An instrument for in situ time-resolved X-ray imaging and diffraction of laser powder bed fusion additive manufacturing processes

Nicholas P. Calta  
*Lawrence Livermore National Laboratory*

Jenny Wang  
*Lawrence Livermore National Laboratory*

Andrew M. Kiss  
*SLAC National Accelerator Laboratory*

Aiden A. Martin  
*Lawrence Livermore National Laboratory*

Philip J. Depond  
*Lawrence Livermore National Laboratory*

*See next page for additional authors*

Follow this and additional works at: [https://lib.dr.iastate.edu/ameslab_manuscripts](https://lib.dr.iastate.edu/ameslab_manuscripts)

Part of the Manufacturing Commons, Materials Science and Engineering Commons, and the Physics Commons

**Recommended Citation**

Calta, Nicholas P.; Wang, Jenny; Kiss, Andrew M.; Martin, Aiden A.; Depond, Philip J.; Guss, Gabriel M.; Thampy, Vivek; Fong, Anthony Y.; Nelson Weker, Johanna; Stone, Kevin H.; Tassone, Christopher J.; Kramer, Matthew J.; Toney, Michael F.; Van Buuren, Anthony; and Matthews, Manyalibo J., "An instrument for in situ time-resolved X-ray imaging and diffraction of laser powder bed fusion additive manufacturing processes" (2018). *Ames Laboratory Accepted Manuscripts*. 427.  
[https://lib.dr.iastate.edu/ameslab_manuscripts/427](https://lib.dr.iastate.edu/ameslab_manuscripts/427)

This Article is brought to you for free and open access by the Ames Laboratory at Iowa State University Digital Repository. It has been accepted for inclusion in Ames Laboratory Accepted Manuscripts by an authorized administrator of Iowa State University Digital Repository. For more information, please contact digirep@iastate.edu.
An instrument for in situ time-resolved X-ray imaging and diffraction of laser powder bed fusion additive manufacturing processes

Abstract

In situ X-ray-based measurements of the laser powder bed fusion (LPBF) additive manufacturing process produce unique data for model validation and improved process understanding. Synchrotron X-ray imaging and diffraction provide high resolution, bulk sensitive information with sufficient sampling rates to probe melt pool dynamics as well as phase and microstructure evolution. Here, we describe a laboratory-scale LPBF test bed designed to accommodate diffraction and imaging experiments at a synchrotron X-ray source during LPBF operation. We also present experimental results using Ti-6Al-4V, a widely used aerospace alloy, as a model system. Both imaging and diffraction experiments were carried out at the Stanford Synchrotron Radiation Lightsource. Melt pool dynamics were imaged at frame rates up to 4 kHz with a ∼1.1 μm effective pixel size and revealed the formation of keyhole pores along the melt track due to vapor recoil forces. Diffraction experiments at sampling rates of 1 kHz captured phase evolution and lattice contraction during the rapid cooling present in LPBF within a ∼50 × 100 μm area. We also discuss the utility of these measurements for model validation and process improvement.

Disciplines

Manufacturing | Materials Science and Engineering | Physics

Authors


This article is available at Iowa State University Digital Repository: https://lib.dr.iastate.edu/ameslab_manuscripts/427
An instrument for *in situ* time-resolved X-ray imaging and diffraction of laser powder bed fusion additive manufacturing processes

Cite as: Rev. Sci. Instrum. 89, 055101 (2018); https://doi.org/10.1063/1.5017236
Submitted: 26 November 2017 . Accepted: 09 April 2018 . Published Online: 01 May 2018


ARTICLES YOU MAY BE INTERESTED IN

Laser powder bed fusion additive manufacturing of metals; physics, computational, and materials challenges
Applied Physics Reviews 2, 041304 (2015); https://doi.org/10.1063/1.4937809

Direct measurements of laser absorptivity during metal melt pool formation associated with powder bed fusion additive manufacturing processes
Journal of Laser Applications 30, 032302 (2018); https://doi.org/10.2351/1.5040636

Review of selective laser melting: Materials and applications
Applied Physics Reviews 2, 041101 (2015); https://doi.org/10.1063/1.4935926
An instrument for in situ time-resolved X-ray imaging and diffraction of laser powder bed fusion additive manufacturing processes

Nicholas P. Calta,1 Jenny Wang,1 Andrew M. Kiss,2 Aiden A. Martin,1 Philip J. Depond,1 Gabriel M. Guss,3 Vivek Thampy,2 Anthony Y. Fong,2 Johanna Nelson Weker,2 Kevin H. Stone,2 Christopher J. Tassone,2 Matthew J. Kramer,4 Michael F. Toney,2 Anthony Van Buuren,1 and Manyalibo J. Matthews1

1Physical and Life Sciences Directorate, Lawrence Livermore National Laboratory, Livermore, California 94550, USA
2Stanford Synchrotron Radiation Lightsource, SLAC National Accelerator Laboratory, Menlo Park, California 94025, USA
3Engineering Directorate, Lawrence Livermore National Laboratory, Livermore, California 94550, USA
4Division of Materials Science and Engineering, Ames Laboratory, Iowa State University, Ames, Iowa 50011, USA

(Received 26 November 2017; accepted 9 April 2018; published online 1 May 2018)

In situ X-ray-based measurements of the laser powder bed fusion (LPBF) additive manufacturing process produce unique data for model validation and improved process understanding. Synchrotron X-ray imaging and diffraction provide high resolution, bulk sensitive information with sufficient sampling rates to probe melt pool dynamics as well as phase and microstructure evolution. Here, we describe a laboratory-scale LPBF test bed designed to accommodate diffraction and imaging experiments at a synchrotron X-ray source during LPBF operation. We also present experimental results using Ti-6Al-4V, a widely used aerospace alloy, as a model system. Both imaging and diffraction experiments were carried out at the Stanford Synchrotron Radiation Lightsource. Melt pool dynamics were imaged at frame rates up to 4 kHz with a ∼1.1 μm effective pixel size and revealed the formation of keyhole pores along the melt track due to vapor recoil forces. Diffraction experiments at sampling rates of 1 kHz captured phase evolution and lattice contraction during the rapid cooling present in LPBF within a ∼50 × 100 μm area. We also discuss the utility of these measurements for model validation and process improvement. © 2018 Author(s). All article content, except where otherwise noted, is licensed under a Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).

https://doi.org/10.1063/1.5017236

INTRODUCTION

Laser Powder Bed Fusion (LPBF), also known as Selective Laser Melting or Laser Beam Melting, is a rapidly developing additive manufacturing technology that provides significant design flexibility relative to conventional manufacturing techniques and enables the production of highly complex parts at minimal added cost for low-volume production.1 In a LPBF process, a high power (~100’s of W) continuous wave (CW) laser selectively scans over a thin metal powder layer, generating a melt pool that rapidly solidifies to create a two-dimensional solid layer adhered to the substrate or part beneath it. After each solid, patterned layer is created, the part is lowered, and a new powder layer is spread over the part. The process is then repeated to build a fully three-dimensional part in a layer-by-layer fashion. The significant differences between this process and more established manufacturing techniques, such as casting and forging, lead to different mechanical properties for parts built with LPBF when compared with wrought or cast material.2,3 One approach to understanding and predicting these differences is to carry out time-resolved in situ experiments at the melt pool scale (~10’s of μm) to improve understanding of the fundamental mechanisms that govern the LPBF process. Information gleaned from in situ experiments coupled with ex situ part inspection can inform process models, reduce process development time and costs, and improve part quality by identifying laser scan parameters optimized to minimize defect formation or produce a desired microstructure.

A large body of literature exists, which has focused on in situ process monitoring of LPBF to improve process outcomes in an attempt to understand the process and validate models.4 Recently, high-speed visible imaging has been used to elucidate the evolution of melt pool morphology and droplet movement in and around the melt pool during a build.5–8 For example, Ly et al. resolved the origins of hot droplet ejection or spatter emanating from the melt pool region during LPBF.9 Bidare et al. used Schlieren imaging to understand the interaction of the laser plume with process gas in the build chamber.10 In addition to visible light-illuminated imaging, thermal emission imaging has been used to measure the surface temperature of the melt pool11–14 or the entire build.15 Melt pool morphology has also been investigated via in situ measurements of the pool depth using inline coherent imaging, an interferometry-based technique.16 While these approaches based on optical methods provide important information about the dynamics of the LPBF process, they are limited to surface imaging only and cannot provide information about bulk material behavior. In contrast, X-ray-based probes are highly penetrating and can non-destructively...
capture melt pool dynamics in the bulk as well as provide structural information through diffraction. Taken together, high-speed optical, thermal, and X-ray diagnostics can provide highly complementary information about process dynamics during LPBF.

Many of the rapid solidification phenomena relevant to LPBF are quite similar to those present in welding. Elmer and co-workers carried out numerous X-ray diffraction experiments during welding and quantified the dynamics of solidification and solid-state phase transitions on cooling for Ti alloys 17-19 and stainless steels. 20-22 Yonemura et al. also used time-resolved synchrotron X-ray diffraction to study the dynamics of solidification during welding of stainless steel. 23 While this prior work provides important context for understanding solidification and phase transitions in LPBF, the time scales relevant for welding are much longer than the dynamics of LPBF. Time resolution of 100 ms is sufficient to resolve cooling dynamics in diffraction experiments of Ti-6Al-4V (Ti-64) welding, 17 while cooling in LPBF is expected to occur on time scales of a few milliseconds. 24 Therefore, higher sampling rates are required to completely elucidate the dynamics of the laser-material interaction in LPBF. Zhao et al. have reported high-speed X-ray imaging and diffraction of the interaction between a laser and Ti-64 powder under conditions similar to LPBF. 25 Although their report achieved high temporal resolution (20 µs), which provides valuable insight into the dynamics of the laser-powder interaction, Zhao et al. used a stationary laser spot and so were unable to probe dynamics related to laser scanning present in LPBF. Kenel et al. performed in situ X-ray microdiffraction of rapid solidification in Ti-64 under well-defined cooling conditions with 1 ms time resolution, providing additional insight into the fundamental solidification behavior of this alloy under LPBF-like conditions. 26 They performed cyclic heating and cooling to elucidate microstructural evolution induced by thermal behavior similar to what occurs in a multi-layer build, though the thermal boundary conditions in their experiment represent a somewhat different case than what is present in an LPBF build. In this article, we report a laser melting system designed and built to simulate LPBF conditions of a commercial machine while also accommodating instrumentation to enable high temporal and spatial resolution in situ X-ray probes. We also report initial X-ray imaging and diffraction experiments using this instrument at the Stanford Synchrotron Radiation Lightsource (SSRL).

INSTRUMENT DESIGN

Figure 1 summarizes the experimental approach used for in situ X-ray imaging and diffraction during LPBF. LPBF is a complex process that is not perfectly reproducible in its details, and the stochastic nature of the powder and melt dynamics do not allow for the exact stitching of sequential experiments. Therefore, continuous data collection of a single event is required to gain insight into the process. This precludes most averaging approaches, and the ultimate temporal resolution is limited by the brilliance of the X-ray source.

Figure 2 shows the detailed design and photographs of the LPBF system. The system design mimics the conditions present in a typical LPBF build while still permitting sufficient X-ray transmission to probe the area in and around the melt pool at high sampling rates. A single-mode, 1070 nm, 500 W, continuous wave (CW) fiber laser (IPG Photonics, Oxford, MA, USA, YLR-500-WC-Y14) is coupled directly to a 3-axis galvanometer scanning mirror system (Nutfield Technology, Hudson, NH, USA, 3XB 3-Axis Scan Head). The scan head focuses the process laser onto the substrate surface and steers the laser across the substrate to create the moving melt pool required for the LPBF process. The output from the scan head is directed through an anti-reflective coated window affixed to a 2.75 ConFlat (CF) port into a 168.9 mm internal diameter, 60.96-mm-wide spherical octagon vacuum chamber with eight equally spaced radial 2.75 CF ports between two larger parallel 8 CF ports (Kimball Physics, Wilton, NH, USA, MCF800-SphOct-G2C8). Unused ports around the chamber sides allow for future incorporation of secondary probes to monitor the process with more traditional optical techniques simultaneously with X-ray probes. The two large vacuum ports are sealed with 500-µm-thick, 142.2 mm diameter Be apertures to allow X-ray transmission. The narrow chamber width minimizes the X-ray path length inside the Ar-filled chamber to reduce background signal due to scattering. The process laser is focused to ∼50 µm diameter (D40) circular Gaussian beam shape at the sample surface [Fig. 2(c)]. The vacuum chamber is evacuated by a diaphragm/turbo molecular pump system (Pfeiffer Vacuum, Nashua, NH, USA, HiCube 80), and Ar gas is admitted by a flow control valve (MKS Instruments, Andover, MA, USA, 248A) to allow for control over the atmosphere of the experiments. An Ar gas environment is used as a non-reactive environment for all the experiments described here. A gas-species-independent pressure transducer (MKS Instruments, 910 DualTrans) is used for pressure monitoring. The system is a light-tight, portable, Class 1 enclosed laser system, permitting convenient transportation to various beamlines at SSRL optimized for either X-ray imaging or diffraction. In a typical experiment, the laser power can be set between 20 and 500 W nominal power and scanned over the substrate surface at any scanning rate between 0 and ∼2000 mm/s.

The sample substrate and holder are shown in Fig. 3. Sample design must satisfy two competing constraints: it must
be thin enough to permit sufficient X-ray transmission for adequate signal to noise for both imaging and diffraction in transmission geometry while also being thick enough to reasonably mimic the thermal boundary conditions experienced in a true LPBF environment. A suitable compromise geometry to satisfy these two requirements is a thin substrate plate sandwiched between two 1-mm-thick glassy carbon windows (Alfa Aesar, Tewksbury, MA, USA). In this geometry, the laser scans parallel to the windows while the X-ray beam is perpendicular to the substrate. Thick substrates provide insufficient transmission for imaging and diffraction at the kHz rates required to resolve process dynamics. However, a thin substrate enforces thermal boundary conditions that differ from the bulk conditions present in a normal LPBF build. The substrate width chosen to balance these competing requirements in the experiments described here is 500 \( \mu \)m, although the sample holder allows for an adjustable substrate width between 0 and 2 mm.

To investigate the effect of thermal boundary conditions on the temperature profiles present in these experiments, we used both an analytical estimate of the thermal diffusion length as well as more comprehensive finite element calculations of thermal transport. The thermal diffusion length in a material being heated by a Gaussian heat source can be written as:

\[
L = \sqrt{\frac{4}{D} \tau},
\]

where \( L \) represents thermal diffusion length, \( D \) is thermal diffusivity, and \( \tau \) is thermal diffusion time.\(^{27} \)

For a scanning laser beam, which we treat as a scanning Gaussian heat source, the steady-state thermal diffusion length during the interaction time can be written as:

\[
L = \sqrt{\frac{2aD}{u}},
\]

where \( a \) represents the \( 1/e^2 \) diameter of the laser beam and \( u \) represents scanning velocity.\(^{28,29} \)

For this estimate, we use \( a = 50 \mu m \), the measured beam diameter in our system; \( u = 144 \) mm s\(^{-1} \), the scan speed used for the imaging data reported here; and \( D = 5.6 \times 10^{-6} \) m\(^2\) s\(^{-1} \), the reported value for Ti-64 at 1500 K.\(^{30} \)

Based on these estimates, \( L \approx 62 \) \( \mu \)m, significantly smaller than 250 \( \mu \)m, the distance between the heat source and the glassy carbon thermal boundary condition. The primary limitation of this estimate is that it uses a single value for \( D \), which varies as a function of temperature. To understand the influence of thermal boundary conditions beyond the simple estimate presented above, a finite element model implemented
in COMSOL calculated the thermal transport resulting from the above laser scan parameters in a 500-µm-thick Ti-64 bare substrate. Two systems were compared: a case in which the substrate was supported between two glassy carbon windows, as in the experiments described here, and a case where the two windows were composed of Ti-64 to mimic a bulk LPBF case. Both cases used radiative boundary conditions with $\varepsilon = 0.6$.

The geometry of the model is shown in Fig. 4(a). The measured intensity of the laser spot is shown in Fig. 4(b) on the same scale. The model used temperature-dependent properties of Ti-64 obtained from Boivineau et al. and room temperature density and thermal conductivity for glassy carbon provided by Alfa Aesar, with glassy carbon heat capacity obtained from two literature sources. Optical constants for Ti metal were taken from Johnson and Christy. In addition to heat transfer in the liquid and solid, the model includes evaporative and radiative losses based on kinetic theory accounting for latent heat of vaporization and heat flux. Temperature-dependent vapor pressure and heat of vaporization values for elemental Ti were used as an approximation of the value for the Ti-64 alloy. It neglects Marangoni flow in the melt pool, so thermal gradients within the melt pool will not be accurately calculated, but this should not affect the accuracy of cooling in the solid, far from the melt pool. A two-dimensional thermal profile within the Ti-64 substrate at the laser location is shown in Fig. 4(c). This thermal profile was extracted from the case using glassy carbon windows. The temperature at the glassy carbon windows is below 300 K, suggesting no effect of boundary conditions on melt pool dynamics. Compared thermal profiles of the two cases are shown in Figs. 4(d) and 4(e). The difference between the thermal profiles in the two cases is negligibly small, and we therefore conclude that the thermal boundary conditions of the glassy carbon windows do not substantially affect the melt pool dynamics under these conditions for the case of a bare plate. Heat conduction through the metal plate is at least 1 order of magnitude larger than conduction through the powder layer, so conduction through the plate should dominate in the powder case as well.

We further note that for a laser spot size of $\sim 50$ µm, a melt pool width of $\sim 150$ µm is expected for scan speeds and laser powers typical of the LPBF process, which leaves $> 150$ µm of unmelted material between the melt pool and the glassy carbon windows. As the solidified material approaches room temperature during cooling, for example, in the heat-affected zone (HAZ) probed during diffraction experiments, the effect of thermal boundary conditions imposed by glassy carbon windows will become more pronounced. Therefore, any analysis of lattice dynamics during cooling using diffraction must include thorough modeling to understand the influence of thermal boundary conditions on the measured diffraction patterns.

The glassy carbon windows are supported independent of the substrate plate, allowing an adjustable height differential between the glassy carbon windows and the substrate. The height of the glassy carbon window above the substrate determines the thickness of the powder layer, which was nominally set to $60 \pm 20$ µm. The 500-µm-thick Ti-64 substrates were
machined from sheet (TMS Titanium, Poway, CA, USA). The Ti-64 powder (Additive Metal Alloys, Maumee, OH, USA) had a particle diameter of $30 \pm 10 \, \mu m$. This powder layer was manually applied, but an automated spreader is being developed to improve layer uniformity and reduce spreading time. The entire sample holder assembly is supported by three encoded piezoelectric stages (Attocube Systems AG, Munich, Germany), providing motion in the two directions normal to the X-ray beam (vertically and horizontally) as well as one tilt axis.

The practical limit on temporal resolution at the two beamlines used in this manuscript is overall X-ray flux. With the imaging detection scheme described here, signal-to-noise ratios dropped to unacceptable levels at exposure times shorter than $\sim 250 \, \mu s$, corresponding to 4 kHz imaging. During normal user operations, X-ray pulses at SSRL arrive in bunches of 4 spaced 30 ns apart. Since the time scale of the pulse bunches is 4 orders of magnitude smaller than the shortest exposure time we use in this report, we treat the X-ray source as continuous for the purpose of timing and synchronization. In a typical experiment, the sample is continuously illuminated by the X-ray beam, and a control computer sends a trigger to a delay generator (DG645, Stanford Research Systems, Sunnyvale, CA, USA), which sends a trigger to the scan head controller followed by a trigger to the camera to initiate data recording at a variable delay time after the mirrors begin to scan. The scan head controller then triggers the laser to turn on a time delay empirically optimized to account for finite mirror acceleration.

**X-RAY IMAGING**

X-ray imaging experiments were carried out at SSRL beamline 2-2. The X-rays available at this beamline are generated by a 1.25 T bend magnet with a critical energy of 7.4 keV. Figure 5(a) shows the calculated X-ray spectrum produced by this bend magnet. A 6.35-mm-thick Al plate was used as an X-ray filter upstream from the sample to remove low-energy X-rays and protect the detection system from damage by the full emission of the bend magnet source. The in situ imaging experiments were conducted using this filtered X-ray spectrum to maximize high-energy X-ray flux.

X-rays transmitted through the sample and chamber are recorded using a high-speed imaging setup. The transmitted X-rays first hit a Ti-doped NaI scintillator crystal positioned directly behind the exit Be window of the chamber, which converts the X-ray photons to visible light. This visible light is then redirected by a silver-coated mirror at 45° and collected by an infinity-corrected tube lens (Thorlabs, Inc., Newton, NJ, USA) coupled to a pco.dimax S4 CMOS recording camera (PCO AG, Kelheim, Germany). The recording camera features a 12-bit, 2016 × 2016 sensor with 11 µm square pixels. It stores images locally during data collection, permitting a maximum frame rate of 1.2 kHz at full resolution and up to 27 kHz with a 480 × 240 region of interest. Each image is flat-field reference-corrected and converted from transmission to absorption using Beer’s law. This detection scheme yields an effective pixel size of 1.1 µm and a 2.2 × 2.2 mm field of view when using the full sensor.

Vapor depression depth and shape are very difficult to measure with surface-sensitive optical techniques but are relatively straightforward to determine using X-ray imaging because of the significant density contrast between the solid/liquid Ti-64 and Ar gas present in the vapor depression. Furthermore, the perspective of a 2D projection perpendicular to both the laser beam and its path is an extremely useful one for measuring depression depth and shape. These measurements are therefore of high value for model validation. Melt pool depth information can be readily extracted from ex situ sectioning and metallography, but such experiments provide information only about the total depth of the melt,

![Figure 5](image-url)

FIG. 5. X-ray imaging of a typical melt pool region. (a) Simulated X-ray spectra produced at SSRL beamline 2-2, used for imaging. Both the full emission from the bend magnet and the attenuated spectrum experienced at the sample are shown. (b) A series of frames collected at 4 kHz during a scan. Each frame shows the absorption difference $A(t) - A(t_0)$ such that darker regions represent decreases in absorption (or material) and lighter regions represent increases in absorption (or material). The $t_0$ frame was collected before +0.00 ms and is not shown. The laser moves from right to left, and the approximate location (within $\sim 50 \, \mu m$) is marked with a red line. A void, circled in orange, forms beginning in the +1.00 ms frame.
not the dynamic depth or shape of the depression induced by the recoil pressure from the metal vapor plume.23 Similarly, X-ray imaging is uniquely suited to observing the dynamics of pore formation during LPBF because it can directly detect pore formation and motion.

A montage of frames from a typical video is shown in Fig. 5(b). These images show the change in X-ray absorption relative to the start of the video to the current frame; i.e. \( A(t) - A(t_0) \), where \( A(t) \) is the measured absorption in the frame collected at time \( t \) and \( A(t_0) \) is the absorption measured in the first frame of the video, time \( t_0 \). Therefore, dark regions represent a loss of material density, grey regions represent no change, and lighter regions represent an increase in material density. For this scan, the laser power and scan speed are set to 100 W and 144 mm/s, respectively. These settings supply a higher energy density than is desired for a typical build. Due to the high energy density, the melt pool is relatively wide and the thermal boundary conditions imposed by the glassy carbon windows may influence melt pool behavior. In this montage, the laser scans from right to left across the substrate. The laser scan causes four major features in the images. First, a significant amount of material is lost in the powder layer, a result of powder movement, which causes the well-known denudation effect during LPBF.5,25,39 This appears as a large dark region above the substrate. Second, a track of deposited material is evident behind the laser, which appears as a small light region above the original powder-substrate interface. Third, a narrow dark region protruding into the substrate follows the laser scan, representing a lack of material caused by the depression of the melt pool surface due to recoil pressure. The density change in this region is near the noise level of the measurement and therefore is difficult to resolve, but it is a consistent feature of the data and we therefore conclude that it is not an artifact. Fourth, a small, oval-shaped region of decreased density appears starting at 1 ms due to the formation of a keyhole pore, in which the deep melt depression collapses and traps a void deep in the substrate. The pore is ellipsoidal with major and minor axes of 38 ± 3 \( \mu \)m and 25 ± 3 \( \mu \)m, respectively, and forms 136 ± 10 \( \mu \)m underneath the initial powder-substrate interface and 188 ± 10 \( \mu \)m below the surface of the newly deposited track. The major source of uncertainty in these position values arises from ambiguity in defining the exact location of the interface between the substrate and the powder. These dimensions are consistent with other observations of keyhole-type porosity in LPBF parts.29,40 Such pores are an undesirable feature that is often present in LPBF builds and significantly reduce the fatigue lifetime of as-built LPBF parts relative to wrought material.41 Observing pore formation \textit{in situ} provides valuable data for simulating the melt pool behavior that causes them.

\section*{X-RAY DIFFRACTION}

X-ray diffraction experiments were performed at SSRL beamline 10-2. The X-rays at this beamline are generated by a 33 pole, 1.27 T wiggler source. The X-ray beam is passed through a double crystal Si monochromator to select the photon energy and focused to a ~160 \( \times \) 600 \( \mu \)m spot using Rh-coated bent cylinder mirrors. These experiments used 20 keV (\( \lambda = 0.6199 \) Å) X-rays, the highest energy at which the mirrors can effectively focus the beam without a significant loss in flux. For the \textit{in situ} diffraction experiments, slits were used to further reduce the beam size to 50 \( \mu \)m (vertical) by 100 \( \mu \)m (horizontal), which provides an approximate flux of 10\(^{11}\) photons s\(^{-1}\) over the beam area. The horizontal spot size is also related to the ultimate temporal resolution available. The laser scans through the horizontal beam size over a finite amount of time dictated by the scan speed, so if the horizontal beam size is large relative to the distance traveled in the exposure time, the diffraction pattern obtained will probe regions of the sample in different states, complicating the interpretation of the lattice dynamics. The diameter of the Be window on the back side of the chamber allows for 20\(\mu\)s X-rays.

An Eiger 1M area detector (Dectris Ltd., Switzerland) with a total detection area of 77 \times 79.9 mm\(^2\) and pixel pitch of 75 \times 75 \(\mu\)m\(^2\) was used to record the diffracted X-rays. It was positioned approximately 128 mm behind the sample to provide full azimuthal coverage for data with \(Q_{\text{max}} = \sim 2.98 \text{ Å}^{-1}\), with a \(Q\) resolution \(\Delta Q/Q \sim 0.005 \text{ Å}^{-1}\). This \(Q\) range captures the full powder diffraction ring of the (100), (002), and (101) reflections from hexagonal close-packed (hcp, \(P6_3/mmc\)) \(\alpha\)-Ti and the (110) reflection from body-centered cubic (bcc, \(Im3\bar{m}\)) \(\beta\)-Ti. A movable diode beam stop wrapped in Pb foil was used to monitor the intensity of the transmitted X-ray beam and shield the detector from damage caused by the direct X-ray beam. Detector distance and tilt were calibrated with a LaB\(_6\) standard spread as a powder layer on top of a Ti-64 substrate. The X-ray beam only sampled the LaB\(_6\) powder, not the metal substrate. Data integration and calibration of detector distance and tilt were performed with the GSAS-II software package.42 Figure 6 shows typical diffraction data collected from a 500-\(\mu\)m-thick Ti-64 substrate with a ~60-\(\mu\)m-thick powder layer. Data were collected at a sampling rate of 1 kHz, and the laser was set to a power of 400 W with a scan speed of 576 mm/s. The X-ray beam was positioned with the beam center ~25 \(\mu\)m below the top of the substrate, as illustrated in Fig. 6(a). In this geometry, the entire sampled volume is within the substrate, with minimal contributions from the powder layer. Figure 6(b) shows the sum of 1500 diffraction patterns collected at 1 kHz, the equivalent of a 1.5 s collection. The (100), (002), and (101) \(\alpha\)-Ti reflections are noted with arrows. Figure 6(c) shows a two-dimensional diffraction pattern collected at 1 kHz prior to laser melting. A time series of integrated one-dimensional diffraction patterns collected at 1 kHz is shown in Fig. 6(d). Prior to laser exposure, the three \(\alpha\)-Ti peaks are present in addition to a very small (110) \(\beta\)-Ti peak between the (002) and (101) \(\alpha\)-Ti reflections [Fig. 6(e)]. Immediately after laser melting at \(t = 0\) ms, the sampled volume includes contributions from the re-solidified zone and the heat-affected zone (HAZ) on either side of the track. All the peaks are shifted to lower \(Q\) due to thermal expansion. Furthermore, the three \(\alpha\)-Ti peaks are lower in intensity and the (110) \(\beta\)-Ti peak is much more intense, indicating that the solidified material is cooling in the cubic \(\beta\)-Ti structure and comprises a large fraction of the sampled volume while the HAZ has warmed significantly from room temperature but
remains in the α-Ti structure. After \( t = 10 \) ms, the (110) β-Ti peak has disappeared and once again the diffraction pattern reflects a sample composed primarily of α-Ti with a small β-Ti contribution. Diffraction patterns collected before and after laser melting show differences in the peak shape and position of the three α-Ti peaks, indicating a change in macro- and micro-strain states. These observations of rapid crystallographic changes during cooling provide insight into the unique microstructural features of LPBF-made materials.

CONCLUSIONS

We report the design and implementation of a laboratory-scale LPBF instrument optimized to enable in situ X-ray experiments at a synchrotron radiation source while providing an environment that is representative of a full-scale LPBF machine. Experiments performed at SSRL produced 4 kHz imaging data with an effective pixel size of 1.1 \( \mu \)m and 1 kHz diffraction data from a 50 x 100 \( \mu \)m region. Initial inspection of the data reveals pore formation, melt depression dynamics, temperature-dependent lattice dynamics, and the β-Ti–α-Ti phase transition upon cooling in Ti-64. Ongoing upgrades to the detection scheme and sample holder will enable multi-layer experiments, faster sample preparation for higher throughput, and improved signal-to-noise ratios to increase maximum achievable sampling rates. Future experiments will include integration with optical process monitors to facilitate correlation between sub-surface defect formation and surface-sensitive optical detection approaches. Such correlations will provide direct insight into defect detection by online process monitoring during LPBF, a significant challenge for qualifying and certifying LPBF parts.

ACKNOWLEDGMENTS

This material is based upon work supported by the U.S. Department of Energy’s Office of Energy Efficiency and Renewable Energy (EERE) under the Advanced Manufacturing Office, CPA Agreement Nos. 32035, 32037, and 32038. Lawrence Livermore National Laboratory is operated by Lawrence Livermore National Security, LLC, for the U.S. Department of Energy, National Nuclear Security Administration under Contract No. DE-AC52-07NA27344. Use of the Stanford Synchrotron Radiation Lightsource, SLAC National Accelerator Laboratory, is supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Contract No. DE-AC02-76SF00515. Work at the Ames Laboratory was supported by the Office of Energy Efficiency and Renewable Energy under Contract No. DE-AC02-07CH11358. The authors acknowledge experimental assistance from Doug Van Campen, Matthew Latimer, and Ron Marks as well as helpful discussions with Peter Collins and Ryan Ott.