in their surface chemistry. On the other hand, extrinsic probe studies of this model apolar cyclic aliphatic compound in some confined states are still missing.

Recently, we have investigated *CHX* in the bulk state by a combination of two extrinsic *probe* techniques using ESR and PALS [40]. It was found that  $T_{50G}$  correlates with the *solid*- to-*solid* phase transition at  $T_{ss}$ (bulk) lying deep in the *solid* state about ca. 95 K below the *solid*-to-*liquid* (melting) point  $T_{m}$ (bulk). This agreement indicates that the reorientation of the *TEMPO* is closely related to the reorientation motion of the *CHX* molecules being responsible for the  $T_{ss}$ (bulk) transition. In the light of several afore-mentioned

standard characterization studies of confined *CHX* it is of interest to reveal the information potential of ESR by addressing the following two important apects, namely, molecular shape and mutual interaction aspect in the three*component system* with respect to the two-*component system* in standard techniques. To solve this question the *apolar cyclohexane (CHX) medium* formed by the quasi-spherical molecules which embedded in a series of both unmodified and two differently modified *silica gel (SG) matrices* is investigated by ESR and DSC techniques.

# 2. Experimental

#### 2.1. Materials

Cyclohexane (CHX) with the purity of 99.5% from Sigma–Aldrich, Inc., Germany was used.

A series of unmodified (virgin) and modified (silanized) *silica gels* KROMASIL<sup>®</sup> consisting of spherical particles with outer diameters of 10 µm were obtained from Eka Chemicals, Bohus, Sweden. The pore diameters  $D_{pore}$  in the virgin *silica gels* named as *SG60-SIL*, *SG100-SIL* and *SG300-SIL* were determined by *the supplier* to be 60, 100 and 300 Å. The virgin *silica gel SG100* matrix was chemically bonded by *dimethylalkylsilyl* groups of various length and the modified *SG100 matrices* were designed as *SG100-C4* (*dimethylbutylsilyl*) and *SG100-C18* (*dimethyloctadecylsilyl*). According to *the supplier* the total surface concentration of *silanol* groups is 8 µmol/m<sup>2</sup> in the original *SG100-SIL*, while the degrees of coverage, i.e., the average bonding densities of *modifier* in both the modified *SG100-C4* and *SG100-C18 matrices* are 3.9 or 3.7 µmol/m<sup>2</sup>, respectively.

As extrinsic probe, 2,2,6,6-tetramethypiperidine-1-oxyl, (TEMPO), from Sigma-Aldrich, Inc, Germany, was used by dissolving it in the liquid *CHX* at a very low concentration of  $5x10^{-4}$  M.

A series of the confined *spin systems* for ESR studies was prepared by drop-by-drop filling of a solution of the *CHX* doped with the spin probe *TEMPO* into the *accessible* pores of all the four *SG matrices* to achieve a completely filled (saturated) state of the *filler* (*CHX*) in the pores of the *confiners* (*SG*'s). The capillary forces allowed to fill the *accessible* pores of *SG* with *CHX* with no liquid remainders on the external surface of the *SG* grains. The *theoretical* and *real* mass fractions of the *filler* for the each *filler/confiner system*, as well as the ratio of the latter quantity to the former one are given in Table 1. Before the own filling procedure all the four *SG matrices* were dried at 393 K for several hours, to evaporate the adsorbed *water*.

### 2.2. DSC

DSC measurements were carried out on DSC 8500 from Perkin-Elmer based on a power compensation principle equipped with a CLN2 cooler. Calibration was performed with a series of three different standard substances: *indium, n-dodecane* and *n-heptane*. The sample masses were about 15 mg. All DSC measurements were carried out under nitrogen atmosphere using standard aluminium pans. The samples were first cooled with -10 K/min from room temperature down to 113 K and subsequently, they were measured with heating rate of +10 K/min from 113 K up to 303 K. The *solid-to-solid* and *solid-to-liquid* (melting) transition phenomena, i.e., endothermal effects in the DSC thermogram, were quantified by the onset *solid-to-solid* transformation temperatures,  $T_{\rm ss}(D_{\rm pore})$ , and the onset melting transition temperatures,  $T_{\rm m}(D_{\rm pore})$ , and the corresponding *solid-to-solid* transformation enthalpy,  $\Delta H_m(D_{\rm pore})$ , and *melting* transition enthalpy,  $\Delta H_{\rm m}(D_{\rm pore})$ , were normalized to the mass of *CHX*.

#### 2.3. ESR

ESR measurements of the spin systems *TEMPO/CHX* were performed with the X-band Bruker – ER 200 SRL spectrometer operating at 9.4 GHz with a Bruker BVT 100 temperature variation controller unit. ESR spectra of the doped *TEMPO/CHX systems* were slowly cooled with ca. -4 K/min rate and subsequently were recorded in heating mode

Table 1	
Physical parameters of a series of the used SG matrices according to the supplier and the confined G	CHX/SG systems.

Matrix	Pore diameter D <sub>pore</sub> , Å	Pore size distribution %	Pore volume V <sub>pore</sub> , cm <sup>3</sup> /g	Pore area $A_{\text{pore}}$ , m <sup>2</sup> /g	F <sub>CHX,theo</sub> * -	<i>F</i> <sub>CHX,sat</sub> ** -	%***
SG 60-SIL SG 100-SIL SG 100-C4 SG 100-C18 SG 300-SIL	60 100 300	$80 \pm 15$ $80 \pm 25$ $80 \pm 25$	1.20 0.85 0.92 0.93	546 312 327 112	0.483 0.398 0.417 0.420	0.479 0.381 0.399 0.298 0.411	99.1 96 95.7 97.8

\*  $F_{CHX,theo} = m_{CHX}/(m_{CHX} + m_{SG})$ , the theoretical mass fraction of *CHX medium* with respect to the *CHX/SG system* estimated using the density of *CHX* at room temperature,  $\rho_{CHX}(RT) = 0.779 \text{ g/cm}^3$  under the complete accessibility condition of all the pores for the *CHX medium*.

\*\*  $F_{CHX,sat} = m_{CHX}/(m_{CHX} + m_{SG})$ , the real experimental mass fraction of *CHX* in the *CHX/SG system* corresponding to the fully filled (saturated) situation of the *CHX* in the *CHX/SG system*.

\*\*\* % =  $F_{\text{CHX,sat}}/F_{\text{CHX,theo}} \times 100.$ 

over a wide temperature range from 115 K up to 350 K using temperature steps of 5–10 K. To reach thermal equilibrium, the samples were kept at a given temperature for 15 min before the starting of the spectra accumulations. In these measurement the temperature stability was  $\pm$  0.5 K. The microwave power and the amplitude of the field modulation were optimized to avoid signal distortion. Analysis of the ESR spectra was performed in terms of the extrema separation of the outermost lines of the triplet spectra,  $2A_{zz'}$ , as a function of temperature with subsequent evaluation of the spectral parameter of mobility  $T_{50G}$  [11,13] as well as further characteristic ESR temperatures,  $T_{Xi}$ , within both slow and fast motional regimes:  $T_{Xi}^{fost}$  [14,23].

#### 3. Results and discussion

# 3.1. DSC data

The macroscopic structural-phase-transition temperatures characterization of bulk *CHX medium* as well as of a series of the confined *CHX/SG samples* were characterized by means of DSC are summarized in Fig. 1 and Table 2.

The obtained thermodynamic data of both the phase transitions in the bulk *CHX* are in a plausible agreement with those from the



**Fig. 1.** Phase i.e. *solid-to-solid*,  $T_{ss}$ , (circles) and *solid-to-liquid* (melting),  $T_m$ , (diamonds) transformation temperatures as a function of inverse pore diameter,  $1/D_{pore}$  for *CHX* confined in irregular modified *CPG matrices* [26] (black points), irregular modified *silica gel matrices* [27] (blue points), in regular unmodified *SBA-15 matrices* [33,42] (green points) and irregular unmodified *silica gels SG–SIL matrices* [present work] (magenta points). Initial larger circle and diamond mark  $T_{ss}$  or  $T_m$  of the bulk *CHX* with  $1/D_{pore} = \infty$ , respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

literature:  $T_{\rm m} = 280.1$  K [26],  $T_{\rm ss} = 186$  K,  $T_{\rm m} = 281$  K,  $\Delta H_{\rm ss} = 79.8$  J/g and  $\Delta H_{\rm m} = 31.9$  J/g [27] and  $T_{\rm m} = 279.5$  K,  $\Delta H_{\rm m} = 31.3$  J/g [41]. On the other hand, the DSC characteristics of the *CHX* after insertion of the organic *filler* into the various inorganic *SG confiners* behave in different way depending on the quality of the pores surface. It is therefore usefull to divide and discuss these changes according the two types of surface composition of the used *SG matrices*.

First, in the series of the three virgin *SG300-SIL*, *SG100-SIL* and *SG60-SIL matrices*, both the phase transiton temperatures  $T_{ss}$  and  $T_m$  are decreased significantly with physical restriction, i.e., with the decrease of the mean pore diameter  $D_{pore}$  from 300 Å through 100 Å down to 60 Å. For the largest spatial limitation these reductions in the  $T_{ss}$  and  $T_m$  values reach -36.4 K or -61.5 K, respectively. Similarly, the dramatic reductions in the phase transformation enthalpies  $\Delta H_{ss}$  and  $\Delta H_m$  occur as they will be presented and discussed later. All these changes in the thermodynamic characteristics on confinement of the *CHX medium* reflect its significant structural-phase reorganization towards the increased rotation disorder in the *solid* state as well as the elevated amorphization of the *CHX medium* after its spatial restriction into the mesopores of the virgin *SG-SIL matrices* with respect to the highly rotation and translation ordered (crystalline) structure of *CHX* in the bulk state.

It is of interest to compare the present DSC data on three virgin *silica* gels with previous data for both the phase transitions of variously confined *CHX* in the literature [26,27,33]. Fig. 1 shows such a comparison with sets of data on three irregular *controlled porous glasses* (*CPG*) matrices modified by silylation with hexamethyldisilazane giving very short  $\equiv$ *Si*-*O*-*Si*(*CH*<sub>3</sub>)<sub>3</sub> grafts instead of the original *silanol* groups [26]. Furthermore, it provides  $T_{ss}$  and  $T_m$  of the five irregular *silica* (*Spherosil*) matrices modified by dehydratation and dehydroxylation [27] as well as on the three regular unmodified *SBA-15 matrices* [33,42]. As one can see, in the case the *solid*-to-*solid* transition the  $T_{ss}$  data are quite close to each other with sligthly larger difference for the irregular unmodified *SG-SIL matrices* indicating a rather weak

Table 2

Temperature and enthalpy of solid-to-solid ( $T_{\rm ss}^{\rm DSC}$ , ( $\Delta H_{\rm ss}^{\rm DSC}$ ) and solid-to-liquid (melting) ( $T_{\rm m}^{\rm DSC}$ ,  $\Delta H_{\rm m}^{\rm DSC}$ ) phase transformations and temperature differences  $\Delta T_{\rm i} = T_{\rm i}$ (conf) -  $T_{\rm i}$  (bulk), where  $T_{\rm i} = T_{\rm ss}^{\rm DSC}$  or  $T_{\rm m}^{\rm DSC}$  in the bulk *CHX* and a series of the various confined *CHX/SG systems*. Typical temperature and enthalpy uncertainties are up to  $\pm 1$  K and  $\pm 1.5$  J/g.

Sample	$T_{ m ss}^{ m DSC}$ K	$\Delta T_{ m ss}$ K	ΔH <sub>ss</sub> J/g CHX	T <sub>m</sub> <sup>DSC</sup> K	$\Delta T_{\rm m}$ K	ΔH <sub>m</sub> J/g CHX
CHX	187	$0 \\ -10 \\ -26.8 \\ -24.8 \\ -42 \\ -36.4$	78.5	280	0	32
CHX/SG300Å	177		22.2	264.3	-15.7	7.2
CHX/SG100Å	160.2		20.5	230	-50	2.8
CHX/SG100Å-C4	162.2		21	234	-46	9.4
CHX/SG100Å-C18	145		8.3	225	-55	10.5
CHX/SG60Å	150.6		19.7	218.5	-61.5	4

sensitivity to the type of the confining SiO2-based matrix, perhaps. On the other hand, as for  $T_{\rm m}$ , in addition to the generally larger depression of the melting transition temperature, we find a rather larger dependence on the type of  $SiO_2$ -based matrix. Moreover, the  $T_m$  values seem to group roughly into two classes reflecting the effect of surface composition on the  $T_{\rm m}$  reduction, being weaker for hydrophobic walls of the pores in the modified CPG's and SG's than for the hydrophilic ones in the unmodified SBA-15 and our series of unmodified (virgin) SG-SIL matrices. These findings suggest a complex picture for the confined CHX in various SiO<sub>2</sub>-based confiners. While the former transition is related to a *smaller*-scale rearangement connected with the rotational disordering of the CHX molecules and is only weakly dependent on the pore morphology and chemistry, the latter case is related to a *large*-scale phase transition due to translational disordering and depends sensivitely on various aspects of the pore morphology and chemistry, e.g. not only on pore regularity or irregularity, i.e., pore separability or pore interconnectivity, but also on pore sizes distribution as well as pore surface composition.

In addition, the effect of pore surface modification on the DSC characteristics can be evaluated in detail for the two modified SG100 matrices. One modification included replacement of a part of the original polar silanol  $\equiv$ SiOH groups by apolar ones, such as the shorter dimethylbutylsilyl groups  $\equiv$ Si-O-Si(CH<sub>3</sub>)<sub>2</sub>(CH<sub>2</sub>)<sub>3</sub>CH<sub>3</sub> in SG100-C4 matrix and the other one by the essentially longer dimethyloctadecylsilyl groups  $\equiv Si - O - Si(CH_3)_2(CH_2)_{17}CH_3$  in SG100-C18. In the former case of shorter grafts one observes the slight increase of  $T_{ss}$  and  $T_m$  by ca. +2 or +4 K, respectively, compared to the virgin SG100-SIL matrix. On the other hand, in the case of the significantly (more than 4 times) longer grafts in the SG100-C18, larger reductions of  $T_{ss}$  by about -42 K and of  $T_{\rm m}$  by about -55 K with respect to the bulk *CHX* is found. Interestingly, the latter decrease of  $T_{\rm m}$  is a bit about -5 K larger than the one for the virgin SG 100-SIL matrix. At first sight we could expect a further reduction in both the  $T_{ss}$  and  $T_m$  values with respect to the virgin SG100-SIL matrix due to the smaller effective pore size, but the situation is not so simple.

In the literature one similar DSC study on the *CHX* in the *SG80* matrix modified via dehydratation and dehydroxylation treament and in the two chemically-derivatized *SG80-C1* (trimethylsilyl, -O-Si (*CH*<sub>3</sub>)<sub>3</sub>) and *SG80-C6* (hexyl,  $-O-(CH_2)_5CH_3$ ) matrices exists [28]. It was found that the differences in the melting temperature  $\Delta T_m$ , defined as  $T_m(\text{conf})-T_m(\text{bulk})$  reach -27 K, -44 K and -31 K. These authors concluded that the observed non-monotonous trend in the melting temperature with the alkyl modifier length cannot simply be attributed only to the "shrinkage" of the pore size due to the presence of the short  $-O-Si(CH_3)_3$ , or the longer  $-O-(CH_2)_5CH_3$  groups on the silica gel surface wall as in the case of the aliphatic linear *n-alkane*, i.e., *n-decane*, where this intuitive supposition is found to be valid [28].

Our data on the modified SG100-C4 and SG100-C18 matrices seem to exhibit similar non-monotonous order of  $\Delta T_{\rm m} = -50$  K, -45 K and -55 K as a function of the modifying alkylsilyl group length in the SG100-SIL, SG100-C4 and SG100-C18 matrices, at first sight, with the rather, unexpected smaller reduction  $\Delta T_{\rm m} = -55 \,\text{K}$  for the CHX/ SG100-C18 system compared with the virgin SG60-SIL with  $\Delta T_{\rm m} = -61.5$  K. According to the detailed experimental and modelling study of the geometry of the pores in a series of empty modified SG's, the effective pore size is not given simply by the linear correction of the original pore diameter on the straight alkylsilyl modifier length because of its coiled minimized-energy conformation [43,44]. However, the situation is complicated in the presence of the confined organic medium in the modified confiner due to the possibility that the conformation of longer alkyl modifier may further be altered with respect to that in the empty SG matrix. Consequently, the observed trend results also from additional interaction between the apolar alkyl modifier and apolar CHX medium in the confined state by the modified confiner.

As for the energetic aspect of phase transitions, the *solid*-to-*solid* transformation enthalpy,  $\Delta H_{ss}$ , and the *solid*-to-*liquid* (melting)

transition enthalpy,  $\Delta H_{\rm m}$ , also decrease drastically in accord with the literature [26,28,29]. Thus, for the most restrictive situation in a series of the three virgin *SG-SIL matrices*, the  $\Delta H_{\rm ss}$  value is reduced by 75%, while the  $\Delta H_{\rm m}$  even by 87.5%. These values indicate very essential disordering and amorphization of the *CHX medium* after physical restriction.

Next, the corresponding transformation enthalpies have rather different trends. Thus, the  $\Delta H_{ss}$  value is comparable for the shorter *C4* modifier with the virgin *SG100-SIL* matrix and essentially lower for the longer *C18* alkyl group. On the other hand, the  $\Delta H_{m}$  increases with alkylsilyl chain lenght in comparison with the parent *SG100-SIL* matrix. This suggests that the orientation order of the *CHX* molecules decreases with an introduction of alkylsilyl groups, but in contrary, their translational order increases with respect to the rotation-translation situation in the the virgin *SG100-SIL* matrix.

All these changes in the thermodynamic characteristics indicate the essential structural-phase reorganization of the *CHX medium* towards significant amorphization after its spatial restriction into the mesopores of the virgin *SG-SIL matrices* and the modified *SG100-C4* and *SG100-C18 matrices* with respect to the highly ordered (crystalline) structure in the bulk state.

# 3.2. ESR data

Figs. 2 and 3 show the spectral parameter,  $2A_{zz'}$ , as a function of temperature for all the six investigated *CHX* systems. For the sake of clearness, the ESR data and their discussions are separated into two groups in dependence on the chemical composition of the pore surface of the *SG confiners*.

3.2.1. ESR data for CHX confined in the unmodified (virgin) SG-SIL matrices

Basically quasi-sigmoidal, significantly distinct  $2A_{zz}$ , vs. *T* plots depending on the physical, i.e. bulk or confined state of the *CHX medium* as well as on the pore size of the virgin *SG matrices* are found in Fig. 2. In the bulk *CHX medium*, the  $2A_{zz}$ , values at the lowest temperature of our measurements reach from 67 Gauss to ca. 32 Gauss at *RT*, being



**Fig. 2.** Spectral parameter  $2A_{zz'}$  of the spin probe *TEMPO* as a function of temperature in the bulk *CHX* and the three confined *CHX/SG 60, 100* and *300-SIL* systems demonstrating the effect of pore size  $D_{pore}$  in the unmodified *SG-SIL matrices*. The characteristic ESR temperatures  $T_{X1}^{slow}$ ,  $T_{50G}$  and  $T_{X1}^{fast}$  for the bulk CHX and  $T_{X1}^{slow}$ ,  $T_{X2}^{slow}$  and  $T_{50G}$  for the confined *CHX/SG 60,100* and *300-SIL* systems are marked by arrows. The characteristic DSC temperatures of both the phase transitions are also depicted by dashed lines for the solid-to-solid transitions at  $T_{ss}$  and by full lines for the melting tranformations at  $T_{m}$ .



Fig. 3. Spectral parameter 2Azz<sup>-</sup> of the spin probe *TEMPO* as a function of temperature in the bulk *CHX* and the three confined *CHX/SG100-SIL*, *CHX/SG100-C4* and *CHX/SG100-C18* systems demonstrating the dramatic effect of the pore wall composition in the modified *SG100-CX matrices*.

typical for *nitroxide free radicals* in *apolar* organic *media* [45–49]. The most pronounced change at the characteristic ESR temperature  $T_{50G}$  correlates with  $T_{ss}$  [40]. In addition, two weak changes in  $2A_{zz}$  can be found, one within the slow motion regime at  $T_{X1}^{slow} \sim 160$  K and the other within the fast one at  $T_{X1}^{fast} \sim 280$  K, the latter being related with the *solid*-to-*liquid* transition at the melting point  $T_m(bulk)$ .

Confinement of the spin probe *TEMPO* doped *CHX medium* in a series of three virgin *SG-SIL matrices* lead to very significant changes in both the  $2A_{zz'}$  value at 115 K and 300 K as well as in the characteristic ESR temperatures,  $T_{Xi}^{slow}$  and  $T_{50G}$  which mark numerous changes in the temperature dependencies.

As for the values of  $2A_{zz'}(115 \text{ K})$ , the drastic increase from 67 Gauss to about 80 Gauss in the immobilized state of the *spin probe TEMPO* in all the three confined *CHX/SG-SIL systems* is found. It indicates a strong change in the mutual interaction between the *polar TEMPO* molecules and the *polar* component of the confined *CHX/SG-SILsystem*, i.e., the virgin *SG-SIL matrices* compared to their relative weak interaction with the *CHX medium* alone. Consistently, the  $2A_{zz'}(300 \text{ K})$  values of *TEMPO* in the confined states ranging from 38 up to 43 Gauss in all the three *CHX/SG-SIL systems* are also significantly higher compared to ca. 32 Gauss for *TEMPO* in the *bulk CHX medium* in consistency with the literature data on the adsorbed *nitroxides* on pure *silica* [24].

On increasing temperature, the three confined *CHX/SG60-SIL*, *CHX/SG100-SIL* and *CHX/SG300-SIL* systems reveal the first weak anomalies in  $2A_{zz'}$  close to  $T_{X1}^{\text{slow}} \sim 150-160$  K, in the vicinity of the reduced  $T_{ss}$ (confined) transitions.

On further increase the temperature, the second stronger changes in  $2A_{zz}$ , are shifted from  $T_{X1}^{slow} \sim 160$  K in the bulk *CHX* to  $T_{X2}^{slow}$ (confined)  $\sim 215$ , 220 and 220 K for the virgin *SG300-SIL*, *SG100-SIL* and *SG60-SIL matrices*. Further enhancement of the temperature leads to the most pronounced transition from the slow to the fast motion regime that occur at  $T_{50G} = 273$ , 274 and 278 K, respectively. Comparison of the characteristic ESR temperatures  $T_{X1}^{slow}$  and  $T_{50G}$  with those of the phase transitions reveals a correlation of  $T_{X1}^{slow}$ (confined) with the decreased melting temperature for the smallest pores  $T_{m}(60$  Å) indicating a sensitivity of the surface-bonded reorienting *TEMPO probe* dynamics to the phase transition for the maximal confinement of the *CHX/SG60-SIL system*. On the other hand, the effect of the phase transition for the 100 Å and 300 Å restrictions is

not so apparent suggesting the local character of the change within the slow motion regime. Finally, the most pronounced slow to fast motion transitions lie essentially above the corresponding  $T_{\rm m}$ (confined), but in the relative vicinity also supporting the local character of the phenomenon. Interestingly, the  $T_{\rm 50G}$  values for our set of the confined *CHX/SG60-SIL*, *100-SIL* and *300-SIL systems* are about 25 K below those for another *apolar organics*, i.e., linear *n-alkanes*, such as *n-hexadecane* (*n-HXD*) in the same series of the virgin *SG-SIL confiners* [14] with  $T_{\rm 50G}$  = 298–313 K and *n-undecane* (*n-UND*) [49] in the same virgin *SG100-SIL* with  $T_{\rm 50G}$  = 298 K. This finding seems to indicate that for the *apolar* aliphatic organic *media* the  $T_{\rm 50G}$  value is somewhat dependent on the *cyclic* vs. *acyclic* topology of the *molecular constituents*.

# 3.2.2. ESR data for CHX confined in the modified SG100-C4 and SG100-C18 matrices

Similarly as in the previous case, basically quasi-sigmoidal, significantly distinct  $2A_{zz'}$  vs. *T* plots depending not only on the physical, i.e., bulk vs. confined, state of the *CHX medium*, but also on the pore surface composition of the *SG100 matrices* are observed. Thus, the  $2A_{zz'}$ (115 K) values for both the modified *SG100 confiners* lie in between those for the bulk *CHX medium*, i.e., 67 Gauss, and the virgin *CHX/SG100-SIL matrix*, i.e. ~80 Gauss, while for the *SG100-C4 confiner* is a bit smaller than for the *SG100-C18* one: 76 < 77 Gauss. The fact that both the  $2A_{zz'}$ (115 K) values do not reach the maximal  $2A_{zz'}$ (115 K) ~80 Gauss, being attributed to the *TEMPO* molecules attached to the pore wall, indicates that the *TEMPO* probe reflects the structural-dynamic changes of the various confined *CHX medium systems*.

In the modified *SG100-C4 matrix* with the short<u>er</u> *alkyl silyl* groups, the slow to fast transition with respect to the virgin *SG100–SIL matrix* with  $T_{50G} = 274$  K is shifted by 36 K to the lower temperature of  $T_{50G} = 238$  K. On the other hand, the other modified *SG100-C18 matrix* with essentially longer *alkyl silyl* groups causes a slowing down of the slow to fast transition at the  $T_{50G} = 323$  K compared to the the virgin *SG100–SIL matrix*. These mutual relationships between all the three types of *SG100 matrices* can be explained by considering the following two factors: effective pore size and pore surface composition.

Regarding the first factor, the effective pore size is reduced after any chemical modification by adding the alkyl groups [43,44]. It might imply that the dynamics should be slowed down with respect to that in

the *virgin SG100-SIL matrix*. Indeed, although this is found for the *SG100-C18* case, for the shorter modifier in the *SG100-C4 matrix* the opposite situation exists. According to the detailed experimental and modelling study of the geometry of the pores in a series of empty modified *SG's*, the effective pore size is not given simply by the *linear* correction of the original pore diameter on the straight alkylsilyl *modifier* length because of its *coiled* energy-minimized conformation [43,44]. However, in the presence of the confined organic *medium* in the *modified confiner*, the real conformation situation of the *modifier* might be complicated by the possibility that the conformation of longer alkyl silyl *modifier* can further be altered with respect to that in the empty *SG matrix*. Consequently, the observed trend results also from certain additional interaction between the *apolar alkylsilyl modifier* and *apolar CHX medium* in the confined state.

As for the second aspect, we recall that the average bonding density of modifier in both the modified SG100 matrices are quite close being  $3.9 \,\mu mol/m^2$  or  $3.7 \,\mu mol/m^2$ , respectively. This means that about 48.8% or 46.3% of the total surface concentration of the original silanol groups 8µmol/m<sup>2</sup> are substituted by the corresponding *modifier* indicating an almost comparable degree of coverage of the pore surface by the alkylsilyl groups. Evidently, the complete substitution of all the silanol groups at the original pore surface for the sterical reasons is impossible. Both the alkylsilyl groups C4 and C18 increase the hydrophobic character of the pores and this coverage of the pore surface decreases an approach of the TEMPO probes to the remaining unreacted silanol groups. Thus, they reduce the potential interaction the polar spin probe TEMPO molecules with them. However, the shorter alkylsilyl groups of *modifier* reduce the effective pore volume to an essentially smaller extent and this fact together with the similar coverage of the pore surface by apolar dimethylbutylsilyl groups results globally in a decrease of the T<sub>50G</sub> value with respect to the virgin CHX/SG100-SIL case, i.e., an acceleration of the transition from slow to fast motion regime. On the other hand, although the longer and coiled C18 groups [43] seem to change the pore surface to the hydrophobic character to an essentially larger extent with better coverage of the remaining unreacted silanol groups, the smaller effective pore diameter together with the larger probability of the presence of a certain fraction of the CHX molecules as well as the TEMPO ones in between the longer modifier chains leads effectively to the dominance of the latter aspect. As the result, the significant increase in the  $T_{50G}$  value with respect to the virgin SG100-SIL matrix. Evidently, in this case of CHX/SG100-C18 system ESR technique measures and reflect rather the qualitatively different confined system compared to the CHX/SG100-C4 and CHX/ SG100-SIL ones.

#### 4. Conclusions

The spectral behavior of the spin probe TEMPO in the bulk cyclic hydrocarbon cyclohexane (CHX) and its confined states in a series of silica gels (SG) and phase behavior using ESR or DSC, respectively, are reported. The spectral parameter  $A_{zz'}$  at low and high temperatures and the characteristic ESR temperature  $T_{50G}$  of the slow to fast regime transition of TEMPO increases dramatically in the virgin SG-SIL's compared to the bulk CHX medium weakly depending on the pores size. This trend is similar to that found for other linear hydrocarbon, nhexadecane (n-HXD). In contrast, the sensitivity to the pores size for the globular CHX molecules is slighter in comparison to the moderate one for the linear *n*-HXD molecules. On the other side, the  $A_{zz}$ , values and the corresponding  $T_{50G}$  ones are very dependent on the pores composition confirming definitely the hypothesis about the preferential localization of polar molecular probe to polar pores of the SG matrices. These observations result from the complex interrelation between the mutual interaction between the polar TEMPO and the polar or apolar SG's and the altered phase behavior of the CHX as evidenced from DSC. These findings point to very careful choice of the organic *filler* and inorganic confiner in the bulk vs. confinement problem as monitored not only by

ESR technique but also by other molecular probes, such as fluorescence ones. The results of further investigations for organic *medium* of distinct structural type, such as polar protic substance, inserted in the same series of inorganic *SG matrices* which allowing for its preferential interaction with their pores over that with the *polar spin probe* will be presented elsewhere.

# **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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