Elastic Phases of $Ge_xSb_xSe_{100-2x}$ Ternary Glasses Driven by Topology

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Supporting Information

ABSTRACT: Topology offers a practical set of computational tools to accurately predict certain physical and chemical properties of materials including transformations under deformation. In network glasses with increased cross-linking three generic elastic phases are observed. We examine ternary $Ge_xSb_xSe_{100-2x}$ glasses in Raman scattering, modulated DSC and volumetric measurements, and observe the rigidity transition, $x = x_c(1) = 14.9\%$ that separates the flexible phase from the Intermediate phase, and the stress transition, $x = x_c(2) = 17.5\%$ that separate the intermediate phase from the stressed rigid one. Raman scattering provides evidence of the structural motifs populated in these networks. Using size increasing cluster agglomeration, we have calculated the rigidity and stress transitions to occur near $x_c(1)^t = 15.2\%$ and $x_c(2)^t = 17.5\%$, respectively. Theory predicts and experiments confirm that these two transitions will coalesce if edge—sharing Ge-tetrahedral motifs were absent in the structure, a circumstance that prevails in the Ge-deficient $Ge_7Sb_xSe_{93-x}$ ternary, underscoring the central role played by topology in network glasses. We have constructed a global elastic phase diagram of the Ge–Sb–Se ternary that



provides a roadmap to network functionality. In this diagram, regions labeled A, B, and C comprise networks that are flexible, rigid but unstressed, and stressed-rigid, respectively.

1. INTRODUCTION

In soft matter, applications of topology include foams,^{1,2} quasicrystals,³ nonuniform media,⁴ and glasses.^{5–7} Concerning the latter, a topological approach to rigidity theory has provided new insights⁸ into understanding glassy covalent networks. It has led to the recognition of elastic phase transitions⁹ separating flexible glassy networks having internal degrees of freedom that allow for local deformations, from stressed rigid ones which are "locked" by their high connectivity.

In a global approach to rigidity one has found it useful to analyze physical properties of multicomponent systems in terms of their network connectivity, measured in terms of mean coordination number "r".⁹ Characteristic structural motifs composed of tetrahedra (edge and/or corner-sharing), pyramids, and quasi-tetrahedra play an important role in understanding network properties including their elastic phases. The use of the term phase within Rigidity theory^{8,10} is to denote changes in physical properties of network glasses with composition. Recently, it was shown¹¹ that the increase of fragility in tetrahedral glass-forming liquids could be quantitatively understood from an increase in edge-sharing (ES) fraction of group IV (Si, Ge) tetrahedra in binary selenide glasses. Experiments¹² supported by theory^{8,13} have also shown the existence of two elastic phase transitions in network glasses, a rigidity transition followed by a stress transition with increasing r. Glass compositions in between these transitions, the intermediate phase (IP), have evoked special interest both at a basic level and in applications because of their most unusual

physical properties including absence of network stress,¹⁴ nonaging,¹⁵ and space filling property.¹²

In the above respect, the case of group V sulfides and selenides has received less attention than the group IV ones, even though IPs in the P–Se, 16,17 P–S, 18 and As–Se 19 and As– S^{20} binaries are established. The case of the Sb_xSe_{100-x} binary would appear to be of marginal interest given that homogeneous glass formation is restricted²¹ to x < 2 at % of Sb with the larger size and more metallic pnictide. However, experiments reveal⁶ that by alloying a few mol % of Ge, glass formation can be extended over an increased range of Sb in the $Ge_7Sb_rSe_{93-r}$ ternary. More spectacular is the fact that in the $Ge_xSb_xSe_{100-2x}$ ternary, containing equimolar concentrations of Ge and Sb, glass formation extends over a much wider range (0 $\langle x \langle 22\% \rangle$ of Sb or a mean coordination number in the 2.00 \langle r < 2.62 range.²² We take Ge, Sb, and Se to be, respectively, 4-, 3- and 2- fold coordinated, yielding r = 2 + 3x. We have examined the Ge_xSb_xSe_{100-2x} ternary in Raman scattering and modulated DSC experiments, and in this paper show that the rigidity and stress transitions appear respectively, near $r_{c}(1) =$ 2.42(1) and $r_c(2) = 2.52(1)$. Using size increasing cluster approximation⁸ (SICA) and the four structural motifs observed in Raman studies, we attempt to predict these transitions. SICA in its simplest form, i.e., in the first level of approximation,

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already reproduces the salient features observed in experiments. These findings when compared to those reported earlier on corresponding P- and As- bearing ternaries^{15,23} highlight the role of topology at a local level, coordination numbers of structural motifs, and on extended scale, fraction of ES/CS units, to determine these elastic phase transitions. Taken together, these results on $\text{Ge}_x \text{Pn}_x \text{Se}_{100-2x}$ ternaries, with the Pn = P, As, and Sb, provide a unified view of elastic phase transitions observed in these ternary covalent selenides.

2. EXPERIMENTAL SECTION

22%, were synthesized starting from 99.999% Ge, Sb and Se pieces (3 mm diam.) from Cerac Inc. The starting materials, sealed in evacuated (10^{-7}Torr) 5 mm ID quartz tubings, were alloyed at 950 °C for periods ranging from 5 to 10 days, and quenched from 50 $^{\circ}C$ above the liquidus in cold water to realize bulk samples. FT-Raman spectra, taken along the length of a quartz tube encapsulating a melt-quenched glass, were taken over the 5-10 day alloying period, until all line-shapes coalesced into a single one (see Figure S1of the Supporting Information, SI), at which point sample compositions were considered homogeneous. This novel technique was recently introduced^{7,24-26} to ascertain batch homogeneity, and has been remarkably powerful in synthesis of homogeneous chalcogenides melts/glasses. Mass density of the glasses was established using Archimedes' principle. A quartz fiber attached to a digital balance was used to weigh samples in air and in pure ethyl alcohol. A single crystal of Si was used to calibrate the density of alcohol, and a single crystal of Ge used to check accuracy of measurements. Modulated DSC experiments used a model 2920 instrument from TA Instruments, and scans were taken at 3 °C/min ramp rate with a sinusoidal modulation of 1 °C temperature modulation, and 100 s modulation time. After scanning up in temperature to $T_{\rm g}$ + 25 °C, a cool-down cycle was programmed, and the frequency correction to the nonreversing enthalpy made in the usual way.^{27–29} Compositional trends in $T_g(x)$, and nonreversing enthalpy, $\Delta H_{nr}(x)$, were measured on T_{g} -cycled samples both in the fresh (rejuvenated) and aged state. Aging was performed at room temperature $T < T_g$ with samples sealed in hermetically sealed Al pans. Figure 1 provides a summary of compositional trends in $T_g(x)$, $\Delta H_{nr}(x)$, and molar volumes $V_m(x)$. We note that $T_g(x)$ increases monotonically in the 0 < x < 18.2% range, but decreases thereafter, which is signature of demixing of of homopolar bonds (Ge-Ge, Sb-Sb) formed above the chemical threshold (Figure 1, inset). The $\Delta H_{nr}(x)$ term shows a deep and sharp reversibility window in the 14.2% < x < 17.5% range, and the term is found to age outside the window but not in it over months. In the same composition range, molar volumes show a local minimum. T_g 's and molar volumes of the present samples are found to be measurably higher than earlier reports^{22,31} (Figure 1), probably because of their dryness and homogeneity.

FT-Raman scattering measurements made use of Thermo-Nicolet Model Nexus 870 unit with a laser spot size of 250 μ m to profile melts. Raman spectra at select compositions are shown in Figure 2. Observed lineshapes were deconvoluted to superposition of appropriate number of Gaussians to extract variations in mode scattering strength and frequency with *x*. In the Se-rich regime (x < 18.2%), we observe several modes, which were noted earlier in binary Ge_xSe_{100-x} glasses^{7,24,25} that result from Ge centered cross-linking structures (see below)



Figure 1. Filled circle data points show trends in (a) $T_g(x)$, (b) $\Delta H_{\rm nr}(x)$, and (c) $V_{\rm m}(x)$ in the present ${\rm Ge}_x {\rm Sb}_x {\rm Se}_{100-2x}$ ternary. In (a) we compare the present results (\bullet) with those on the ${\rm Ge}_x {\rm Sb}_x {\rm Se}_{100-2x}$ ternary by Mahadevan et al ($\mathbf{\nabla}$)²² and the ${\rm Ge}_x {\rm As}_x {\rm Se}_{100-2x}$ ternary (\bigcirc) by T. Qu et al.²³ The inset in (a) shows the fraction of Ge–Ge/Ge–Se bonds derived from a stochastic agglomeration theory (SAT) based analysis of $T_g(x)$ variation.³² Note that $T_g(x)$ in the present ternary shows a kink near x = 18.2%, which identifies the chemical threshold. Such a kink is not observed in ${\rm Ge}_x {\rm As}_x {\rm Se}_{100-2x}$ glasses. The smooth line is a prediction of $T_g(x)$ based on SAT for the present ternary. In (b) we show the presence of aging of the $\Delta H_{\rm nr}(x)$ term outside the reversibility window but not in it. In (c) we compare the present molar volumes to those reported by Mahadevan et al.²²

formed with polymeric Se-chains. In addition, we observe other modes that are due to pyramidal SbSe₃ units. In the Se-deficient regime (x > 18.2 at %), new modes appear (see the SI) that can be assigned to homopolar bonds, Ge–Ge and Sb–Sb, the former as part of ethanelike units (Ge₂Se₃) and the latter as part of ethylenelike units (Sb₂Se₂(Se_{1/2})₄. We are guided in Raman mode assignments by first principles cluster calculations^{33,34} and by observed variations in mode scattering strengths with x (see the SI).

The systematic blue-shift of CS mode frequency (ν) with increasing *x* in the Se-rich regime confirms the blue-shift to be the result of increased network connectivity. From these data,



Figure 2. Observed Raman spectra of glass samples at indicated compositions in $\text{Ge}_x\text{Sb}_x\text{Se}_{100-2x}$ ternary. Raman line shape at x = 14.2%, is deconvoluted into five Gaussian components, and these are identified with modes of CS GeSe₄ tetrahedra (195 cm⁻¹), ES tetrahedral (219 cm⁻¹), Pyramidal Sb(Se_{1/2})₃ units (215 cm⁻¹), Se_n-chain mode (260 cm⁻¹), PYR umbrella mode of Sb(Se_{1/2})₃ (115 cm⁻¹), and an E mode of Se_n chains (135 cm⁻¹) New modes identified with Sb–Sb and Ge–Ge bonds are observed in the Se-deficient phase (x > 18.2%), and are commented upon in the SI.

we deduce an elastic power-law of p = 1.20(5) in the IP for the corner-sharing mode frequency (ν) increase with x using,

$$v^2 - v_c^2 = A(x - x_c)^p \tag{1}$$

Here ν_c represents the mode frequency at the rigidity threshold x_c and find the optical elastic power-law, p, to be in reasonable accord with earlier results.^{15,23}

3. DISCUSSION

3.1. Topology and Elastic Phases. The observation of the sharply defined reversibility window (Figure 1b), permits fixing the *rigidity* and *stress* transitions near $x_c(1) = 14.9\%$ and $x_c(2) = 17.5\%$, from the walls of the window. The feature of a vanishing nonreversing enthalpy at T_g in the IP phase has been noted earlier in many experiments,¹² and recently modeled in theory.²⁸ Thus, one views Se-rich glasses at x < 14.9% to be in the flexible phase, and Se-deficient ones at x > 17.5% to be in the stressed-rigid phase, while those in the intervening region to be in the IP. The local minimum of molar volumes (Figure 1c) in the IP is consistent with the compacted or space filling nature³⁵ of networks formed in this phase.

A salient feature of the present results on the $Ge_xSb_xSe_{100-2x}$ ternary is the onset of rigidity near $r_c(1) = 2.45$, significantly upshifted in r (Figure 3) in relation to the cases of $Ge_xP_xSe_{100-2x}$ ($r_c(1) = 2.28$) and $Ge_xAs_xSe_{100-2x}$ ($r_c(1) =$ 2.28) ternaries. There is strong evidence that P¹⁵ and As²³ cations occur in pyramidal (PYR) (r = 2.40) as well as in quasitetrahedral (QT) structural motifs (r = 2.28) in respective ternary selenides, but Sb takes only the PYR configuration in the present ternary as no specific Raman signature for the QT species has been found for the present investigated system. The



Figure 3. Intermediate phases observed in titled binary and ternary Selenides. The vertical lines drawn at r = 2.28 and 2.40 are for reference. Results on the titled glass systems are taken from (a) Selvanathan et al.,⁴² (b) Bhosle et al.,⁷ (c) Georgiev et al.,⁴³ (d) Georgiev et al.,¹⁶ (e) S. Chakravarty et al.,¹⁵ (f) T. Qu et al.,²³ (g) present work, and (h) Madhu et al.⁶

onset of the rigidity transition in the P- and As-based ternaries at lower r in relation to the Sb-based ternary is traced to the presence of QT motifs^{23,36} that possess a low mean coordination number of r = 2.28. Onset of rigidity in these ternary glasses is determined by ring-like configurations⁸ of the most Se-rich isostatic structural motifs formed. Network stress is a nonlocal property and much of the challenge to understanding IPs comes from the multiply connected nature of structure determined by network topology. The IP library (Figure 3), along with the structural motifs present in the backbone provides important bounds on realistic topological structural models of these phases. The excitement and challenge of IPs is that such networks adapt,³⁷ reconnect to lower their free energy, i.e., self-organize during melt quench.

The stressed-rigid phase forms at x > 17.5%, and glassy networks in this phase are largely demixed (see below), a feature also found³⁸ in the 225 phase change material (Ge₂Sb₂Te₅ or x = 22.2%) in corresponding Ge_xSb_xTe_{100-2x} Tellurides as well as other systems.³² The nature of chemical bonding in the present ternary, as suggested by the observed elastic phases is covalent as atoms conform to the 8-N bonding rule. This is an important result in its own right given that in corresponding amorphous-Tellurides, the bonding rule is intrinsically broken³⁹ as Pauling resonant bonding effects prevail as found in their crystalline counterparts.⁴⁰ Knowledge of the elastic phases in the Ge_xSb_xTe_{100-2x} ternary tellurides⁴⁰ will undoubtedly help better understand their functionality as phase-change materials.

3.2. Glass Structure and Onset of a Chemical Threshold. Our Raman scattering results suggest that Se-rich glasses (x < 18.2%) largely consist of polymeric Se_n chains that are progressively cross-linked by 4-fold Ge and 3-fold Sb, thus locally satisfying their chemical bonding valence requirements. The local structures formed at the cross-links consist of CS-and ES- GeSe₄ tetrahedra and pyramidal SbSe₃. Stoichiometrically, one can write,

$$Ge_x Sb_x Se_{100-2x} = Ge_x Se_{2x} + Sb_x Se_{3/2x} + Se_{100-2x-2x-3/2x}$$
(2)

Table 1. SICA Structures with Their General Probabilities (Not Normalized), Their Estimate in Case of Self-Organization, And the Associated Number of Constraints n_c

Cluster Structures	Probability (not	Probabilities (not normalized) using equs.	n _c
	normalized)	(ba, ob) in the self-organized case	
	$4P_S^2 e_{flex}$	$\frac{\frac{8(2-11x)(72xe_{ring}+4-31x)}{3(2-3x)^2}}{\text{if } x < x_c}$	2.0
		0 if x>x _c	
- James	16P _S P _G e _{iso}	$\frac{64(2-11x)x}{(2-3x)^2}$	3.0
-	$12P_SP_{Sb}e_{flex}$	$\frac{\frac{32x(72xe_{ring}+4-31x)}{(2-3x)^2}}{\text{if } x < x_c}$	2.56
<i>×</i>		0 if x>x _c	
	16P _G ² e _{stress}	0 if x <xc< td=""><td>3.67</td></xc<>	3.67
		$\frac{16x(31x-4-72xe_{ring})}{(2-3x)^2} \qquad \text{if } x > x_c$	
	$72P_G^2 e_{ring}$	$1152x^2e_{ring}$	3.25
		$(2-3x)^2$	
	24P _G P _{Sb} e _{stress}	0 if x <xc< td=""><td>3 36</td></xc<>	3 36
		$\frac{\frac{24x(31x-4-72xe_{ring})}{(2-3x)^2}}{\text{if } x > x_c}$	5.50
	$9P_{Sb}^2 e_{iso}$	144 <i>x</i> ²	3.0
Å 👎		$\overline{(2-3x)^2}$	

with the last term on the right-hand side of eq 2 representing Se atoms left over in a polymeric chain configuration. The chemical threshold (ct) would then occur when the last term vanishes, i.e., $11x_{ct}/2 = 100$, or $x_{ct} = 18.2\%$. As $x > x_{ct}$ homopolar (Ge-Ge, Sb-Sb) bonds can be expected to appear in the network. Thus, glass compositions at x < 18.2% are viewed as Se-rich, while those at x > 18.2% as Se-deficient. In the Se-deficient glasses, a pertinent issue is- do the heteropolar bonds form part of the network backbone or do they segregate as Ge-rich and Sb-rich clusters? The $T_g(x)$ variation (Figure 1a filled circles) in the present ternary reveals a kink near $x = x_{ctr}$ an observation that constitutes signature of homopolar bonds demixing from the backbone.³⁰ If these homopolar bonds were formed as part of the backbone, then one would expect $T_{\sigma}(x)$ to monotonically increase past x_{ct} as predicted by Stochastic Agglomeration Theory⁴¹ (Figure 1a smooth line), and as observed in the $Ge_x As_x Se_{100-x}$ ternary.³⁰ Note that the covalent radii of Ge, As and Se are nearly the same in the $Ge_xAs_xSe_{100-x}$ ternary. The demixing of the $Ge_xSb_xSe_{100-2x}$ ternary at $x > x_{ct}$ is thought to be driven by atomic size mismatch (covalent radii, $r_{\rm cov}$ (Sb) = 1.40 Å, $r_{\rm cov}$ (As)= 1.20 Å, $r_{\rm cov}$ (Ge) = 1.22 Å, and $r_{\rm cov}$ (Se) = 1.17 Å).

In Raman scattering existence of Sb–Sb and Ge–Ge bonds is observed in Se-deficient glass compositions ($x > x_{ct}$). The growth in Raman activity in the 160–190 cm⁻¹ range, starts

already at x = 18.2% (compare spectrum at x = 17.0% with the one at x = 18.2% in Figure 2). Furthermore, there is growth in scattering in the high frequency (240 cm⁻¹ to 330 cm⁻¹) range as can be seen by comparing the spectrum of a glass at x = 22%with the one at x = 18.2% (Figure 2). We identify these systematic changes in the observed lineshapes with composition as resulting from growth of Ge-Ge bonds present in an ethanelike cluster and Sb-Sb bonds in an ethylenelike cluster in these glasses. There are three characteristic vibrational modes $(187 \text{ cm}^{-1}, 216 \text{ cm}^{-1}, \text{ and } 234 \text{ cm}^{-1})$ of ethylenelike clusters and two $(179 \text{ cm}^{-1} \text{ and } 288 \text{ cm}^{-1})$ of ethanelike clusters that are predicted.^{33,34} from first principles calculations, and also observed in the spectra. We discuss these findings including Raman spectral deconvolution results in the SI. Another feature of the Raman results is that vibrational mode frequency of these clusters steadily red-shift as mode scattering strength increases at $x > x_{ct}$. The behavior apparently stems from cluster geometries changing as they grow in size and decouple from the backbone.

3.3. Size Increasing Cluster Approximation. SICA combined with rigidity theory permits predicting the elastic threshold for the rigidity and stress transition.^{8,44} We build on the successful modeling of Group IV selenides and derive in a similar fashion clusters for the present ternary glass $Ge_xSb_xSe_{100-2x}$ which can be rewritten as (Ge-

 $Se_{4/2})_x(SbSe_{3/2})_xSe_{100-11x/2}$. In these calculations, one agglomerates the basic structural motifs observed (GeSe_{4/2} tetrahedra, SbSe_{3/2} pyramids, and Se₂-chains) whose probabilities are given by stoichiometry, i.e.:

$$P_{\text{Se}_2} = \frac{2 - 11x}{2 - 3x} = 1 - P_{\text{G}} - P_{\text{Sb}} = P_{\text{S}}$$
(3a)

$$P_{\text{GeSe}_{4/2}} = P_{\text{SbSe}_{3/2}} = \frac{4x}{2 - 3x} = P_{\text{G}} = P_{\text{Sb}}$$
 (3b)

for compositions $x < x_{cv}$ i.e., compositions for which homopolar Sb–Sb or Ge–Ge can be safely neglected.

From these initial probabilities P_{i} , one computes the probability of finding larger clusters.

We obtain a series of 7 different clusters (including ES $GeSe_{4/2}$ connections) represented in Table 1 with their corresponding probabilities and constraint count (see below). The probabilities of such clusters are given by the following:

$$P_{ij} = \frac{W_{ij}}{Z} P_i P_j e^{-E_{ij}/k_{\rm B}T}$$

$$\tag{4}$$

where the normalizing factor Z ensures that $\sum_{i,j} P_{ij} = 1$. The term W_{ij} is a statistical factor which depends on the number of equivalent ways to connecting the starting species to yield a given cluster. For simplicity, we reduce the number of energy gains E_{ij} to three, depending on the elastic nature (flexible, intermediate, stressed rigid) of the considered cluster: E_{flex} , E_{iso} , E_{stress} . An additional energy, E_{ring} , is introduced for ring closure leading to ES connections. Due to the normalizing factor Z, the whole construction depends only on three parameters $e_i = \exp[(E_{iso} - E_i)/k_{\rm B}T]$ with i = (flex, stress, ring). Two of these parameters can be exactly determined by writing a conservation equation for the species, i.e.,

$$P_{\rm GG} + \frac{1}{2}P_{\rm GS} + \frac{1}{2}P_{\rm GSb} = x = P_{\rm SbSb} + \frac{1}{2}P_{\rm SbS} + \frac{1}{2}P_{\rm GSb}$$
(5)

which can be solved in the chalcogen-rich ($e_{\text{stress}} = 0$) and chalcogen-poor ($e_{\text{flex}} = 0$) region of the phase diagram, leading respectively to the following:

$$e_{\text{flex}} = \frac{2(72xe_{\text{ring}} + 4 - 31x)}{3(2 - 11x)}$$
(6a)

$$e_{\rm stress} = -\frac{(72xe_{\rm ring} + 4 - 31x)}{16x}$$
(6b)

One also remarks that to be valid $e_i > 0$ (i = stress, flex), and thus one has a constraint on the compositional domain bounded by the concentration:

$$x_{\rm c} = \frac{4}{31 - 72e_{\rm ring}} \tag{7}$$

The composition x_c also represents $x_c(2)^t$ as discussed below. Equations 6a and 6b allow us to have the probabilities depend only on a single parameter e_{ring} (Table 1).

Note that the condition $e_i = 1$ (i = flex, ring, stressed) leads to the usual random bond distribution with probabilities only given by their statistical weight. In the latter case, CS or ES GeSe_{4/2} connections have non-normalized probabilities, which are given by $16P_G^2$ and $72P_G^2$, respectively. Among this population of generated clusters, one can compute the probability of finding flexible ($n_c < 3$), isostatically rigid ($n_c = 3$) and stressed-rigid ($n_c > 3$) clusters. Here n_c represents the count of bonding constraints due to bond-stretching and bondbending forces per atom. Various situations^{8,44} can then be investigated: random bonding case, and self-organized case which amounts to avoid forming stressed- rigid CS connections as far as possible as x is steadily increased. This amounts requesting $e_{\text{stress}} = 0$ (eq 6b) as long as possible.

One identifies the *rigidity* transition by requiring the vanishing of the count of floppy modes $(f = 3 - n_c = 0)$, and the stress transition x_c given by equ. 7 when CS stressed-rigid links can no longer be avoided as *x* increases. In the case of random connections, theory predicts a *single* transition near r = 2.40 or x = 13.3% (broken blue curve, Figure 4). This also



Figure 4. SICA results for $Ge_xSb_xSe_{100-x}$ glasses. Calculation of the number of floppy modes $f = 3 - n_c$ from the local structures (broken blue line) and from self-organized clusters having a finite fraction of ES motifs (solid blue line). The latter defines the Rigidity Transition (RT) at 15.3%. Right axis: probability (black solid lines) of finding flexible, isostatic (which peaks at the Stress Transition near 17.2%) and stressed rigid clusters as function of composition. The pink zone defines the IP.

corresponds to the result obtained from a direct constraint count on the initial local structures ($\text{GeSe}_{4/2}$ tetrahedra, $\text{SbSe}_{3/2}$ pyramids and Se_2 -chains). In the case of self-organized networks (avoidance of CS connections as long as possible, see Table 1), and with a finite fraction of ES motifs as the only adjustable parameter in the theory, calculations predict (i) the rigidity transition (RT) to upshift in *x* or *r* (solid blue curve, Figure 4), and (ii) the RT and the stress-transition (ST) to split, and to occur near $x_c(1)^t = 15.3\%$ and near $x_c(2)^t = 17.2\%$ (right axis, Figure 4) for, e.g., at an ES/CS fraction η of 1.4 at a fixed composition of x = 18.18%. In fact, e_{ring} can be expressed as a function of the ES to CS ratio defined by the following:

$$\eta = \frac{p^{\rm ES}_{\rm GG}}{p^{\rm CS}_{\rm GG}} = \frac{9e_{\rm ring}}{e_{\rm stress}}$$
(8)

and using eq 6b, one has the following:

$$e_{\rm ring} = \frac{13\eta}{72(\eta+1)} \tag{9}$$

at $x = x_{ct} = 18.18\%$.

Our SICA predictions are is excellent accord (Figure 5) with the experimental results if the ES/CS fraction is taken to be about 1-1.7, of the same order to what is obtained (1.17) from



Figure 5. SICA results for the location of the rigidity (red curve) and stress transition (black curve) as a function of the ES/CS ratio. The broken horizontal lines correspond to the experimentally determined transitions $x_c(1)$ and $x_c(2)$ and the intercept with the SICA result provides an estimation of η (between 1 and 1.7, gray zone).

Molecular Dynamics simulations⁴⁵ for a similar system (GeSe₂). The experimental results of the ES/CS ratio come from Raman scattering and requires knowledge of the scattering cross-section of the ES and CS vibrational modes to convert the observed scattering strength ratio of about 1/2 to reliably determine ES/CS fraction. At present, there are no reliable estimates of these matrix elements effects to deduce the ES/CS fraction. It is also possible that if the agglomeration steps in SICA are extended to higher level, a smaller ES/CS fraction would suffice to understanding the observed thresholds of the rigidity- and stress-transitions in the present Ge_xSb_xSe_{100-2x} ternary. We plan to explore these possibilities in future studies.

3.4. Global Elastic Phase Diagram of the Ge–Sb–Se ternary. Given the present results on the $Ge_xSb_xSe_{100-2x}$ ternary, and those on the Ge_tSe_{100-t} binary,^{24,25} it is possible to construct a global elastic phase diagram of the present Ge–Sb–Se ternary as shown in Figure 6. It is useful to recall that in



Figure 6. Global elastic phase diagram of bulk Ge–Sb–Se ternary glasses constructed from the established elastic phases in the Ge_xSe_{100-x} binary,⁷ Ge₇Sb_xSe_{93-x} ternary,⁶ and the present work on $Ge_xSb_xSe_{100-2x}$ ternary. Phases are indicated as flexible (A, yellow), intermediate (B, green), and stressed rigid (C, indigo). The chemical threshold phase boundary is shown by the black dashed line and resides in the Stressed-rigid phase. The glass forming region is taken from Sreeram et al.³¹ and Borisova et al.⁴⁷

the Sb_ySe_{100-y} binary, homogeneous glass formation is restricted to a rather small concentration, y < 0.02 of Sb as noted earlier.^{21,46} At higher concentrations, Sb tends to cluster as a nanocrystalline Sb₂Se₃ moiety in Se–rich glasses, as suggested by the glass transition temperature remaining essentially close to 40 °C, the value characteristic of Se-glass, while displaying a slope $dT_g/dy = T_g(y = 0)/\ln(3/2)$ for y < 1% in agreement with SAT predictions.²¹ The glass forming tendency of binary Ge_xSe_{100-x} glasses and their elastic phases have been extensively studied over the years, and the recent results^{24,25} on homogeneous glasses are included in Figure 6.

Madhu et al. reported⁶ observing a narrow ($\Delta r = 0.02$) IP centered near x = 26% or r = 2.40 in the Ge-deficient Ge₇Sb_xSe_{93-x} ternary (Figure 3). In this ternary, as in the case of the base Ge₇Se₉₃ binary, Raman scattering unambiguously shows absence of ES GeSe₂ tetrahedral units. These structural motifs are first manifested when the Ge concentration x exceeds about 12% in binary Ge_xSe_{100-x} glasses. Thus, theory and experiment come together in revealing that the narrowing of the IP width in the Ge₇Sb_xSe_{93-x} ternary stems from absence of ES motifs.

From the above cited reports one can now construct a global elastic phase diagram of the present Ge-Sb-Se ternary (Figure 6). Se-rich bulk glasses composed of Se_n chains are elastically floppy⁴⁸ and comprise the Flexible phase of the present ternary shown as the A (yellow)-region in Figure 6. Upon progressive cross-linking by alloying both Ge and Sb, a rigid but unstressed elastic phase, Intermediate phase, manifests in the B (green)region. This phase is of special interest in applications because of its unstressed nature by virtue of topology. And as one crosslinks the network further by alloying more Sb and Ge, the stressed-rigid elastic phase C(indigo) is manifested. Stressedrigid glassy networks eventually demix,³² a feature observed in the present glasses as well. A parallel elastic phase diagram in corresponding ternary sulfides, Ge-Sb-S, probably exists although studies⁴⁹ to date have not been directed toward that goal.^{31,47}

4. CONCLUSIONS

In summary, topology and connectivity determined by local structural motifs (Figure 4) provides a useful basis to quantitatively understand the three elastic phases of ternary $Ge_xPn_xSe_{100-2x}$ glasses, with Pn = P, As, Sb. In these polymerized networks, the ring topology and connectivity of the most Se-rich isostatic local structural motif determines the Rigidity transition, while percolation of the most Se-deficient structural motif the Stress transition. Presence of a multitude of isostatic local structural motifs with variable stoichiometry can be used to design wide IPs in multicomponent network glasses for applications. In this work, we present the first example of a global phase diagram in Sb bearing chalcogenides established experimentally and supported by theory. Such materials are the active element of phase change memories.⁵⁰

ASSOCIATED CONTENT

Supporting Information

Details on synthesis of bulk glasses, and particularly a check on their structural homogeneity by FT-Raman profiling experiments. Raman vibrational modes characteristic of homopolar Ge–Ge and Sb–Sb bonds. Results of numerical simulations of IR and Raman active modes of ethane-like (Ge₂Se₆) and ethylene-like (Sb₂Se₄) clusters containing homopolar bonds of interest. Predictions from first principles cluster calculations

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compared to the observed modes. Observed Raman spectra of Se-deficient glasses at x > 18.2% are deconvoluted, and observed modes identified. Comparison of the predicted and observed mode frequencies and particularly the scattering strength variation of the observed modes with x providing for consistency of mode assignments. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

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