Optical Identification of Materials Transformations in Oxide Thin Films

Duncan R. Sutherland, Aine Boyer Connolly, Maximilian Amsler, Ming-Chiang Chang, Katie Rose Gann, Vidit Gupta, Sebastian Ament, Dan Guevarra, John M. Gregoire, Carla P. Gomes, R. Bruce van Dover,* and Michael O. Thompson*

ABSTRACT: Recent advances in high-throughput experimentation for combinatorial studies have accelerated the discovery and analysis of materials across a wide range of compositions and synthesis conditions. However, many of the more powerful characterization methods are limited by speed, cost, availability, and/or resolution. To make efficient use of these methods, there is value in developing approaches for identifying critical compositions and conditions to be used as a priori knowledge for follow-up characterization with high-resolution techniques, such as micrometer-scale synchrotron-based X-ray diffraction (XRD). Here, we demonstrate the use of optical microscopy and reflectance spectroscopy to identify likely phase-change boundaries in thin film libraries. These methods are used to delineate possible metastable phase boundaries following lateral-gradient laser spike annealing (lg-LSA) of oxide materials. The set of boundaries are then compared with definitive determinations of structural transformations obtained using high-resolution XRD. We demonstrate that the optical methods detect more than 95% of the structural transformations in a composition-gradient La-Mn-O library and a Ga2O3 sample, both subject to an extensive set of lg-LSA anneals. Our results provide quantitative support for the value of optically detected transformations as a priori data to guide subsequent structural characterization, ultimately accelerating and enhancing the efficient implementation of micrometer-resolution XRD experiments.

KEYWORDS: laser spike annealing, reflectance spectroscopy, high-throughput experimentation, materials transformations, metastable phases

INTRODUCTION

Combinatorial materials science research employs high-throughput experimentation to rapidly probe and analyze large regions of synthesis and processing space.1,2 The exploration of metastable phases is of particular interest in the context of designing new materials,3 and synthesis of metastable materials requires precise control of transformation kinetics. Structural characterization during metal oxide synthesis has revealed a complex interplay between thermodynamics and kinetics as a function of composition, temperature, and time,4 which is consistent with the general understanding of phase behavior/evolution in materials. The composition–temperature–time synthesis phase space is rich in structure and largely unexplored due to its complexity and the limited ability to process materials on fast time scales. Conventional methods for annealing thin film samples include hot plate, furnace, and rapid thermal annealing,5 but the 1–100 K/s cooling rates available with these techniques are insufficient to kinetically trap many high-temperature metastable phases for room-temperature characterization. Methods for studying metastable materials include characterization of thin films quenched during thin film deposition, both with and without subsequent annealing,6,7 and nanocalorimetric processing that provides quench rates on the order of 104 K/s.8

Recently, lateral gradient laser spike annealing (lg-LSA) has been developed as a high-throughput technique in which thin film samples are annealed in the sub-millisecond9 time domain. The specific Gaussian-like laser beam profile in lg-LSA generates a reproducible spatial temperature gradient across the scan direction, typically ~1 mm in width (fwhm). With an accessible domain of peak temperatures (300–1410 °C) and dwell/quench rates (200 μs to 10 ms), a wide range of anneal conditions can be probed rapidly in a single thin film sample. We have demonstrated that lg-LSA can form and quench metastable phases in functional oxide materials, for example, by stabilizing high-temperature polymorphs under ambient...
conditions.9 Due to the large thermal gradient across an lg-LSA stripe, micrometer-scale spatial resolution is required to analyze the processed thin film properties. Structural characterization using X-ray diffraction (XRD) allows the identification of the crystalline phases present in addition to providing information on the overall degree of crystallinity.10 However, typical lab-source diffractometers are unsuitable for resolving structural transformations across an lg-LSA stripe in a high-throughput fashion since they (a) produce spot sizes on the order of 1 mm or larger and (b) provide insufficient X-ray intensity to account for the weak scattering of thin films, demanding long integration times. Using slits to improve spatial resolution of the X-ray source further degrades the throughput. On the other hand, synchrotron facilities produce spatial resolution of the X-ray source further degrades the throughput. On the other hand, synchrotron facilities produce features that are readily available, our methods can speed up the iterative feedback cycle between characterization and synthesis/processing by orders of magnitude.

In Methods we discuss the two spatially resolved optical analysis methods developed in this work in detail, particularly in Spatially Resolved Techniques. The application of these methods to two materials systems is discussed in the Results and Discussion, which are then assessed for accuracy using two metrics in Absolute Gradient Sum and Completeness of the Materials Transformations. Finally, we summarize our results in Conclusions.

■ METHODS

Thin Film Deposition and lg-LSA Processing. Thin film libraries were deposited onto thermally oxidized (20 nm), highly doped (p-type, 0.02 Ω cm), lithographically patterned Si wafers with gold alignment marks. The Ga2O3 sample was deposited via reactive sputtering from a Ga2O3 target in an atmosphere of 4.5 mTorr Ar and 0.5 mTorr O2 in an AJA sputter system. The target was operated at an RF power of 210 W while the substrate was rotated to create a 160 nm thick film with a <5% thickness variation.

The La-Mn-O system was deposited by reactive cosputtering from La and Mn targets offset to generate a lateral composition gradient using a custom deposition system described previously.11 In an atmosphere of 5.4 mTorr Ar and 0.6 mTorr O2, the La and Mn targets were energized with RF powers of 83 and 140 W, respectively. XRF measurements, calibrated using thin film standards, revealed that the La/(La + Mn) concentration in the as-deposited film varied from 12 to...
78% across the library with a metals loading at the substrate center of 1.4 μmol/cm², corresponding to a thickness of approximately 250 nm.

The beam of a CO₂ laser operating at \( \lambda = 10.6 \) μm and a maximum power of 125 W was shaped to provide a 320 μm wide (fwhm) lateral beam profile and used to generate lg-LSA stripes on the two materials system samples. Each anneal stripe was 5 mm long, with peak temperatures ranging from 600 to 1410 °C and processing dwell times between 250 μs and 10 ms. The dwell is defined as the fwhm of the laser in the scan direction divided by the scan velocity; this is approximately the time the temperature is within 5% of the peak temperature.\(^9\)

The anneal locations were randomized across the thin film library to avoid location bias on the wafer and to evenly distribute the dwell-time and temperature conditions with respect to the composition gradient in the La-Mn-O sample. A 100 mm diameter wafer offers space for a total of up to 625 stripes with distinct anneal conditions.

**Spatially Resolved Techniques. Optical Microscopy.** Optical micrographs were collected using a Thorlabs CMOS camera (RGB channels with 1024 × 1280 pixels) aligned normal to the sample. The sample was illuminated coaxially with white light over a spot size approximately 1 mm in diameter. The RGB color channels were summed using the weights of (0.3, 0.59, 0.11) and (0.33, 0.33, 0.33) for the Ga₂O₃ and La-Mn-O samples, respectively, to create high-contrast grayscale images. The magnification was set to provide a 0.79 μm spacing between pixels in the final image; this produced images with an approximately 1 mm wide horizontal field of view (FOV). A representative RGB micrograph is shown in Figure 1a, which is horizontally cropped to be consistent with the displays associated with spectroscopic and diffraction techniques.

**Reflectance Spectroscopy.** For reflectance spectroscopy, the sample was illuminated with a white light source (wavelength 400 < \( \lambda < 900 \) nm). A seven linear fiber bundle was imaged onto the sample to collect reflected light from an area of approximately 10 μm × 70 μm, aligned along the laser scan direction. The fiber bundle (a) improves the signal to noise ratio and (b) reduces erroneous signals from defects and particles. A reference dark spectrum and a spectrum from an undisturbed beam of Ga₂O₃ were measured for each sample. The fiber image was scanned across laser stripes at 10 μm increments for 201 samples, each with an optimized integration time of ~250 ms. Reflectance traces from three representative scans are shown in Figure 1c.

**X-ray Diffraction.** The data from the two optical techniques were compared to reference data collected using the ID3B beamline at the Cornell High Energy Synchrotron Source (CHESS). A 9.7 keV beam was focused to yield a spot size with a 20 μm × 40 μm footprint on the sample at a 2° angle of incidence. The diffracted signal was captured with a Pilatus 300 K detector. The X-ray beam was scanned along the same path as in the reflectance spectroscopy technique, collecting data every 10 μm across the anneal stripe with a 50 ms integration time. The 2D detector data were integrated along the \( \chi \) direction using pyFAI.\(^12\)

**Analysis.** Systematic data processing protocols were developed in order to compare the different techniques. Since they vary in their spatial extent and resolution, a key step involved carefully aligning the resulting data maps across the techniques to allow a quantitative comparison. The optical micrographs were cropped to conform to the dimensions of the region probed by reflectance spectroscopy. Each map was smoothed with the same spatial degree of filtering to decrease noise. An unsharp mask filter was applied to the X-ray diffraction maps to enhance sharpness and emphasize featuring.

### RESULTS AND DISCUSSION

To demonstrate the value of the novel optical methods presented in subsection Optical Microscopy and Reflectance Spectroscopy, we apply them to study phase transitions of (meta)stable phases in two materials systems: (a) single-composition Ga₂O₃ and (b) composition-gradient La-Mn-O.

Ga₂O₃ has four known accessible crystalline polymorphs, \( \alpha, \beta, \gamma, \) and \( \epsilon \), with distinct temperature regimes of thermodynamic stability, allowing a clear interpretation of the phase space regions. \( \beta \)-Ga₂O₃ is of particular interest due to its commercial application in electronic and power devices as a wide-band-gap semiconductor.\(^16\) The \( \alpha \)- and \( \epsilon \)-Ga₂O₃ polymorphs have, for example, applications in functional heterostructures, as transparent conductive oxides, and exhibit tunable band gaps.\(^14,15\)

The La-Mn-O system has been the subject of experimental interest for many years due to its application in solid oxide fuel cells\(^13\) and the colossal magnetoresistance found in lanthanum manganite perovskites.\(^18,19\) The equilibrium pseudobinary phase diagram in air was reviewed by Grundy et al.,\(^17\) who identified four polymorphs of MnO₃. Two polymorphs of La₂O₃ and the perovskite LaMnO₃. However, nonequilibrium metastable phases of the system have not been studied extensively. The richness of the chemical system along with the interest in potential metastable phases makes it an excellent candidate for a combinatorial high-throughput study.

**Absolute Gradient Sum.** We commence our analysis of the data arising from the three different experimental sources by deriving key quantities that can be compared across the sampling techniques. The images and heat maps, as described in Spatially Resolved Techniques, share a common horizontal axis corresponding to the spatial location along the temperature-gradient (lateral) direction of a given lg-LSA stripe. The maps are treated as images, where \( v(x, y) \) is the signal strength at position \( x, y \) of the map, and \( v \) represents either the physical location (microscopy image, reduced to gray scale), the photon energy (spectroscopy), or Q value (XRD). Figure 1 gives a visual comparison of the resulting images/maps for a representative sample from the La-Mn-O library, where we align the \( x \) axis with respect to the center of the lg-LSA stripe. To quantitatively compare the images and maps across the different measuring techniques, we compute the absolute value of the gradients along \( x \) for every row \( i \), \( g(x) = \frac{dy}{dx} \). We then reduce the dimensionality by summing up every column along \( y \) and obtain the absolute gradient sum (AGS) \( a(x) = \sum g(x) \), the key quantity for our further analysis. Note that for the microscopy images there is essentially no variation along the \( y \) axis in the region imaged since the annealing conditions have reached steady state (i.e., along the longitudinal axis). \( a(x) \) is therefore simply a sum over redundant information along that direction.

The AGS represents a measure for changes in the overall signal intensity across a stripe. Regions with a high AGS value, e.g., pronounced peaks, correspond to rapid variations in the underlying measured properties associated with materials transformations. For the two optical methods, the physical
source of these variations in the transparent films stems from varying interference due to changes in the film thickness \( d \) and/or the refractive index \( n \) of the film, and possible diffuse scattering of the light due to formation of a non specular surface. In particular, small changes in either \( n \) or \( d \) markedly alter the optical reflectance of the material. In regions where the phase mixture is uniform, the optical properties are uniform as well. However, when a new phase is introduced, it changes the optical properties, even if slightly. The ability to detect such subtle systematic changes in optical behavior through the AGS allows us to detect changes from one phase or phase mixture to another. Changes in the AGS in the XRD patterns are directly associated with structural changes, including local ordering of the amorphous film and changes in the crystalline phase of the material. A core hypothesis of this work is that structural changes detectable by XRD will also give rise to optical transitions identified by peaks in the gradient signal, and conversely that detected optical transitions will typically correspond to an XRD-detectable structural transition.

Figure 2 shows how the images/maps compare with each other for the same representative lg-LSA stripe as in Figure 1.

![Maps and Gradients](image)

**Figure 2.** Optical micrograph, reflectance spectroscopy, and XRD maps aligned and stacked over the same spatial extent \( x \). The teal vertical lines indicate the peaks in the corresponding AGS shown in the panels on the right.

and includes the corresponding AGS in the right-hand panels. The height and widths of the peaks in the AGS vary from technique to technique due to the sensitivity and spatial resolution of each method. The location of the peaks in the AGS are indicated by vertical teal lines in the signal maps; these align well across the three experimental methods. With the XRD patterns of known LaMnO\(_3\) phases at hand\(^{20,21}\) we can identify the structural transformations in the film and associate them with the processing conditions on the stripe. As the maximum temperature experienced by the film increases from room temperature at the position \( x = -400 \) \( \mu m \) toward the peak temperature of the entire lg-LSA stripe at the position \( x = 0 \) \( \mu m \), the film undergoes two definable transformations (note that there are two definable transformations for this particular lg-LSA stripe, but films can exhibit a fewer or greater number of transformations, varying in nature, depending on the processing parameters and film composition). First, the amorphous as-deposited \( \text{La}_{0.7}\text{Mn}_{0.3}\text{O}_3 \) film begins to change to an optically denser condition at \( x = -250 \) \( \mu m \). Second, the fully dense amorphous film transforms to a crystalline cubic LaMnO\(_3\) structure at \( x = -100 \) \( \mu m \). The symmetrical nature of the lateral thermal gradient leads to the same set of transformations in reverse order on the right-hand half of the stripe.

Ideally, any materials transformation should be captured with all three methods, leading to a 1:1 mapping of the associated AGS. In general the same symmetries and transformations are observed in the optical micrograph and the reflectance spectroscopy map, with AGS peak locations consistent with those observed in the XRD map. To quantify the difference of the AGS across the techniques for all of the experimental conditions, we use the Pearson correlation coefficient (PCC) between each pair of techniques as a measure of the similarity in the overall material transformation characterization. A PCC value of 0 equates to no linear correlation, whereas a value 1 or \(-1\) equates to total positive or negative linear correlation, respectively. For the PCC calculation the AGSs are interpolated with a cubic spline function spanning the same spatial extent. Note that the comparison is sensitive to the difference in spatial resolution of each technique (the effective probe sizes in Figure 1), which can cause a sharp transformation boundary to appear indistinct or could create or obscure the appearance of a multiphase region. Further, inaccurate calibration of the lateral anneal space can compress or expand the AGS and significantly alter the calculated correlation coefficients. Hence, we carefully calibrated our spatial scales to minimize the effect of such artifacts.

The subplots (a)–(c) and (d)–(f) in Figure 3 summarize the PCC values between each pair of techniques (optical to spectroscopy, optical to XRD, and spectroscopy to XRD) for the 612 and 603 LSA conditions analyzed in the Ga\(_2\)O\(_3\) and La-Mn-O samples, respectively. The right-hand panels in each subplot show the scatter of PCC values between each pair of techniques (optical to spectroscopy, optical to XRD, and spectroscopy to XRD) for the 612 and 603 LSA conditions analyzed in the Ga\(_2\)O\(_3\) and La-Mn-O samples, respectively. The left panels in each subplot show the histogram of PCC values between the specified techniques over all LSA conditions. For the Ga\(_2\)O\(_3\) system, the variation in dwell conditions are encoded with different opacities (light and dark for short and long dwells, respectively), while for the La-Mn-O system the color encodes the compositional variation of the La/(La + Mn) atom fraction. The left panels in each subplot show the histogram of the PCC from the corresponding panel on the right. The red horizontal line denotes the median value of the PCC.

The histograms for the Ga\(_2\)O\(_3\) system clearly show a bimodal distribution, which indicates the presence of distinct, systematic features in the PCCs over the investigated LSA conditions. The first mode exhibits a peak with a median of \( \approx 0.1 \), while the peak of the second mode is located at a value of \( \approx 0.7 \), with some variation depending on the compared techniques. A comparison with the corresponding scatter plot shows that the two modes stem from samples that are separated distinctly at a critical temperature \( T_c \approx 600 \) °C that is independent of the anneal time, as observed in Figure 3a–c. We ascribe this sharp increase in the PCC to the onset of crystallization from the amorphous Ga\(_2\)O\(_3\) film, consistent with the findings reported previously.\(^{22}\) Below the threshold value \( T_c \), there is essentially
no variation in the AGS, since the signal stems from purely amorphous samples, leading to correlation values close to zero. Above $T_c$, the crystallization events lead to clear changes in the AGS that correlate strongly across the techniques. Note that the correlations between the spectroscopy and XRD characterization have the tightest distribution for the second mode (above $T_c$), which we attribute to the very similar spatial resolutions of the two techniques.

A similarly sharp onset is neither expected nor observed for the La-Mn-O system due to the increased complexity associated with compositional variation across the wafer. The PCC gradually increases as a function of anneal temperature between 500 and 800 °C, leading to a unimodal, skewed distribution in the histograms. The median value is $\sim$0.6, which is lower than the second mode of the Ga$_2$O$_3$ system but still indicates a strong correlation. Thin films of La$_2$O$_3$ and MnO$_2$ crystallize from amorphous as-deposited films at 600 and 450 °C$^{23,24}$ respectively. While various lanthanum manganate materials have been investigated, they typically incorporate different substitutions$^{25}$ that complicate comparative measurements or are grown on a heated substrate at 900 °C to synthesize as-deposited crystalline films.$^{26}$

Overall, we observe a high degree of correlation (>0.5) in the PCC analysis, which clearly indicates that the optical and spectroscopic measurements are sensitive enough to identify the material transformations observed by XRD. For both materials systems, the strongest correlation is observed when spectroscopy is compared to XRD. This result is not surprising, since these two methods contain a broader spectrum of information (photon energy and diffraction angles) in comparison to microscopy (single value from the three color channels). Nevertheless, microscopy imaging is a simple and powerful method to rapidly assess the AGS features, since the time scale to capture a single RGB image is 2–3 orders of magnitude shorter than that needed for reflectance spectroscopy or XRD (when available).

**Completeness of the Material Transformations.** The PCC serves as a statistical method to assess and compare the AGS but does not reflect whether or not the optical methods are sensitive to a particular type of crystallographic transformation occurring in the LSA processed thin films, i.e., it does not clarify what kind of transformations are identified or missed. To address this issue, we employ a metric complementary to the PCC by analyzing the completeness of the material transformations identified by XRD to those identified by either of the optical techniques. Of particular concern are the cases where crystalline structure changes occur (based on the complete XRD data) without detectable changes in the optical techniques: i.e., “false negatives”. To assess the presence of such cases, we manually compile a list of transformations detected in any or all sets of techniques in a given stripe through visual inspection and identify the false negatives. “False positives” correspond to transitions observed in either of the optical methods corresponding to putative phase transformations that do not appear in the XRD measurements. The completeness of the observed structural transformations is best visualized with a Venn diagram (Figure 4). The subsets represent the portion of the transformations identified by one, two, or all three characterization techniques. The total number of identified transformations, independent of transformation type, within each set are given in Table 1.

In contrast to what one would expect from the PCC analysis, the anneal stripes in the Ga$_2$O$_3$ system display very gradual materials transformations across the stripe that lead to a relatively large false negative rate of approximately 4.3% for each of the optical techniques. This value is substantially higher than for the La-Mn-O system but is low on an absolute scale. A detailed investigation of this subset of false negatives reveals that they occur primarily when the $\gamma$ phase transforms to the $\beta$ phase with increasing temperature. This result confirms that the two optical methodologies may be insensitive to certain types of materials transformation, particularly when it occurs gradually between two phases with similar optical
XRD areas are colored in red, yellow, and blue, respectively. False positive rate, 25.1%, higher than that in Ga₂O₃. In general, such morphological changes, or other defects in the thin film, respectively. The optical microscopy, reflectance spectroscopy, and XRD techniques for the La-Mn-O and Ga₂O₃ samples in the panels a and b, respectively. The optical microscopy, reflectance spectroscopy, and XRD measurements.

For the La-Mn-O system, 0.7% of the structural transformations are demonstrably false negatives (X\(\backslash(S\backslash O)\)); see the tiny blue sliver at the bottom of Figure 4b). Investigation into these false negatives reveals the crystallographic transformations not detected in optical measurements because the optical signals were obscured by laser-induced roughening of the film. Such film damage occurs at the highest temperatures, where the introduction of substantial nonspecular reflection from the roughening lowers the sensitivity to optical transitions from microscopic structural changes. As a result these false negatives do not counter the hypothesis that optical transitions are a useful proxy for structural transitions but rather illustrate a limitation of the technique that the transitions are only well characterized up to a material-dependent maximum temperature and dwell time where macroscopic film damage occurs. For further automation of the analysis workflow, such film damage can be readily detected in the microscopy data. The sensitivities of each optical method applied to the La-Mn-O system are similar to those for Ga₂O₃ with regard to capturing identical material transformations. The La-Mn-O system shows a larger false positive rate, 25.1%, higher than that in Ga₂O₃. In general, such false positives can be attributed to dust, delamination, film morphological changes, or other defects in the thin film that do not pertain to the underlying crystallographic transformations.

As a means for prescreening phase transitions prior to XRD experiments, both optical methods are effective despite the large number of false positives. Such false positives are undesirable because they motivate redundant sampling but they do not lead to any loss of information about crystallographic material transformations. For example, exhaustive XRD sampling would require 125,000 individual measurements in the La-Mn-O composition spread, while focusing only on spectroscopy-detected transitions would reduce the number of required measurements to as few as 960. Thus, the false positive rate of 23% nevertheless enables substantial resource savings in terms of the total number of required XRD measurements.

On the other hand, false negatives may result in loss of information in the synthesis phase diagram and should be minimized. The low false negative rate of approximately 0.7% in the La-Mn-O sample demonstrates the capability to effectively prescreen transition boundaries prior to XRD experiments. The higher false negative rate of 4.3% observed in the Ga₂O₃ sample stems from a single type of phase transformation that is especially difficult to detect optically: i.e., in which the optical absorption and optical thickness (the product of n and d) both remain approximately constant. Such instances are coincidental and are expected to be rare in higher order composition spaces.

**Accelerating Experimental Throughput.** To assess the value of this work in accelerated exploration of materials synthesis spaces, we consider the demonstrated workflow of (i) composition library deposition, (ii) LSA-based processing with approximately 600 LSA stripes (peak temperature and dwell time conditions) per composition library, and (iii) characterization within each LSA stripe to identify transitions in temperature, dwell time, and composition.

The throughput of these processes in terms of number of composition libraries is 1 and 10 per day for deposition and IgLSA, respectively. For transition characterization, the alternate approaches described in the present work — optical microscopy, reflectance spectroscopy, and micrometer-resolution XRD — have throughputs of 1000, 2, and 10 per day, respectively, although the XRD throughput can be substantially lower for thin films that require larger acquisition times due to weak scattering.

With these nominal throughputs, the deposition step appears to be approximately rate limiting until one considers resource availability, which is in the order of 9 days per year for the specialized synchrotron XRD measurements and 300 days per year for the other, purpose-built optical instruments. Limited XRD availability makes this technique rate limiting for annual workflow throughput, and this limitation is exacerbated when the value of feedback is considered — the ability to choose the deposition conditions on the basis of a consideration of all prior data.

Typical synchrotron XRD throughput is up to 90 libraries per year corresponding to 90 traversals of the workflow, although deposition selection informed by characterization of all prior libraries only occurs in up to 9 of those traversals because, during the 9 days of XRD experiments, the decision

<table>
<thead>
<tr>
<th>library</th>
<th>S(\backslash O\backslash U\backslash X)</th>
<th>(S(\backslash O))(\backslash X)</th>
<th>(O(\backslash U))(\backslash S)</th>
<th>(S(\backslash U))(\backslash O)</th>
<th>O(\backslash(S\backslash U))</th>
<th>S(\backslash(O\backslash U\backslash X))</th>
<th>X(\backslash(O\backslash U\backslash S))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ga₂O₃</td>
<td>848</td>
<td>78</td>
<td>5</td>
<td>12</td>
<td>2</td>
<td>18</td>
<td>39</td>
</tr>
<tr>
<td>La-Mn-O</td>
<td>943</td>
<td>279</td>
<td>8</td>
<td>2</td>
<td>30</td>
<td>3</td>
<td>7</td>
</tr>
</tbody>
</table>

**The critical false negatives are X\(\backslash(O\backslash U\backslash S)\), while false positives leading to unnecessary XRD measurements are (S\(\backslash O\))\(\backslash X\).**

---

**Figure 4.** Venn diagrams illustrating the intersections among all three techniques for the La-Mn-O and Ga₂O₃ samples in the panels a and b, respectively. The optical microscopy, reflectance spectroscopy, and XRD areas are colored in red, yellow, and blue, respectively. False negatives are represented by a small blue sliver at the bottom of each Venn diagram.
throughput is limited by the deposition throughput. The ability to detect transitions with optical techniques increases the decision throughput from 9 to 300 per year, enabling optimized allocation of the XRD resource to structural characterization of the optically identified transitions.

This scheme is particularly effective if the XRD-based (structural) transitions are consistently identified with the optical techniques; Figure 4 provides the most extensive demonstration to date that optically detected transitions are most often due to structural transitions. While the throughput of the optical microscopy technique is much higher than that of reflectance spectroscopy, in the context of the annual workflow analysis the increased throughput is not the essential benefit. The value of the microscopy lies in its spatial resolution. The spectral resolution of reflectance spectroscopy facilitates transition detection, but its spatial resolution is limited in practice to \( \sim 5 \) \( \mu \)m. If multiple transitions occur within that length scale (which corresponds to a range in temperature of up to \( \sim 25 \) K for our lg-LSA system), optical microscopy becomes a critical complementary transition detector. Narrowly clustered transitions are not observed in the data presented herein, but the concurrence of the transitions identified by microscopy and spectroscopy indicate that the combination of these techniques provides a robust approach to transition detection in synthesis parameter space.

**CONCLUSIONS**

The methods described here enable high-throughput experimental identification of materials phase transitions across an lg-LSA stripe. Using rapid, low-cost optical techniques such as optical microscopy and spatially resolved reflectance spectroscopy, locations of materials transitions can be identified and may serve as prior knowledge for experimentally expensive techniques such as synchrotron XRD studies. For both single composition and composition-gradient samples, structural transitions were identified via optical methods. An anomalous transition behavior was noted in \( \text{Ga}_2\text{O}_3 \), where a structural change—the introduction and growth of a secondary phase—was not always identified via optical means, demonstrating the possibility of false negatives under certain circumstances. The appearance of unidentified transitions and the common occurrence of optical changes not corresponding to a structural change (false positives) suggest that further development could improve the effectiveness of the methods and extend the properties that can be identified optically. Collectively, our results demonstrate the utility of using in-lab optical characterization methods to accelerate iterative feedback between processing and thin film deposition in the discovery and development of new materials. The results also motivate further development of algorithms that leverage the optical characterization techniques to design synchrotron XRD experiments that enhance the efficient utilization of that limited resource.

**AUTHOR INFORMATION**

**Corresponding Authors**

R. Bruce van Dover – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States; Email: rbv2@cornell.edu

Michael O. Thompson – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States; Email: mot1@cornell.edu

**Authors**

Duncan R. Sutherland – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States

Aine Boyer Connolly – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States

Maximilian Amsler – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States; Department of Chemistry and Biochemistry, University of Bern, CH-3012 Bern, Switzerland; orcid.org/0000-0001-8350-2476

Ming-Chiang Chang – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States

Katie Rose Gann – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States

Vidit Gupta – Department of Materials Science and Engineering, Cornell University, Ithaca, New York 14853, United States

Sebastian Ament – Department of Computer Science, Cornell University, Ithaca, New York 14853, United States

Dan Guevara – Joint Center for Artificial Photosynthesis, California Institute of Technology, Pasadena, California 91125, United States

John M. Gregoire – Joint Center for Artificial Photosynthesis, California Institute of Technology, Pasadena, California 91125, United States; orcid.org/0000-0002-2863-5265

Carla P. Gomes – Department of Computer Science, Cornell University, Ithaca, New York 14853, United States

Complete contact information is available at: https://pubs.acs.org/10.1021/acscmbsci.0c00172

**Author Contributions**

D.R.S. and A.B.C. contributed equally to this work.

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

The authors acknowledge the Air Force Office of Scientific Research for support under award FA9550-18-1-0136. This work is based upon research conducted at the Materials Solutions Network at CHESS (MSN-C), which is supported by the Air Force Research Laboratory under award FA8650-19-2-5220. This work was also performed in part at the Cornell NanoScale Facility, a member of the National Nanotechnology Coordinated Infrastructure (NNCI), which is supported by the National Science Foundation (Grant NNCI-2025233). M.A. acknowledges support from the Swiss National Science Foundation (project P4P4P2-180669).

**REFERENCES**


(20) Sayagüés, M. J.; Córdoba, J. M.; Gotor, F. J. Room temperature mechanosynthesis of the La1−xSr xMnO3δ (0 ≤ x ≤ 1) system and microstructural study. J. Solid State Chem. 2012, 188, 11–16.


