

Crystallinity and perfection in ethylene vitrimers probed by combined calorimetry, scattering, and time-domain NMR

1 Kay Saalwächter^{1*}, Bhaskar Soman^{2,3}, Christopher M. Evans^{2,3*}

2 ¹Department of Materials Science and Engineering, Urbana, Illinois 61801, USA

3 ²Frederick Seitz Materials Research Laboratory, Urbana, Illinois 61801, USA

4 ³Institut für Physik – NMR, Martin-Luther-Universität Halle-Wittenberg, 06099 Halle (Saale),
5 Germany

6 * Correspondence:

7 Corresponding Authors

8 kay.saalwaechter@physik.uni-halle.de

9 cme365@illinois.edu

10 **Keywords: covalent adaptive networks, polymer crystallization, crystallization kinetics,**
11 **polymorphism, memory effect, superheating**

12 Abstract

13 The kinetics of crystallization and crystal-crystal transformations in ethylene vitrimers are studied by
14 time-domain NMR. These vitrimers previously exhibited polymorphic transition of crystal structures,
15 which are shown here to be distinguishable by NMR via their dipolar line widths based upon
16 different proton densities. The conditions under which the polymorphs are formed and interconvert
17 are identified via time-resolved NMR experiments, with a focus on recrystallization after full and
18 partial melting. The DSC experiments are used to clarify an unexpected superheating effect, which
19 challenges the determination of actual melting points. We further identify a strong memory effect in
20 isothermal (re)crystallization. Implications of the dynamic nature of the vitrimers in relation to the
21 kinetics of crystallization are discussed.

22 1 Introduction

23 Covalent adaptable networks (CANs) are polymer networks crosslinked using dynamic covalent
24 bonds (Kloxin and Bowman, 2013). Their ability to be reprocessed and recycled have made them
25 prime candidates for the next generation of sustainable plastics. Vitrimers are a sub-category of
26 CANs where the network strands are crosslinked by associative covalent bonds. (Montarnal et al.,
27 2011; Guerre et al., 2020; Van Zee and Nicolay, 2020; Zheng et al., 2021) These bonds undergo
28 exchange reactions such that the overall crosslink density of the networks is conserved during the
29 exchange. The bond exchange adds a degree of freedom on the molecular scale, that has a strong
30 impact on macroscopic properties. A core question in the study of vitrimers has been to understand
31 how parameters like molecular weight, molecular weight dispersity, junction chemistry, backbone
32 chemistry, temperature, and crosslink density impact macroscopic properties such as viscosity, stress
33 relaxation, creep, reprocessability, and self-healing ability (Denissen et al., 2015; Denissen et al.,
34 2017; Röttger et al., 2017; El-Zaatari et al., 2020; Hubbard et al., 2021; Ishibashi et al., 2021; Porath
35 and Evans, 2021; Ricarte and Shanbhag, 2021; Hubbard et al., 2022; Krishnan et al., 2022; Luo et al.,

2022; Porath et al., 2022). In recent years, using the knowledge generated from numerous fundamental studies, functional vitrimers have been developed for wide-ranging applications such as solid polymer electrolytes, durable coatings, shape-memory materials, and additive manufacturing (Jing and Evans, 2019; Yan et al., 2020; Chen et al., 2021; Ma et al., 2021; Niu et al., 2021; Rossegger et al., 2021; Gu et al., 2022; Lin et al., 2022; Van Lijsebetten et al., 2022). Despite the advances in vitrimers design and processing, a good understanding of how an exchange event that takes place on an Angstrom scale influences macroscopic events such as crystallization, rheological flow and self-assembly is lacking. Little work has focused on how adding dynamic bonds will affect the ability of polymer chains to crystallize (Soman et al., 2022a; Kuenstler and Bowman, 2023), which is a key attribute of polyolefins and many commercial plastics.

Most work on vitrimers has used rheology or dynamic mechanical analysis to study the moduli and flow properties of the networks (Yan et al., 2020; Chen et al., 2021; Ma et al., 2021; Niu et al., 2021; Gu et al., 2022; Lin et al., 2022; Van Lijsebetten et al., 2022). Some work has also utilized dielectric spectroscopy to probe segmental scale dynamics (Soman et al., 2022b; Pascual-Jose et al., 2023). While these methods can probe materials which are crystallizing, they are not routinely used to understand such processes and to infer quantities such as percent crystallinity or the degree of crystal perfection (Wurm et al., 2005; Soccio et al., 2007; Ezquerro et al., 1994). Solid-state NMR spectroscopy provides a plethora of techniques which can provide a great deal of insight regarding crystallization physics in soft materials (Chen and Kurosu, 2007). This holds in particular for proton NMR, possibly performed on cost-efficient low-field low-resolution instruments, which allows for a quick mobility-based determination of the absolute crystallinity (in terms of proton signal fraction, equaling mass fraction), provides access to domain sizes via spin diffusion, and importantly, has sufficient sensitivity to allow for time-resolved experiments with a resolution of less than a minute (Tanaka and Nishi, 1986; Maus et al., 2006; Schäler et al., 2015). Work along these lines is relevant beyond the field of vitrimers, as crystallization in networks, in particular highly crosslinked ones, is little researched and understood in general (Golitsyn et al., 2019). The cited paper reports on one of the few systematic studies of model networks with short but defined network chain length, where the normally high crystallinity of polyethylene oxide (PEO) linear polymers (typically 80% or higher) is significantly reduced in end-linked networks with chains of less than a few kDa. Tightly crosslinked thermosets are mostly amorphous, as are the covalent equivalents of the samples studied herein (Soman et al., 2022a). A peculiar exception seems to be urea/formaldehyde resins, which can reach crystallinities in the 50% range (Wibowo and Park, 2020). This, however, seems to be a special case where for the given small-molecule-based resin formulation, crystallization of linear stretches and crosslinking compete already during curing.

To sum up our recent work relevant for the given follow-up study, the role of dynamic bond exchange on crystallization in networks has been investigated in ethylene vitrimers (Soman et al., 2022a). They are comprised of monodisperse telechelic alkane diols $(\text{CH}_2)_n(\text{OH})_2$, with n being 10 and 12 in this work, crosslinked by boric acid, $\text{B}(\text{OH})_3$, samples with exact linker spacing were synthesized. The samples are denoted C_{10} and C_{12} , respectively. In comparison to analogous, permanently crosslinked networks, which do not show any evidence of crystallization, ethylene vitrimers did crystallize with local packing motifs analogous to those of polyethylene (PE). At room temperature, crystallization was previously followed over 30 days, and the melting temperature of the crystals continuously increased by ~ 25 degrees, while crystallinity as reflected in the DSC-detected melting enthalpy remained nearly constant. While the absolute crystallinity was not yet quantified, wide-angle X-ray scattering (WAXS) suggested rather high crystallinity indicated by the virtual absence of an amorphous halo and high thermal conductivity. WAXS further revealed an unexpected crystal-crystal transition which indicated a second crystal polymorph. It is interesting to note that

these changes in properties take place in quiescent samples. We attributed the initial crystallization and the subsequent evolution in properties to the dynamic bonds. Local rearrangements of the network strands facilitated by bond exchange are hypothesized to allow crystal perfection or an increase in crystallinity, and to enable a crystal-crystal transformation into a more stable polymorph. Even more recent work by Kuenstler and Bowman (2023) have also investigated crystallization in dynamic networks, where different catalysts allowed for control of bond exchange kinetics. A similar long-time evolution in T_m was observed, and the actual T_m values were found to be inversely correlated with the exchange kinetics.

Here we investigate in more detail the crystallization of the recently reported high-crosslink-density ethylene vitrimer (Soman and Evans, 2021; Soman et al., 2022a; Soman et al., 2022b; Lv et al., 2023). We focus on the kinetics of isothermal crystallization as well as the annealing/recrystallization kinetics of two different samples with C_{10} and C_{12} spacer length. Along the lines of earlier work on the confined crystallization of pharmaceuticals (Sonnenberger et al., 2016), we are able to distinguish two different crystal polymorphs by simple ^1H NMR via their dipolar line width, allowing us to monitor the interconversion in real time. Interestingly, we could identify an unexpected superheating effect, which challenges the determination of the melting points reported previously for our samples, as well as a prominent memory effect on crystallization.

2 Materials

The samples investigated in this work are prepared via the condensation of alkane diols with boric acid. The synthetic procedure follows exactly from our prior work (Soman et al., 2022a)

3 Results and Discussion

3.1 Samples and rheological characterization

The previously investigated and carefully dried C_{10} and C_{12} ethylene vitrimers were prepared in sealed glass tubes flushed with Argon for NMR measurements. We refer to our previous publication for details (Soman and Evans, 2021). The structure of the dynamic networks is shown as in inset in **Figure 1a**, where we have compiled the terminal relaxation times of these samples. The rheological measurements were already done in the context of our first publication from which the Arrhenius slopes were determined, but the relaxation times of C_{10} and C_{12} were not explicitly published there. The data shows that the networks relax on a 0.1 – 1 s timescale in the temperature range relevant for the present study. This will be taken up in the discussion below.

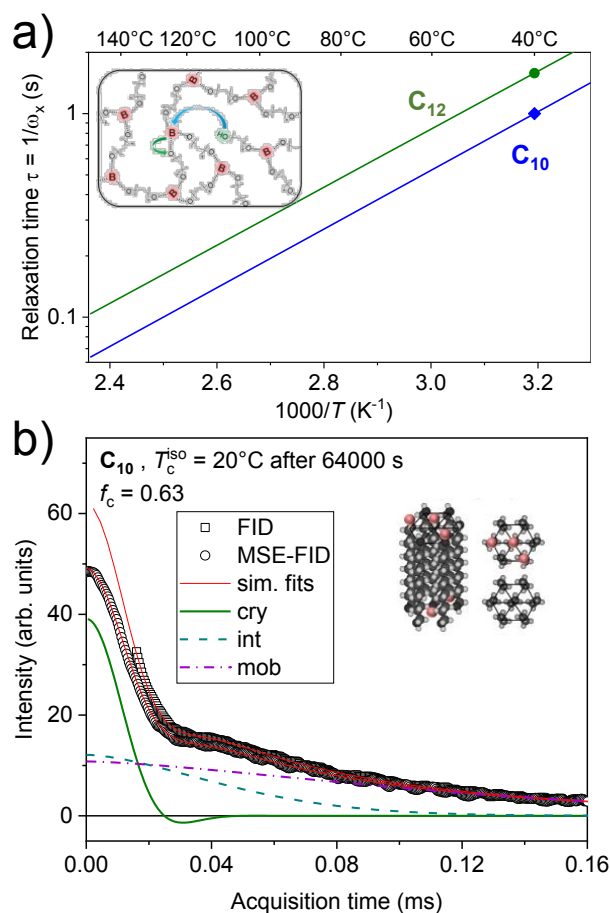


Figure 1. a) Arrhenius plot of vitrimer relaxation times calculated as the inverse of the rheological crossover frequency of samples C₁₀ and C₁₂ measured at 40°C along with extrapolations based upon shift factors (data taken from Soman et al., 2021). The inset shows the network structure. b) Simultaneous three-component fit of ¹H FIDs, detected during isothermal crystallization at 20°C, both directly after the excitation pulse and after an MSE to avoid the dead time. The fits provide the crystalline fraction (f_c) and second moments of the corresponding crystalline lineshapes (M_2), as well as T_2 relaxation times characterizing the restricted mobility of the intermediate and amorphous fractions. The inset shows the hexagonal crystal structure formed under the given condition (Soman et al., 2022a)

3.2 Differential scanning calorimetry (DSC)

Samples (2-5 mg) were melted into hermetically sealed pans in an Argon glovebox. Crystallization proceeded in the glovebox at ambient temperature. A TA Instruments DSC 25 was used to control the heating and cooling rates as described in the text.

3.3 Combined small- and wide-angle X-ray scattering (SAXS/WAXS)

Samples were prepared for X-ray analysis by sealing between two Kapton sheets using epoxy and a spacer. A Xenocs GeniX3D Cu K α source generated 1.54 Å X-rays and scattering was recorded on a Pilatus 1M detector. Sample-to-detector distance was calibrated using silver behenate, and 2D data was azimuthally averaged using a MATLAB-based GIXSGUI software to determine intensity as a function of wavevector q .

3.4 NMR spectroscopy and data analysis

NMR experiments were conducted on a Bruker minispec mq20 instrument at a Larmor frequency of about 20 MHz. Pulse lengths were in a range of 2.5 and 5 μ s for flip angles of 90° and 180°, respectively. A recycle delay of 1 s was sufficient for full T_1 relaxation of both semicrystalline and amorphous/molten samples. The free induction decay (FID) of the proton signal is measured both directly and after a magic-sandwich echo (MSE-FID) to address the dead-time problem (Maus et al., 2006; Schäler et al., 2015). The latter suffers some intensity loss, which means that we fit the two datasets (restricted to an acquisition time interval of 200 ms) simultaneously to a three-component model accounting for the crystalline (index c), amorphous (a), and interfacial (i) regions (Schäler et al., 2015):

$$I(t) = A_c e^{-(at)^2/2} \sin(bt)/(bt) + A_i e^{-(t/T_{2i})^{\beta_i}} + A_m e^{-(t/T_{2m})^{\beta_m}}$$

The MSE leads to a small reduction of the amplitude prefactors, which are the only fitting parameters that are allowed to vary between FID and MSE-FID signals. Sample fits are shown in **Figure 1b**. Most importantly, these fits provide the crystalline fraction ($f_c = A_c/(A_c + A_i + A_m)$), using the amplitudes of the undistorted FID. Since NMR intensities are strictly proportional to proton fractions, f_c reflects crystallinity by weight. Information on the crystal packing is obtained in terms of the second moment of the crystalline lineshape ($M_2 = a^2 + b^2/3$), which depends on the overall strength of multi-proton dipole-dipole interactions as governed by their proximity, i.e. the density. It thus reflects the crystalline perfection of the sample.

The β exponents for the intermediate and mobile amorphous components are stretching parameters reflecting in the one hand dynamic heterogeneity, and on the other hand a partial Gaussian character typical for dipolar dephasing originating from the dominance of residual dipolar couplings in the short network chains (a discussion is given below). This explains why the β values were always ranging between 1 and 2. The T_{2i} was around and below 0.1 ms in the network melt down to 0.03 ms for fully crystallized samples, while T_{2m} was around 0.2, staying the same or even increasing somewhat during crystallization. For the given short network chains, T_2 values are in the range of 0.1 to 0.2 ms, which means that the mobile and intermediate components together reflect inhomogeneities of the network structure. As network chains become part of the crystal, dangling defects with longer T_2 are left over, thus it increases in some cases.

4 Results and Discussion

4.1 High crystallinity and increase of T_m being related to crystal perfection

Our NMR-based crystallinity determination, applied to monitor isothermal crystallization of both samples in **Figure 1a,b** confirms the initial WAXS-based observation that the samples reach high crystallinity of ~90% within a reasonably short time. This is an astonishingly high crystallinity for polymer samples that are quiescently kept at room temperature, in particular considering the disordered network structure and the short chains. The boric acid ester linkers are thus expected to be part of an ordered fold surface or to form intracrystalline layers, along the lines of recent studies of linear precision polymers (Trigg et al., 2017; Marxsen et al., 2021). To reach such high crystallinity, long-chain polymers are usually annealed at higher temperatures or shear aligned, and have to feature fast chain mobility in the crystals (Schulz et al., 2022). For highly crosslinked networks, it is in fact a peculiarity, as mentioned in the Introduction.

The crystallization kinetics $f_c(t)$ were evaluated by fits to the Avrami function (black solid lines in **Figure 2a,b**), which revealed Avrami exponents in the range between about 1 and 2. Since the variations were non-systematic across all our data (see also **Figure 5** below), arising from uncertainties (noise) in the data at short times and possible secondary crystallization at longer times, we refrain from reporting and discussing the fitted exponents, but just report crystallization half times $t_{1/2}$. The values are 830 and 63 min for C₁₀ and C₁₂, respectively, with the large difference mainly attributed to the different undercooling at 20°C. In both samples, the second moment continues to grow during crystallization, indicating a gradual densification or a suppression of fast small-angle motions. We thus interpret the observation in terms of an increase in crystal perfection, which, notably, takes place without further significant increase in crystallinity. The increase in enthalpy of melting previously reported (Soman et al., 2022a) can in this new light be attributed to the increasing crystal perfection observed from NMR rather than percent crystallinity.

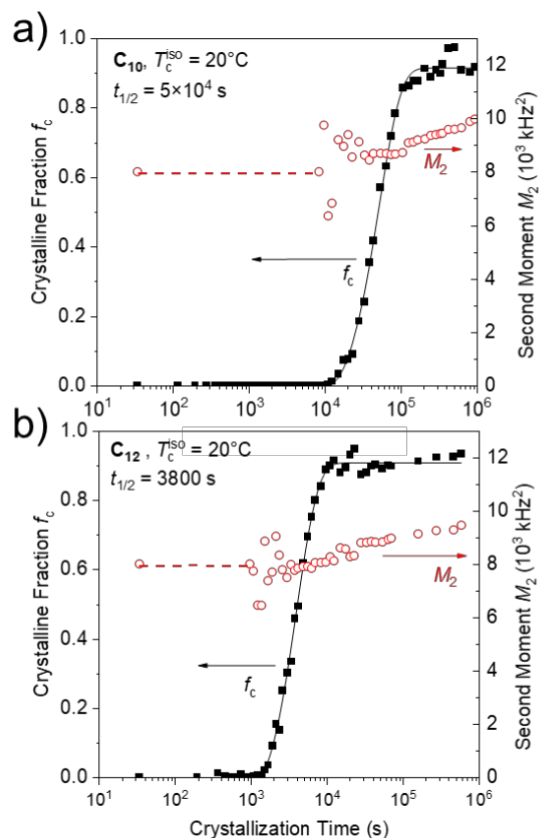


Figure 2. Crystalline fraction (f_c) and second moments of the crystalline lineshapes (M_2) plotted as a function of annealing time at 20 °C for the a) C₁₀ and b) C₁₂ ethylene vitrimers. The red dashed line indicates the time range where M_2 was fixed since stable fitting of a finite crystalline fraction was impossible. The black solid lines are fits to the Avrami function, from which we take the crystallization half time.

The crystallization of the networks was also tracked by WAXS over a 150 day crystallization period (**Figure 3a,b**). After 1 day of crystallization, the WAXS patterns already indicate a high crystallinity as inferred from the absence of an underlying amorphous halo which is observed in the scan of the amorphous network (0 min scan). In addition to the high crystallinity, the FWHM of the peaks does not show significant variation after the first day indicating a stable crystal size. At room temperature, the C₁₀ eventually undergoes a spontaneous transformation from hexagonal to orthorhombic but not

within the first 30 days. In the C_{12} network, the crystal structure appears to be hexagonal over the entire 150 day window and only transitions to orthorhombic at elevated temperatures of 65 °C (to be discussed later).

The M_2 values for hexagonal and orthorhombic crystals are distinct, and the NMR measurements reveal only hexagonal crystals in the first month consistent with the prior WAXS data. This is evidenced by the comparison of M_2 values for a C_{10} sample crystallized isothermally at 20°C during heating and cooling with a rates of 1 K/min in **Figure 3c**. The first heating scan to 40°C reveals a slight decrease of M_2 in a range below 10^4 kHz² (accompanied by only a slight decrease in crystallinity from about 0.92 to 0.86). The sample was then cooled again to room temperature, upon which the variation of M_2 is fully reversible, as evidenced by the second-heating data. Crystallinity actually diminishes above 48 °C, and annealing at 50 °C leads to a recrystallization into the orthorhombic phase (details to be discussed below in the context of **Figure 6a**). Importantly, this is directly reflected in the considerably larger M_2 value of around 1.2×10^4 kHz², which is now typical for similar crystal structures such as those of PE or aliphatic polyesters (Schäler et al., 2013). The orthorhombic crystallinity in the shown T range varies between 79 and 85%. The T -dependence of its M_2 is even weaker, and depends mostly on thermal expansion. The stronger dependence and lower values of the initial hexagonal phase may be related to enhanced dynamics such as increasing amplitude of local librations or conformational fluctuations (Schäler et al., 2013). It is noted that the M_2 values for the two phases are too close to allow for a separation of the fractions of either phase by fitting in samples where they coexist. However, in-between values can be interpreted qualitatively in terms of coexistence.

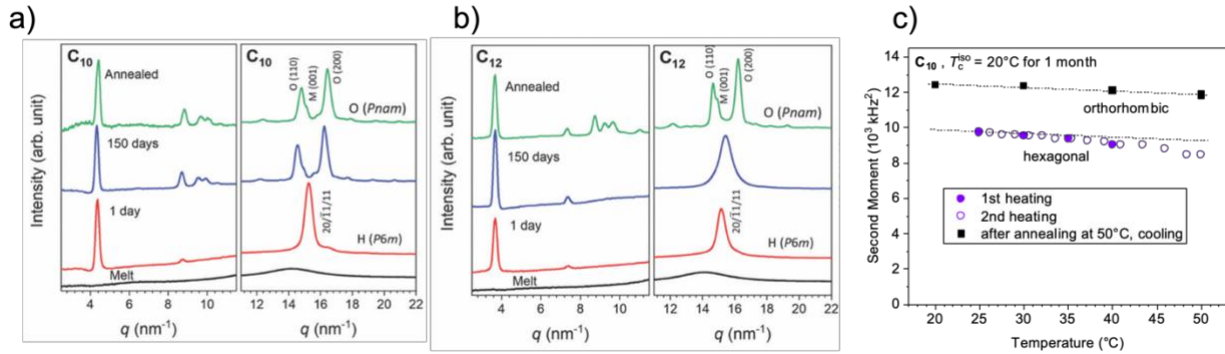


Figure 3. Time evolution of SAXS/WAXS spectra in the a) C_{10} and b) C_{12} networks at 23 °C revealing the transformations from hexagonal to orthorhombic unit cells which occur with time (in the C_{10} network) or heating to 65 °C (in the C_{12} network). Data reproduced from Soman et al., 2022a with permission. c) Second moment of the crystalline lineshape for the two relevant crystal structures as a function of temperature, recorded upon heating or cooling at 1 K/min. The parallel dashed lines indicate the typical weak decay due to thermal expansion of the lattice; the stronger and reversible decrease in the hexagonal phase indicates increased dynamics, e.g. increasing concentration of fluctuating conformational defects or librational motions.

Our previous report of increasing T_m with crystallization time corresponded to samples heated at a rate of 20 °C/min in the DSC (Soman et al., 2022a). In the present work, the time evolution of T_m was examined using different heating rates spanning from 0 to 20 °C/min. The DSC traces are shown in the Appendix (**Figure A1**). In all cases an increase is observed with essentially the same slope (**Figure 4**). Surprisingly, the value of T_m decreases significantly with slower heating rates, a phenomenon also observed in conventional linear PE (Hellmuth and Wunderlich, 1965). This puts the previous T_m results (Soman et al., 2022a), taken to be not too far from equilibrium, into

perspective. In that study, the increase in T_m was attributed to increasing crystallinity due to the change in the enthalpy of melting. Here, it is shown that increasing perfection leads to the increase in T_m . The higher apparent T_m values at 20 °C/min are thus attributed to the superheating of crystals, which refers to the ability to be heated above the equilibrium melting temperature (Hellmuth and Wunderlich, 1965). For polyethylene and polyoxymethylene used in that study, the superheating effect resulting in an ~4-5 °C shift for a range of heating rates from 0.6 to 20 °C/min. The effect is much larger in ethylene vitrimers. In the present work, we deduce from **Figure 4** that while the vitrimeric rearrangements and enhanced crystal perfection result in systems which are more stable and melt at higher T_m , the superheating effect, apparent from the large differences between the different heating rates, is affected only little. We hypothesize that the melting kinetics are governed by the timescale of vitrimeric rearrangements, which could be slower in the organized environment between the crystalline alkyl sheets. Note that the total time for a heating cycle ranges from 5 – 100 minutes depending on the ramp rate, and the rheological crossover time in the melt state is ~ 0.1 sec (**Figure 1a**), indicating ample time for relaxation events and bond exchange to modify the samples during this ramp.

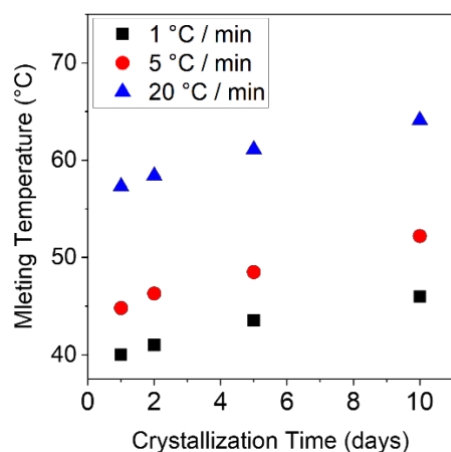
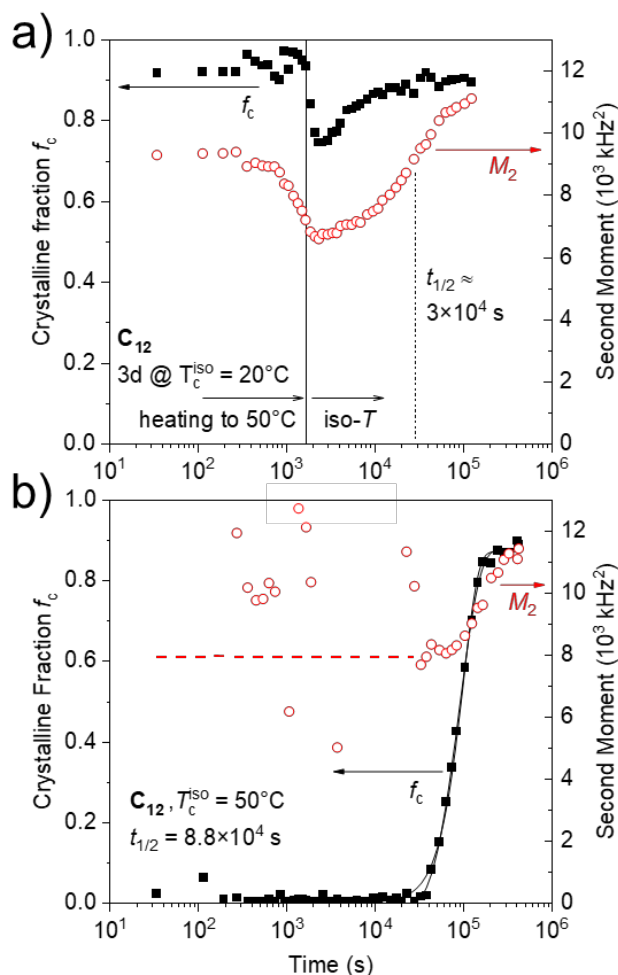


Figure 4. Melting temperatures of the C₁₀ network as a function of crystallization time with different heating rates.

As noted in the Introduction, Kuenstler and Bowman (2023) found that thiol/thioester-based vitrimers with crystallizable C₁₀ alkyl-dithiol linkers also show a long-time increase in T_m , the absolute value which is of the order of 20°C, thus much lower than in our case. T_m was higher for systems with faster bond-exchange kinetics. The annealing time dependence of T_m was not studied explicitly. The effect was attributed to potential lamellar thickening – which was not measured. This explanation must be reconciled with a system in which the disordered thioether/(thio)ester moieties are likely part of the lamellae and whose thickness is further bound by the crosslinks. In that work, the extended C₁₀ units probably also crystallize in PE-type fashion similar to polymers with alkyl sidechains. Such a system, namely atactic poly(*n*-octadecyl methacrylate) was studied by Beiner and coworkers (Hempel et al., 2006), where a similar long-term increase of T_m was also observed. In that case, the C₁₈ side-chains form interdigitated layered nanodomains separated by the disordered main chains, within which a hexagonal crystalline packing develops and was hypothesized to perfection over time, as detected by a slow increase of both T_m as well as the heat of melting. It remains an open question whether a thickening of an initially shorter crystallized stretch of the C₁₈ (as depicted by Hempel et al.), or an improving organization/packing at the crystallite surface, is responsible for the

266 observed increases in T_m and melting enthalpy. In our system with its much shorter C_{10} chains, the
 267 second option appears more likely, also in view of the largely invariant sharp SAXS/WAXS peaks
 268 (Figure 3).



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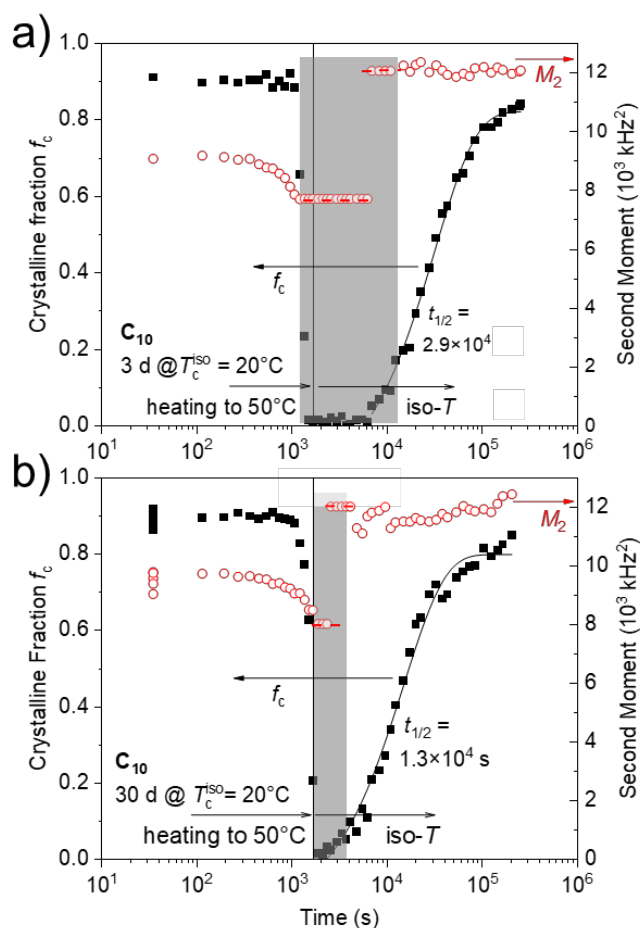
270 **Figure 5.** a) C_{12} sample undergoes a crystal-crystal transition when annealed at 50 °C after isothermal
 271 crystallization at RT for 1 day. b) The C_{12} network directly cooled from the melt (150°C) transitions to the
 272 orthorhombic phase during isothermal crystallization at 50°C. The two black solid lines are Avrami fits
 273 excluding or assuming an onset time t_0 of 2.8×10^4 s. The two scenarios, providing Avrami exponents of 2.6 or
 274 1.6, respectively, cannot be distinguished on the basis of the data/fitting quality.

275 4.2 Higher-temperature recrystallization and melting

276 As observed in the SAXS/WAXS data of Figure 3, the C_{12} sample can transition from hexagonal to
 277 orthorhombic when annealed at 65 °C. This transition was tracked using NMR on a C_{12} network first
 278 crystallized for 1 day followed by heating to 50 °C in 1700 s (rate of ~ 1 °C/min) as shown in Figure
 279 5a, where we used the lower temperature so as to be able to compare with isothermal crystallization
 280 at the same temperature. This procedure leads to a minor partial melting, and then recrystallization
 281 into a denser phase (orthorhombic) but with almost the same crystallinity and a higher perfection.
 282 Notably, the M_2 values decrease first on heating from 20 to 50°C (see also Figure 2c), suggesting
 283 fast (but effectively small-angle) segmental fluctuations with increasing amplitude. Then, they
 284 steadily increase to the level expected for the orthorhombic crystal. The increase of M_2 can be

285 interpreted in terms of increasing crystalline perfection, or simply by a converted fraction. Taking the
 286 latter perspective, a conversion half time of 500 min is estimated.

287 This can be compared with the isothermal crystallization of the same network that is cooled directly
 288 from the melt to 50 °C, revealing a half time of about 1500 min (**Figure 5b**). This temperature is
 289 apparently below the equilibrium melting point of the hexagonal phase, but is above the annealing
 290 temperature required to form the orthorhombic phase. The hexagonal phase forms first (M_2 of around
 291 8000 kHz²), and then gradually converts to the orthorhombic phase during the crystallization ($M_2 > 11$
 292 kHz²). The crystallinity still reaches $f_c \sim 0.9$. The orthorhombic phase does not seem to directly
 293 nucleate and we note that this denser phase is effectively at a lower degree of undercooling than the
 294 hexagonal phase.



295

296 **Figure 6.** Memory effect on isothermal recrystallization at 50°C observed upon annealing of C₁₀ a sample
 297 crystallized isothermally at 20 °C for a) 3 days and b) 1 month. The latter has a stronger memory effect and
 298 thus faster crystallization (including a shorter apparent “induction time”). Note that isothermal crystallization
 299 at 50°C after cooling from the virgin melt is not observed on the given timescale.

300 4.3 Memory effect in vitrimer crystallization

301 Polymer crystals are known to have memory effects which can take long times to anneal out and
 302 fully erase thermal and processing history (Cheng et al., 1986; Bastiaansen et al., 1990; Hamad et al.,
 303 2015; Sangroniz et al., 2020). The C₁₀ network was first examined by crystallizing for 3 days or 1

month, followed by heating to 50°C (**Figure 6**). This temperature is above the melting point of the hexagonal phase initially formed on crystallization at lower temperatures (see **Figure 4**), and no crystallization directly into the orthorhombic phase was observed with several days at 50°C after cooling from the melt. As expected, both 20°C pre-crystallized samples show essentially complete melting upon heating to 50°C, yet this is now followed by recrystallization directly into the orthorhombic phase (as evidenced by the high and constant M_2). The 3-day sample (**Figure 6a**) shows a notable induction period, while the 1-month sample has an even stronger memory effect (**Figure 6b**) with an apparently shorter induction period and shorter overall crystallization half time. The orthorhombic phase apparently does not nucleate well (or not all all) and requires some “memory” facilitating its formation

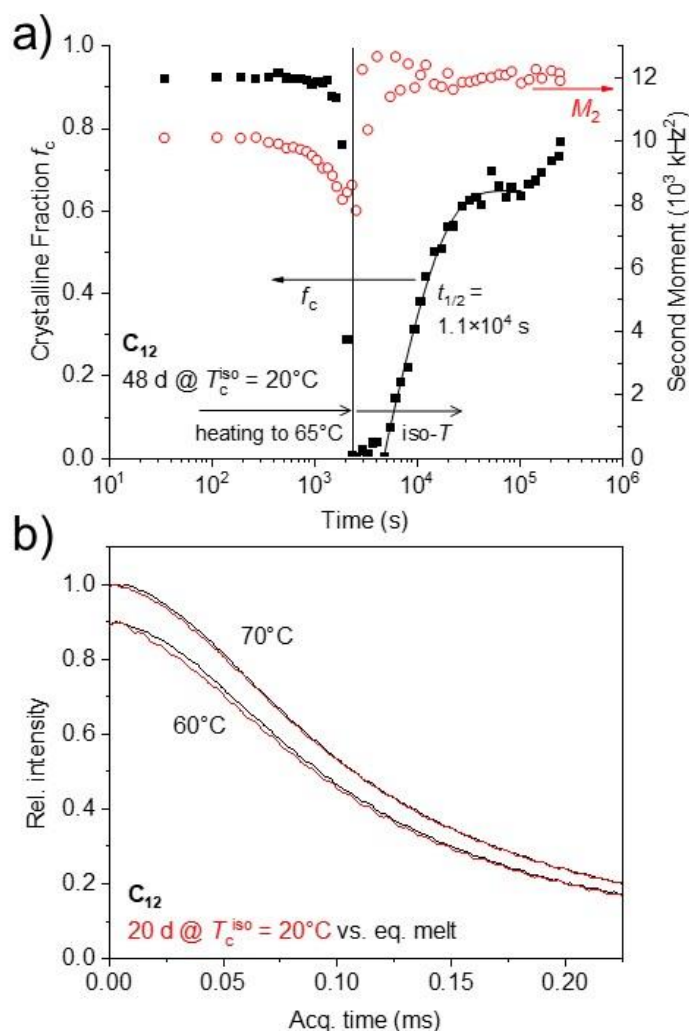


Figure 7. a) Memory effect of C_{12} on isothermal recrystallization at 65°C . **b)** MSE-FIDs at 60°C and 70°C; the latter was taken after 40 min. hold time at 60°C. The intensity data at 60°C is scaled by a factor of 0.9 for visual separation.

The C_{12} networks were also examined for memory effects, but heating to 50 °C is not sufficient to induce the same phenomena as in the C_{10} networks, as isothermal crystallization from the virgin melt and formation of (or transition into) the orthorhombic phase is already observed at this temperature (see **Figure 5 a,b**). We have thus focused on higher temperatures, where a sample crystallized at 20

°C for 1.5 months showing the hexagonal modification melts completely at 65 °C and recrystallizes directly into the orthorhombic phase after a lag time of about 1.5 hours (**Figure 7a**). Qualitatively the same behavior was observed for a sample crystallized at 20°C for 20 days after heating to 60°C, where crystallization sets in already after about 20 min. (data not shown). In contrast, a virgin melt cooled to the same temperatures did not crystallize.

As to the origin of this memory effect, one may consider an adapted network mesh topology providing a “melt memory” with pre-arranged chains. This option is, however, unlikely, as the memory is stable for hours, while the terminal (flow) time of the vitrimers, as governed by the bond-exchange process, is well below the second scale (see **Figure 1a**). To test this hypothesis, or check for the possible remainder of a small crystalline fraction, we compare in **Figure 7b** two (MSE-)FIDs of melt states at 60°C and 70°C for a sample crystallized for 20 days with and without memory (the latter taken after complete melting at 150°C for about 15 min.). For this experiment, the pre-crystallized sample was held at 60°C for 40 min, during which time crystallization set in visibly. It was then further heated to 70°C and held. The shown data are averages of the FIDs taken during the induction times, where the signals did not change visibly and where no stable fitting of a finite crystallinity was possible. Nevertheless, the direct comparison at 60°C with the data of the equilibrated melt shows that the signal from the samples “with memory” is always decaying somewhat more quickly, the data being compatible with a crystallinity degree on the percent level. The effect is very small but still visible at 70°C, while at 80°C (with data taken after another 40 min holding at 70°C) showed signals that are identical to those of the equilibrated melt within the noise. Thus, the memory is most likely related to the existence of a very small fraction of high-melting crystallites, possibly such ones for which the perfectioning has progressed more. We note that it is expected that the observed differences pertain not just to a minute fraction of crystalline signal but a somewhat decreased T_2 relaxation time of a part of the sample (i.e., the intermediate fraction in our three-component fit), arising from some degree of immobilization of the chains surrounding the small crystallites.

We finally add some comments on the shape of the melt-state FIDs, which could hold information on a “melt memory”. It is approximately Gaussian, as expected for very short network chains featuring rather large residual dipolar coupling constants arising from the strongly anisotropic motions of the short end-fixed chains (Saalwächter and Heuer, 2006). In other words, the transverse relaxation process (normally characterized by a T_2 relaxation time) is rather pure dipolar dephasing, where an orientational (“powder”) average combined with multiple multi-spin couplings leads to the well-known Gaussian shape. In case of a homogeneous system, is described by $I(t) = \exp\{(9/40)D_{\text{res}}^2 t^2\}$, where D_{res} is the residual dipolar coupling, which is inversely proportional to the length of the network strands (Saalwächter and Heuer, 2006). Since in our case D_{res} is large due to the short strands (corresponding to a high crosslink density typical for thermosets), direct analysis of the FID, which is normally precluded by field inhomogeneities and additional contributions of segmental dynamics, becomes possible. Topological structures such as loops would contribute lower D_{res} and the relaxation function would become modified/stretched, yet again, no differences are apparent between the cases with and without memory.

Another relevant piece of information in this context is a nearly exponential long- T_2 tail, for which fitting (not shown) provides a relative fraction of less than 10%. This corresponds to network defects, whose rather low amplitude confirms the high conversion of the networks with its mostly 3-functional crosslinks. More in-depth analysis into the network structure and a more stable fitting is possible by using multiple-quantum (MQ) NMR, along the lines of our previous work (Saalwächter

367 and Heuer, 2006). Results along these lines are deferred to a future publication, where we focus on
368 stoichiometrically imbalanced, defect-containing samples.

369

5 Conclusions

The crystallization of ethylene vitrimers was investigated by time-domain NMR to probe the kinetics, perfection, and polymorphic transitions of these systems. NMR analysis confirms that the crystallinity is very high, of order 90%, in agreement with prior WAXS data. A notable and significant long-time increase of T_m upon annealing was shown to not be related to an increase in crystallinity (reaching its final level comparably quickly) but rather to an increased crystal perfection, enabled by a reorganization of the dynamic bonds on the surface of the crystalline alkyl layers. The dynamic bonds thus become part of a layered crystal structure, as also evidenced by a sharp SAXS peak reflecting the boric ester layer spacing.

These results may be of general relevance for systems in which alkyl nanodomains can crystallize. Both in a recent study of somewhat more complex C₁₀-based vitrimers (Kuenstler and Bowman, 2023) and in a linear methacrylate polymer with C₁₈ side-chains (Hempel et al., 2006), similar T_m increases were reported. In the latter case, the melting enthalpy change over time was also studied by DSC, and shown to increase along with T_m , in the same way as in our ethylene vitrimers (Soman et al., 2022a). We could here show that this increase is not related to increased crystallinity but is rather related to perfectioning of the layers containing the crosslinks, meaning that the surface energy is reduced. This emphasizes the relevance of our NMR investigations, providing a robust measure of the degree of crystallinity.

NMR also allows us to follow the polymorphic transition from hexagonal to orthorhombic based on the dipolar second moment of the crystalline lineshape, originating from slightly different proton densities and being detected in terms of differences in transverse relaxation times. We further identified a strong superheating effect, where higher heating rates lead to significantly higher apparent T_m values as the samples are heated beyond their equilibrium melting temperature. The related sluggish melting kinetics may be tied to a slower dynamic bond exchange in the crystalline interlayers. This finding suggests that it is generally difficult in the given system, as well as in related ones (Kuenster and Bowman, 2023; Hempel et al., 2006), to measure – and draw conclusions from – the actual melting temperature of the given structure.

The vitrimers are finally shown to exhibit strong memory effects, where our NMR observations are compatible with a small remaining high-melting fraction of well-optimized crystals that can nucleate the crystallization directly into the orthorhombic phase. In summary, our study has provided key new insights into the role of dynamic bonds on crystallization in dense networks, that would typically not crystallize despite being made up of monodisperse building blocks. Future work is directed towards introducing controlled defects and studying their impact on terminal flow and on semicrystalline structure formation.

6 Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

7 Author Contributions

Synthesis of vitrimers, X-ray scattering, rheology, and calorimetry were performed by B.S. All NMR experiments, fitting, and analyses were performed by K.S. The project idea, development, and writing of the manuscript was contributed by all authors.

8 Funding

This work was partially supported by the National Science Foundation through award CBET-2029928 (B.S. and C.M.E.). Partial funding was also provided by the Deutsche Forschungsgemeinschaft (DFG) in the framework of the SFB-TRR 102 (project-ID 189853844), project A1 (K.S.).

9 Acknowledgments

We acknowledge the use of the facilities at the Materials Research Laboratory at the University of Illinois, Urbana-Champaign.

10 Data Availability Statement

The datasets [GENERATED/ANALYZED] for this study can be found in the [NAME OF REPOSITORY] [LINK]. [will be added upon final submission]

11 Appendix

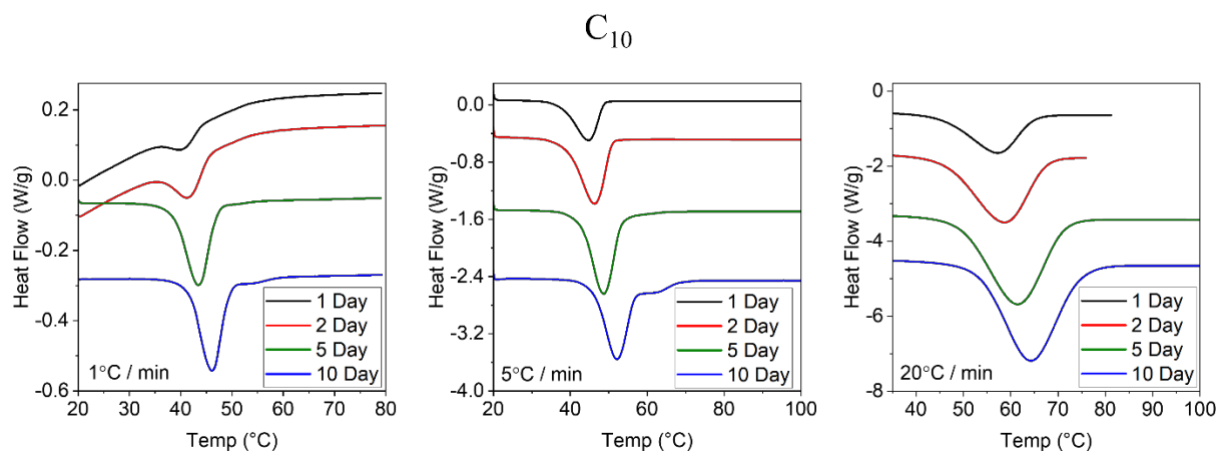


Figure A1. DSC heating traces of C_{10} crystallized isothermally at 20°C for different durations (1-10 days), using different heating rates of 1 (left), 5 (middle) and 20 °C/min (right).

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