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ABSTRACT
We present a multi-purpose radiation furnace designed for x-ray experiments at synchrotrons. The furnace is optimized specifically for dark-field x-ray microscopy (DFXM) of crystalline materials at beamline ID06 of the European Synchrotron Radiation Facility. The furnace can reach temperatures above 1200 °C with a thermal stability better than 10 °C, with heating and cooling rates up to 30 K/s. The non-contact heating design enables samples to be heated either in air or in a controlled atmosphere contained within a capillary tube. The temperature was calibrated via the thermal expansion of an α-iron grain. Temperature profiles in the y and z axes were measured by scanning a thermocouple through the focal spot of the radiation furnace. In the current configuration of the beamline, this furnace can be used for DFXM, near-field x-ray topography, bright-field x-ray nanotomography, high-resolution reciprocal space mapping, and limited powder diffraction experiments. As a first application, we present a DFXM case study on isothermal heating of a commercially pure single crystal of aluminum.

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I. INTRODUCTION

The microstructure of a material often governs its performance in engineering applications. Heat treatments are used in industry to produce the desired mechanical properties in metals and alloys, as the microstructure depends on the material’s thermal history.1 Indeed, heat treatments are used to change the microstructure of a host of different materials, including ceramics and optical materials, to improve the material’s performance for diverse applications, including aerospace, electronics, communication, and medicine. To understand how temperature drives the microstructural changes in materials, this wide range of applications requires in situ studies to understand the microscopic processes during heat treatment to refine designs and develop new engineering materials.

Many industrial materials have hierarchical structures, with grains and domains that span several length scales; high-temperature studies of these materials thus require non-destructive measurements that can accurately measure structures over a range of 10 nm to 1 mm with a high angular resolution. Electron-based microscopy methods such as electron back-scatter diffraction (EBSD) and transmission electron microscopy (TEM) can probe several length scales with a high spatial resolution. However, the low penetration depth of electron beams limits these measurements to surfaces or very thin samples (often including invasive sectioning processes).2 Synchrotron diffraction methods such as
3DXRD and its near-field derivative, diffraction contrast tomography (DCT), can measure 3D information in a non-destructive manner. However, these tools cannot map highly deformed samples or nm-sized domains because of overlapping peaks and insufficient spatial resolution, respectively.

Dark-field x-ray microscopy (DFXM) is a new technique that non-destructively collects three-dimensional (3D) information about the material’s strain and orientation at different length scales.\cite{1,2} Analogous to dark-field electron microscopy, DFXM uses an objective lens to magnify features in a crystalline sample that diffract. Using high-energy synchrotron x rays, the beam penetrates the crystal, resolving deeply embedded features. Full 3D mapping can be performed in minutes, with changes to the field of view and spatial resolution requiring only simple reconfiguration of the x-ray objective lens. The spatial resolution has reached 100 nm with an angular resolution of 0.001°. The high resolution and high sensitivity coupled with the deep penetration depth makes DFXM a unique tool to study the in situ behavior of crystalline engineering materials at high temperatures.

Many different types of heating systems are used for synchrotron experiments, including resistive heating,\cite{3,4} laser heating,\cite{5} inductive heating,\cite{6} and radiation furnaces.\cite{7,8} In radiation furnaces (also called optical or mirror furnaces), reflective mirrors focus radiation from a lamp (e.g., halogen) onto the sample. This design makes radiation furnaces compact, with high heating and cooling rates and high thermal stability for experiments in both transmission and reflection geometries. Most existing radiation furnace designs have been developed for in situ x-ray diffraction (XRD) studies; however, the aforementioned advantages also present advantages for in situ diffraction imaging\cite{9,10,11}, which also require a compact design and high thermal stability. Here, we present a non-contact radiation furnace, designed specifically to be compatible with DFXM experiments at the European Synchrotron Radiation Facility (ESRF).

II. SPECIFICATIONS

The initial goals of this furnace focused on metals, so we focus the calibrations on the temperature range relevant to process steel, namely, from room temperature to above 1200 °C. Within this range, we also explored fast rates of heating and cooling, which are important for industrial applications that require isothermal studies and additive manufacturing. Cooling rates of 25–30 K/s are typical for austenite to pearlite and martensite transition in many steel alloys.\cite{12} Therefore, the furnace is needed to achieve these cooling rates or more.

A well-defined thermal stability is essential to interpret the material’s response to temperature. The microstructure of a material (especially ductile metals) is also very sensitive to additional stresses from thermal fluctuations, which can produce results that are not representative of an isothermal material. This means that temperature fluctuations must be avoided such that they do not alter the state of the sample during a DFXM scan for isothermal studies. As DFXM experiments use objective lenses that have small effective apertures, they are highly sensitive to changes in the diffraction angle (varies with the d-spacing) that originate from thermal expansion or strain.

In addition to the primary application, DFXM, the furnace also needs to be compatible with the other x-ray techniques available at ID06, including bright-field tomography, near-field diffraction topography, and high-resolution reciprocal space mapping. For use over this wide range of experimental capabilities, the furnace thus needs to have clear apertures for the incident, transmitted, and diffracted x-ray beams with scattering angles up to 2θ ≈ 30° in the vertical scattering plane (i.e., the maximum scattering angle of the current instrument on ID06).\cite{13} Finally, the furnace needs to permit a full 360° sample rotation about the y axis to allow experiments to rock the crystal along one rotational axis for diffraction in the vertical geometry.\cite{14} At the same time, the apertures need to be as small as possible to prevent heat from leaking out of the system.

As DFXM images embedded crystalline regions beneath the material’s surface, the x-ray measurements are performed in transmission (Laue) geometry, with photon energies covering 15–35 keV. Typical samples have a size of 0.5–2 mm in the x and z directions with a length of several mm along the y axis. The samples are mounted inside a quartz capillary that can be filled with either air or an inert atmosphere.

The primary design constraint for the furnace was the limited space available on the sample stage at ID06, specifically, the distance between the sample mounting surface and the goniometer’s center of rotation (i.e., the sample position). The sample mounting surface of the goniometer is on the negative y side [right side in Fig. 1(b)]. Furthermore, the furnace needs to be small enough to allow a near-field camera to be positioned 50–100 mm downstream from the sample for alignment or imaging experiments (e.g., near-field x-ray topography).

![FIG. 1. General layout of the furnace: (a) side view and (b) front view. The incident beam travels along the x-axis, the y-axis is to the left in (b), and the z-axis is up. (c) Photograph of the radiation furnace. The opening for mounting the sample is on the left and the slot for the scattered beam is on the right.](https://example.com/furnace_image)
III. IMPLEMENTATION

To accommodate the specifications and constraints listed above, we constructed a non-contact radiation furnace, where the furnace and sample are mounted independently. This has several benefits:

- Heat transfer from the furnace to the goniometer is reduced, thus minimizing long-timescale thermal drift of the sample position.
- The furnace and the associated power cables and cooling water lines do not increase the mechanical load on the goniometer stages.
- Potential vibrations of the furnace (e.g., due to water cooling) are not transmitted to the sample.
- The furnace remains stationary during x-ray experiments even when the sample is rotated or moved, minimizing the necessary size of the openings for the incident and diffracted beam. This reduces power loss, increasing the efficiency of the furnace and minimizing inhomogeneities in the heating.
- As the furnace is mounted on a separate motorized y translation stage, the furnace may be retracted without repositioning the sample. This can be helpful to access a wider range of scattering angles, as has been performed in interrupted annealing measurements with 3DXRD.
- The sample's temperature can be changed rapidly, as only the lamps and sample are "hot" (i.e., the thermal inertia of the system is very small). In principle, the rapid temperature response makes proportional integral derivative (PID) regulation easy.

A schematic layout of the furnace is shown in Fig. 1. Two lamps are used as the heating elements in this design, as preliminary tests with a single bulb did not reach the desired temperatures. The bulbs are positioned in a reflective cavity with a shape defined by two intersecting ellipsoids. The sample is positioned near the shared central focus, and the bulbs sit at the other two foci, as shown in Fig. 1.

The furnace was constructed by AS Special Devices, Echirolles, France. The reflectors were machined from copper, then polished, and gold plated. Integrated cooling channels regulate the temperature in the furnace, circulating either air or water. Before machining, the copper casing was annealed at 850 °C for 12 h to relieve any built-up stresses in the material, preventing the material from deforming as the outer shell and water connections are brazed in the subsequent machining.

Machining of the furnace was carried out in two steps. First, the internal shape was machined to the rough-shape of an ellipsoid, with axes 1 mm smaller than the desired shape. The outer shell and water connections were then brazed, and the ellipsoid was subsequently machined to its final size and shape.

The cavities were then electropolished, refining the surface roughness to \( R_a \approx 0.1 \) μm. Loss of material during the electropolishing was partially compensated by a surface layer of nickel, which prevents diffusion between the copper casing and the gold layer that was deposited on the surface.

The integrated cooling channels keep the furnace's outer surface below 20 °C, even when the sample is heated to internal temperatures above 1000 °C.

The sizes and flow rates for the cooling channels were calculated to allow for locations for the input and output feeds that could minimize forces on the furnace as much as possible. Ultimately, reducing the net force would mitigate vibrations from the water circulation that could reduce blur of the experiment's images.

Parasitic heating of the surroundings is thus limited to black-body radiation that escapes through apertures for the x-ray beams, the sample, and near the bases of the two lamps. The cooling channels allow for safe operation of equipment surrounding the furnace, including the goniometer, the sample stage, and the near-field camera.

The sample is positioned 3 mm upstream of the furnace radiation's focal spot. This gives an illuminated circular area with a diameter of 5 mm through the depth of the sample, which heats the sample more homogeneously than it would if the sample were in the exactly focal spot. The defocused radiation therefore improves the precision of the temperature with respect to possible misalignment of the sample's position in the furnace.

The casing's apertures for its heat lamps were designed to reduce air convection at the sample position and—as much as possible—the resulting temperature inhomogeneities that they could also generate around the sample. Its diameter was optimized using a flow simulation program in order that 75% of the air enters through the lower lamp's aperture and exits through the upper lamp's aperture.

The heating elements in the furnace are two identical 400 W halogen light bulbs. The lamps are connected in parallel to a Delta Elektronika DC power supply, operated at constant power via a software feedback loop. The required power for a given set-point temperature was obtained from a lookup table calculated from calibration measurements with a thermocouple at the sample position. The beamline control software can control the sample's temperature, and thus, data can be acquired automatically with slow, step-wise temperature ramps or interrupted annealing over a precise duration. Different samples may require a different power to achieve the same set-point temperature, based on the surface reflectivity and absorption cross section over the radiation furnace's energy spectrum.

Samples can be inserted horizontally (perpendicular to the scattering plane) through an opening with \(~\approx 5\) mm diameter. The distance from the outside edge of the furnace to the sample position is \(~\approx 27\) mm. The size of this opening allows the samples to be sealed in capillaries that are backfilled with a desired atmosphere and, if necessary, mounted to a Huber pin with a temperature-resistant ceramic sleeve to ensure stability. The aperture for the incident x rays has diameter of 2.5 mm, which matches the field of view of the near-field camera. On the downstream side, a vertical slot with 2.5 mm width allows x-ray beams with vertical scattering angles of up to 30° to exit the furnace unobstructed. The blackbody radiation within the furnace is focused by double-ellipsoid mirrors. The axis or rotation of the double-ellipsoid is inclined 45° toward the incident x-ray beam, which maximizes the accessible range of scattering angles, as shown in Fig. 1.

There are some drawbacks of the chosen design geometry:

- The radiation's focal point within the furnace is located 27 mm from the outer edges of the furnace (along the radius of the cylinder). The sample support must, therefore, be long...
and thin to position the sample at the center of the furnace, with supporting material that is also heated by the blackbody radiation. Temperature changes and thermal expansion within the supporting materials may require adjustments to the sample position after large changes in temperature. The support should therefore be made from a material with low thermal expansion (e.g., quartz) and ideally with a high infrared reflectively or low infrared absorption.

- The small diameter of the sample aperture (5 mm) strongly limits the range of sample rotations about the x (incident beam) and z (vertical) axes. Increasing the accessible angular range would require a larger lateral opening in the furnace, which would leak significantly more heat onto the goniometer, potentially destabilizing the thermal uniformity at the sample position. Furthermore, these changes could also cause asymmetry and inhomogeneity in the radiation and result in temperature gradients across the sample.

- The sample’s temperature is not measured directly. The equilibrium sample temperature depends on a combination of parameters, including the sample’s absorption cross section for the relevant blackbody radiation spectrum and its emissivity. At a given power, a strongly absorbing and low-emissivity sample may be much hotter than a highly reflective or transmissive sample. This means that the relationship between the power to the lamps and the temperature at the sample must be independently calibrated using, for example, thermal expansion measurements.

- The small mass of heated samples (thermal mass) and the capability for rapid temperature changes both reduce the system’s short-term temperature stability.

### IV. RESULTS AND DISCUSSIONS

#### A. Characterization by thermocouple

Figure 2 shows the temperature response of a K-type thermocouple located at the center of the furnace at different power values. We used a fitting function that contains the sum of two exponential functions to describe the time–temperature behavior of the furnace, with the sum of two exponentials couple located at the center of the furnace at different power values.

\[
T = T_0 + A_{\text{slow}} \exp\left(-t/\tau_{\text{slow}}\right) + A_{\text{fast}} \exp\left(-t/\tau_{\text{fast}}\right).
\]

The results of the fits are reported in Table I. For the first temperature step, we compute the thermal stability at 5 min to be 4.63 K based on the exponential fit, which yields \(\tau_{\text{slow}} = 137 \pm 6.7\) s with an amplitude of \(A_{\text{slow}} = -41 \pm 0.6\) K. For the fourth temperature step, which cools the sample to room temperature and, as radiative heat transfer becomes considerably small when the shell and the sample are nearly at room temperature, is therefore the slowest of these steps, the thermal stability at 5 min is calculated as 7.72 K from the fit (Table I).

We note that the two exponentials in each fit correspond to two different stages for each heating step, each with their own inherent timescales. We interpret the fast heating stages to be equilibration between the furnace and the sample, while the slow heating stage corresponds to the equilibration of the sample holder to the new temperature (in this case, the stainless steel housing of the thermocouple). The diffraction-based calibration measurements used a quartz capillary as the sample holder and showed better thermal stability after 5 min, indicating less lag time from the slower process. Further improvements to the timescales of thermal equilibration and stability could be made by implementing a closed-loop feedback system.

We also note that the measured temperatures are different at the two plateaus of \(P = 37\) W. This result is caused by the blackening of the thermocouple (shown by the black arrow in Fig. 2) during the heating at \(P = 69\) W, which increases its absorption efficiency for the furnace’s radiation and therefore increases the temperature reached the second time when the thermocouple was heated with \(P = 37\) W.

The heating and cooling rates were calculated by fitting the temperature–time plots using exponential functions, as shown in Fig. 2. Heating to a set temperature always results in a faster rate than that of the corresponding temperature decrease, as there is no active cooling element to drive the cooling rates faster than thermal diffusion of the sample. The cooling rates could be increased by blowing room temperature air or nitrogen gas through the cavity of the furnace, although this would also introduce thermal gradients, as it would be a surface-driven cooling element. Note that in this experiment, the internal copper elements of the furnace were cooled.

Figures 3(a) and 3(b) show the spatial profile of temperature along the y and z directions for different lamp powers, as measured

#### TABLE I. Fit parameters of the temperature calibration using the K-type thermocouple.

| Temperature step | \(A_{\text{fast}}\)(K) | \(\tau_{\text{fast}}\)(s) | \(A_{\text{slow}}\)(K) | \(\tau_{\text{slow}}\)(s) |
|------------------|---------------------|----------------|-------------------|--------------|
| 1                | -78.6 ± 0.8         | 8.91 ± 0.05    | -41 ± 0.5         | 137.6 ± 6.7  |
| 2                | -264.6 ± 0.9        | 4.80 ± 0.03    | -40.9 ± 0.34      | 111.1 ± 3.4  |
| 3                | -298.4 ± 1.5        | 6.53 ± 0.06    | 33.7 ± 0.7        | 85.2 ± 4.2   |
| 4                | 419.9 ± 1.6         | 12.8 ± 0.1     | 90.3 ± 2.5        | 122.1 ± 2.5  |

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![FIG. 2. Temperature calibration using a K-type thermocouple showing heating and cooling rates for given set temperatures.](image-url)
with a K-type thermocouple. The measured temperature increases as a function of power [Fig. 3(c)]; however, the spatial temperature gradient (ΔT from the edges to the center) also increases—meaning that the edges of the sample are significantly cooler than the center. Both temperature distributions are expected to be symmetrical around the center; we interpret asymmetry in this case to be caused by misalignment of the lamps. The temperature variation near the center was also below 1200 °C at 185 W, compatible with most of the industrially relevant heat treatments of steel alloys as they take place well below 1200 °C. We note that higher temperatures could be achieved within the power limits of the lamps, as we did not use the full 400 W power range for the two halogen lamps in the furnace. For example, we measured a temperature of 1450 ± 20 °C at 250 W using a blackened type-S thermocouple.

**B. Characterizing temperature from lattice parameters**

We used a recrystallized grain of α iron within an Fe-3%Si sample with dimensions of 0.17 × 0.17 × 8 mm³ inside a quartz capillary to calibrate the furnace based on thermal expansion within the sample. We measured shifts in the (110) Bragg peak along the vertical direction, using 17 keV x rays that were diffracted onto a 2D FReLoN CCD detector that was placed 5090 mm behind the sample. Using the motorized translation stages at ID06, we were able to translate the detector to the appropriate 20 angle across the entire temperature range required for the calibration. The d-spacing of (110) measured for each furnace power was used to calculate the temperature of the sample using the equation described in Ref. 21. This approach of using the lattice parameter and thermal expansion relation to measure temperature is ideally suited for the x-ray measurements at ID06, so long as care is taken to ensure that the crystal’s orientation will diffract at least one beam through the apertures of the furnace (Fig. 4).

**C. Usage during a DFXM experiment**

To demonstrate the utility of the radiation furnace in DFXM-based studies, we include initial data from experiments investigating the evolution of dislocations with temperature (annealing) in single-crystalline aluminum. The 0.5 × 0.5 × 20 mm³ crystals were used as purchased (no further purification or heat treatments) from Surface Preparation Laboratories in this experiment. The samples were cut from a larger ore and polished by chemical etching to mitigate residual surface strains. This experiment performed slow-ramp heating of the samples, with long equilibration times at each temperature, spanning 6 h. The microstructure of the crystal was then measured at a series of temperatures using the dark-field x-ray microscope at the ESRF beamline ID06-HXM. A Si(111) double crystal monochromator selected x rays at photon energy 17 keV from the undulator source.

Before reaching the sample, the incident x rays were passed through a 2D trans focator with compound refractive lenses (CRLs) totaling 8 Be lenses (an apex radius of curvature R = 200 μm), followed by a 1D condenser with 58 Be lenses (R = 200 μm). A 400 μm × 600 nm (horizontal × vertical) area of the sample was illuminated by the x rays. The diffracted x rays were then imaged with an x-ray objective lens comprised of 88 2D Be parabolic lenses (R = 50 μm) that was placed 274 mm behind the sample, producing an effective focal length of 260 mm. A far-field CCD camera was positioned 5364 mm from the sample to capture the magnified image. The effective magnification was 18.5×, resulting in a final effective pixel size of 75 nm/pixel when combined with the scintillator-based optical components used with the detector. The incident x-ray line beam illuminated a 5600 nm high layer of the sample to capture section topographs. To map the relative axial strain, longitudinal (θ − 2θ) scans were performed by collecting dark-field images across scans of the sample tilt and scans of the objective and camera positions. We probed the (002) Bragg reflection of the aluminum crystal at 2θ = 20.84°. The temperature was calibrated for the aluminum sample using the thermal expansion coefficients described in Ref. 22.

Figure 5 reveals the refinement of the microstructure with the increase in temperature, covering the temperature range required...
FIG. 4. Temperature calibration curve as measured based on thermal expansion of the lattice parameter in $\alpha$ iron. The (110) Bragg reflection of a grain was traced on a CCD located $\approx 5$ m downstream of the sample. The plots show (a) the sample temperature as a function of applied power and (b) the measured lattice parameter for each temperature upon heating.

FIG. 5. Reconstructed DFXM relative strain maps of single-crystalline aluminum during in situ isothermal heating at (a) $155$ °C showing mostly positive strain in the interior of the crystal (b) at $252$ °C with some negative strain present in the interior (c) at $416$ °C where the strain fluctuations start to become prominent and linear structures appear (d) at $501$ °C showing higher strain fluctuations with more evident linear structures. The color scale represents relative axial strain.

V. CONCLUSION

We demonstrated a radiation furnace developed for DFXM studies at ID06 beamline at the ESRF. The furnace was calibrated up to $1050$ °C using two distinct methods: by means of a K-type thermocouple and by resolving thermal expansion through the lattice parameter in $\alpha$ iron. Extrapolating the calibration curve, we expect that temperatures above $1200$ °C can be reached—potentially higher for samples with very high infrared absorption. This makes the furnace ideal to study steel alloys and ceramics. While originally designed for DFXM studies, this furnace is compatible with the other methods available at ID06 such as bright-field tomography, near-field diffraction topography, and high-resolution reciprocal space mapping. We showed a case study using DFXM of isothermal heating of a pure single-crystal aluminum. Axial strain maps showed subtle structural modifications as a function of temperature, which are characteristic of the annealing that was performed. The limitations of the furnace result from the restricted apertures for the sample and beam. Mosaicity scans are difficult due to the small tilt range orthogonal to the rocking, and full 3DXRD scans are impossible due to the limited size of the beam exit. Nevertheless, these studies are still possible for interrupted...
heating schemes by translating the furnace out without moving the sample.

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DATA AVAILABILITY

Raw data were generated at the ESRF large scale facility. Derived data supporting the findings of this study are available from the corresponding author upon reasonable request.

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