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1. Introduction

Spectroscopic ellipsometry as a route to thermodynamic characterization †‡

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Strategies for synthesizing molecularly designed materials are expanding, but methods for their thermodynamic characterization are not. This shortfall presents a challenge to the goal of connecting local molecular structure with material properties and response. Fundamental thermodynamic guantities, including the thermal expansion coefficient, α , can serve as powerful inputs to models, yielding insight and predictive power for phenomena ranging from miscibility to dynamic relaxation. However, the usual routes for thermodynamic characterization often require a significant sample size (e.g. one gram), or challenging experimental set-ups (e.g. mercury as a confining fluid), or both. Here, we apply spectroscopic ellipsometry, which is an optical technique for thin film analysis, to obtain thermodynamic data. We clarify issues in the scientific literature concerning the connection between ellipsometric and volumetric thermal expansion coefficients for substances in both the glass and melt states. We analyze temperature-dependent data derived using both ellipsometry and macro-scale dilatometric techniques for ten different polymers. We find superb correlation between the α values obtained via the two techniques, after considering the effects of mechanical confinement by the substrate for a glassy thin film. We show how the ellipsometric α can serve as input to the locally correlated lattice theory to yield predictions for the percent free volume in each polymer as a function of temperature. We find that the ellipsometric α at the glass transition temperature, T_{gr} is not only material dependent, but it is linearly correlated with T_q itself. Spectroscopic ellipsometry, which requires only very small quantities of sample and is straightforward to perform, will significantly expand the range of systems for which thermodynamic properties can be characterized. It will thus advance our ability to use theory and modeling to predict the miscibility and dynamic relaxation of new materials. As such, ellipsometry will be able to underpin materials synthesis and property design.

Thermodynamic data are necessary for the modelling of material phenomena, including miscibility and dynamic relaxation. One essential parameter is the volumetric thermal expansivity (α_v). When a substance is subjected to an isobaric temperature shift, it will equilibrate by changing its volume. The fractional volume change with temperature defines α_v . Typically, α_v is obtained *via* the gold standard method of pressurized dilatometry, which collects accurate pressure-volume-temperature (*PVT*) data.¹⁻³ If a substance is in the form of a thin film on a substrate, volume changes with increasing temperature are not easily measured; changes in the thickness are more easily quantified. For instance, ellipsometry is a non-invasive optical technique⁴ that measures film thicknesses with sub-nm accuracy. It has been used to measure the thermal expansivity in the direction normal to the substrate, α_N . Here, we are interested in the relationship between α_N and α_V , and whether the connection between the two depends on the state (solid or liquid). As we describe below, there is inconsistent understanding of how an α_N value (derived from ellipsometry) relates to the familiar, well-defined α_V and to the one-dimensional linear expansivity. In this paper, we use new extensive experimental data to confirm the relationships explicitly for both the melt and glass states.

A key motivation of our research arises because important aspects of a material's dynamic, thermodynamic, and mechanical behavior can be correlated with α_V using models such as theory-based equations of state (EOS).^{5–12} For a three-parameter

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EOS, a complete characterization requires an α_V value together with the compressibility, κ , and a single value of specific volume. However, as we demonstrate in this work, useful correlations can be made using α_v values alone. In our previous work using the EOS derived from the Locally Correlated Lattice (LCL) model, we were able to predict and compare the free volume (V_{free}) that exists in a range of melt and glassy materials.¹³ V_{free} is the amount of space a solid or liquid has in excess of its limiting state at close molecular packing. It is strongly connected to both the material's thermodynamic behavior (one example being its compatibility, or miscibility, with other components),^{14,15} and to its dynamic properties,^{16,17} including viscosity, diffusion, segmental relaxation times, and ultimately the glass transition temperature, $T_{\rm g}$. Thus, thermodynamic characterization powerfully unlocks an understanding of a wide range of properties.

Although α_V (obtained *via PVT* measurements) is essential, in most laboratories, α_N (e.g. via ellipsometry) is far more accessible. While carefully applied volume dilatometry can also yield atmospheric V(T) data,¹⁸ many substances, such as polymers, can be challenging to work with, and hence data availability is limited. Pressurized or ambient volume dilatometry requires confining fluids, commonly mercury. If mercury is avoided by the use of another type of confining fluid, then the PVT behavior of that fluid must be accurately known, and further, there must be no uptake of that particular fluid into the sample.³ With gas pycnometers, uptake of He into the sample is a similar concern in some cases (e.g. fluorocarbon polymers).¹ Other methods to obtain ambient density, such as weighing bottles and hydrostatic weighing techniques, can be complicated by sample viscosities that are too high (former) or too low (latter).^{1,2} Further, most of these approaches have the disadvantage of requiring a relatively large sample size,1-3,18 typically 1 to 2 g, compared to <1 mg for a thin film for ellipsometry. Deuterium-labelled molecules¹⁹ and some novel materials are synthesized in small (<1 g) quantities. Synthetic efforts continue to expand the range of interesting polymers and copolymers, with the goal of optimizing properties for desired applications. Currently, aside from perhaps standard thermal analysis, many new materials have not been thermodynamically characterized at all. This objective would be significantly advanced by the ability to determine volumetric thermal expansion coefficients using the more readily available measurements of α_N via ellipsometry. The resulting expanded data set would lead to a better understanding of how to link molecular structure to properties and function.

We note that a comparison between volumetric and ellipsometric data is only relevant when the films are thick enough that the behavior can be considered bulk-like. There is substantial evidence, including in the ellipsometry studies of Singh *et al.*,²⁰ and the X-ray reflectivity work of Miyazaki *et al.*,²¹ that for film thicknesses of at least 100 nm, α_N becomes independent of thickness. In the work reported here, the mean thickness of the samples is *ca.* 450 nm, which is well into the bulk regime.

In this work we present experimental evidence, with appropriate data analysis, that shows ellipsometry can be a convenient and accessible route to fundamental thermodynamic characterization. We focus on polymers, but the methods and theory apply to any optically-transparent amorphous substance. An outline of the remainder of the paper is as follows. In Section 2, we define the relevant quantities, and provide physical background on the expected quantitative relationships between the measured expansivities, α_N and α_V . In Section 3, we verify these relationships as we present results and discussion that covers our ellipsometry measurements and analysis performed on 10 different polymers, including a comparison with corresponding results from *PVT* data. Following that, we use the α_N values from these ellipsometry experiments to characterize and compare the free volume of all of the polymers by application of the LCL EOS. We present a summary in Section 4 and provide experimental details in Section 5.

2. Theoretical background

Using ellipsometry measurements of the height, *h*, of a thin film as a function of temperature, *T*, leads to results for the expansivity $\alpha_N = (1/h)dh/dT$, where N indicates that the expansion is in the direction normal to the substrate. However, α_N is not the same quantity as the material's coefficient of linear expansion, $\alpha_L = (1/L)dL/dT$ which characterizes expansion in the length, *L*, in a single direction when the conditions are isotropic and unconstrained. In other words, α_L is defined for the situation in which the material is allowed to expand freely in all three directions and therefore (for an isotropic material) $\alpha_L = \alpha_V/3$, where $\alpha_V = (1/V)dV/dT$ is the familiar volumetric thermal expansion coefficient of the bulk material.²² (Maintaining simplicity in notation, it is understood in the present context that α_N , α_L , and α_V are all defined by derivatives taken at a constant ambient pressure.)

In ellipsometry experiments, the material being analyzed, here a polymer film, adheres to a substrate that remains essentially at a fixed area, A, because the coefficient of thermal expansion for any typical substrate, *e.g.* Si, is two orders of magnitude lower than for polymers.²³ Consequently, expansion is only possible in the direction normal to the substrate, *i.e.* only the film height, h, changes. See Fig. 1. When a polymer film is in the melt state, we make an important assumption that its behavior should be analogous to that of a liquid expanding inside a piston of fixed area. For such a case, with a constant A, we expect the following relationship:

$$\alpha_{\rm V} = \frac{1}{V} \frac{\mathrm{d}V}{\mathrm{d}T} = \frac{1}{Ah} \frac{A\mathrm{d}h}{\mathrm{d}T} = \frac{1}{h} \frac{\mathrm{d}h}{\mathrm{d}T} = \alpha_{\rm N} \tag{1}$$

This physical picture wherein an area of a substrate is assumed to hold a fixed mass of polymer due to lateral constraints coincides with the view advanced several decades ago in works by Beaucage *et al.*²⁴ and Wallace *et al.*²⁵ in studies of polystyrene.

In contrast to the melt case, a polymer in a solid or glassy state cannot expand in just a single direction without accruing internal elastic stress. Thus, as temperature increases, a glassy sample will achieve a final volume that is less than the



Fig. 1 How a thin film expands with increasing *T* when constrained by a non-expanding substrate. It is assumed here that there is no slipping at the interface.

equilibrium volume of the free sample at that temperature. In this case, one should expect that $\alpha_{\rm N} < \alpha_{\rm V}$. The elastic model proposed by Wallace *et al.*²⁵ (below) predicts this trend analytically. Additionally, in the ESI,† we walk the reader through a simple thought experiment that can illustrate this inequality in a more pictorial way.

For the case of an isotropic solid or a glass, α_N can be connected to α_V through the Poisson ratio, v, which is the negative of the ratio of the lateral strain to the longitudinal strain.²⁶ Based on this concept, the elastic model of Wallace *et al.*²⁵ gives the following:

$$\alpha_{\rm N} = \left(\frac{1+\upsilon}{1-\upsilon}\right) \alpha_{\rm L} = \frac{1}{3} \left(\frac{1+\upsilon}{1-\upsilon}\right) \alpha_{\rm V} \tag{2}$$

For glassy polymers, typical values of v are around 0.3 to 0.4. This implies that one might expect ellipsometry results to yield $\alpha_N \approx (2/3)\alpha_V$ when the thin films are glassy.

For the case of the melt, eqn (2) will reduce to the eqn (1) result, $\alpha_N = \alpha_V$, because the Poisson's ratio logically must become v = 0.5; tabulated values for the Poisson ratio are typically quoted for the glassy state. Note that v = 0.5 is sometimes associated with the system being mechanically "incompressible" (with *P* fixed). However, a liquid or melt is certainly compressible (*e.g. via* a pressure change), and in this case the value of 0.5 simply represents the reality that, upon constant pressure compression in one direction, an unrestricted liquid will expand in the other directions so as to retain its original volume.

Several examples in the literature exhibit some confusion or at least a lack of consensus - in using ellipsometry data to determine thermal expansion coefficients. A recent review article²⁷ implied that α_N from ellipsometry (treated as a linear expansion coefficient, α_L) should equal $\alpha_V/3$ and did not describe quantitatively the effects of mechanical confinement. Somewhat less directly, reported measurements²⁸ have been implicitly connected to the linear expansion coefficient by the choice of notation, but with no relation to the given α_v . There have also been several instances²⁹⁻³⁵ where the intent is to equate α_N for the glass directly with α_V ; the mechanical confinement effect leading to the $\alpha_N < \alpha_V$ prediction of the Wallace et al. elastic model is not considered. There are other examples^{36,37} that apparently apply the Wallace et al. model without changing v to 0.5 at temperatures above T_{g} ; such an approach leads to predicted α_V values for the melt that will be larger by roughly 50%.

In summary, according to the simple model picture summarized in this section, α_N , calculated by monitoring film thickness changes in a carefully executed ellipsometry experiment on a melt sample constrained by a non-expanding substrate should give the same result as α_{v} , extracted from experimental PVT data collected on a corresponding unrestricted melt sample. However, for a glass, one should expect $\alpha_{\rm N}$ < $\alpha_{\rm V}$. While the implications for the analysis of melt and glass data (eqn (1) and (2)) have been accepted and applied by multiple researchers, there nevertheless exists some significant confusion in the literature. This paper is the first to provide a substantive test of the model result, which we accomplish through careful analysis of ten different polymers. This level of scrutiny, verifying the relationship between $\alpha_{\rm V}$ and $\alpha_{\rm N}$ in the melt (eqn (1)) and in the glass (eqn (2)), is essential for promoting spectroscopic ellipsometry as a viable route to the data needed for EOS modeling, as will be demonstrated.

3. Results and discussion

Table 1 shows the polymer names and acronyms for the ten investigated polymers, as well as their molecular weights, T_g , and references to experimental *PVT* data to which we will compare our expansivity results obtained *via* ellipsometry. The systems cover a range of polymer chemistries and include one copolymer. Spectroscopic ellipsometry data were collected on thin films of these polymers supported on an Si substrate (with $\alpha_L = 2.6 \times 10^{-6} \,^{\circ}\text{C}^{-1}$),²³ deposited from solution *via* spincoating (see Materials and methods). The films had a mean thickness of about 450 nm (all ≥ 150 nm). At these thickness values, we expect the relative contribution of the interfacial regions (*e.g.* α -relaxation times vary within ~ 5 nm of an interface^{38,39}) to be negligibly small compared to the amount of more bulk-like material in the film.

By analyzing the reflection of polarized light from a sample, ellipsometry measures – as a function of wavelength – two parameters: Ψ (related to the change in the amplitude of light) and Δ (related to its change in phase). There are two primary methods reported in the literature for using ellipsometry to determine α_N values in thin films. In a dynamic scan, 28,35,45,46 the temperature is increased at a constant heating rate while ellipsometry data are collected continuously. In a step-wise scan, 20,24,30,34 the temperature is raised in increments, and data are collected at fixed temperatures. The data collection rate in a dynamic scan usually must be faster than in a stepwise scan. Typically, only a few wavelengths are used in the

 Table 1
 Polymers and their characteristics

Acronym	Chemical name	$M_{ m w}{}^a ({ m g}{ m mol}^{-1})$	$T_{g}^{\ b}$ (°C)	Ref. for <i>PVT</i> data
PS	Polystyrene	96 000	100	1
PMMA	Poly(methyl methacrylate)	100 250	105	1
PCHMA	Poly(cyclohexyl methyl methacrylate)	65 000	104	40 and 41 ^c
PVME	Poly(vinyl methyl ether)	23 070	-31	42^d
PVAc	Poly(vinyl acetate)	100 000	32	1
ΡαΜS	Poly(α-methyl styrene)	97 600	168	43
PBMA	Poly(butyl methacrylate)	182 600	20	1
PEMA	Poly(ethyl methacrylate)	340 000	63	1
PSAN	Poly(styrene-co-acrylonitrile) (75%/25%)	165 000	107	1
TMPC	Tetramethyl bisphenolA polycarbonate	54000	196	44

^a The molecular weight (M_w) values tabulated here correspond to the samples studied here via ellipsometry. Literature sources for PVT data and T_g
values correspond to different samples. ^b The T_g literature values are from the compilation in Table 1 of ref. 13. The T_g values apparent from the
present ellipsometry measurements (for those polymers with T_{g} > room temperature) are within 5 °C of these values. ^c The PCHMA melt data
points shown were regenerated from the published fit to the Tait equation of state (parameters available in ref. 12 and 40); given the accuracy of the
Tait equation these values are equivalent to the actual measured data. Data for the PCHMA glass are from ref. 41. ^d The original data set contains
<i>PVT</i> data for $P \ge 10$ MPa; by fitting the LCL equation of state to this set we have predicted $V(T)$ data points at $P = 1$ atm, which are expected to be
very accurate. See ESI.

analysis, but fast spectroscopic scans over a wide spectrum are possible with modern instruments.^{46,47} Hereafter, we will compare the results obtained *via* dynamic scans and step-wise scans for PCHMA thin films, as an example.

Dynamic data are presented in Fig. 2(a and b) showing results obtained with four wavelengths measured as the temperature was raised at a constant rate of 2.5 °C min⁻¹. The ellipsometry parameters change (increasing or decreasing) in approximate direct proportion to the change in the thickness (Δh) and refractive index (Δn).⁴⁵ Following a quick glance at these data, two linear regions are apparent, corresponding to the glass in the lower temperature region and the melt state in the higher region. These data were analyzed using commercial software to obtain *h* and refractive index (*n*) of the PCHMA film as a function of temperature, as is shown in Fig. 2(c and d). The thickness increases weakly with temperature in the low temperature region, which is attributed to the glass. Similarly, there is a weaker dependence of *n* on temperature in this region.

The dynamic scans are very effective in identifying the temperature above which the thermal expansivity increases, which is used to define the $T_{\rm g}$. For the purposes of illustration, our data analysis uses the room-temperature value of the complex refractive index as a function of wavelength, $\tilde{n}(\lambda)$, for the silicon substrate, across the wide range of temperatures. (Here, $\tilde{n} = n + ik$, where k is the extinction coefficient.) This model is chosen frequently in the literature.^{28,32,35,45,48} Although neglecting the temperature dependence of $\tilde{n}(\lambda)$ is not expected to affect the values obtained for the inflection point, and hence the $T_{\rm g}$, it has a pronounced effect on the value of $\alpha_{\rm N}$ obtained. For this reason, dynamic scans are not suited for the determination of α_N , unless the data analysis explicitly considers the changing $\tilde{n}(\lambda)$ in the substrate.^{20,46,47} This point has been made previously by Kahle et al.46 In the analysis presented hereafter, we will show quantitatively the implications of using the fixed $\tilde{n}(\lambda)$ model.

Continuing the example of PCHMA, we next present in Fig. 3(a) representative spectra obtained from a thin film at

two different temperatures in a step-wise scan. The best-fits to the spectra were made by using $\tilde{n}(\lambda)$ for Si at the corresponding temperatures. See Fig. S2 in ESI[†] for example data. The results of this analysis for a PCHMA thin film across the entire range of temperatures are shown in Fig. 3(b and c). Analysis of the h(T)obtained α_N to be 1.86 (±0.07) × 10⁻⁴ °C⁻¹ in the temperature range from 30 to 95 °C (attributed to the glass) and $\alpha_{\rm N}$ = 5.49 $(\pm 0.03) \times 10^{-4}$ °C⁻¹ from 100 to 180 °C (attributed to the melt). These values differ substantially from what were found using the dynamic analysis (reported in the caption for Fig. 2(c)). The difference between the two methods can be explained by the fact that the analysis of the step-wise data uses a different $\tilde{n}(\lambda)$ for the Si substrate at each temperature, which is more difficult to include analytically in the dynamic modeling when the temperature is constantly changing during the measurement. Note that some commercial software does not offer $\tilde{n}(T)$ in the analysis of dynamic data, which has resulted in some poor published results.

To provide a comparison of the models, the same ellipsometry spectra used to construct Fig. 3 were re-analyzed without considering the temperature dependence of $\hat{n}(\lambda)$ for the Si substrate. (See Fig. S3 and the comparison of α_N values in Table S1 of the ESI.†) With inaccurate values for $\hat{n}(\lambda)$ of Si, the values of α_N are noticeably higher. This same pattern was found in the experiments on other polymers reported in Table S1 (ESI†).

Moreover, we note that in a dynamic scan both *T* and *h* are changing during the course of a measurement ("on the fly"), naturally leading to uncertainty in both, which will be greatest when heating rates are fast and spectral acquisition is slow. The dynamic scan performed in Fig. 2 measured ψ and Δ at only four wavelengths (λ), and hence $n(\lambda)$ was determined by fitting to four measurements (as described in the Methods), rather than across the entire spectrum, as was the case for the step-wise scans. (Data could have been collected at more wavelengths, but then the time of data acquisition would be greater, and the measurement would be made over a greater temperature interval, thereby adding uncertainty to *T*.) There is thus greater uncertainty in the *n* (and *h*) values obtained by the



Fig. 2 (a) Ellipsometric data obtained from a dynamic heating scan for a thin film of PCHMA using four wavelengths as are shown in the legend: (a) Ψ data and (b) Δ data. (c) Thickness of this film as a function of temperature, obtained from the analysis of the data shown in (a and b). The α_N value in the temperature range from 28 to 101 °C (attributed to the glass region) is 2.12 (± 0.02) × 10⁻⁴ °C⁻¹, whereas in the temperature range from 105 to 170 °C (attributed to the melt region) $\alpha_N = 7.52 (\pm 0.02) \times 10^{-4} °C^{-1}$. The glass transition temperature is indicated by the change in the gradient. (d) Refractive index, *n* (at a wavelength of 0.55 µm) as a function of temperature for the same film.

dynamic scan method, which explains the lack of agreement of values for PCHMA when comparing data in Fig. 2d and 3c.

Given these findings, all of the results presented hereafter will use measurements from step-wise temperature scans analyzed using temperature-dependent values of $\tilde{n}(\lambda)$ for the substrate.

We aim to develop a robust method for measuring α_N using ellipsometry, and to compare the results with α_V from *PVT* data. We continue here to use PCHMA as an example to demonstrate a few of the finer points in calculating the α values. To calculate $\alpha_N = (1/h)(dh/dT)$ (eqn (1)), one could evaluate the slope dh/dTover a chosen linear region and then divide it by the initial *h* value, or preferably, by the mean *h* value over that region. The selection of the particular value of *h* (or equivalently, the associated reference temperature that defines it) will have an impact on the value of α_N , *e.g.* by about 2 or 3%. In the literature, there are examples of using an initial thickness at an arbitrary reference temperature^{35,45,47} and other examples using the mean thickness value for the temperature range.²⁹ A more convenient approach is illustrated in Fig. 4(a),⁴⁹ where we plot the natural log of *h* vs. *T*; here the slope will correspond directly to α_{N} .

To draw the best comparison with PVT measurements, we have been careful to match the temperature ranges of the data used in both methods, because for some polymer melts the α value depends on temperature. (In these cases, typically, α will gradually increase with increasing T.) In deciding on the temperature range, it is important to avoid points that cross the glass transition temperature. Further, for PEMA and PBMA, we avoid choosing a temperature range that could lead to possible changes in partial melt crystallinity; the latter can be detected via calorimetry and have been discussed previously in Walsh et al.⁵⁰ Finally, we excluded any ellipsometry data in the high T extreme of the range that show evidence of developing negative curvature in h vs. T. Pronounced negative curvature of h (or V) against T on an ambient pressure isobar would not be typical of standard single-phase expansion. Such a reversal in h(T) dependence might be caused by significant sample



Fig. 3 (a) Ellipsometry spectra (Ψ data (open symbols) and Δ data (filled symbols)) for a thin PCHMA film from a step-wise heating experiment at 30 °C (square symbols) and 170 °C (triangles). The best fits to the data, shown as the solid lines, take the substrates' complex refractive index, $\tilde{n}(\lambda)$, change with temperature into account. Each arrow points to the relevant axis denoting the parameters. (b) Thickness as a function of temperature graph from a step-wise heating scan performed on this PCHMA film showing the glass (blue) and melt (red) regions. The glass transition temperature can be found from the change in the gradient. (c) Refractive index, *n* (at a wavelength of 0.55 µm) varying with temperature for the same film as in (b).



Fig. 4 Comparison of ellipsometry and *PVT* data for PCHMA. Experimental measurements of the temperature dependence of (a) the natural logarithm of *h* from Fig. 3b and (b) the natural logarithm of volume (taken from the literature^{40,41}). (All data correspond to P = 1 atm.) The regions over which the linear best fits were taken are show by the red (melt) and blue (glass) symbols; the slope of each line gives the α value. Note that for consistency the regimes of the linear fits of the ellipsometry and *PVT* data are selected to cover matching *T* ranges.

Table 2 α values via ellipsometry and via PVT data: results for the melt

Polymer	$\begin{array}{l} Ellipsometry \\ \alpha_N \times \ 10^4 \ (^\circ C^{-1}) \end{array}$	$\begin{array}{l} \textit{PVT}\alpha_V\times10^4 \\ \left(^\circ C^{-1}\right) \end{array}$	α_N/α_V	Ellipsometry T range (°C)	<i>PVT T</i> range (°C
PS^a	5.88 ± 0.29	5.87	1.002	100-165	100-170
PMMA	5.33 ± 0.38	5.97	0.893	115-180	121-179
PCHMA ^a	5.69 ± 0.12	6.03	0.943	105-160	103-163
PVME	6.39 ± 0.45	6.70	0.954	50-100	44-110
PVAc	6.85 ± 0.48	7.02	0.976	55-105	54-103
PaMS	5.05 ± 0.36	5.26	0.960	165-225	190-209
PBMA ^a	6.50 ± 0.26	6.21	1.047	40-100	43-101
PEMA	5.51 ± 0.39	6.01	0.917	70-110	70-112
PSAN ^a	5.43 ± 0.11	5.69	0.954	105-170	110-170
TMPC^{b}	8.20 ± 0.58	7.69	1.066	180-205	218 - 242

^{*a*} For PS, PCHMA, and PBMA, the α_N value is the average from five separate trials and three separate trials in the case of PSAN; the α_N result for all other polymers is from a single trial. ^{*b*} There was a negative curvature in h(T) for TPMC in the melt state at higher temperatures. Hence, only a narrow range of temperature could be used to estimate the α_N for TMPC, and the results for this polymer melt are less reliable than for the others.

Table 3 α values via ellipsometry and via PVT data: results for the glass^a

Polymer	$\begin{array}{l} Ellipsometry \\ \alpha_N \times \ 10^4 \left({^\circ C}^{-1} \right) \end{array}$	$\begin{array}{l} \textit{PVT}\alpha_V\times10^4 \\ \left(^\circ C^{-1}\right) \end{array}$	α_N/α_V	Ellipsometry T range (°C)	<i>PVT T</i> range (°C)
PS^b	1.68 ± 0.14	2.46	0.681	40-90	40-91
PMMA	2.33 ± 0.45	2.65	0.879	40-105	40-101
$PCHMA^{b}$	1.84 ± 0.10	3.27	0.562	40-85	40-85
PαMS	1.47 ± 0.28	2.14	0.687	120-160	120-160
PEMA	2.01 ± 0.38	2.33	0.863	40-45	30-38
$PSAN^{b}$	1.50 ± 0.26	2.08	0.712	30-95	30-90
TMPC	1.34 ± 0.26	1.60	0.838	100-175	103-177

^{*a*} Results were not obtained for three of the polymers (PVAc, PBMA, and PVME) in their glassy states because of their lower T_g values. Our ellipsometry stage does not cool below standard room temperature. ^{*b*} The α_N value is the mean from five separate trials for PS, four separate trials for PCHMA, and three separate trials for PSAN; the α_N result for all other polymers is from a single trial.

changes, for example, in the degree of partial crystallinity (*e.g.* ref. 50) or decomposition. Using this set of practical guidelines, we identified a viable T range for each set of ellipsometry data, and then (data availability allowing) have taken the same range from the *PVT* data. For completeness, the temperature ranges for all polymers are summarized in Tables 2 and 3.

Continuing with our example of PCHMA in Fig. 4(b), we show PVT data plotted as $\ln V$ over the relevant range in T (P = 1 atm). This is analogous to the ellipsometry $\ln h vs$. T plot in panel (a). Both the PVT and ellipsometry data show a regime corresponding to the melt and another regime corresponding to the glass. ($T_{\rm g}$ occurs around 100 °C.) As discussed in the guidelines above, the figure shows how matching data ranges were selected for the determination of α , and how the corresponding linear fits of ln h vs. T for the ellipsometry experiments and ln V vs. T for the PVT measurements are obtained for both the melt and glass regimes. The ellipsometry results for this particular replicate PCHMA sample (more on replicate trials below) thus yield α_N = 5.63 (±0.03) × 10⁻⁴ °C⁻¹ for the melt and $\alpha_{\rm N}$ = 1.83 (±0.07) × 10⁻⁴ °C⁻¹ for the glass, while the *PVT* measurements show corresponding results of α_V = 6.03 × 10^{-4} °C⁻¹ for the melt and $\alpha_V = 3.27 \times 10^{-4}$ °C⁻¹ for the glass.

For the case of the melt, it is important to note that the α_{N} value for PCHMA thin films is in good agreement with its PVTderived α_{v} . This general agreement is evident from the summary of our results from a survey of ten different polymers in Table 2. The corresponding results for the polymers in their glassy states are summarized in Table 3. These tables report uncertainties for the α_N values, which are estimated from random errors obtained from the analysis of replicates. Five independent measurements were made for three of the polymers (PCHMA, PS and PBMA), and three independent measurements for PSAN. For these polymers, the α_N listed in Table 2 corresponds to the mean of the replicates, and we take the standard error in the mean to define the random error. The four polymer sets showed an average standard deviation of 7.1%, which led to an average standard error of 3.3% in the value of their mean α_N . Using PBMA as example, the standard error of the mean was found to be 4.0% whereas the linear fitting error of ln h vs. T was less than 0.1%. The random error likewise dominated the analysis for PCHMA and PS. Given the average standard deviation of 7.1% found for the α_N of the four sets of replicates, we then applied this percentage error to estimate the error in α_N for the remaining six polymers in Table 2, which have only a single measurement.

For polymers in their melt states (Table 2), we compare the α_N and α_V values by plotting them in Fig. 5(a). The best-fit slope (forcing a zero intercept) of 0.977 falls very close to the value for a perfect diagonal, illustrating excellent agreement between the two methods. There is one partial outlier, TMPC, for which the melt analysis is considered less reliable than the others. This is because the onset of a negative curvature in h(T) began at approximately 20 °C above the T_g . Consequently, data points over a very narrow T range were used in approximating the TMPC melt expansivity. Keeping or disregarding this approximated result will not change our overall conclusions.

Our result that $\alpha_N \approx \alpha_V$ for the melt supports our previously noted physical assumption, where a film maintaining a fixed mass on a substrate with a constant area expands only in the normal direction such that, if it is a melt, it can reach the expected volume for any *T* (without accruing stress) much like a fluid in a piston. More broadly, the data reported here, on a large set of polymers, with multiple replicates, demonstrates the feasibility of using ellipsometry experiments to approximate *PVT* properties.

We next consider results for the polymers in their glassy state. The data in Table 3 clearly show that the α_N from ellipsometry for the glass is systematically lower than the corresponding α_V from *PVT* measurements. In Fig. 5(b) the α_N values are plotted against α_V , and the line of best fit reveals that the α_N for the glass falls short of the α_V by an average factor of $\alpha_N/\alpha_V = 0.72$. Note that here we report this fitted slope because it is informative in average assessment, however, we should not expect, for the polymers in their glassy forms, that $\alpha_N vs. \alpha_V$, should form a perfect point-by-point linear relationship because glassy polymers do not all have the same Poisson ratios and hence the conversion factor in eqn (2) will change (more below).



Fig. 5 (a) Correlation of α_N (obtained from ellipsometry) and α_V (derived from literature *PVT* data) for (a) polymer melts and (b) polymer glasses. The line on each is a linear fit (with intercept forced through the origin). The fit in (a) includes consideration of the TMPC values (outlier in the upper right) which is derived from very limited data (see footnote in Table 2). The error bars represent the standard error in the mean for four polymers with replicate measurements or otherwise the mean standard deviations. The gradient is 0.98 for the melts, and 0.72 for glasses.

The results for multiple polymers shown in the plot of Fig. 5(b) support the prediction of the Wallace *et al.* elastic model (eqn (2)) and the physical picture presented in the Theoretical Background, that, at least qualitatively, there is a significant difference between the α_N and α_V for a polymer glass. As for the melt, a glassy film is expected to be restricted to thermal expansion only in the normal direction. However, unlike a melt, a glass cannot expand solely in one direction without accruing an elastic stress, and so the observed α_N is lower than the *PVT*-based α_V .

We can further analyze the results in terms of the Wallace *et al.* model. Since eqn (2) proposes the relationship between α_N and α_V using the Poisson ratio, we therefore use our measured α_N and α_V values to back-calculate a value for v that would be "implied" by the model. The results are tabulated in Table 4. Notice that for the polymer glasses we obtain an average implied Poisson ratio of 0.38 ± 0.03 , which is in fairly good agreement with typical values of v for polymer glasses. For instance, PS has a literature value of 0.33,⁵¹ and PMMA has a value of 0.40.⁵² Though the predictions may not always be quantitative, the elastic picture provides a sensible way to understand, at least qualitatively, why the polymer glasses are the systems for which α_N is noticeably less than α_V .

As discussed in the assumptions described in the Theoretical Background, we expect polymer melts to expand without accruing elastic stress. In this limit the elastic model must reduce to give $\alpha_N = \alpha_V$, which follows from eqn (2) whenever v = 0.5, which is the expected value for a liquid. The results of our back-calculations using eqn (2) for the polymer melts, presented in Table 4, show an average implied Poisson ratio of 0.49 \pm 0.01, which equals 0.5 within uncertainty. The implied value of v for TMPC in the melt state is >0.5, which is physically unrealistic. However, as already noted, our measurement of α_N is less reliable for this polymer.

Table 4 Implied Poisson ratio values from elastic model: melt and glass

Polymer	v melt	v glass
PS	0.50	0.34
PMMA	0.46	0.45
PCHMA	0.48	0.26
PVME	0.48	_
PVAc	0.49	_
PaMS	0.49	0.35
PBMA	0.52	_
PEMA	0.47	0.44
PSAN	0.48	0.37
TMPC	0.52	0.43
Mean	0.49 ± 0.01	0.38 ± 0.03

We now present some examples of how the present ellipsometry results can be used for thermodynamic characterization and analysis, an endeavor that would have normally required PVT data. In particular, we analyze and compare the free volume content of the polymer melts using the LCL EOS.¹³

In our work, free volume, V_{free} , is defined as a quantity that depends on thermodynamic properties only, as follows

$$V_{\rm free} = V - V_{\rm hc} \tag{3}$$

It is the total system volume, *V*, minus the system's limiting (hard-core) volume at random close-packing, V_{hc} , as is illustrated in Fig. 6a. These quantities can be calculated once the LCL molecular level parameters are known. V_{hc} is taken to be a constant for a particular system, where *V* varies with both *P* and *T*. More background on this physical picture of a limiting state at random (amorphous) close-packing and its connection to the system's volume and expansivity behavior can be found in recent studies of both experimental and simulated systems.^{17,53,54}

The LCL model has three molecular parameters, and solving uniquely for all of them requires, at a minimum, three single



Fig. 6 (a) An illustration of the free volume, V_{tree} in two dimensions. The blue circles represent molecular segments that are contained in a total system volume (*V*), which is marked by the dark purple boundary. To the right, the yellow boundary shows the same system's corresponding volume at its limiting state of random close-packing (the hard-core volume, V_{hcc}). The difference between these volumes (marked by the green boundary) is the free volume (V_{free}). (b) LCL free volume predictions derived from α_{N} obtained *via* ellipsometry, which shows the percent free volume (= $100 \times V_{\text{free}}/V$) as a function of temperature (P = 1 atm) for polymers in their melt state. The points correspond to the value of the percent free volume at that polymer's experimental T_{g} . (Note the portions of the curves extending below the experimental T_{g} thus correspond to a theoretical extrapolation of the equilibrium melt.) The heavy line is a best fit to the points and marks the average boundary where upon cooling, the polymer melt curves intersect and thus transition from melt into glass. See text for details.

experimental quantities: α_V , the compressibility (κ), and a specific volume value measured at some particular given (T, P). (A complete PVT data set would of course effectively contain all of this information and such data sets are commonly used to characterize the LCL model.) Relevant to the application here, we emphasize that if we are interested only in properties at ambient pressure, then the requirements lessen, and we can do without κ . Going further, if our goal is only to predict the fractional free volume V_{free}/V , rather than absolute values of $V_{\rm free}$ itself, then we only need an experimental value for α , and do not even require the specific volume. Further, α values lead directly to one of the key LCL characteristic parameters, viz. the non-bonded nearest-neighbor interaction energy operating between segments, *ɛ*. We have shown in previous work¹⁵ that both ε and the polymer V_{free} are very strongly correlated with α . Thus, we can proceed using α_N values determined from ellipsometry for the polymer melts where we have shown that we can now approximate that $\alpha_V = \alpha_N$.

In Table 5 we summarize modeling results for the polymer melts; details on these calculations are provided in the ESI.† The table lists the α_N values determined *via* ellipsometry and the resulting fitted values of the characteristic LCL energetic parameter, ε . The right-hand columns in Table 5 list predictions for the percent free volume ($%V_{\text{free}} = 100 \times V_{\text{free}}/V$) of each polymer compared all at a single common temperature ($T = 125 \,^{\circ}$ C, and $P = 1 \,$ atm), and, compared when at each polymer's corresponding T_{g} . The free volume present in a material is intimately connected with its coefficient of thermal expansion. For example, the table shows that when they are compared at a common temperature, the higher the α , the higher is the $%V_{\text{free}}$.

As reflected in eqn (3), the system volume increases with T, therefore so must the free volume. In Fig. 6b we plot the

Table 5LCL fitting to ellipsometry results for the melt: parameters and $V_{\rm free}$ predictions

Polymer	T_{g}^{a} (°C)	$lpha_{ m N} imes 10^{4a} \ \left(^{\circ} { m C}^{-1} ight)$	$\frac{\text{LCL}}{(\text{J} \text{ mol}^{-1})}$	LCL %V _{free} at 125 °C	LCL % $V_{\rm free}$ at $T_{\rm g}$
PS	100	5.88	2119	12.38	11.12
PMMA	105	5.33	2223	11.44	10.51
PCHMA	104	5.69	2154	12.05	11.02
PVME	-31	6.39	2037	13.22	6.01
PVAc	32	6.85	1972	13.96	9.02
PαMS	168	5.05	2283	10.96	12.97
PBMA	20	6.50	2021	13.40	8.15
PEMA	63	5.51	2187	11.75	8.92
PSAN	107	5.43	2204	11.61	10.75
TMPC	196	8.20	1816	16.05	21.33

^{*a*} See Tables 2 and 3 for details on T_{g} and α_{N} values.

temperature dependence of the percent free volume, $VV_{\text{free}}(T)$, at ambient pressure for each of the polymer melts. Over a *T* span of about 200 °C, the VV_{free} of a polymer melt changes by about 10%. This may not seem like a large amount, however, small changes – even a fraction of a percent – can be important in determining both the dynamic response of a polymer, *e.g.* its segmental relaxation times near T_g ,^{16,17} and its thermodynamic behavior, *e.g.* its miscibility with other components in blends.^{14,15}

Fig. 6b also shows that the curve for any one polymer will not cross that of another, so the ranking between polymers persists over the entire temperature range. Comparing polymers among our set, we see that PVAc has the highest $\% V_{\rm free}$, followed by PBMA and PVME; at the other end we have P α MS, with the lowermost value.

Related to this, P α MS has the highest T_g , whereas PVAc, PBMA, and PVME have the lowest T_g s among the set. We have noted in previous work¹³ that, comparing polymers at any

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single temperature, those with the lowest free volume have a tendency, on average, to be the ones with the highest T_g value and *vice versa*. The glass transition is a kinetic event where a material's molecular level dynamics slow down to the point that they are effectively slower than the practical experimental observation time scale. A polymer that tends to have more free space at any given *T* will tend to require less cooperativity for segmental motion, leading to lower activation barriers compared to other polymers. This serves to delay the glassy dynamic slowdown as *T* is further decreased. The connection between $V_{\rm free}$, cooperativity, and activation energy is discussed in detail in ref. 16 and 17.

In addition to comparing polymers at a common temperature, it is also very revealing to examine the trend in %V_{free} across the series when calculated at each polymer's corresponding experimental $T = T_g$. In Fig. 6b, these $\% V_{\text{free}}(T = T_g)$ values are marked as the large points, one on each polymer's $%V_{\text{free}}(T)$ curve. (Note the portion of any curve at temperatures below the experimental $T_{\rm g}$ would thus correspond to the theoretical extrapolation of the equilibrium melt.) These points, determined by applying our LCL model through ellipsometric characterization of each polymer, show the same striking linear trend we uncovered in earlier work¹³ for a large set of 50 polymers via analysis of their PVT data. Here, both the slope and intercept of the best-fit line drawn through the points are in excellent agreement with those of the line found from that previous PVT-based analysis, e.g. the slopes are 0.034 °C⁻¹ and 0.033 $^{\circ}C^{-1}$ respectively.

Note that, while this is a strong correlation, it will not be "exact" because the glass transition temperatures themselves are not exact. The kinetic nature of the transition is reflected in the fact that the observed $T_{\rm g}$ point depends on the experimental timescale of observation (*i.e.* the heating or cooling rate), and on the experimental technique. It is common for reported $T_{\rm g}$ values to vary by ± 5 °C (or more). However, this is still reproducible enough for our purposes. The range of system $T_{\rm g}$ values considered here spans hundreds of degrees; if one were to hypothetically alter each of the values randomly by ± 5 °C, it would only have a small effect on the slope of the correlation line; none of the main conclusions would be affected.

The physical significance of the $\% V_{\text{free}}(T = T_g)$ vs. $T = T_g$ correlation line in Fig. 6 is that it marks the boundary between the melt regime (above) and glass regime (below). In other words, following along any particular melt's $\% V_{\text{free}}(T)$ curve from high *T* as *T* decreases, the curve will intersect this boundary, and that intersection temperature is where, on average, the LCL theory predicts the polymer will become glassy. For a more complete description of the melt/glass boundary, see the discussion of Fig. 3 in ref. 13. In addition, we note explicitly that the correlation line in Fig. 6 demonstrates that the %free volume of a polymer at its T_g varies considerably, depending on its nature. This is counter to the historical view of free volume at T_g having a universal constant value, based on models^{55–57} that incorrectly assumed that polymer dynamics follow the Doolittle equation.⁵⁸

 $T_{\rm g}$ is an interesting quantity, a thermodynamic signature that reflects one aspect of dynamic response. The results above, *i.e.* that V_{free} is not simply a constant at T_{g} , but rather carries with it a T-dependence, demonstrate that free volume plays an important role in dynamics but is not the only important variable. We have used our model for dynamic relaxation of melt materials and shown for many systems that segmental relaxation times (τ) follow a general form $\ln \tau \sim f(T) \times (1/V_{\text{free}})$.^{16,17} In other words, independent contributions to dynamics, both from temperature and specific volume, are important in understanding this material response. This form also reflects our discovery that the volume-dependent contribution is most effectively and analytically captured by using the free volume, not the total volume. Thus, we have found an essential role for V_{free} , which can only be independently evaluated using thermodynamic information. Some alternative approaches can only operate by extracting V_{free} values from dynamic data, however, that precludes using the results to understand dynamic response, since such a route would be circular. With the view that V_{free} results can be illuminating for deeper understanding of both thermodynamic and dynamic properties, in this work we have extended this characterization route to the convenient ellipsometry approach.

4. Summary and conclusions

This work advances the soft matter community's ability to link material dynamic phenomena with thermodynamic data. Firstly, we have demonstrated spectroscopic ellipsometry as an ideal method to obtain vital thermodynamic information *via* the analysis of a material in the form of a thin film. Secondly, we combined the new results with thermodynamic analysis using the Locally Correlated Lattice (LCL) model to calculate the temperature dependence of the percent free volume, which is a quantity that has been used to explain trends in both miscibility and local dynamic relaxation, among other behavior.^{13,15,17} This is the first application of ellipsometry to find $\%V_{\text{free}}(T)$ in this way.

Ellipsometry offers distinct advantages over other techniques. In contrast to many dilatometric approaches, only a very small quantity of material is required for analysis, such as a thin film over a square centimeter (having a mass <0.1 mg). In experiments using bespoke deuterium-labelled molecules, often only small quantities are available.⁴⁹ In contrast to other thin film analysis techniques, such as neutron or X-ray reflectivity,^{21,25} ellipsometry is often readily available within a laboratory and relatively inexpensive.

Our ellipsometric characterization of ten different polymer films on a substrate with a low thermal expansion coefficient demonstrates conclusively that the measurement of α_N in the melt state is equal to the corresponding α_V . However, in the glass state, a value of the Poisson ratio of the glass is needed to recover α_V from a measurement of α_N *via* ellipsometry. Our new data strengthen previous LCL observations that the percentage of free volume at the T_g varies between 5% and 12%, depending on the particular polymer, and increases linearly with the T_g itself. Ellipsometry has been used widely in the literature for decades to characterize solids and liquids. However, this work is the first to provide comprehensive and conclusive data testing the relationship between α_N and α_V for both the glass and melt states and thereafter obtaining the percent free volume as a function of temperature. Until now, the full usefulness of ellipsometry had not been exploited. Our research has thereby opened up the possibility of reliable essential thermodynamic characterization using this commonly available, straightforward technique, even when there is a small quantity of sample. We envisage that our methods will in the future underpin molecular design and synthesis to achieve target properties.

5. Materials and methods

5.1 Materials

The polymers, which are listed in Table 1, were obtained from either Sigma-Aldrich (Gillingham, Dorset, UK) or Polymer Laboratories (Church Stretton, Shropshire, UK). Laboratory reagent-grade, anhydrous toluene with low sulfur (Sigma-Aldrich) was used as the solvent to prepare solutions (typically 5 wt%). The only exception is that tetrahydrofuran (Sigma-Aldrich) was used as the solvent for PSAN. No unexpected or unusually high safety hazards were encountered. Pieces of single-side polished silicon (100) wafers (PI-KEM Tamworth, UK) were used as the substrate. Wafers were cut into pieces with typical dimensions of 2 cm \times 2 cm.

5.2 Sample preparation

The Si substrates were placed into a UV ozone cleaner (Bioforce Nanosciences, Inc.) for 15 min to remove any organic contaminants. Thin films were spin-cast onto the substrates from 0.5 mL of the polymer solutions using a photoresist spinner (PWM32 Series, Headway Research, Garland, Texas, USA). Samples were spun at 2000 rotations per min for 30 s. The thin films were then annealed in a vacuum oven (Technico) at 50 °C above the specific polymer's literature value for its $T_{\rm g}$ (Table 1) to remove residual solvent and to allow structural relaxation. Films were then cooled naturally while still under vacuum at a rate of approximately 0.25 °C min⁻¹.

5.3 Spectroscopic ellipsometry measurements

Ellipsometry analyzes the change in the state of the polarization of light upon reflection from interfaces. In our experiments, the thickness and refractive index of films were determined as a function of temperature using a variableangle spectroscopic ellipsometer (V-VASE, J. A. Woollam Co., Lincoln, NE, USA). The films were secured to a vertical heating stage (HS-190) that was controlled *via* software (WVASE, J. A. Woollam Co.). A thermally-insulated cover (HTC-100 HeatCell) was placed over the vertical heating stage to stabilize and maintain the temperature of the sample holder. The cover has two ports on either side to allow light into and out of the sample cell, which contained the ambient atmosphere. Light from a 75 W Xe lamp passes through a linear polarizer before reflecting from the sample.

Two different ellipsometry techniques were used: dynamic scans and step-wise scans. In a dynamic scan, four wavelengths of light (in the range between 400 nm and 800 nm) were selected. Data (consisting of ψ , Δ pairs) were collected over time consecutively at each of these wavelengths using an angle-of-incidence of 70° (measured from the normal to the plane of the substrate) as the sample was heated at a constant rate of 2.5 °C min⁻¹.

In a step-wise scan, the temperature of the thin film was increased from room temperature in 5 °C increments using a heating rate of 2.5 °C min⁻¹. The sample was allowed to equilibrate for 30 s before a spectroscopic scan commenced. Spectroscopic ellipsometry spectra were obtained at each temperature over the visible wavelength range from 400 nm to 800 nm, in increments of 10 nm, at an incident angle of 70°.

5.4 Ellipsometry data analysis

To determine a thin film's *h* and *n* at each temperature, a model was fit to the data using commercial software (WVASE). The model for the analysis, which is presented in Table 6, consisted of a substrate, thermal oxide layer (SiO₂), and a polymer film layer. Complex refractive index values, $\tilde{n}(\lambda)$ for the Si for a given temperature were obtained from the WVASE software. Preliminary ellipsometry measurements on uncoated Si wafers found that the mean thickness of the native oxide layer was 2.5 nm. This value was used in all of the models. The model used the refractive index values of thermal oxide obtained by Herzinger *et al.*⁵⁹

The refractive index of the polymer layer was modelled using a Cauchy equation of the form:

$$n(\lambda) = A + \frac{B}{\lambda^2} + \frac{C}{\lambda^4} + \dots, \qquad (4)$$

where *A*, *B* and *C* are Cauchy coefficients, and λ is the wavelength in units of μ m. The extinction coefficient, *k*, was set to 0 because the ten polymers are transparent in the visible range of λ . There was some variation in the thickness across the footprint of the light beam. The thickness non-uniformity was included in the model and typically had a value <5%. The unknown parameters were obtained by iteratively reducing the mean-squared error using a Levenberg–Marquardt algorithm

I able 6 Model used for ellipsometry data analysis	
Layer in the model	Varied parameters
 Polymer film Thermal silicon dioxide layer (SiO₂) Silicon substrate 	Cauchy parameters (A, B); $h(T)$; thickness non-uniformity None Refractive index (n, k)

within the WVASE software. Typically, an approximate thickness was first obtained by fitting when using an estimate of A and B. (C was typically set to 0, unless an acceptable fit could not be obtained.) Then in a second step, values of A and B were obtained by fitting simultaneously with h. In a final step, the thickness non-uniformity was fitted while refining the values of h, A and B.

Author contributions

R. P. W.: conceptualization, formal analysis, methodology, visualization, writing original draft; D. B.: methodology, investigation, visualization, writing original draft; A. M. J. M. Beale: investigation; I. G.: investigation; J. L. K.: conceptualization, funding acquisition, supervision, writing – reviewing and editing; J. E. G. Lipson: conceptualization, funding acquisition, supervision, writing – reviewing and editing.

Conflicts of interest

There are no conflicts of interest to declare.

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