Photo-activated raster scanning thermal imaging at sub-diffraction resolution

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Active thermal imaging is a valuable tool for the nondestructive characterization of the morphological properties and the functional state of biological tissues and synthetic materials. However, state-of-the-art techniques do not typically combine the required high spatial resolution over extended fields of view with the quantification of temperature variations. Here, we demonstrate quantitative far-infrared photo-thermal imaging at sub-diffraction resolution over millimeter-sized fields of view. Our approach combines the sample absorption of modulated raster-scanned laser light with the automated localization of the laser-induced temperature variations imaged by a thermal camera. With temperature increments ~0.5–5 °C, we achieve a six-time gain with respect to our 350-μm diffraction-limited resolution with proof-of-principle experiments on synthetic samples. We finally demonstrate the biological relevance of sub-diffraction thermal imaging by retrieving temperature-based super-resolution maps of the distribution of Prussian blue nanocubes across explanted murine skin biopsies.
Active photo-thermal imaging over submillimeter-sized to centimeter-sized fields of view with tunable spatial resolution in the 10–100 μm range would be a valuable tool for the nondestructive characterization of the morphology and functional state of both biological tissues and synthetic materials. Differently from passive techniques, which measure temperature changes spontaneously occurring in the sample due to intrinsic heat generation, active approaches rely on monitoring the peculiar sample thermal response upon application of an external heating (or cooling) stimulus \(^1\text{-}^4\). In the clinical and pre-clinical setting, for example, active medical thermography allows identifying the modified blood perfusion and metabolic activity associated to the presence of inflammations, cancer masses or physiological dysfunctions by accessing the resulting alterations in the tissues thermo-physical properties (e.g., the thermal conductivity \(^3\), or the sample temperature during the thermal relaxation phase \(^6\)). Externally induced temperature variations also provide direct access to the spatial distribution of bio-markers, as demonstrated with melanin for the identification of age-related macular degeneration in the retina \(^1\text{-}^3\) or the discrimination of in-situ and invasive malignant skin lesions in the context of melanoma screening and diagnosis \(^6\text{-}^7\). In nanomedicine and nanotechnological research, photo-induced heat release events allow characterizing the distribution of nanoparticles following systemic administration in animal model systems, and the development of related photo-thermal therapy protocols \(^8\text{-}^9\).

Overall, active medical thermography finds promising applications ranging from the development of quantitative screening tools for the (early-stage) detection of pathologies, to the characterization of disease progression and therapy response in forefront pre-clinical research. High resolution active thermal imaging is equally relevant to monitor the dissipation efficiency of all the measured temperature peaks. Provided the technique sensitivity \(^3\text{-}^4\), may limit the applicability to biological systems.

More straightforward access to temperature values, coupled to imaging over wider areas, can be achieved by infrared thermal imaging \(^10\text{-}^12\). Upon calibration of the sample emissivity \(^3\text{-}^4\text{-}^33\text{-}^36\), temperature increments can be quantified based on Stefan-Boltzmann’s law by the measurement of the infrared radiance in the gray-body approximation. However, the typical \(~\sim100 °C\) range of interest corresponds to peak radiances in the \(~7–11 μm\) band \(^13\text{-}^4\): even though expensive thermo-cameras with pixel size as low as 5–20 μm on the sample plane exist in commerce, the effective resolution is limited to \(~\sim0.1–1 mm\) \(^15\text{-}^40\) due to the thermal waves diffusion in the sample and to the diffraction of the far-infrared radiation at the low numerical aperture of Germanium or Zinc-Selenide optics (Abbe’s law \(^41\text{-}^43\)). Higher spatial resolution (\(~100 μm\) and \(~0.1 °C\) sensitivity have been reached in specially designed military equipment only \(^44\text{-}^45\).

It is therefore our purpose here to develop and validate a non-contact super-resolution infrared thermal imaging approach capable of quantitatively measuring temperature increments, as allowed by conventional thermography, and of simultaneously achieving the \(~50 μm\) resolution required by forefront medical, electronic and nano-technical applications on variably extended (sub-millimeter to centimeter-sized) fields of view. Similarly to what is performed in PALM \(^46\) (Photo-Activated Localization Microscopy) and STORM \(^47\) (STochastic Optical Reconstruction Microscopy), our strategy exploits the automated sub-diffraction centroid localization of sparse temperature increments primed by modulated raster-scanned focused laser light and imaged by a thermal camera. The super-resolution image of the light-absorbing centers in the sample is reconstructed and color-coded by the localized centers and amplitudes of all the measured temperature peaks. Provided the fit localization precision is only limited by the shot-noise of thermal emission \(^48\), the spatial resolution of the rendered image can in principle be tuned down to the \(~1 μm\) diffraction-limited laser spot size at the excitation visible to near-infrared wavelength. \(<60-μm\) resolution is initially demonstrated with temperature increments as low as \(~0.5–5 °C\) by proof-of-principle experiments on synthetic samples. We thereby achieve 6-fold and 20-fold enhancements with respect to the diffraction-limited prediction and the effective resolution of our thermo-camera in default operation. Validated here and particularly advantageous on a low-cost (\(~\sim5k\) $) thermo-camera, our super-resolved imaging procedure could be adopted to enhance the resolution of any, even expensive (\(~\sim70k\) $), infrared thermography setup. We finally demonstrate application to biological samples, and provide super-resolution maps of the distribution of 30-nm Prussian blue nanocubes \(^49\text{-}^50\) across millimeter-sized expanded murine skin biopsies. This is a necessary step for the development of a photo-thermal therapy protocol to be subsequently applied in-vivo to the tissue in pathological conditions. Demonstrated here with exogenous nanostructures, exemplary application of super-resolution thermal imaging can be envisioned in the characterization of the melanin distribution in pigmented skin lesions, aimed at the objective discrimination of benign tissue and malignant melanoma both in-vivo and ex-vivo in a non-invasive label-free approach.

**Results**

Photo-activated thermography at sub-diffraction resolution. The radiance of a gray-body of emissivity \(ε (0 < ε < 1)\) at temperature \(T\) is \(R = εσT^4\), where \(σ = 5.67 \times 10^{-8}\) W m\(^{-2}\)K\(^{-4}\) is Stefan-Boltzmann’s constant. Non-contact thermal imaging produces thermographic sequences based on the intensity of emitted
A Gaussian laser beam (red spot in a) is focused on the sample at \((x_0, y_0)\) with a square pulse of duration \(\tau_{on}\). Light absorbing entities (blue) induce a local temperature increase, which is measured by the thermal camera and as an evolving 2D Gaussian peak. A Gaussian fit of the frame collected at time \(\tau_{on}\) (b, c) provides the temperature variation \(\Delta T_{max} = \Delta T_{max} - T_0\) and the peak coordinates \((x_c \pm \sigma_x, y_c \pm \sigma_y)\) that localize the center of the distribution of the absorbing objects within the laser spot size (magnified region in a). The procedure is repeated over sets of isolated points as exemplified in e, f. The center coordinates provided by the Gaussian fit of temperature peaks identify the position of the absorbing entities on the scan grid (d, g) with uncertainty \(\sigma_{xy}\) and a color code assigned by the best-fit \(\Delta T_{max}\). A maximum projection of the signal across the raw thermo-camera image sequence provides a low-\((\sim \text{mm})\) resolution image of the sample (i), whereas a maximum projection of the stack containing all the localized absorptive centers provides the super-resolution image (j) of the scanned region (h). Scale bars (100 \(\mu\text{m}\) in the magnification of a, 1000 \(\mu\text{m}\) elsewhere) indicate the typical imaged fields of view; note that the -50-\(\mu\text{m}\) laser spot and the whole magnified area in a lie within a single -400-\(\mu\text{m}\) typical thermo-camera pixel on the sample plane. All panels have been derived from experimental data acquired on an explanted murine skin biopsy treated with 30-nm Prussian blue nano-cubes analogous to the one employed for Fig. 3.

Fig. 1 Photo-activated thermal imaging at sub-diffraction resolution. A Gaussian laser beam (red spot in a) is focused on the sample at \((x_0, y_0)\) with a square pulse of duration \(\tau_{on}\). Light absorbing entities (blue) induce a local temperature increase, which is measured by the thermal camera and as an evolving 2D Gaussian peak. A Gaussian fit of the frame collected at time \(\tau_{on}\) (b, c) provides the temperature variation \(\Delta T_{max} = \Delta T_{max} - T_0\) and the peak coordinates \((x_c \pm \sigma_x, y_c \pm \sigma_y)\) that localize the center of the distribution of the absorbing objects within the laser spot size (magnified region in a). The procedure is repeated over sets of isolated points as exemplified in e, f. The center coordinates provided by the Gaussian fit of temperature peaks identify the position of the absorbing entities on the scan grid (d, g) with uncertainty \(\sigma_{xy}\) and a color code assigned by the best-fit \(\Delta T_{max}\). A maximum projection of the signal across the raw thermo-camera image sequence provides a low-\((\sim \text{mm})\) resolution image of the sample (i), whereas a maximum projection of the stack containing all the localized absorptive centers provides the super-resolution image (j) of the scanned region (h). Scale bars (100 \(\mu\text{m}\) in the magnification of a, 1000 \(\mu\text{m}\) elsewhere) indicate the typical imaged fields of view; note that the -50-\(\mu\text{m}\) laser spot and the whole magnified area in a lie within a single -400-\(\mu\text{m}\) typical thermo-camera pixel on the sample plane. All panels have been derived from experimental data acquired on an explanted murine skin biopsy treated with 30-nm Prussian blue nano-cubes analogous to the one employed for Fig. 3.

In order to reconstruct a super-resolution image across extended areas, the procedure of illumination and localization of absorbing centers is sequentially repeated (Fig. 1e). A minimum distance \(\Delta x\) between pairs of consecutively irradiated points and/or a minimum time interval \(\Delta t\) in between their illumination have to be adopted to ensure that the two temperature variations appear as spatially and/or temporally separate peaks in the thermal camera images. This is crucial to enable the correct localization of the center of each detected temperature increase by Gaussian fitting. The minimum values for \(\Delta x\) and \(\Delta t\) depend on the laser activation time and on the sample thermal diffusivity, as detailed in Supplementary Note 2. Any illumination scheme satisfying this constraint on \(\Delta x\) and \(\Delta t\) can be exploited. We adopt here a modulated laser illumination (Supplementary Fig. 3 and Methods section), where the sample is raster scanned by the focused laser beam while a synchronized shutter allows modulating the laser activation and de-activation. The scan grid, \(N_xN_y\) pixels in format, has a pixel size \(\delta x \sim 10 \mu\text{m}\) and is oversampled relatively to the \(\sim 400-\mu\text{m}\) thermo-camera pixel size on the sample plane. During each raster-scan,
illumination (open-shutter condition) is only allowed on the limited subset of isolated pixels lying on a square lattice with characteristic spacings \( \Delta x \) and \( \Delta y \) along the horizontal and vertical directions, respectively (Supplementary Fig. 3). Complete coverage of the investigated area is achieved when the raster scan is repeated \( \Delta x \Delta y \) times, since for each scan only \( N_x N_y / (\Delta x \Delta y) \) pixels get illuminated. The laser pixel dwell time coincides with the shutter opening time (~100–1000 ms) during laser activation, whereas a ~2 ms dwell time is selected during shutter closure (see Methods section). This results in a total acquisition time \( t_{\text{on}} = N_x N_y \tau_{\text{on}} + N_x N_y \tau_{\text{off}}(\Delta x \Delta y - 1) \sim 10–100 \) min over millimeter-sized areas depending on \( \tau_{\text{on}}, N_x, N_y, \Delta x, \) and \( \Delta y \) (Supplementary Fig. 3).

During the scanning, the thermal camera acquires frames at rate \( f_{\text{rate}} \), providing a typical \( 10^4–10^5 \)–frames sequence (Fig. 1b, f). If a maximum projection of the signal is performed along the time axis of the raw image stack, a low-resolution thermal image—resembling the one that would be produced under wide-field illumination of the sample—is obtained (Fig. 1i and Supplementary Note 3). By contrast, when the center coordinates and amplitudes of all the detected temperature peaks are determined by Gaussian fitting and are associated to the corresponding pixel along the scan grid (Fig. 1d, g), the center coordinates provide much better resolved topological information on the absorptive centers in the sample. Best-fit amplitudes encode the information on the sample absorption and local induced temperature variations, and provide the color code for the final super-resolution image that is obtained by superimposing all the color-coded pixels where absorptive centers have been localized (Fig. 1j and Methods section). Significantly, if the scan pixel size and the localization uncertainty are smaller than the thermal wavelengths of the detected signal only, this is an estimate of \( \sigma_x, \sigma_y \), and the spatial resolution \( \Delta x, \Delta y \) is systematically minimized when the fit of temperature peaks is performed as a two-step Gaussian fit: a first fit to a symmetric function with fixed variance provides the peak coordinates, which are fixed in a second step to recover the variance \( \sigma^2 \) and \( \Delta T_{\text{max}} \). Similarly, a symmetric Gaussian trial function outperforms asymmetric and skewed fit surfaces when the laser beam hits on a microfluid ink stripe with 30-μm width, which only partially covers the excitation spot with 1/2\( \mu \)m diameter and 56 ± 2 μm (Supplementary Fig. 5g). Exploitation of the same fitting routine is justified therefore for any a priori unknown distribution of absorptive entities.

Plotting the uncertainty \( \sigma_{\text{xy}} \) resulting from the two-step Gaussian fit as a function of the best-fit \( \Delta T_{\text{max}} \) for both the ink square and the ink stripe further reveals that all data points lie along the expected curve\(^{46,48}\)

\[
\sigma_{\text{xy}} \approx \sqrt{\left(\alpha^2 + \alpha^4/\mathcal{N} + 4x/\pi \sqrt{C}\right)/\Delta \mathcal{N}^2} \approx \sqrt{\alpha/\Delta T_{\text{max}}^2 + \beta/\Delta T_{\text{max}}^4} \tag{a}
\]

The fit of the trend of \( \sigma_{\text{xy}} \) versus \( \Delta T_{\text{max}} \) (Supplementary Fig. 5e, best-fit parameters \( \alpha = 599 \pm 221^\circ \text{C} \mu^2 \) and \( \beta = 634 \pm 133^\circ \text{C}^2 \mu^2 \)) allows extrapolating the localization uncertainty at any measured \( \Delta T_{\text{max}} \) thereby providing the estimate of \( \sigma_{\text{xy}} \) for the following experiments on synthetic samples.

Microfluidic samples have been subsequently exploited to perform proof-of-principle experiments aimed at validating the proposed super-resolution imaging approach.

A LABS-shaped structure (acronym for Laboratory of Advanced Bio-Spectroscopy—Fig. 2a) has been reconstructed first. Following experimental measurement of the sample emissivity (Supplementary Note 5), the temperature variations induced by modulated illumination and localized by two-step Gaussian fitting of the thermo-camera frames have been exploited to render the final super-resolution image of the sample (Fig. 2c). Visual inspection of Fig. 2c already suggests proper reconstruction of the features of the printed pattern, and the 0.14°C standard deviation of the measured temperature variations confirms that the sample is remarkably uniform in terms of the absorption properties (Supplementary Fig. 6). None of the letters of the LABS pattern would instead be recognized by the thermal camera in conventional operation, as can be seen on the temporal maximum projection of the whole stack of the raw thermo-camera images (Fig. 2b).

Quantitatively, the accuracy of the image reconstruction algorithm is demonstrated in Fig. 2d by the comparison with the transmitted-light image of the same sample. The spatial profile of \( \Delta T_{\text{max}} \) values drawn along the space between the
letters B and S of the ink pattern correctly superimposes to the corresponding intensity profile in the transmitted-light image, within the 30–50 μm localization error $\sigma_{xy}$ expected from the values of $\Delta T_{\text{max}} = 0.6–0.9 \degree\text{C}$ (Supplementary Fig. 5e). The agreement between temperature variations and transmitted-light intensity profiles should further increase at increasing observed $\Delta T_{\text{max}}$ (increasing laser intensity or activation time and decreasing $\sigma_{xy}$). Still, the localization error $\sigma_{xy}$ of the reported data-set allows discriminating absorbing structures (in Fig. 2d, the letters B and S) 100 μm apart.

The 100 μm resolution achieved in Fig. 2c, d, which already demonstrates a 3.5-time gain with respect to the theoretical 345-μm diffraction-limited resolution of our thermal imaging setup (Supplementary Note 3), is limited by the 56 ± 2 μm laser spot 1/e² diameter (Supplementary Fig. 7), combined with the 30–50 μm localization uncertainty. It has been pushed further by both increasing the excitation laser power (and the detected $\Delta T_{\text{max}}$ values) and reducing to 22 ± 1 μm the spot 1/e² diameter. A microfiche sample reproducing a grid of uniform ink stripes, 60 μm in width and relative distance (Fig. 2f), has been imaged by modulated illumination yielding the super-resolution image in Fig. 2g. Again, the structures that would be unresolved in conventional thermal imaging (as shown by the temporal maximum projection over the raw image stack, Fig. 2e) appear instead discernible in the super-resolved frame. The comparison with the transmitted-light (200 nm resolution) image of the same sample quantitatively confirms that the size of the imaged ink stripes is correctly retrieved (Fig. 2h). Furthermore, the average spatial profile of temperature increments detected orthogonally to the ink stripes satisfies the 26.4% contrast threshold required by the well-known Rayleigh’s criterion55, thereby quantifying in ≤60 μm the achieved resolution (Fig. 2h). This corresponds to a 5.8-time gain relatively to the diffraction-limited prediction, and a 20-time gain with respect to the (1200 ± 180)-μm thermo-camera resolution in conventional operation (Supplementary Note 3 and Supplementary Fig. 8).

**Sub-diffraction thermal imaging on biological samples.** Sub-diffraction thermal imaging has been finally tested on biological systems with sparse distributions of visible-light absorbers (Fig. 3). Explanted murine skin biopsies have been treated with 30-nm Prussian blue nanocubes (PB-NPs), which exhibit a strong absorption band centered at 700 nm and a highly efficient thermal relaxation upon VIS/nearIR irradiation49,50. After experimental characterization of the sample emissivity and of its heterogeneity (Supplementary Note 5 and Supplementary Fig. 9), the temperature increments induced by modulated illumination and localized with the two-step Gaussian fitting procedure have been exploited to provide a super-resolution spatial map of the distribution of the nanostructures inside the tissue (Fig. 3b). Details as close as ~40 μm appear resolved under the adopted imaging conditions, as highlighted by the magnified area in Fig. 3c. It is to be noted that the lateral size of the magnified region corresponds to the thermo-camera theoretical 345-μm diffraction-limited resolution. The achieved resolution gain appears even larger when the reconstructed images (Fig. 3b, c) are compared to the conventional thermographic image obtained as the temporal maximum projection over the raw thermo-camera image stack (Fig. 3d).

A high degree of co-localization has been observed between the temperature variations in the super-resolution image and the lowest-intensity pixels in the corresponding transmitted-light image of the same sample, ascribed to PB-NPs (Fig. 3a–c and Supplementary Fig. 10). This highlights the specificity of thermal detection and proves the possibility of imaging NPs at sub-diffraction resolution in a heterogeneous environment. This conclusion is reinforced by the results obtained on nanoparticle-untreated samples under identical imaging conditions (Fig. 3f–h). In the absence of exogenous nanoparticles, nearly no temperature variation is detected. Only if the $\Delta T_{\text{min}}$ threshold is lowered from 0.3 °C (as adopted for treated samples in Fig. 3b) to 0.1 °C (corresponding to the thermal camera sensitivity), a few spurious temperature increases get identified (Fig. 3g and Supplementary Fig. 10).
We have proposed a photo-modulated thermal imaging technique that we have showed to overcome Abbe’s resolution limit for the emitted thermal wave on a conventional thermo-camera. We take advantage of a compact, easily aligned and low-cost (hardware cost ~20k€) benchtop microscope, where a continuous-wave laser beam is raster-scanned and synchronously modulated by a programmable shutter. By relying on the localization of the isolated induced temperature variations, our approach reconstructs the distribution of the absorptive centers in the sample and combines this morphological information with the quantitative measurement of induced temperature variations.

Sub-diffraction 60-μm resolution has been demonstrated by proof-of-principle experiments, thereby enabling a 5.8-enhancement with respect to the 345-μm diffraction-limited prediction and a 20-time enhancement with respect to the 1200 ± 180 μm effective resolution of our thermal camera in conventional operation. The achieved resolution already proves sufficient to accurately map the distribution of light-absorbing nanostructures injected into explanted murine skin biopsies.

We have shown that, with single-spot scanning, a 3.7 × 1.8 mm² area can be scanned with 40-μm pixel size in 90 min, and this time reduces to 20 min with a 75-μm scan pixel size. Extended acquisition times are common to all the super-resolution techniques (e.g., PALM and STORM) that rely on a stochastic molecular switching and readout\cite{46,47,56}. A number of bio-relevant applications would benefit from the achievement of super-resolution spatial information while not necessarily requiring transient thermal imaging at high temporal resolution.
we can envision exemplary applications for our imaging approach in the characterization of the distribution of metallic nanoparticles in explanted bioplates for the optimization of photothermal therapies, or in the development of active thermography ex-vivo pre-clinical protocols in the context of melanoma screening and diagnosis. Whenever beneficial for in-vivo biological applications, our current imaging time can also be shortened by multi-spot illumination (via a Spatial Light Modulator) and/or by relaxing the constraint on the observation of individual, isolated temperature peaks. Classes of algorithms allow handling multiple and overlapping signal peaks, based on deep-learning, sequential fitting, sparsity, or maximum-likelihood estimation, and could be adopted to speed up the imaging process\cite{89, 90}. Methods developed for ghost imaging\cite{91} could be borrowed as well, especially for those applications (e.g., in cryptography) that require the recognition of an encrypted but known pattern.

By taking advantage of the full analytical description of the space-time dependence of the photo-activated temperature spots imaged on the thermal camera (Supplementary Note 5), and by properly treating both the sample transparency to infrared wavelengths and its heterogeneity in terms of emissivity and thermal diffusivity (Supplementary Note 5), we should also be able to combine the reconstruction of super-resolved images to the quantitative characterization of the sample thermal properties. The possibility of producing 2D maps of (spatially heterogeneous) diffusivity values is currently being investigated.

The voltage dependence of the path length has been quantified with a relative 60% absorbance at the 633-nm wavelength we employ for photo-thermal imaging. The laser source consists in a 633 nm He-Ne laser beam (power P = 10 μW on the sample plane), which is focused on the sample by a 20 x 0.5 NA, air objective (HCX PL Fluotar, Leica Microsystems, D). Images have been acquired by detecting the transmitted light signal with a non-spectral dedicated photo-multiplier tube, with no confocal pinhole along the detection optical path. A 400-Hz raster scan frequency per line has been adopted, and millimeter-sized sample regions have been imaged using a tile-scan acquisition mode.

**Methods**

**Optical setup.** Supplementary Fig. 3a shows a schematic of the experimental photo-thermal imaging setup. Sample absorption is primed by a He-Ne laser beam (λ_{max} = 633 nm, Thorlabs, NJ, USA) with 30-mW maximum output power. Gray filters are employed to adjust the laser power on the sample plane and achieve a minimum 0.1 °C temperature variation with a relatively short illumination time t_{on} (10–100 ms). The excitation spot size on the sample plane may be adjusted by a two-lens Keplerian beam reducer with focal lengths f1 and f2: the 1/e^2 beam diameter ranges from 56 ± 2 μm in the absence of the beam reducer down to 22 ± 1 μm with f1 = 3 cm and f2 = 10 cm (Supplementary Fig. 7).

A scanning unit with servo-board electronics and a pair of galvanometric mirrors (Cambridge Technology Inc., MA, USA) is coupled to a two-lens scan system (f1 = 4 cm, f2 = 10 cm), followed by beam fociation (f1 = 3 cm) onto the sample plane. The mirror positioning system is driven by an Arduino Uno microcontroller board and interfaced with a custom written Python code. Mirrors are operated in conventional raster-scanning mode (typical pixel size, 10–50 μm on the sample plane), with user-defined voltage values applied to the mirrors to regulate the scan path length along the horizontal and vertical axes. The voltage dependence of the path length has been quantified following the calibration procedure described with Supplementary Note 6 and Supplementary Fig. 13.

The Arduino board and Python code used to drive the scan system are also exploited to synchronize the galvanometric mirrors with the electronic shutter (Oriel 76992, Newport, CA, USA; 150 Hz maximum operation frequency) aimed at modulating the laser illumination in time. During laser de-activation (i.e., shutter closure), a default 2-ms pixel dwell time of the laser beam along the scan grid is usually followed by an activation, the time showning is the time desired by the user, and the laser pixel dwell time is temporarily increased to the same value to ensure the sample illumination occurs at the same location. Since neither the thermal relaxation time nor the sample characteristic heating time is significantly affected by the sample thermal diffusivity (Supplementary Fig. 1a), a simple practical criterion can be adopted to choose t_{rel} irrespectively of the sample thermal properties: based on the desired temperature peaks amplitude (i.e., the desired signal-to-noise ratio, which will impact in turn on the peaks localization uncertainty and the achievable resolution), we adopt the minimum laser activation time and the maximum laser power that ensure the achievement of such a desired signal amplitude while avoiding any sample photo-damage. Once the laser activation time has been set, the spatial distance between pairs of consecutive illumination events is specified according to the sample thermal properties following the criteria we detail in Supplementary Note 2.

The detection of thermal radiation is performed by an uncooled microbolometer-based thermal camera (FLIR E40, FLIR Systems Inc., OR, USA) with 30-Hz frame rate. The detector provides 320 × 240 images, with 25° x 19° FOV and ~400 μm pixel size on the sample plane. As detailed with the measurement procedure in Supplementary Note 5, the thermo-camera automatically corrects for the atmospheric attenuation and the sample reflection of the thermal radiation emitted by the surroundings\cite{92}. To this aim, the sample distance d from the camera front lens, the sample emissivity ε, and the relative humidity h and temperature of the atmosphere have to be provided to the camera software. h = 50% has been assumed for all the experiments of the present work, whereas the sample distance d (~30 cm) and the ambient temperature have been measured in each imaging experiment. The emissivity ε of the sample has been characterized by the black-tape method\cite{93} for both microfiche ink samples (ε = 0.8) and biological specimens (ε = 0.95) as described in Supplementary Note 5. Finally, imaging of a known-size object at constant temperature has been exploited before each calibration to calibrate the image and will take small variations associated to the thermo-camera positioning into account.

All the experiments have been performed with a maximum 20° vertical tilt of the thermal camera with respect to the sample (Supplementary Fig. 14). Such a thermal camera tilt does not impact on measured ΔT max values (Supplementary Note 5), nor it impacts on the size of imaged objects, the foreshortening effect of view being excluded by the experimental results reported in Supplementary Fig. 15.

**Transmission microscopy setup.** A Leica TCS SPS STED-CW scanning confocal microscope (Leica Microsystems, D) has been employed for transmitted-light imaging. The laser source consists in a 633 nm He-Ne laser beam (power P = 10 μW on the sample plane), which is focused on the sample by a 20 x 0.5 NA, air objective (HCX PL Fluotar, Leica Microsystems, D). Images have been acquired by detecting the transmitted light signal with a non-spectral dedicated photo-multiplier tube, with no confocal pinhole along the detection optical path. A 400-Hz raster scan frequency per line has been adopted, and millimeter-sized sample regions have been imaged using a tile-scan acquisition mode.

**Prussian blue nanoparticles.** PB nanoparticles have been prepared according to the synthesis protocol reported in refs. 95, 96, starting from 10 mM FeCl3 (Fe(III)), 10 mM K4[Fe(CN)6] (FeII) and 0.025 M citric acid reagents. Solutions have been heated to 60 °C during the synthesis and then allowed to cool at room temperature. Purification has been performed by ultracentrifugation for 25 min at 13,000 rpm, followed by pellet re-suspension in half the original volume.

The synthesis leads to nanocrystals with cubic shape and an average side of (29 ± 8) nm, measured by TEM imaging. The absorption band peaks at 700 nm, with a relative 60% absorbance at the 633-nm wavelength we employ for photo-thermal imaging.

**Ink samples.** Microfiche printing has been employed to produce reference ink patterns with convenient shape and known sub-resolved size for the initial experimental demonstration of super-resolution photo-thermal imaging. Patterns have been designed (PowerPoint, Microsoft Office) as white shapes over a black background (microfiche printing produces the negative of the original drawing). The pattern linear dimensions have been selected based on the desired size on the final image (10 × 8 cm), taking into account the contribution of the reduction factor to the final image size, as well as the thermo-camera positioning into account. Imaging by conventional transmission microscopy at 633 nm allows inspecting the uniformity of the printed ink layers as in Fig. 2a, f.

**Biopsies.** Murine skin biopsies have been explanted from the flank derma of sacrificed Balb/c mice at 6 weeks of age. Explanted tissue sections have been embedded in OCT freezing medium (Bioptica, I), cut in 10-μm sections on a cryostat and adhered to glass slides (Superfrost Plus, Thermo Fisher Scientific, MA, USA) for photo-thermal imaging. When necessary, Prussian blue nanoparticles (100 μL, 10x concentration of the initial stock) have been injected into the biopsies right after the explant. All the experiments have been performed under protocols approved by the Institutional Animal Care and Use Committee of the University of Milano-Bicocca and the Italian Ministry of Health.

**Image reconstruction.** Raw data for super-resolution thermography consist in an image sequence of ~10^6 - 10^7 frames acquired by the thermal camera during the modulated laser illumination of the sample. Due to the ~10 × 8 cm² field of view of the thermal camera employed here, the analysis is restricted to a Region Of Interest (ROI) covering the scanned area and easily identified by computing the maximum projection over the time axis of the acquired image stack. For each pixel i within the ROI, the temperature-versus-time profile T(t) of the laser spot may contain N temperature peaks with approximately exponential rise and decay. Each peak is produced by a laser square pulse impinging on light-absorbing and heat-releasing entities located within the i-th pixel or in pixels nearby (due to heat diffusion). Since the scan pixel size is typically several times the thermal camera pixel size (i.e., δx ~ 10–100 μm, δy ~ 400 μm), tens to hundreds of peaks can be detected in a single T(t) plot; peak amplitudes vary according to the distance of the illuminated heat-releasing objects from pixel i and from the center (highest-intensity portion) of the excitation laser spot at the time of illumination. A pattern-recognition algorithm can be exploited to locate the peaks, fit the experimental data on the theoretical model for the short- and long-time behavior of the temperature rise and decay for the short adopted t_{rel} = 0.3–1 s, and perform a fit to a 1D symmetric Gaussian function that (reasonably) approximates the exponential decay for both the short and long-time behaviors.
Data analysis software. Photo-thermal imaging raw data have been acquired with the thermo-camera’s software (FLIR Tools +, FLIR Systems Inc., OR, USA), exported in .csv file format and entirely processed by a custom written Python code. Both non-linear 1D fits of temperature time profiles and the symmetric, asymmetric and skewed Gaussian surface fits of detected 2D temperature peaks have been performed by the code with the curve fitting routine of the Scipy open source Python tool. Reconstructed super-resolution images have been saved as .txt files and rendered with ImageJ (US National Institutes of Health, MD, USA) for visualization and look-up table adjustment. Their comparison with the results of transmitted-light imaging has been accomplished by the Origin Pro 8 software (Origin Lab Corporation, MA, USA) based on temperature and intensity profile plots.

Data availability statements prepared and analyzed in the present study are available from the corresponding author on reasonable request.

Code availability. The custom code employed for the analysis of the datasets acquired in this study is available from the corresponding author on reasonable request.

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Author contributions

Ma.B. designed the experiments, analyzed the data, and wrote the paper. Ma.B. and M.M. acquired the data, conceived, and wrote the data analysis code. A.Z., M.M., and Ma.B. built and tested the experimental setup. M.C. and G.C. conceived the technique, supervised the project and wrote the paper. P.P. and My.B. provided Prussian blue nanoparticles. F.M. and F.G. prepared and provided biological samples. L.S. helped with biological samples and related data analysis. I.D. wrote the paper and discussed the project.

Competing interests

The authors declare no competing interests.

Additional information

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