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	1 2	Microstructural characterization of a star-linear polymer blend under shear flow by using rheo-SANS
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Abstract

We present an investigation into the dynamic relaxation mechanisms of a polybutadiene blend composed of a four-arm star (10 wt. %) and a 14 linear polymer matrix in the presence of an applied shear flow. Our focus was the response of the star polymer, which cannot be unambiguously 15 assessed via linear viscoelastic measurements since the signature of the star polymer can barely be detected due to the dominant contribution of 16 the linear matrix. By utilizing small-angle neutron scattering (SANS) coupled with a Couette shear device and a deuterated matrix polymer, we 17 investigated the dynamics of the minority star component of the blend. Our results confirm that the stars deform anisotropically with an increas-18 ing shear rate. We have compared the SANS data with predictions from the well-established scattering adaptation of the state-of-the-art tube 19 model for entangled linear polymer melts undergoing shear, i.e., Graham, Likhtman, Milner, and McLeish (GLaMM) approach, appropriately 20 modified following earlier studies in order to apply to the star. This modified model, GLaMM-R, includes the physics necessary to understand 21 stress relaxation in both the linear and nonlinear flow regimes, i.e., contour length fluctuations, constraint release, convective constraint release, 22 and chain retraction. The full scattering signal is due to the minority star component and, although the contribution of the linear chains is hidden 23 from the neutron scattering, they still influence the star polymer molecular dynamics, with the applied shear rate ranging from approximately 24 8 to 24 s^{-1} , below the inverse relaxation time of the linear component. This study provides another confirmation that the combination of rheology 25 26 and neutron scattering is an indispensable tool for investigating the nonlinear dynamics of complex polymeric systems. © 2020 The Society of

27 Rheology. https://doi.org/10.1122/1.5121317

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28 I. INTRODUCTION AND THEORETICAL29 BACKGROUND

Over the past decade, new applications and processing 30 methods have led to a renewed interest in the mechanical prop-31 erties of polymer blends comprising both linear and branched 32 architectures [1-11]. There is both a practical and theoretical 33 interest in understanding how the properties of a linear 34 polymer matrix might be altered via blending with a small 35 amount of polymer possessing a different, well-defined, 36 complex architecture. An enormous advantage would be 37 gained, for example, if there were a predictable recipe for 38 blending polymers of different well-defined complex architec-39 tures to achieve and optimize mechanical properties of a 40 polymer melt, e.g., extension hardening [12–21]. 41

42 An active area of research for polymer solutions and melts 43 containing both branched and linear polymers is to understand

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how the presence of a linear chain matrix influences the relax-44 ation of branched chains. In the linear viscoelastic regime, 45 significant progress has been made and different theoretical 46 models describe accurately the effects of adding branched 47 polymers into a linear matrix [7-10]. Actually, a recent study 48 by Hall et al. has shown that when the linear polymer has a 49 relaxation time longer than the star component, the terminal 50 relaxation time dependence has nonmonotonic dependence on 51 the blend composition, and this result is obtained both theoret-52 ically and experimentally [11]. However, in the nonlinear flow 53 regime (relevant to industrial processing conditions), the theo-54 retical understanding of this problem is still in its infancy, and 55 conventional rheological techniques are not always well suited 56 for probing the response of branched polymers, especially 57 when their fraction is small [22-25]. 58

The tool necessary to predict both molecular scale structure 59 and macroscopic flow in realistic processing configurations is 60 a quantitatively accurate microscopic theory. The macroscopic 61 properties of a bulk polymer in the fluid state depend on the 62 chain configurations and their modification in flow, and these, 63 in turn, depend on the topological interactions between the 64 polymer molecules. The primary topological effect in concentrated solutions or melts is the entanglements, namely, the 66

0148-6055/2020/64(3)/1/10/\$30.00

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67 long-lived topological constraints that form a physical network
68 that hinders the chains from translating normal to one another,
69 thus impacting their dynamics [26–29].

The most significant advance in the field was the tube 70 model theory developed originally by de Gennes, Doi, and 71 Edwards [28,29] and subsequently improved to provide more 72 accurate predictions of the linear and in part nonlinear visco-73 elastic behavior, based on the molecular characteristics of 74 entangled polymers, mainly molar mass, plateau modulus, and 75 Rouse time of an entanglement segment [4, 14-16, 19, 30]. 76 Although the tube model was originally intended to describe 77 entangled melts (or solutions) of monodisperse, linear poly-78 mers, it has since been successfully extended to polydisperse 79 80 linear chain melts [19,31,32] and complex macromolecular architectures [33–36]. However, at present, the understanding 81 of (linear and nonlinear) dynamics of mixed architectures 82 (branched and linear polymers) remains an open issue, as will 83 be discussed below. 84

According to the tube model, topological constraints in 85 the surrounding environment restrict the lateral motion of a 86 polymer chain but not its curvilinear motion. To the extent 87 that the polymer chain has some limited freedom for lateral 88 motion, the complexities of the surrounding topological envi-89 ronment can be replaced by a smooth tubelike structure 90 [26–29]. The polymer cannot pass through the tube walls 91 (which are not fixed in reality, as discussed below), but it is 92 free to fluctuate within this tube by Rouse-like rearrange-93 ments of its configuration. Rearrangements taking place over 94 distances comparable to the tube diameter occur very fast, on 95 96 a time scale τ_e (Rouse relaxation time of an entanglement segment). Rearrangements within the tube taking place over 97 distances comparable to the full chain contour length are 98 slower, taking a chain Rouse time $\tau_R = Z^2 \tau_e$, where Z is the 99 number of entanglements per chain. To fully relax its stress, 100 the polymer must evacuate the tube via different mecha-101 102 nisms, namely, curvilinear diffusion (reptation), contour length fluctuation (CLF), thermal or convective constraint 103 release (TCR/CCR), and constraint release Rouse (CRR) 104 relaxation [3,19,28,37]. The characteristic time scale for 105 relaxation by reptation is $\tau_{d,\text{lin}} = 3Z^3 \tau_e$. Depending on the 106 details of the polymer architecture and the applied deforma-107 tion (linear or nonlinear), the relaxation processes (CLF, 108 TCR/CCR, and CRR) may play an important, or in some 109 110 cases a dominant, role.

CLFs reflect the fact that chain ends can escape the con-111 112 fining tube (and thereby relax the associated stress) as they fluctuate back toward their preferred unstretched contour 113 114 length [14–26]; it also applies to star arms that do not reptate. TCR and CCR are complementary mechanisms that account 115 for the release of entanglements due to Brownian or exter-116 nally induced motion of chains, respectively. When a 117 constraint is removed by reptation or CLF, we have TCR, 118 since reptation and CLF are driven by thermal fluctuations. 119 When a constraint is removed by convection due to flow, 120 CCR takes place [19,30,38,39]. CRR relaxation is the mecha-121 nism by which stress relaxes following TCR or CCR. When 122 123 TCR and CCR are present, the tube is better described as a fluctuating field instead of a fixed object. To a first approxi-124 125 mation, one expects that CRR and TCR allow the whole tube to undergo the same kind of Rouse-like rearrangements but 126 on a much longer time scale than the Rouse time, τ_R . In the 127 present work, we refer to TCR and CCR as stress relaxation 128 mechanisms, but in such cases, the CRR mechanism is also 129 implied. 130

With this wealth of relaxation mechanisms, we can 131 describe the distinctive features for melts of linear and star 132 polymers, as well as star/linear blends where constraint 133 release effects can be important. Upon applying a certain 134 shear rate less than $(\tau_R)^{-1}$, chain retraction can be viewed as 135 an instantaneous process for both the linear matrix and the 136 star arms. Here, we focus on the processes that lead to relaxa-137 tion of chain orientation. For linear polymers, this includes 138 reptation, TCR, and CLF at low shear rates and CCR at 139 higher shear rates. For star polymers, we have CLF and TCR 140 at low shear rates and CCR at high shear rates. The longest 141 relaxation time for star arm orientation due to CLF, τ_{arm} , is 142 predicted to be [26]

$$\tau_{\rm arm} = \frac{\pi^3}{2} \tau_e Z_{\rm arm}^2 \exp\left[\frac{15}{24} Z_{\rm arm}\right],\tag{1}$$

where Z_{arm} is the number of entanglements along each star 144 arm. While the numerical factor inside the exponent may 145 change slightly depending on the approximations used, all 146 approaches agree that it is O(1) and this will not be further 147 discussed hereafter. 148

For star/linear blends, the relaxation mechanisms of the 149 linear chains remain unchanged by the presence of a small 150 fraction of star arms, whereas on the other hand, the stars 151 dynamics is affected by the linear matrix. The orientation of 152 their arms is still primarily relaxed by CLF and TCR, but the 153 nature of these processes is altered when star/linear entangle-154 ments are "short-lived" compared to star/star entanglements 155 for blends having span molar mass of the arms similar to that 156 of the linear chains. It is helpful for the subsequent discus- 157 sion to introduce the concept of "thin tube" and "fat tube" 158 surrounding the star arms [31,40]. The thin tube describes 159 the restriction on lateral motion of the star arm due to entan- 160 glement with both other star arms and with the linear matrix 161 molecules. The "fat tube," on the other hand, describes the 162 restriction to lateral motion that remains after all of the star- 163 linear entanglements are removed, leaving only star-star 164 entanglements. Depending on the molar masses of the linear 165 chains and the star arms, the latter may relax at least partially 166 within their fat tubes. For a given star/linear blend, we must 167 then consider whether CLF occurs in the thin tube or the fat 168 tube. The thin tube CLF time, $\tau_{\rm arm}$, is essentially unchanged 169 from the case of pure star polymers (without the linear 170 polymer background but with a number of entanglements 171 equal to the sum of the star-star and star-linear entanglements 172 of the blend). The fat tube CLF time, $\tau_{\rm arm}^{\rm Fat}$, can be estimated 173 by accounting for entanglement dilution. First, we replace 174 $Z_{\rm arm}$ by $Z_{S/S} \approx \phi_S Z_{\rm arm}$, where ϕ_S is the volume fraction of 175 stars and the value of the dilution exponent is taken to be 1 176 [7,23]. Second, we replace τ_e by τ_e^{CRR} , which is the Rouse 177 time of the thin tube that is envisioned as undergoing 178 Rouse-like fluctuations within the star/star entanglements 179

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180 (i.e., the fat tube)

$$\tau_e^{\text{CRR}} = \frac{2}{3\pi^2} \tau_{d, \text{lin}} \left[\frac{Z_{\text{arm}}}{Z_{\text{S/S}}} \right]^2$$

181 When the star polymers are not self-entangled or only 182 weakly self-entangled, $Z_{S/S} \sim 1$, and thus the fat tube CLF 183 time can be derived as $\tau_{arm}^{Fat} \approx \tau_e^{CRR}$. The star arm can be 184 approximated as no longer self-entangled, and consequently, 185 there is no separation between time scales for relaxation and 186 orientation in the fat tube.

The state-of-the-art molecular constitutive model, which 187 describes the structure and stress of entangled linear polymer 188 melts in flow, is the Graham, Likhtman, Milner, and McLeish 189 (GLaMM) model [19]. It has also been adapted for entangled 190 star polymers in a solution by Tezel et al. [25], simply by 191 switching-off the terms that describe reptation (GLaMM-R is 192 GLaMM without the inclusion of reptation). Although it might 193 seem at first sight as though the GLaMM-R approach could 194 also be applied for star-linear blends, there is an important 195 196 potential limitation for adapting GLaMM to mixed architecture blends of monodisperse polymers. The GLaMM model is a 197 description of single-component monodisperse polymer melts, 198 and it is, therefore, not possible to incorporate the thin/fat tube 199 physics that arise from constraints with different characteristic 200 rates of release and renewal. Significant progress has been made 201 202 toward this end for bidisperse linear/linear blends [12,40,41]. Such considerations would also be necessary in our system at 203 higher concentrations of the star polymer, and we should not 204 expect the GLAMM approach to work in that case, but the gen-205 eralization to mixed architecture systems remains a challenge. 206 However, we show here that if we limit the system to a range of 207 molar masses and concentrations where there are no star self-208 entanglements, the star/linear blends can be described within 209 the GLaMM-R framework, which effectively treats the arm 210 relaxation as an apparent one-component star-linear network. 211 212 modified to follow the approach of Tezel *et al.* 213

214 In particular, we address the coupling of flow and constraint release with the aim to establish an experimental pro-215 tocol for identifying component contributions and improving 216 our understanding of nonlinear stress relaxation processes 217 in mixed architecture blends. We combine the existing 218 219 GLaMM-R model with rheo-small-angle neutron scattering (SANS) in order to measure and predict the structure of star 220 polymers in a star/linear blend in the nonlinear shear flow 221 regime. We choose a star/linear blend for which the star arms 222 223 are roughly half the size of the surrounding linear chains. In this way, we ensure that both the star and linear polymers 224 share virtually the same Rouse relaxation spectrum within 225 their respective thin tubes. We also chose the blend composi-226 tion such that the star polymers are weakly self-entangled, 227 hence there is virtually no fat tube around the star arms. 228 The linear viscoelastic properties of similar systems are 229 fairly well understood [7,20], which forms a good basis for 230 investigating the nonlinear flow regime. 231

Through this systematic investigation, we wish to achieve three goals: (i) reaffirm the predictive power of tube-based models for mixed architecture blends which allow tuning constraint release effects, hence the dynamics; (ii) demonstrate 235 the power of rheo-SANS (simultaneous measurement of bulk 236 rheology and probe microstructure) as a tool uniquely suited 237 for studying nonlinear rheology in mixed architecture blends; 238 (iii) show that whereas the linear viscoelasticity of the examined star/linear polymer blend is hardly distinguishable from 240 that of the linear matrix, rheo-SANS provides distinct signatures of star and linear polymer deformation in flow. 242

II. MATERIALS AND METHODS

A. Sample preparation

The polymer blend used consisted of a fully protonated 245 polybutadiene (h-PBd) symmetric four-arm star polymer at 246 10 wt. % in a matrix of fully deuterated linear with weight 247 average molar mass $M_w = 89 \text{ kg/mol}$ and polydispersity 248 $M_w/M_n = 1.08$ The overlap concentration of the star is 249 estimated to be $c^* = 3.42$ wt. % [42]; hence, the star at about 250 3c* is semidilute albeit not entangled (the estimated entan- 251 glement concentration is about 15 wt. % [42]). This mixture 252 provides adequate contrast versus the background of deuter- 253 ated material and the star is at low enough concentration to 254 avoid phase separation due to isotope effects, which is 255 known to occur in solutions with 35-65 wt. % deuterated 256 materials [43]. The linear and star PBd polymers, with 1,4 257 rich microstructure [cis-1,4, 70%; trans-1,4, 23%; and 3, 4, 258 7% determined from nuclear magnetic resonance (NMR) 259 spectroscopy], were synthesized using high vacuum anionic 260 polymerization techniques. In the case of the four-arm 261 polybutadiene star, an excess of the living polybutadiene 262 (~30%) was reacted with 1,2-bis (dichloromethylsilyl) ethane 263 (linking agent) in order to drive the reaction to completion. 264 After completion of the reaction, the excess of the linear 265 PBd was removed by repeated fractional precipitation 266 (toluene/MeOH), until highly pure four-arm star PBd was 267 obtained, as monitored by size exclusion chromatography 268 (SEC). All intermediates and final product were analyzed 269 by SEC and NMR. The total molar mass of the final fraction- 270 ated star was $M_w = 176 \text{ kg/mol}$ with $M_w/Mn = 1.08$ as obtained 271 from SEC with PS standards and appropriate correction for 272 PBd. With this choice of molar masses, both the linear matrix 273 and the star arms are significantly above the entanglement 274 molar mass of 1,4-PBd (2.17 kg/mol, extracted from the fit of 275 the linear PBd data with the Likhtman-McLeish (LM) model, 276 see Sec. III A below). Each blend was made by first dissolving 277 the individual components into good solvent tetrahydrofuran 278 until they were fully dispersed. Then the ingredients were 279 mixed together thoroughly before driving off the solvent. The 280 latter was initially evaporated at room temperature and atmo- 281 spheric pressure before being subjected to progressively higher 282 vacuum at 30-40 °C. The samples also contained small 283 amounts (0.1 wt. %) of butylated hydroxytoluene antioxidant 284 to reduce the risk of degradation. The polymers were then kept 285 under high vacuum for 24 h before testing, in order to remove 286 any possible remaining solvent. 287

B. Rheological measurements

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Linear viscoelastic measurements were performed with an 289 ARES (TA, USA) strain-controlled rheometer. The rheometer 290 4



FIG. 1. The horizontal and vertical shift factors (a_T and b_T , respectively) as functions of the temperature. For the horizontal shift factor, the WLF fit for the pure components and the star/linear blend is also reported (solid black line; $C_1 = 3.92$ and $C_2 = 180 \pm 1$ K). For the vertical shift factor, the calculated vertical shift reflecting the temperature density compensation is also reported (dashed black line; the density was calculated according to the formula $\rho = 1.0547 - 5.6 \times 10^{-4}$ T, where T is the temperature in Kelvin) [33].

is equipped with a 2 K FRTN1 force rebalance transducer, 291 292 and temperature control $(\pm 0.1 \,^{\circ}\text{C})$ is achieved by means of a nitrogen gas convection oven and a liquid nitrogen Dewar. 293 Dynamic oscillatory shear experiments were performed from 294 -75 to 75 °C in steps of 25 °C. A single measurement was 295 also performed at -80 °C in order to detect the high-296 frequency crossover of the viscoelastic moduli. The strain 297 amplitude was in the range 2-7%, ensuring measurements in 298 the linear viscoelastic regime at all temperatures investigated. 299 It was ensured that residual stresses due to loading were fully 300 relaxed and the polymer samples were allowed to equilibrate 301 at the test temperature. 302

303 Time-temperature superposition was used in order to obtain master-curves at a reference temperature $T_0 = 25$ °C. The hori-304 zontal shift factor a_T is described empirically by the Williams-305 Landel--Ferry (WLF) [44] equation $\log(a_T) = -C_1(T - T_0)/$ 306 $(C_2 + T - T_0)$, where C_1 and C_2 are material constants that 307 depend on T₀. In our specific case, the polybutadiene was 308 found to have $C_1 = 3.9 \pm 0.1$ and $C_2 = 180 \pm 1$ K as also shown 309 in Fig. 1. The vertical shift factor was $b_T = \rho_0 T_0/(\rho T)$, where ρ 310 is the polymer density and T the absolute temperature (in 311 Kelvin). Its values at different temperatures are of the order of 312 unity. These parameters are comparable to the values reported 313 in the literature [45]. 314

315 C. Rheo-SANS measurements

The Rheo-SANS measurements were performed at the 316 Manuel Lujan, Jr. Neutron Scattering Center, Los Alamos 317 National Laboratory in Los Alamos, NM, using a specially 318 constructed Couette flow device (CLNR) on the low-q diffrac-319 tometer beam-line [46]. The scattering vector range 320 $q = 4\pi \sin(\theta/2)/\lambda_N$ covered $2 \times 10^{-2} < q < 1 \times 10^{-1} \text{ Å}^{-1}$, where 321 the scattering vector was defined by the neutron wavelength 322 λ_N and the scattering angle θ . The 8 mm diameter neutron 323 beam passes through the sample in the radial or shear-gradient direction and thus probes the polymer conformation along the 325 velocity-vorticity plane. 326

The Couette device (CLNR) was originally developed as 327 a strain-rate controlled neutron scattering device for studying 328 the behavior of highly viscous systems [46]. Its main features 329 include high torque (up to 200 Nm), high shear rate 330 (900 s⁻¹), and high temperature (over 260 $^{\circ}$ C) under a nitro- 331 gen environment. The CLNR geometry consists of two alu- 332 minum concentric cylinders: an inner stationary bob and an 333 outer rotating cup. The inner bob has a diameter of 334 53.34 mm and has a hollowed core to provide an unimpeded 335 neutron path. The bottom of the bob is machined to have a 336 1° cone to mitigate edge effects and produce a constant shear 337 across the sample. The gap is small, 0.375 mm, in order to 338 provide a stable Taylor-Couette flow (free of elastic instabili- 339 ties) well into the nonlinear flow regime. The maximum attain- 340 able Weissenberg number prior to instability, Wimax, is 341 calculated from $Wi^{\text{max}} = K_{\text{crit}}(r_1/d)^{1/2}$, where d is the gap sep- 342 aration, r_1 is the radius of the inner cylinder, and $K_{crit} = 5.92$ 343 [47]. Internally housed heating elements maintained the tem- 344 perature at 25 °C and an insulating blanket surrounded the 345 apparatus in order to reduce thermal fluctuations. After sample 346 loading, a rest time of about 10 min was allowed for the poly- 347 mers to reach the set temperature and release residual stresses 348 from loading. Details of the first CLNR prototype are reported 349 in the literature [48]. However, the system used here was rede- 350 signed to allow stable flow up to $Wi^{max} = 30$ (based on the 351 linear chain terminal time) and also to reduce the sample 352 volume required to fill the device. 353

The isotropic SANS data in the absence of shear were 354 reduced to 1D scattering intensity using a radial average. Due 355 to the geometry of the experimental setup, the scattering pro- 356 files of the blend in the presence of flow are elongated in the 357 vorticity direction, i.e., perpendicular to the flow direction. 358 The two-dimensional (2D) scattering intensities were calcu- 359 lated using software from the host institute and normalized to 360 the absolute scale via calibration with the incoherent scattering 361 and application of background corrections [46]. The degree of 362 anisotropy in these 2D scattering patterns was characterized by 363 determining a sectorial average of the scattering intensity 364 along the vorticity and flow directions with an opening angle 365 of $\pm 2.5^{\circ}$ from each axis. The anisotropy ratio was then defined 366 by taking the integrated data $2 \times 10^{-2} < q < 1 \times 10^{-1} \text{Å}^{-1}$ and 367 dividing the scattering intensity in the vorticity direction by 368 the data along the flow direction. 369

The rheo-SANS measurements were used to investigate 370 how the single chain structure factor evolves for the star/ 371 linear polymer blend under shear flow in the Couette device. 372 A direct comparison of the experimental data, obtained at dif- 373 ferent flow rates, was made with the GLaMM-R model. 374

SANS has been performed by previous investigators for 375 various polymer architectures [2,8,37]. *In situ* SANS of poly- 376 meric melts under deformation is of particular interest in 377 studying the relaxation and deformation processes. Recent 378 work has been performed via neutron scattering for a Pouseille 379 geometry, using data taken at several points along a 4:1 con- 380 traction, and subsequently, a 1:4 expansion with a recirculating 381 polymer flow [18]. In the present work, the use of a low- 382 curvature Couette geometry simplifies the data analysis in that 383

changes in the velocity gradient across the gap are small 384 enough that the flow may be approximately modeled as 385 simple shear flow in the plane of interest (in our case, it is the 386 flow/vorticity plane). The capability of the Couette shear cell 387 to operate at high torque and high shear, discussed later in 388 further detail, allows for the investigation of entangled poly-389 meric systems, which are too viscous to be studied in conven-390 tional rheometers with this geometry. The blend investigated 391 in this work (10 wt. % four-arm star polybutadiene in 90 wt. % 392 linear deuterated polybutadiene with the same span molecular 393 weight) offers a large operational time-window to study the 394 molecular dynamics of the minority star component by SANS. 395 In particular, our aim is to consider rates of deformation that 396 397 orient the star polymers but leave the linear matrix mostly relaxed. 398

399 III. RESULTS AND DISCUSSION

400 A. Linear rheology

401 Figure 2(a) depicts the frequency-dependent storage and loss moduli, G' and G'', respectively, at a reference tempera-402 ture of 25 °C as well as the $tan(\delta)$ versus shifted frequency. It 403 can be seen that the relaxation process of the pure star 404 polymer is characterized by a broad and smooth peak of G", 405 followed by a strong decay through the crossover between 406 G'' at lower frequencies associated with the fluctua-G' 407 tions of the arms. On the other hand, the linear chain matrix 408 exhibits a single loss peak (at the moduli crossover), which 409 marks its terminal regime. In the case of the polymer blend, 410 the linear chain dominates the terminal relaxation and there is 411 very little distinction between the linear viscoelastic (LVE) 412 spectra of the linear chain and the blend, with the exception 413 of the low-frequency G' data. We have confirmed that the 414 data are not an artifact associated with phase angle resolu-415 tion, hence this small deviation of the blend G' data marks 416 the effect of star-linear topological coupling which can be 417 also appreciated in the loss angle plot in Fig. 2(b) and are 418 419 further discussed in the context of the scattering data below. However, over most of the frequency range, it is evident that 420 the dynamic oscillatory measurements are not sensitive 421 enough to clearly separate, at least quantitatively, the contri-422 bution of the star polymer from the linear matrix in the 423 polymer blend. 424

The estimated values for τ_d shown in Fig. 2(a), and also 425 listed in Table I, are based upon the crossing points for the G' 426 and G" data. Also shown in Table I are the number of entan-427 glements per chain Z (per arm for the pure star). In the case of 428 the blend, we get $Z_{\text{blend}} = \phi_s \frac{1}{2} (1 - \phi_s) Z_{\text{linear}} = 45$, consider-429 ing both star and linear chain entanglements, although only a 430 few entanglements come from the star polymer. Finally, we 431 give the estimated value of the plateau modulus G_e evaluated 432 as the value of G' corresponding to the minimum of the loss 433 factor, and the estimated values of the Rouse time based 434 upon the measured values of τ_d and the expected relationship 435 $\tau_R = \tau_d/3Z$ for the linear chain, and the approximation, 436 $\tau_{R,\text{Star}} \longrightarrow 2Z_{\text{arm}} \approx \tau_{R,\text{linear}}(Z)/4$. It should be noted that 437 the data stered in Table I for the blend reflects values for the 438 439 linear chain within the blend. The relaxation for the star in the



FIG. 2. (a) Linear viscoelastic master-curves (shifted storage and loss moduli versus shifted frequency) of the linear matrix, the star polymer, and the star/linear blend at a reference temperature of 25 °C, along with fits (solid lines) using the tube-based Likhtman–McLeish (LM) model for the linear matrix and the BoB model for the star polymer, and the star/linear polymer blend (see text). The respective characteristic times at the moduli crossover are indicated by arrows. (b) Respective data are represented in the more sensitive plot of tan(δ) versus shifted frequency.

blend cannot be measured directly but can be estimated by 440 means of Eq. (2). 441

We can also obtain estimates of parameters for the linear 442 chain applying the LM mesoscopic model [14] for the entan- 443 gled linear polymer, including the characteristic relaxation 444 times. The LM model takes into account CLF, CR, and 445 longitudinal stress relaxation along the tube. The only 446

TABLE I. Linear viscoelastic parameters of the linear matrix, the star polymer, and the star/linear blend at a reference temperature of 25 °C. The LVE parameters were obtained directly from the data (see text).

	τ_d (s)	Z (-)	$\tau_R(s)$	G_{a} (MPa)
Linear	4.5×10^{-2}	41	3.7×10^{-4}	\mathcal{L}
Star	16.6	20	8.7×10^{-5}	1.65
Star/linear blend	6.5×10^{-2}	45	4.8×10^{-4}	1.65

447 parameters are M_e , G_e , τ_e , and c_v (constraint release factor). 448 Their values, obtained from a fitting procedure, are 449 $\tau_e = 3.9 \times 10^{-7}$ s, $M_e = 2.17$ kg/mol (from the high-frequency 450 moduli crossover), and $G_e = 1.65$ MPa. This value of M_e 451 corresponds to Z = 41 for the linear matrix and $Z_s = 80$ for the 452 entire four-arm star. The model predictions are shown as the 453 solid lines in Fig. 2(a).

Similarly, the generalized tube-based branch-on-branch 454 455 model (BoB) has been used for analyzing the dynamics of the entangled star polymer and the star/linear blend [15,16] 456 (to this end, the open source program RECTORE has been 457 employed). A dilution exponent of 1 has been used for the 458 star polymers and the blend. For the BoB fit of the star/linear 459 460 blend, we have considered the molecular characteristics of the components presented in Sec. II A above and the same param-461 eters from the LM fit ($\tau_e = 3.9 \times 10^{-7}$ s, $M_e = 2.17$ kg/mol, and 462 $G_e = 1.65$ MPa), with $Z_{\text{blend}} = 45$ for the blend, considering 463 both star and linear chain entanglements. The density value 464 used in BoB is 1450 kg/m³, making the above G_e and M_e 465 values consistent through the rubber elasticity relationship. 466 467 The BoB fitting results are also shown in Fig. 2(a). It is evident that the BoB model is able to capture the small low-468 frequency deviation of G' due to the relaxation of the star 469 polymer in the linear matrix. The loss factor plot of Fig. 2(b) 470 is suggestive of a decoupled two-mode relaxation (linear and 471 star) of the blend; however, the BoB model in the present 472 form may not account entirely correctly for the dilution effect 473 of the linear chain's CLF on the star. In view of this as well as 474 the lack of experimental data at lower frequencies, definite 475 conclusions about the role of the star cannot be drawn. 476

Note that the values of the parameters obtained from the 477 LM and BoB model fits are slightly higher than those esti-478 mated directly from the experimental data. The latter values 479 are consistent with the literature, where the respective reported 480 values are also slightly lower than those obtained from the 481 fitting with the LM model [49,50]. The value of τ_e is consis-482 tent with that used by Kapnistos et al. [33] and the monomeric 483 friction coefficient of PBd. We shall not explore this further 484 485 and consistently apply the GLaMM-R model for predicting both rheology and structure of the stars with the same values. 486

The above analysis of the LVE spectra shows that discriminating between the pure linear matrix and the star/linear blend in the present situation with similar Z_{linear} and Z_{blend} is a subtle issue, hence, complementary evidence from scattering is highly desirable.

Since a direct measurement of the terminal relaxation time 492 of the star component within the blend is not possible given 493 the lack of a distinct feature in the LVE spectrum, we 494 obtained estimates of τ_{arm}^{Fat} (obtained through τ_e^{CRR}). By using 495 $Z_{S/S} = 2$ for our system, we can use Eq. (2) to estimate that $\tau_e^{\text{CRR}} \approx 0.3 \text{ s.}$ It thus follows that $\tau_{arm}^{\text{Fat}} \approx \tau_e^{\text{CRR}} (Z_{S,S})^2 = 1.2 \text{ s.}$ 496 497 We recognize the approximate nature of this estimate. 498 Nevertheless, it does suggest that we have approximately one 499 order of magnitude separating the characteristic terminal 500 relaxation time of the star (τ_{arm}^{Fat}) and linear $(\tau_{d,lin})$ polymers, 501 and this provides a reasonable window in terms of shear 502 rate for interrogating the star component in the star/linear 503 polymer blend. In fact, this means that the star arms will 504 505 fully relax only after the linear matrix has fully relaxed.

A clear message from the linear rheological experiments 506 is that the linear polymer matrix dominates the star/linear 507 polymer blend, hence, the star signature cannot be easily discerned. The rheo-SANS experiment is the most viable alternative tool to uniquely extract the information on the 510 conformational changes and dynamics of the minority component under shear. 512

513

B. Rheo-SANS results

In the rheo-SANS measurements, the deformation process 514 is controlled by the applied shear rates, i.e., $\dot{\gamma} = 0$ (equilib- 515 rium), 8.1, 16.1, and 24.2 s⁻¹ at 25 °C and probed through the 516 scattering patterns. We can use the crude estimate of $\tau_{\rm arm}^{\rm Fat}$ to 517 convert these shear rates to Weissenberg numbers, $Wi \approx 9.86$, 518 19.6, and 29.45, respectively. It should be noted, however, 519 that these values do not appear in the GLaMM calculations 520 and thus they are given here mainly to provide a qualitative 521 frame of reference for estimating the strength of the flow. The 522 use of $\tau_{\rm arm}^{\rm Fat}$ is motivated by the fact that our aim is to explore 523 the dynamics of the star polymer in the blend (see also discus- 524 sion on LVE above). Despite this approximate approach, we 525 note that the Weissenberg number is actually not necessary in 526 our analysis. What is important to appreciate is that the 527 applied rates are lower than the inverse relaxation time of the 528 linear component and larger than the estimated inverse relaxa- 529 tion component of the star component in the blend (see Fig. 2 530 and Table I) Although we do not perform any direct structural 531 measurement of the linear matrix chains, we expect that they 532 remain virtually unstretched in this range of shear rates, which 533 correspond to Weissenberg numbers based on the terminal 534 time of the pure linear polymer (τ_d in Table I) of 0.36, 0.72 535 and 1.09, respectively. 536

The scattering data were collected in the q-range 537 2×10^{-2} -1 $\times 10^{-1}$ Å⁻¹ for 20 min. This chosen time allowed 538 good statistics and at the same time ensured a stable thermal 539 environment. In fact, the two principle limitations of the 540 Couette device are the difficulty to perform experiments 541 below room temperature (six heater cartridges give a good 542 thermal control from room temperature up to 240 °C with 543 small thermal fluctuations ±2 °C) and the difficulty to run 544 experiments under these conditions at steady state for times 545 longer than 45 min. 546

Therefore, the rheo-SANS procedure to scan and enlarge 547 the investigated q-range [3] was not applicable here. An 548 optimum compromise between short time scales (less than 549 45 min) and good data statistics (times long enough to have 550 high resolution) was used, and some very interesting structural features were still detectable in the chosen q-range. Due 552 to the experimental conditions, the lowest scattering q-values 553 are missing and thus the chain configuration could not be 554 determined with high precision. Instead, we focused our 555 attention on length scales where chain orientation is accurately determined, i.e., the star arm length scale. 557

The total intensity scattering in the perpendicular and parsize allel directions is plotted versus q in Fig. 3. The scattering size data have been scaled with the reference length of the tube so diameter, which is $a \sim 44$ Å (q ~ 0.023 Å⁻¹) as reported in so the literature [50]. As can be observed from Fig. 3, the so



FIG. 3. Experimental 1D perpendicular and parallel components (symbols) versus GLAMM-R calculations (lines).

563 scattering intensity is in good agreement with the predicted 564 values from the GLAMM-R/SANS theory at mid-to-high 565 q-values. However, it fails at low q-values where probably 566 some high-order structuring (nonhomogeneity) as well as the 567 curvature of the Couette cell may contribute.

In the original application of the GLAMM-R model for 568 stars in solution, five parameters were needed: three are well-569 tabulated material properties and two are universal characteris-570 tic constants for stress relaxation processes. They are G_e , τ_e , 571 572 and M_e as well as the dimensionless parameters for retraction, c_{ν} , and the constraint release term, R_s . We also need the 573 applied shear rate, $\dot{\gamma}$. The first three parameters are obtained 574 from LVE. The other two parameters, c_v and R_s , are universal 575 constants associated with constraint release (CCR and TCR) 576 and chain retraction [18,19]. Recommended values for these 577 constants can be found in the literature, $c_v = 0.1$, $R_s = 1$ and 578 were selected based on an extensive comparison with experi-579 mental data [18,19,25]. Finally, the shear rate can be deter-580 581 mined directly from the experimental protocol. In our predictions with the GLaMM-R model, we have used Z = 20582 for the probe chain. We recall that the probe chain (star 583 polymer) is surrounded by and entangled with the matrix 584 (linear polymer) in a semidilute regime. Although the neutron 585 scattering investigation is focused on the star polymer behavior, 586 the contribution of the linear matrix is yet present and implicit. 587

In our previous works [2,3], with data over a larger 588 q-region (i.e., lower q-values corresponding to the radius of 589 synation, R_{e}), we used the ratio of the scattering intensity 590 along the perpendicular (vorticity) and parallel (velocity) 591 directions, $S(q)\perp/S(q)\parallel$ as a simple metric of microstructural 592 anisotropy at varying q. With this anisotropy metric, we con-593 sistently find a q for which star arm anisotropy is maximized. 594 Again, a comparison with the modified GLaMM-R model is 595 possible in the q-range where the orientation of the probe 596 chain is observed. Indeed, in Fig. 4, the ratio $S(q)\perp/S(q)\parallel$ at 597 different Weissenberg numbers distinguishes between 598 599 deformed and undeformed configurations. This provides a measure of the degree of deformation (in terms of chain ori-600 601 entation) for a range of length scales $2\pi/q$ along the star arm.



FIG. 4. Plot of the $S(q)\perp/S(q)\parallel$ (perpendicular-parallel ratio) with SANS data corresponding to Weissenberg numbers of 0, 9.86, 19.6, and 29.45. Lines represent the modified GLaMM-R model predictions (for the isotropic case the line is horizontal through the data). The data used to generate this figure is the same as the data shown in Fig. 3. The maximum experimental uncertainty is 10%.

The isotropic data ($\dot{\gamma} = 0$) shows a very small, nearly neg- 602 ligible upturn at the lowest q-values; this might be due to 603 some parasitic forward scattering for $q < 1/R_g$ or residual 604 voids (air) after the loading process. We note, however, that 605 the SANS experiments were performed after a rest time 606 longer than $10\tau_{d,linear}$, as discussed above. 607

To better appreciate the response of the star in the studied 608 star-linear system (10 wt. % star with Z=20 and 90 wt. % 609 linear), we compare the calculated $S(q)\perp/S(q)\parallel$ ratio from 610 GLAMM-R with the respective scattering response of a 611 linear polymer mixture (10 wt. % linear with Z=20 and 612 90 wt. % linear) based on GLAMM. The results are shown in 613 Fig. 5, where we have selected arbitrary values of the shear 614



FIG. 5. GLAMM-based calculations of $S(q)\perp/S(q)\parallel$ for star (solid lines) and linear (dashed-dotted lines) polymers with Z = 20, using GLAMM-R (10 wt. % star with Z = 20 and 90 wt. % linear) and GLAMM (10 wt. % linear with Z = 20 and 90 wt. % linear) models, respectively.

rat (12), 270, 402, and 540 s⁻¹ (at lower rates the anisotropy 615 of the linear chain would overlay on the isotropic profile). 616 617 Due to the branching nature of the star polymer (and the omission of reptation in GLAMM-R), the anisotropy of the 618 star is more than twice larger (note the higher peak and 619 broader shape of the scattering curves representing the q-620 dependent $S(q)\perp/S(q)\parallel$ ratio). It is clear from these results 621 that while the star-modified GLAMM model is not fully 622 capable of capturing the anisotropy of the scattering data in 623 Fig. 4, it does much better than a linear chain prediction. A 624 further comparison between the GLaMM-R modeling and 625 experimental data is shown in Fig. 6, where 2D scattering 626 contours of the scattering intensities are reported. In the 627 investigated system, the applied flow causes the probed star 628 polymer arm to orient in the flow direction. As mentioned 629 above, at the shear rates considered here, the linear matrix is 630 almost completely relaxed and thus the anisotropic 2D scat-631 tering patterns show that the star polymer is weakly oriented 632 along the velocity direction. The measured profiles in the 633 2D-detector plane are in good qualitative agreement with the 634 calculated results obtained via the GLaMM-R model. 635

The isointensity contours of 2D anisotropic patterns show 636 a transition from circular (isotropic) to a more ellipsoidal 637 (anisotropic) shape with the increasing rate of deformation 638 (see Fig. 6). 639

The details of the full scattering pattern provide a more 640 complete picture of chain orientation and deformation than 641 the simple ratio $S(q)\perp/S(q)\parallel$ presented in Fig. 4. At zero 642 shear rate, we obtain the well-defined isotropic pattern, 643 possessing a q-invariant value of the scattering ratio 644



FIG. 6. Experimental (black lines) and modeled (red lines) 2D scattering intensities in the vorticity-velocity plane of the star/linear polymer blend corresponding to applied shear rates of 0 (top left), 8.1 s⁻¹ (top right), 16.1 s⁻¹(bottom left), and 24.2 s⁻¹ (bottom right). The calculations were performed in the $2 \times 10^{-2} < q < 1 \times 10^{-1} \text{ Å}^{-1}$ range. The 2D q \perp and q \parallel ranges reported are between -1×10^{-1} and 1×10^{-1} . The deformation direction is horizontal.

 $S(q)\perp/S(q)\parallel = 1$ and a perfectly circular 2D profile, as 645 reported in Figs. 4 and 6, respectively. At an applied shear 646 rate of 8.1 s^{-1} (*Wi* = 9.86), a weak anisotropy is observed, 647 highlighted by a smooth leveling-off from the isotropic 648 profile (see Fig. 4). In this region, GLaMM-R provides 649 a good description of the data accounting for the previously 650 mentioned assumption. Further, at 16.1 s^{-1} (*Wi* = 19.6), we 651 still observe an increased anisotropic behavior and GLaMM-R 652 predicts fairly well the scattering pattern. Finally, at the 653 highest applied rate of 24.2 s^{-1} (*Wi* = 29.45), the trend is 654 still qualitatively consistent with the previous shear rates in 655 the scattering event and the anisotropy increases even more, 656 but the GLAMM-R model now seems to underestimate the 657 anisotropy as also seen earlier in Fig. 4. The reason is that 658 the complex star-linear polymer coupling is not considered 659 in the current formulation of one-component system with a 660 simple CCR term. Higher anisotropy with strong appear- 661 ance of lozenge-shaped scattering patterns (which can be 662 barely evidenced at the low-q regime of the highest rate in 663 Fig. 6) are expected to appear over a wide q-range at much 664 higher shear rates exceeding 100 s^{-1} (not accessible in the 665 present experiment). 666

IV. CONCLUSIONS

With a combined experimental (rheo-SANS) and modeling 668 (GLaMM-R) approach, we probed the structural change of 669 stars immersed in a linear polymer matrix that had the same 670 molar mass as the sum of two arms and was being sheared at 671 moderate shear rates (8.1 $< \dot{\gamma} < 24.2 \text{ s}^{-1}$) between the inverse 672 relaxation times of the linear and star components of the blend. 673 The respective range of approximate Weissenberg numbers 674 was 9.86 < Wi < 29.45, based on the estimated value of τ_{arm}^{Fat} . 675 While the star contribution is barely perceptible in the linear 676 viscoelastic spectrum at low frequencies, their structural anisot- 677 ropy when weakly sheared is clearly evident in the SANS data, 678 albeit the effect is small. The orientation of the star polymer in 679 the flow is clearly observed in 2D anisotropic patterns. At this 680 time scale, the star-star entanglements are effectively absent, 681 and thus can be neglected in the treatment of the molecular 682 mechanism of the star relaxation. The neutron scattering 683 experiments allowed precise discrimination of the probe star 684 microscopic behavior in the transition from linear to nonlin- 685 ear regime. 686

Finally, this work shows that the combination of rheology 687 and neutron scattering is uniquely capable for selectively 688 probing the nonlinear configurations of branched polymers in 689 a bimodal polymer mixture. This protocol is particularly 690 useful for architectural blends and can be extensively applied 691 to other modes of deformation (e.g., extensional) in order to 692 obtain the material's full response to deformation. A more 693 detailed theoretical approach may still be needed for fully 694 interpreting the linear chain contribution in these bimodal 695 polymer melts. Nevertheless, the present set of approxima- 696 tions works well for moderate deformation rates but would 697 not be expected to describe polymer-melt behavior subjected 698 to stronger strain fields. Here, we have provided a platform 699 that can be expanded upon in future works to develop more 700 robust models. 701

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702 ACKNOWLEDGEMENTS

Professor Evelyne van Ruymbeke is thanked for helpful discussions. This project was primarily supported by the IMMS/LANL program at UCSB. The authors also acknowldege partial support by the European Commission (ITN "Supolen," No. FP7-607937, and Horizon 2020-INFRAIA-2016-1, EUSMI Project No. 7310219).

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