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ABOUT THERMO-OXIDATIVE AGEING AT MODERATE TEMPERATURE OF CONVENTIONALLY VULCANIZED NATURAL RUBBER

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ABSTRACT

This article is dedicated to the understanding of the mechanisms involved in the evolution of the structure of a “conventionally” vulcanized rubber during thermo-oxidative (TO) aging. Based on swelling measurements, WAXS and DSC, a scenario of these mechanisms is proposed. Our results show that the crosslinking reactions are far from being complete at the end of the vulcanization process. During TO aging carried out at a moderate temperature 77 °C (350 K), the creation of long bridges first takes place via the consumption of residual sulfur; this mechanism is mainly responsible of the increase in the density of elastically active chains. The presence of residual antioxidants inhibits in the early stages of aging the chemical reactions involving oxygen. For longer aging time (here 7 days), these reactions can then occur and create "unidentified" bridges whose formation may also involves sulfur grafted onto the polymer chains. Finally, all these cross-linking reactions seem to enhance the heterogeneity of the spatial distribution of the crosslinks - already existing in the initial material - and to create highly crosslinked domains.

Keywords: Natural Rubber, thermo-oxidative, ageing, vulcanization, network architecture

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1. INTRODUCTION

For most of its applications, Natural Rubber is submitted to a crosslinking reaction, which consists in the formation of bridges between the polymer chains. They are obtained through curing of the polymer blended with different ingredients. The most common recipe contains sulfur, which will form the bridges, and additives necessary to control the kinetics of the different involved crosslinking reactions. Among these additives, one distinguishes accelerator molecules. Their type and the ratio of sulfur to accelerator pilot the length distribution of the sulfur crosslinks. Networks rich in long polysulfidic crosslinks, called hereafter Conventionally Vulcanised NR (CV), obtained when this ratio is largely above 1, are said to provide better fatigue and tear resistance [1], [2] than networks rich in mono- and disulfide crosslinks. These ones, so called hereafter Efficiently Vulcanised NR (EV) system, are obtained when this ratio is below 1.

However, compared to EV systems, CV systems are said to have a lower resistance to ageing. Indeed, polysulfidic crosslinks should be thermally less stable than disulfide and monosulfide bridges [3]. When submitted to thermal ageing (anaerobic), depending on the temperature and the ageing time, non–reversible change can occur in the crosslink structure [4]–[7]. It leads for instance to a non-reversibility of the material deformation when this ageing is performed on a deformed material [8], [9]. When oxygen is also present, which is common in most applications, a reduction of the molecular weight of the elastomer through chain scissions (which can produce a decrease of the Young Modulus) is also observed [10], [11]. This reduces the strength properties and possibly adds nucleation sites for crack initiation and propagation. Oxidation can nevertheless also contribute to additional crosslinks formation [12]–[14], a behavior mainly depending on the matrix chemistry. Thus, thermo-oxidative ageing is the result of different mechanisms, whose consequences on the material properties can be contradictory, and strongly depend on the ageing temperature [12].

For tire application, the use temperature is below 80°C [15]–[17]. The studies to understand these mechanisms are often performed at higher temperatures [7], [10], [18], [19]. Indeed, this choice enables to accelerate the processes. Nevertheless, the activation energies of the involved chemical reactions can be different [10], [20]. For this reason, accelerated ageing tests may induce erroneous lifetime prediction, especially in the case of CV systems where likely more mechanisms occur [12], [14]. Some studies display experimental results at low ageing
temperatures for CV NR [8], [21]. However, to the knowledge of the authors, only Ahagon et al. [12], [14] independently analyzed thermal ageing and oxidative ageing, which can help to discriminate the chemical mechanisms involved. He reported that thermal ageing, performed below 80°C does not significantly change elastic active chains (EAC) density while oxidative ageing strongly increases it. All the samples studied were filled, but the possible influence of the fillers on the oxidation rate [22], [23] was not taken into account in the discussion, which hinders any definitive conclusion on the mechanisms involved in the matrix alone. Besides, using only mechanical tests, he could not dissociate them, neither he evaluated their different contribution. Thus, the aim of this paper is to go further in the understanding of the multi-steps thermo-oxidative ageing of CV system. To do so, model unfilled samples are processed and analyzed by combining rheological, DSC, WAXS and swelling measurements.

2. EXPERIMENTAL

2.1 MATERIALS AND SAMPLES PROCESSING

| Sample code | NR1.5 | NR2.5 | NR2.5_Eff |
|-------------|-------|-------|-----------|
| NR\textsuperscript{a}, phr\textsuperscript{b} | 100   | 100   | 100       |
| 6PPD\textsuperscript{c}, phr | 3     | 3     | 3         |
| Stearic Acid, phr | 2     | 2     | 2         |
| Zinc Oxide, phr | 5     | 5     | 1.5       |
| CBS\textsuperscript{d}, phr | 0.5   | 1     | 2.52      |
| Sulfur, phr | 3     | 6     | 1.6       |
| $v$ ($\times 10^{-4}$ mol.cm$^{-3}$) | 1.5   | 2.5   | 2.5       |
| $t_{98}$, min | 55    | 55    | 13        |

\textsuperscript{a} Natural rubber TSR20
\textsuperscript{b} Parts by weight for hundred parts of rubber.
\textsuperscript{c} N-(1,3-dimethylbutyl)-N'-phenyl-p-phenylenediamine
\textsuperscript{d} N-Cyclohexyl-2-benzothiazole sulfenamide.
\textsuperscript{e} EAC Average density of estimated from swelling measurement with an affine assumption.

Table 1: Samples recipes and initial EAC density

The materials are unfilled natural rubber, obtained by gum vulcanization (Technically Specified Rubber TSR20) according to the recipe given in Table 1. NR1.5 and NR2.5 have a Sulfur/CSB ratio of 6, i.e. above 1, leading to
a network crosslinked mainly by polysulfidic bridges. In spite of different sulfur content, their same Sulfur/CBS ratio should ensure a same average length of the sulphur bridges. NR2.5_Eff with a Sulfur/CBS below 1 is crosslinked by an “efficient” recipe leading to a network mainly crosslinked by monosulfides bridges. First, the gum is introduced in an internal mixer and sheared at 50 rpm for 2 min at 50 °C. The material is subsequently sheared at 70 rpm in an open mill for 5 minutes at 70 °C. Then the vulcanization ingredients are added and the so-formed mixture is sheared for 5 minutes at 50°C. Thin sample sheets (0.8 mm thickness) are finally obtained by hot pressing under nitrogen flux at 150°C, for conventional system and 170°C for efficient system, 30 bars.

The cure times were estimated from the torque measurements performed with a Monsanto analyzer as illustrated in Fig. 2. As usually, they are defined as the time needed to reach 98% of the maximum torque value (t_{98}). It is worth mentioning that NR2.5 and NR2.5_Eff exhibit the same EAC density with different sulphur/CBS ratios, whereas NR1.5 and NR2.5 have a different EAC density with the same sulphur/CBS ratio.

2.2 RHEOMETRY ANALYSIS

Cure characterization was carried out with an Alphamodel ODR2000 oscillating disk rheometer in accordance with ASTM D2084-92. Evolution of torque has been followed on samples sheets (2 mm) at 150°C from 0 to 100 minutes. Results were provided by Michelin Company.

2.3 DSC ANALYSIS

DSC data were acquired using a Pyris Diamond calorimeter (Perkin Elmer, U.S.A) beforehand calibrated with an indium standard. The samples weight was 10 mg encapsulated in an aluminum pan. They were heated from 20 to 150°C at 50°C.min\(^{-1}\). Measurements are performed when the samples are maintained at 150°C, 160 min in order to follow the vulcanization process. This one is evidenced by the appearance of an exothermic peak on the thermogram (cf. Fig. S2 in SI). Then, samples were quenched at 20°C and submitted again to the same thermal program to ensure the end of curing. The DSC thermogram obtained during this second isothermal treatment at 150°C provides the baseline needed to calculate the previous peak enthalpy.

2.4 WAXS ANALYSIS

Wide-Angle X-ray scattering measurements were carried out on an apparatus equipped with a copper rotating anode (\(\lambda = 1.54 \text{ Å}\) (Rigaku Corporation, Japan), Gobel’s mirrors collimation system 160 (Exeliance, France) and a 2D detector (CCD camera from Princeton Instruments, USA). The acquisition time of the diffraction pattern is
Each pattern was integrated azimuthally and corrected from the background scattering. The corrected scattering intensity was normalized by the thickness and the absorption of each sample. The deconvolution of the curve \( I = f(2\Theta) \) was performed using Lorentzians functions (cf. Fig. S1) in order to extract the ZnS peak. This method was used to follow the evolution of the ZnS peak area during the vulcanization process.

### 2.5 AGEING CONDITIONS

All materials were submitted to thermo-oxidative (TO) and thermal ageing at 77 °C from 0 to 21 days. To fulfill thermal conditions (absence of oxygen), the samples were first submitted to a degassing step. This specific protocol used an home-made device and a succession of 3 steps described in Fig. 1.

![Fig. 1: Schematical view of the under vacuum device and the procedure to remove oxygen from the sample.](image)

The samples are sealed in a multilayer envelop (aluminum + cardboard) impermeable to air. They are put in a dedicated system made of two chambers. First, the vacuum pump extracts the air from the vacuum pack (Step ①), then the valve of this pack is closed (Step ②). Afterwards the vacuum is made inside the glass chamber (Step ③). At the end of the third step, the vacuum pack swells due to desorption of air from the sample. These three steps are repeated until the swelling of the vacuum pack becomes unnoticeable, indicating a total gas desorption. Finally, the vacuum pack is sealed before returning to atmospheric pressure. When not tested, all the samples are then stored at 7 °C to avoid any degradation. When tested, they are first maintained at 60°C during 5 min to avoid any presence of crystallites.

### 2.6 SWELLING MEASUREMENTS
Swelling measurement provides elastomer network chain density (from the swelling ratio \( Q \)) and the fraction of the chains and soluble species that does not belong to the elastically active network (from the soluble fraction \( F_s \)). The swelling procedure is as follows: samples with an initial mass \( m_i \) is introduced in the solvent (these experiments are performed with toluene and cyclohexane) for 5 days in order to reach the swelling equilibrium at room temperature. Then, the swollen material having a weight \( m_s \) is dried under vacuum at 50 °C during one day resulting in a final sample mass \( m_d \). The swelling ratio \( Q \) of the polymer and the soluble fraction \( F_s \) are then calculated through the Eq. 1 and 2:

\[
Q = \frac{1}{V_2} = 1 + \frac{\rho_p}{\rho_s} \times \left( \frac{m_s}{m_d} - 1 \right)
\]

\[
F_s = 100 \times \left( \frac{m_i - m_d}{m_i} \right)
\]

Where \( \rho_p \) and \( \rho_s \) correspond to the density in g.cm\(^{-3} \) of the polymer and the solvent respectively (0.92 for natural rubber, 0.87 for toluene and 0.78 for cyclohexane). From the volume fraction \( V_2 \) of polymer in the swollen mass (inverse of \( Q \)), the elastic active chain (EAC) density \( v_{\text{swelling}} \) is estimated using the Flory-Rehner equation [24]. \( v_{\text{swelling}} \) is the result of two contributions: EAC created by trapped entanglements and by the chemicals crosslinks formed during the vulcanization process:

\[
v_{\text{swelling}} = -\frac{\ln(1 - V_2) + V_2 + \chi \times V_2}{V_1 \times \left( V_2^\frac{1}{f} - 2 \times \frac{V_2}{f} \right)}
\]

Where \( \chi \) the Flory-Huggins interaction parameter is \( V_1 \) is the molar volume of the solvent (107.9 cm\(^3\).mol\(^{-1} \) for cyclohexane and 106.3 cm\(^3\).mol\(^{-1} \) for toluene) and \( f \) is the crosslink functionality. The low soluble fraction of the processed material (below 5% weight, this fraction being mainly curative residuals) suggests that most chains belong to the rubber network; thus \( f \) can be taken equal to 4. As recently mentioned by Valentin et al, the Flory-
Huggins interaction parameter $\chi$ has a strong impact on the calculated $v_{swelling}$ [25]. A widely used equation to estimate this parameter is based on the Hildebrand solubility constants:

$$\chi = \chi_s + \frac{V_s}{RT} (\delta_s - \delta_p)^2$$

(4)

Where $\delta_s$ and $\delta_p$ are the solubility parameters for solvent and polymer respectively, $R$ the ideal gas constant and $T$ the absolute temperature. The first factor $\chi_s$ is an entropic contribution and was found empirically equal to 0.34 for non-polar systems [26]. During thermo-oxidative ageing, oxygen modifies the chemical nature of the elastomer network by oxidation (i.e. there is formation of polar functions on the chains). Thus, the chemical affinity between the solvent and the polymer and therefore the value of the interaction parameter $\chi$ changes. Considering that the concentration of elastically active chains $v_{swelling}$ is identical regardless of the nature of the good solvent used, the observation of different swelling ratio with different solvent is inevitably linked to a modification of $\chi$ (and $\delta_p$). Thus, swelling measurements with two good solvents make possible the estimation of the solubility parameter of the polymer $\delta_p$ via Eq. 4. The calculations associated with these measurements are detailed in the SI (cf. S6).

2.7 THERMOPOROSIMETRY

Thermoporosimetry is a characterization techniques used to measure the distribution of pore size in porous material from calorimetry measurements. This is based on the modification of the solvent melting temperature by its confinement in the pores. In swelled crosslinked network, a phase segregation in small domains is observed during cooling. Like in porous materials, the idea is that the size of the segregated domains is related to the space available for this segregation, in other words to the local extensibility of the chains around the domain. Thus, this size is related to the local active chain density. The samples were swollen in hexadecane, which is the best solvent for this experiment, as its crystallization can be complete [27]. They were then introduced into an aluminum pan. The DSC apparatus was a Perkin-Elmer Pyris Diamond. The samples were first cooled at 1°C/minute from 25°C to -10°C, and then maintained at -10°C for 5 minutes. They were then heated at 1°C/minute to 25°C. While increasing temperature from -10°C to 25°C, a peak is first observed corresponding to the melting of the hexadecane domains crystallized in the network. The largest this melting temperature, the largest are the related melting domains (following the Gibbs Thomson description). The first melting peak is followed at around 19°C by a second, much sharper peak corresponding to the melting of the free hexadecane. The signal is normalized by the mass of dried sample.
2.8 DYNAMIC MECHANICAL ANALYSIS

DMA measurements were carried out on a homemade torsional pendulum previously detailed [28]. This apparatus measures the values of the real and imaginary parts of the shear modulus (storage modulus $G'$ and loss modulus $G''$). Parallelepiped samples (Length*Width*Thickness: 30*6*0.8 mm$^3$) were tested. The sample chamber was filled with inert gas (helium at 600 mbar) and the measurements were performed between 200 K and 320°K at a heating rate of 1 °C.min$^{-1}$ and a strain frequency of 1 Hz.

2.9 RESIDUAL CONCENTRATION OF ANTI-OXIDANTS (6PPD)

Residual concentration of anti-oxidants was measured using HPLC and ultraviolet absorption spectrum. Results were provided by Michelin Company only for non-aged and aged NR2.5.
3. RESULTS

3.1 VULCANIZATION AT 150°C

Fig. 2 shows the evolutions of $\nu_{\text{swelling}}$ and torque for NR2.5 and NR1.5 during the vulcanization process at 150°C. As expected, the maximum torque increases with the sulfur content. Moreover, similar evolutions are observed for $\nu_{\text{swelling}}$ and torque as a function of cure time for both materials. These parameters continually increase during vulcanization until optimal mechanical properties are reached after $t_{98} = 55$ minutes. Beyond this time, EAC densities and torque weakly decrease. This behaviour has been ascribed to the cleavage of polysulfidic crosslinks. This decreases the crosslink density and therefore the EAC density [6]. At $t_{98}$, average EAC densities of NR1.5 and NR2.5 are found equal to $1.5 \times 10^{-4}$ and $2.5 \times 10^{-4}$ mol.cm$^{-3}$ respectively.

![Fig. 2: Torque and EAC density as a function of cure time at 150°C - Lines are plotted as a guide for the eyes.](image)

The progress of the vulcanization reactions is generally associated with the conversion of the crosslinks precursors (i.e. active sulfuring agent) into crosslinks. According to the generally accepted – and simplified – theory of accelerated sulfur vulcanization (cf. Fig. 3) : the first step (1) in the reaction sequence involves interaction between the accelerator (CBS), ZnO and a ligand (stearic acid and/or an amine, like cyclohexylamine freed by CBS decomposition) to form an accelerator-zinc complex [6][29], [30]. Then the latter is assumed to react with free sulfur $S_8$ (2) to form an active sulfuring agent with variable $S_a$ and $S_b$ arm lengths: the higher the S/CBS ratio, the longer these arms, and consequently the sulfur bridges finally created.
This Zn species is found at an early stage of vulcanization and is likely to react with a carbon-carbon double bond (3) forming a polysulfidic pendant group along the polymer chain, and a new complex which may react with this newly formed pendant group and help it link (4) with another C=C bond, and thus finalize the desired crosslink (either active or inactive). Following this scheme, it will eventually produce some ZnS byproduct: at first sight, formation of two crosslinks, i.e. two polysulfidic bridges during vulcanization process should be indirectly associated to the formation of one ZnS molecule. Nevertheless, the authors acknowledge that the mechanical scheme is far from being clearly understood, as demonstrated by new evidences from literature which suggests for example that the Zn complex may be a bimetallic one with a 1:1 ratio between Zn and the created pendant groups [31]. Anyhow, looking at the relative ZnS content evolution through the vulcanizing process remains interesting as a potential ongoing vulcanization marker when one follows its characteristic peak ($2\theta = 27^\circ$ [32], [33]) through WAXS measurements (cf. Fig. 4a). We define the following $\xi_{\text{WAXS}}$ index:

$$\xi_{\text{WAXS}}(\%) = \frac{A_{\text{ZnS Peak}}(t)}{A_{\text{ZnS Peak}}(t_{\infty})}$$ (5)

Where $A_{\text{ZnS Peak}}(t)$ is the ZnS peak area of compound cured for a time $t$, and $A_{\text{ZnS Peak}}(t_{\infty})$ the ZnS peak area for the same compound in which all the chemicals reactions producing ZnS molecules are assumed to be finished.
(i.e. when the peak does not evolve anymore with time). $\xi_{WAXS}$ is plotted as a function of the curing time in Fig. 4b.

Fig. 4 : (a) Evolution of ZnS and ZnO peaks for NR2.5 along the vulcanization process (In blue : Curing time in min) (b) $\xi_{WAXS}$ as a function of curing time at 150°C.

Fig. 5 : (a) $\xi_{DSC}$ as a function of Curing time at 150°C (b) $\xi_{DSC}$ as a function of $\xi_{WAXS}$. The dashed line is a linear fit.

Isothermal DSC measurements were recorded a 150°C as a function of time for an uncured compounded rubber. All involved reactions are exothermic ($\Delta H$, enthalpy); the observed exotherm depends on the nature of the polymer, curatives and other additives in the matrix [34], [35] (Fig. S2). Its evolution is associated with the progress of all the chemicals reactions involved during the vulcanization process. It can be followed through the index:

$$\xi_{DSC}(\%) = \frac{\Delta H (t)}{\Delta H (t_{\infty})}$$  \hspace{1cm} (6)
Where $\Delta H (t)$ is the enthalpy of the compound cured for time $t$ and $\Delta H (t_{\text{m}})$, the total enthalpy for which all the chemicals events are finished (estimated when $\Delta H (t)$ is stabilized). $\xi_{\text{DSC}}$ is plotted as a function of the curing time and of $\xi_{\text{WAXS}}$ in Fig. 5a and 5b respectively. The evolution of $\xi_{\text{DSC}}$ with time is consistent with the one observed for $\chi_{\text{WAXS}}$ in Figure 5b. A linear dependency is found between both indexes indicating that ZnS could be considered as a probe of the vulcanisation progress. Additionally, $\xi_{\text{WAXS}}$, i.e the ZnS content, still increases beyond $t_{98}$ for both materials. Such finding is unexpected and indicates that the optimal mechanicals properties (i.e. the maximum EAC density measured at $t_{98}$) may not correspond to the end of the crosslinking process.

### 3.2 AGEING AT 77°C

Once cured at $t_{98}$, the materials have been aged from 0 to 21 days at 77°C, in thermal (no oxygen), and thermo-oxidative conditions (so called TH ad TO conditions respectively).

#### 3.2.1 THERMAL AGEING

Dashed lines on Fig. 6 report the evolution of the EAC densities during TH ageing from 0 to 21 days for NR2.5, NR1.5 and NR2.5_Eff. At 7 days, conventionally crosslinked rubbers display an increase of $v_{\text{swelling}}$ while no evolution is noticed for NR2.5_Eff. Variation of EAC density is more important for NR2.5 (+ 32 %) compared to NR1.5 (+ 19 %). Beyond 7 days, $v_{\text{swelling}}$ does not longer evolve for all the samples. By the use of two solvents (cyclohexane and toluene) and of Eq. 4 (Calculation presented in S6), the solubility parameter of the polymer $\delta_p$ can be deduced and indirectly provides an estimate of the evolution of the polar component of our samples. Fig. 7 displays its evolution as a function of the ageing duration. As expected, $\delta_p$ is found constant, which is consistent with the fact that no oxygen is involved in the chemical reactions occurring during TH ageing of NR1.5 and NR2.5.

#### 3.2.2 THERMO-OXIDATIVE AGEING

Evolutions of EAC densities during TO ageing (solid lines) are compared to the ones obtained during thermal ageing (dashed lines) in Fig. 6. For NR1.5 and NR2.5, the curves superimpose between 0 and 7 ageing days. Meanwhile, $\delta_p$ stays roughly constant (cf. Fig. 7a). These similar trends for $v$ and $\delta_p$ are quite unexpected, as oxidative mechanisms should normally take place. An explanation may be that during these first 7 days, oxygen
is consumed by the residual antioxidants (6PPD, used to hinder oxidative reactions [18],[29]) unconsumed during the vulcanization process.

Fig. 6: EAC density versus ageing time at 77°C - Lines are plotted as a guide for the eyes

Fig. 7: Solubility parameter of (a) NR2.5 and (b) NR2.5_Eff as a function of ageing time at 77°C - Lines are plotted as a guide for eyes

| TO ageing time                  | 0 day | 7 days | 14 days |
|---------------------------------|-------|--------|---------|
| NR2.5 previously TO aged 7 days | 3.1   | 3.9    | 4.2     |
| NR2.5 previously TH aged 7 days | 3.1   | 3.1    | 3.9     |

Table 2: \(v_{swelling} (\times 10^{-4} \, \text{mol.cm}^{-3})\) for NR2.5, as a function of TO ageing time
To address this question, a specific study was performed on the most crosslinked sample NR2.5. It was first thermally aged for 7 days at 77°C, during which residual anti-oxidant should be not consumed. Then, it was aged for additional 7 days and 14 days, in presence of oxygen. The sample EAC densities deduced from swelling measurements are presented in Table 2. EAC density of the sample previously thermally aged during 7 days is constant during the following 7 days of TO ageing. Then it increases again to reach, at 14 days of TO ageing, the value obtained when the sample is only submitted to TO ageing during 14 days (In other words, there is a shift of the EAC((time) curve). Additionally, some experiments were performed to measure the concentration of residual anti-oxidants after TO ageing (cf. Fig. 8). As expected, residual antioxidants are still present after vulcanization. Their concentration rapidly decreases during the first 7 days and much slowly the following days. This confirms our assumption that the apparent similarity between TO and TH ageing during the first 7 days is due to the initial presence of antioxidant in the sample, whose consumption prevents the oxygen from playing a role in the degradation mechanisms. This explanation should also be valid for NR1.5 if one makes the reasonable assumption that the remaining quantity of antioxidant after vulcanization is the same as in NR2.5 (reasonable as the initial 6PPD quantity and the curing time are the same).

Fig. 8 : Concentration of residual antioxidants after TH and TO ageing at 77°C for NR2.5 - Lines are plotted as a guide for the eyes

For TO ageing longer than 7 days, \( \delta_p \) (cf. Fig. 7) and EAC densities (cf. Fig. 6) increase for NR2.5 and NR1.5 whereas they remain both constants for NR2.5_Eff. Such finding suggests that presence of oxygen in conventionally crosslinked rubbers leads simultaneously to supplementary crosslinking and to the formation of polar groups. Note that, after 21 days TO ageing, the \( \delta_p \) variation is the same for NR1.5 and NR2.5, whereas the EAC density evolution is twice higher for NR2.5 (+33%) than for NR1.5 (+18%).
Competitive processes should be involved here: chain scissions, chain crosslinking and possibly sulfur bridges reorganization/scission. The consequences of chains crosslinking predominates for NR1.5 and NR2.5 (cf. Fig. 6). Constant weight soluble fractions, around 5% (results presented in S3) means that scissions are not numerous enough to create supplementary free chains. Nevertheless, a significant amount of scission occurs as TO ageing at 77°C of pure isoprene rubber (no vulcanization system) leads to a decrease of its $M_n$ (cf. Fig. 9a).

Fig. 9 : (a) Average molar mass of the non-vulcanized isoprene rubber during thermal ageing (dash line) and thermo-oxidative ageing (solid line) at 77°C - Lines are plotted as a guide for the eyes
(b) Loss modulus $G''$ (filled dots) and $\tan(\phi)$ (unfilled dots) as a function of temperature for non-aged and thermo-oxidative aged NR2.5 (f=0.1 Hz, $\gamma$ below 0.001).

Consequently, TO ageing necessarily leads to the formation of dangling chains in the aged rubber network. This is evidenced by the increase in $G''$ and $\tan(\phi)$, measured in the rubbery plateau, with the ageing time increase (cf. Fig. 9b). Fig. 9a indicates that 21 days of TO ageing should lead to around a maximum of $0.36 \times 10^{-5}$ mol.g$^{-1}$ of chain scissions (“maximum” as the presence of vulcanisation system may decrease the number of chain scissions in the elastomer). Considering NR1.5, the less initially crosslinked material, its EAC density increase is around $0.27 \times 10^{-4}$ mol.g$^{-1}$ after 21 days of TO ageing. This is much larger than the previous chains scissions estimate ($\approx$ at least 10 times lower) and suggests that these scissions can be considered as negligible compared to crosslinking.

$\xi_{WAXS}$ is plotted as a function of ageing time in Fig. 10. First, it is worth recalling that at $t_{0b}$, the ZnS initial concentration is not the maximum one : $\xi_{WAXS}$ is equal to 0.71 for NR1.5 and to 0.60 for NR2.5. More importantly, $\xi_{WAXS}$ does not evolve from 0 to 21 days which suggests that the Zn species involved in the
vulcanization mechanism illustrated in Fig. 3 are not involved in the EAC density increase at 77°C (cf. Fig. 6). It means that a different crosslinking mechanism should take place during this ageing time at 77°C.

![Figure 10: \( \xi_{WAXS} \) as a function of the ageing time measured during TO ageing at 77°C from 0 to 21 days - Lines are plotted as a guide for the eyes](image)

### 3.2.3 STABILITY OF THE NEW BRIDGES FORMED DURING AGEING

![Fig. 11: (a) ln(Crosslink density) versus thermal ageing time for different ageing temperature for NR2.5 previously TO aged at 77°C for 7 days (b) Arrhenius plot and the deduced thermal activation energy](image)

Evolution of the crosslinks density is studied during TH ageing for a previously 7days-TO aged NR2.5. The temperature chosen for this experiment are 140°C, 150°C, 160 °C, and 170 °C, i.e. consistent with curing temperature. In this temperature range, it is known that polysulfidic crosslinks dissociate and form mostly non-elastically active sulfur chains while di or mono-sulfides are relatively stable [3], [12]. Indeed, sulfur bridges of lengths greater than 4 are more easily broken owing to their lower bond dissociation energies of 150 kJ/mol [36]. Conversely, decomposition of di- and monosulfide bridges are more difficult due to their higher dissociation...
energy, which are 270 and 320 kJ.mol$^{-1}$ respectively [37], [38]. Considering that thermolytic chain scission is not involved during thermal ageing up to 170°C along the polymer backbone [8][30] (and assuming that this is also true in presence of sulfur species), variations of the crosslinks density are mainly due to the destruction of sulfur bridges. Assuming a first order kinetic, the concentration of [S-S] bridges may be expressed as:

\[ [S - S] = [S - S]_0 \cdot \exp(-kt) \]  

(7)

Where \( k \) is the rate constant in s$^{-1}$. As shown by Fig. 11a, two regimes are highlighted by these experiments and the transition between both regimes occurs, whatever the ageing temperature, when crosslink density has roughly decreased by around \( 7 \times 10^{-5} \) mol.cm$^{-3}$, i.e. when the crosslink density of the sample has returned to the value of the non-aged sample. From these results, two scissions constant rates \( k_1 \) and \( k_2 \), corresponding to these regimes, can be deduced, as a function of temperatures (from 140°C to 170 °C). Arrhenius extrapolation assumes that a reaction rate \( k \) is proportional to \( \exp(-E_a/RT) \), where \( E_a \) is the Arrhenius activation energy, \( R \) the gas constant (8.314 J/mol.K$^{-1}$), \( T \) the absolute temperature and \( A \) the pre-exponential factor [14], [40] :

\[ k = A e^{-E_a/RT} \]  

(8)

Arrhenius plots give two straight lines for \( k_1 \) and \( k_2 \) (with a correlation coefficient \( R^2 = 0.992 \)) whose slope corresponds to activation energies of 99.1 kJ.mol$^{-1}$ and 177kJ.mol$^{-1}$ (Fig. 11b). Such result indicates the successive destruction of two different sulfur networks in NR2.5. Indeed, the former value is found lower than bond dissociation energy typically found for length much larger than 4 [36] while the latter corresponds to dissociation energy of sulfur bridges of length around 4 [37], [38]. Thus, the first bridges destroyed during the post curing of the aged sample should corresponds to long polysulfur bridges. The variation of the EAC density due to this destruction, obtained after 30 minutes at 150°C, is around three times larger than the EAC decrease observed when the initial material is cured 30 min more than \( t_{98} \). This indicates that the majority of these long polysulfur bridges have been created during the material ageing at 77°C.

4. DISCUSSIONS
From the previous results, let us now discuss the network structure evolution of conventionally crosslinked rubber during TO ageing at 77°C (up to 21 days). Assuming that ZnS is a probe of the sulfur bridges concentration during curing, we have plotted EAC density as the function of $\xi_{WAXS}$ (i.e ZnS) in Fig. 12 for different curing times, from 0 to 100 minutes at 150°C.

![Fig. 12: EAC density as a function of $\xi_{WAXS}$ at 150°C. Curing time in minutes is indicated for each data point - Lines are plotted as a guide for eyes](image)

First, EAC densities increase as a function of $\xi_{WAXS}$ for both samples until $t_{eq}$ is reached and then they decrease slightly beyond this time. Additionally, evolution of EAC density is always 1.6 times higher for NR2.5 than the one for NR1.5. This suggests that the kinetics of all the involved chemical reactions are proportional to the initial amount of sulfur (and CBS). The decrease of EAC with $t$ above $t_{eq}$ cannot be ascribed to thermolytic chain scission, as it was shown in literature that chain scission mechanisms are negligible during thermal ageing of pure isoprene, even at 170°C (within the reasonable assumption that the presence of bridges and vulcanisation system has no influence on this) [10], [39]. Besides, ZnS concentration still increases, which indicates that formation of crosslink precursors still occurs, and therefore that sulphur bridges are likely to be formed. Nevertheless, these additional sulphur bridges, and those possibly formed via the transformation of long bridges in two shorter bridges [18] does not hinder the decrease of the EAC density. An explanation of this result is given by the literature which suggests that a significant part of the polysulfidic bridges can be transformed as non-elastic cyclic sulphides groups along the polymer chains.

Conversely to what is observed at 150°C, the EAC crosslink densities of the optimally cured samples increase during TO or TH ageing at 77°C. Thus, more numerous active bridges are created than destroyed (destruction via
the formation of polysulfidic cycles). As shown by the initial value of $\xi_{WAXS}$ (cf. Fig. 10) for both materials, the ZnS concentration does not evolve. Thus, as said previously, other mechanisms than the ones involved during the vulcanization process at high temperature are responsible for the formation of new crosslinks. To confirm this, the evolution of EAC density and $\xi_{WAXS}$ has been measured during the TH ageing at 77°C of a non-cured NR2.5 (cf. Fig. 13). The EAC density increases up to $5.10^{-4}$ mol.cm$^{-3}$ while $\xi_{WAXS}$ stays nul, showing that crosslinking can occur at 77°C without the involvement of the active sulfurating agent appearing in the vulcanization process at high temperature (cf. Fig. 3). The stabilization of the EAC after 7 days may be explained by the total consumption of the free sulfur. Some supplementary experiments measuring the residual sulfur concentration during ageing would be necessary to confirm this assumption. This total consumption of residual sulphur could also explain the absence of supplementary crosslinking during the TH ageing of NR2.5_Eff (cf. Fig. 6).

![Evolution of $\xi_{WAXS}$ (black) and active chain density (red) during thermal ageing of a non-cured NR2.5 from 0 to 21 days](image)

The crosslinks formed after 7 days in NR2.5 are weak bridges (cf. Fig. 11). They may be formed by the process often named “unaccelerated sulfur vulcanization” in which only the consumption of sulfur ($S_8$) is involved. As reviewed by Krejsa [29], $S_8$ is able to undergo homolytic (radical) or heterolytic (polar/ionic) ring opening reactions to form long polysulfidic bridges (as observed by NMR studies [41]). There is no difference between TH and TO ageing at 77°C during the first 7 days as the consumption of residual antioxidants inhibits the role of the oxygen. For conventionally crosslinked rubber NR2.5 and NR1.5, and for longer ageing duration, TO ageing leads to supplementary crosslinks (cf. Fig. 6 & 7). It is known that ageing by oxygen at high temperature
(140°C) can lead to crosslinks made of peroxide bridge [10]; nevertheless, the $M_n$ decreases of pure polyisoprene during TO ageing at 77°C (cf. Fig. 9a) suggests that such mechanism is negligible at this temperature. Thus, the formation of crosslinks needs here reaction(s) involving both oxygen and sulfur. This might be a reaction between oxygen and cyclic and pendant sulfides groups which are numerous in conventionally crosslinked NR [42], and are randomly distributed along the polymer chain. Additionally, the non-evolution of EAC density and $\delta_p$ for NR2.5_Eff (cf. Fig. 6 and 7) in the presence of oxygen from 7 to 21 days can be explained by the low concentration of reactive polysulfidic groups along the backbone (related to the lower initial sulphur concentration) [43].

The formation of supplementary crosslinks during ageing raises the question of their spatial distribution. From thermoporosimetry experiments, Fig. 14 presents the comparative distribution of the solvent melting temperature for the aged and non-aged NR1.5 and NR2.5. Heterogeneities in the spatial distribution of the crosslinks already exist in the samples prior thermo-oxidative ageing, particularly for non-aged NR2.5, for which the shape of the peak suggests the presence of domains with a high EAC density. TO ageing leads to a shift of the distribution towards low temperature and the magnitude of this transformation is more important for NR2.5. This is consistent with EAC density evolution. In addition, ageing leads to appearance of a second peak at low temperature indicating that the crosslinking is spatially heterogeneous. This phenomenon is more pronounced in the case of NR2.5, in other words, aged NR2.5 is expected to have a network more heterogeneous than NR1.5. It is worth mentioning that the same comment can be made from the observation of the solvent crystallization curves (cf. Fig. S4).
Fig. 14: Heating curves from thermoporosimetry experiments for (a) NR1.5 and (b) NR2.5 aged from 0 to 21 days during TO at 77°C.

To summarize, a schematic vision of the network evolution in conventionally crosslinked natural rubber during TO ageing is given in Fig. 15. First, it is often claimed that weakly and densely crosslinked phases are formed after the vulcanization process [44]–[46]. The latter consist of ZnO/ZnS clusters surrounded by a high crosslinked phase. Such vision is partly confirmed by the thermoporosimetry curves of unaged NR1.5 and NR2.5 (cf. Fig. 14, and Fig. S5 in SI) which show a small shoulder at around 16°C and 15°C respectively. Besides, cyclic sulfide pendant groups are formed along the polymer chain when NR is conventionally crosslinked [43]. Some crosslinks precursors (as $\xi_{\text{WAXS}} \neq 1$) and residual antioxidants also remain in the network after the vulcanization process (cf. Fig. 15a). During the first 7 days of TO ageing, whereas vulcanization reactions are not activated and $\xi_{\text{WAXS}}$ stays constant, residuals antioxidants and free sulfur are consumed, leading mainly to the formation of long polysulfidic bridges (cf. Fig. 15b). Once antioxidants are consumed, oxygen leads to the formation, via reaction(s) involving the oxygen, of new crosslinks, likely from polysulfidic cycle pendant groups (cf. Fig. 15c). Such mechanisms are also spatially heterogeneous, and decrease the fraction of rubber which can be highly swollen, in other words they suppress the weakly crosslinked domains and may create domains too crosslinked to swell. To be complete, in spite of the large predominance of crosslinking mechanisms, TO leads also to relatively few chain scissions, and to the formation of pendants groups along the polymer chain. The schemes in Fig. 15 summarize these interpretations.
Fig. 15: Proposed schemes to highlight network structure (a) after the vulcanization process and its evolution during TO ageing at 77 °C (b) after 7 days and (c) 14 days.
4. CONCLUSIONS

This work aims at understanding the chemicals mechanisms involved during thermo-oxidative ageing at 77°C on a conventionally crosslinked natural rubber. The main findings are the following:

- It is important to keep in mind that in conventionally crosslinked system, the vulcanization process is far to be finished at \( t_{90} \). An important consequence is that kinetics models cannot be built only from the mechanical properties evolution during the vulcanization process when it stopped at that time.

- TO ageing is similar to TH ageing as far as the concentration of antioxidants remaining form the vulcanization process, is large enough.

- TH ageing at 77°C involves chemical mechanisms which are different from the ones met during the material processing.

- At 77°C, TH ageing leads to the formation of weak long polysulfidic bridges from the residual free sulfur remaining in the unaged material. This process last as long as all the free sulfur is consumed.

- Shortening of the polysulfidic bridges does not seem to be a predominant mechanism during TH at 77°C.

- During TO at 77°C, chemical reactions involving the oxygen take place to lead to supplementary crosslinking in conventionally crosslinked materials, which suggests that these reactions need the presence of cyclic sulfide pendant groups (crosslinking being negligible, compared to chain scission mechanisms, in materials in which these groups cannot be present). However, some question remains on the chemical nature of the bridges created during TO ageing. A specific study devoted to this question would be undoubtedly interesting…even though it is likely very difficult, given the complexity of NR and the vulcanization recipe, to be able to associate among all the chemical species formed during TO, those specifically forming these supplementary crosslinks.

- In spite of the large predominance of crosslinking mechanisms, TO leads also to relatively few chain scissions, and to the formation of pendants groups along the polymer chain.

- The supplementary crosslinks induced by ageing seem to enhance the heterogeneity of the spatial distribution of the crosslinks - already existing in the initial material - and to create highly crosslinked domains.

To conclude this work clearly proves that the use, for conventional NR, of accelerated ageing procedure (at temperature for instance above 100°C), to address the question of its durability, is highly questionable.
5. SUPPORTING INFORMATIONS (SI)

Fig. S1 : Deconvolution and fitting of a WAXS pattern.

Fig. S2 : Exothermic peak during the vulcanization process at 150°C.
Fig. S3: Evolution of the soluble fraction $F_s$ during thermo-oxidative ageing at 77°C.

Fig. S4: Cooling curves from thermoporosimetry experiments for (a) NR1.5 and (b) NR2.5 aged from 0 to 21 days swelling in hexadecane.
Fig. S5: Heat flow normalized by the dried sample weight as a function of \(1/(T^0_m - T_m)\) described by the Gibbs-Thomson equation

According to the Gibbs-Thomson equation (Eq. S1), a “pore” size in the polymer network can be calculated from the melting temperature of the solvent it contains:

\[
T^0_m - T_m = \frac{2\sigma_{SL}T^0_m}{L_c\Delta H_f}
\]  

(S1)

Where, \(\Delta H_f\) is the solvent melting enthalpy, \(L\) the crystal radius within the assumption of a spherical pore, \(\sigma_{SL}\) is the solid-liquid interfacial energy and \(T^0_m\) the free solvent melting temperature which is around 21°C for hexadecane. \(\sigma_{SL}\) is not known for the hexadecane-NR couple. However, the crystal radius can be considered proportional to \(1/(T^0_m - T_m)\). Normalized heat flow is plotted as a function of \(1/(T^0_m - T_m)\) for NR1.5 and NR2.5 in Fig. S5. For both samples, the formation of large crystals is less pronounced after TO ageing. As the consequence, crosslinking mechanisms decrease the fraction of rubber which can be highly swollen (i.e. corresponding the formation of larger crystals when solvent crystallize). In other words they suppress the weakly crosslinked domains and may create domains too crosslinked to swell.
Supplementary information S6 : Calculation of δ_p by the use of two good solvents

We assumed equality between \( v_{\text{sweeling}} \) measured by the use of cyclohexane \( v_{\text{sweeling}} \) measured by the use of toluene:

\[
v_{\text{sweeling}}(\text{Toluene}) = v_{\text{sweeling}}(\text{Cyclohexane})
\]  
(S6.1)

Considering the Flory-Rehner approach [24]:

\[
\frac{1}{V_{\text{toluene}}} \left( \frac{\ln(1 - V_{r_{0,\text{tol}}} + V_{r_{0,\text{tol}}} + X_{\text{toluene}} V_{r_{0,\text{tol}}}^2)}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}} \right) = \frac{1}{V_{\text{cyclohexane}}} \left( \frac{\ln(1 - V_{r_{0,\text{cyclohexane}}} + V_{r_{0,\text{cyclohexane}}}) + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}} \right)
\]  
(S6.2)

\[
\frac{V_{\text{cyclohexane}}}{V_{\text{toluene}}} \left( \frac{\ln(1 - V_{r_{0,\text{tol}}} + V_{r_{0,\text{tol}}} + X_{\text{toluene}} V_{r_{0,\text{tol}}}^2)}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}} \right) \times \left( \frac{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}}}{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}} + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2} \right) = \ln(1 - V_{r_{0,\text{cyclohexane}}}) + V_{r_{0,\text{cyclohexane}}}) + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2
\]  
(S6.3)

\[
X_{\text{toluene}} = \frac{V_{\text{cyclohexane}}}{V_{\text{toluene}}} \times \frac{\ln(1 - V_{r_{0,\text{tol}}} + V_{r_{0,\text{tol}}} + X_{\text{toluene}} V_{r_{0,\text{tol}}}^2)}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}} \times \left( \frac{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}}}{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}} + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2} \right) = \ln(1 - V_{r_{0,\text{cyclohexane}}}) + V_{r_{0,\text{cyclohexane}}}) + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2
\]  
(S6.4)

\[
X_{\text{cyclohexane}} = \frac{V_{\text{cyclohexane}}}{V_{\text{toluene}}} \times \frac{\ln(1 - V_{r_{0,\text{tol}}} + V_{r_{0,\text{tol}}} + X_{\text{toluene}} V_{r_{0,\text{tol}}}^2)}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}} \times \left( \frac{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}}}{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}} + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2} \right) = \ln(1 - V_{r_{0,\text{cyclohexane}}}) + V_{r_{0,\text{cyclohexane}}}) + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2
\]  
(S6.5)

With some constants defined by:

\[
A = \frac{V_{\text{cyclohexane}} \times \left( \frac{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}}}{\frac{1}{V_{r_{0,\text{cyclohexane}}} - \frac{V_{r_{0,\text{cyclohexane}}}^2}{2}} + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2} \right)}{V_{\text{toluene}} \times V_{r_{0,\text{cyclohexane}}^2} \times \left( \frac{\frac{1}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}}}{\frac{1}{V_{r_{0,\text{tol}}} - \frac{V_{r_{0,\text{tol}}}^2}{2}} + X_{\text{toluene}} V_{r_{0,\text{tol}}}^2} \right)}
\]  
(S6.6)

\[
C_1 = A \times \ln(1 - V_{r_{0,\text{tol}}})
\]  
(S6.7)

\[
C_2 = A \times V_{r_{0,\text{tol}}}
\]  
(S6.8)

\[
C_3 = A \times V_{r_{0,\text{tol}}^2}
\]  
(S6.9)

\[
C_4 = - \frac{\ln(1 - V_{r_{0,\text{cyclohexane}}}) + V_{r_{0,\text{cyclohexane}}}) + X_{\text{cyclohexane}} V_{r_{0,\text{cyclohexane}}}^2}{V_{r_{0,\text{cyclohexane}}}^2} - \frac{1}{V_{r_{0,\text{cyclohexane}}}}
\]  
(S6.10)
We have:

\[ C_1 + C_2 + C_4 + C_3 \times X_{\text{NR-tol}} = X_{\text{NR-cyclo}} \quad (S6.11) \]

Considering the widely used equation to estimate the interaction parameter:

\[
\begin{align*}
X_{\text{NR-tol}} &= 0.34 + \frac{V_{\text{tot}}}{RT} \cdot (\delta_p - \delta_{\text{tot}})^2 \\
X_{\text{NR-cyclo}} &= 0.34 + \frac{V_{\text{cyclo}}}{RT} \cdot (\delta_p - \delta_{\text{cyclo}})^2
\end{align*}
\]  

(S6.12)

With (S5.11) and (S5.12), we have:

\[
C_1 + C_2 + C_4 + C_3 \times \left[ 0.34 + \frac{V_{\text{tot}}}{RT} (\delta_p^2 - 2\delta_p\delta_{\text{tot}} + \delta_{\text{tot}}^2) \right] = 0.34 + \frac{V_{\text{cyclo}}}{RT} \cdot (\delta_p^2 - 2\delta_p\delta_{\text{cyclo}} + \delta_{\text{cyclo}}^2) 
\]  

(S6.13)

So,

\[
c_1 + c_2 + c_4 + c_3 \times (c_3 - 1) + \frac{\delta_p^2}{RT} \cdot \left[ C_3 V_{\text{tot}} - V_{\text{cyclo}} \right] + \frac{\delta_p}{RT} \cdot \left[ -2C_3 V_{\text{tot}}\delta_{\text{tot}} + 2V_{\text{cyclo}}\delta_{\text{cyclo}} \right] \\
+ \frac{1}{RT} \left[ C_3 V_{\text{tot}}\delta_{\text{tot}}^2 - V_{\text{cyclo}}\delta_{\text{cyclo}}^2 \right] = 0
\]  

(S6.14)

Finally, (S6.14) is in the form of a second-degree equation:

\[ a\delta_p^2 + b\delta_p + c = 0 \quad (S6.15) \]

With:

\[
a = \frac{C_3 V_{\text{tot}} - V_{\text{cyclo}}}{RT} \quad (S6.16) \\
b = \frac{2(V_{\text{cyclo}}\delta_{\text{cyclo}} - C_3 V_{\text{tot}}\delta_{\text{tot}})}{RT} \quad (S6.17) \\
c = c_1 + c_2 + c_4 + 0.34 \times (c_3 - 1) + \frac{(C_3 V_{\text{tot}}\delta_{\text{tot}}^2 - V_{\text{cyclo}}\delta_{\text{cyclo}}^2)}{RT} \quad (S6.18)
\]
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