Letter

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Gold Nanoparticles as Absolute Nanothermometers

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- 4 Supporting Information

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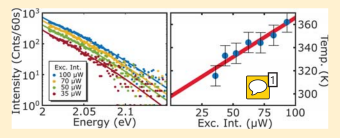
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ABSTRACT: Nanothermometry is a challenging field that can open the door to intriguing questions ranging from biology and medicine to material sciences. Gold nanorods are excellent candidates to act as nanoprobes because they are reasonably bright emitters upon excitation with a monochromatic source. Gold nanoparticles are commonly used in photothermal therapy as efficient transducers of electromagnetic radiation into heat. In this work we use the spectrum of the anti-Stokes emission from gold nanorods irradiated in resonance to measure the absolute temperature of the



nanoparticles and their surrounding medium without the need for a previous calibration. We show a 4 K accuracy in the determination of the temperature of the medium with spectral measurements of 180 s integration time. This procedure can be easily implemented in any microscope capable of acquiring emission spectra, and it is not limited to any specific shape of nanoparticles.

KEYWORDS: Gold nanorods, plasmon, anti-Stokes, sensing, temperature

ost physical, chemical, and biological processes depend on temperature. Together with the miniaturization of devices and the advent of nanotechnology, the need for measuring temperature with high spatial accuracy started to emerge. Notably in biology^{1,2} and medicine³ measuring and controlling temperature at a subcellular scale are the challenges that must be overcome to achieve better understanding and control of new therapies such as photothermal tumor ablation⁴ or controlled drug delivery.^{5,6}

Nanometer-size probes with distinctive spectral features are 30 ideal candidates for temperature measurements since they 31 provide high spatial accuracy while far-field optics allow a 22 noncontact readout. Some of the proposed strategies include 33 structures that undergo a conformational change upon an 34 increase in temperature, 7 thus inducing variations in 35 fluorescence intensity of a dye molecule embedded in them.

Also cleverly designed lanthanide-based fluorescent probes in which the ratio of particular emission peaks depends on temperature provide a high accuracy and can be used as nanothermometers even in biological samples. Photobleaching is often an important limitation of these approaches. Recently, surface enhanced Raman spectroscopy (SERS) allowed to measure spectral changes induced by temperature down to single molecules, to but a careful calibration of the measurements is crucial.

45 Gold nanoparticles continue to receive a fair amount of 46 attention because of their unique optical properties. ¹¹ The 47 collective oscillation of conduction electrons, also known as 48 surface plasmon, shows a resonance in the visible to near-49 infrared wavelengths. This resonance can be tuned by changing 50 the shape of the particles ¹² and is responsible for a large 51 absorption and scattering cross section at the resonance wavelength. These cross sections can be calculated by solving 52 the Maxwell equations numerically employing different 53 computer packages, 13-15 providing a good agreement between 54 calculations and what is experimentally achievable. 55

Thanks to their high absorption and scattering cross section 56 (several times higher than their geometrical cross section), it is 57 relatively simple to detect nanoparticles in a dark-field 58 scattering configuration 16 or via photothermal imaging. 17,18 59 Alternatively, detecting gold nanoparticles through their 60 luminescence 19 is also possible; their low quantum 61 yield, 20–23 around 10⁻⁶, is compensated by the enhanced 62 cross section at the surface plasmon resonance (SPR). The 63 luminescence signal is stable over time; gold nanoparticles do 64 not blink nor bleach and therefore are useful labeling agents for 65 processes that require extended periods of observation. 24

Different metallic nano-objects are being introduced as 67 agents for photothermal therapy^{5,25} or drug delivery.²⁶ One of 68 the advantages of gold nanoparticles is the possibility of tuning 69 their resonance to the near-infrared range, where the 70 penetration of light into tissues can be of several 71 centimeters.^{3–5,25,27,28} Moreover, the particles can be used 72 not only for treatment, but also for imaging.^{5,29} In the case of 73 photothermal therapy, nanoparticles are used as heat 74 sources^{4,27} to locally increase the temperature in order to 75 induce the death of specific cells in a tissue.^{5,25} However, the 76 temperatures reached³⁰ can only be estimated from models²⁹ 77 or from an ad-hoc calibration. Therefore, a method to 78 simultaneously increase and monitor the local temperature

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80 will be of great interest in a broad range of fields. 1 this Letter,
81 we show that the anti-Stokes luminescence of single gold
82 nanorods can be used to measure their temperature upon
83 resonant CW irradiation. The temperature error for a single
84 measurement is less than 10 K with an acquisition time of a
85 few minutes; with a set of such measurements, the temperature
86 of the surrounding medium can be determined with 4 K
87 accuracy or better.

Luminescence of metallic nanoparticles has been the subject 89 of extensive study in recent years. Since the first observation of 90 luminescence from bulk gold,³¹ different groups have tried to 91 quantitatively describe the observed properties, ^{32–36} such as 92 the quantum yield^{20–23,37} and the emission spectrum. ³⁸ In 93 particular, gold nanorods present two distinct resonance 94 energies, namely, the transverse and the longitudinal plasmon 95 resonances. These particles can therefore be excited efficiently 96 at one of those energies; the transverse resonance corresponds 97 to a wavelength of about 530 nm and will give rise to a broad luminescence emission with a peak at the longitudinal plasmon energy. Conversely, it is possible to excite the particles with a 100 wavelength matching the longitudinal plasmon resonance. In 101 this case, the excitation benefits from an enhanced absorption 102 cross section, but the emission that overlaps the plasmon 103 resonance will be mostly blocked by the filters needed to 104 prevent direct excitation light from reaching the detectors.

In this work, we call "photoluminescence" any secondary 106 light emission³⁹ at energies different from the excitation laser 107 energy, $\hbar\omega_{\rm L}$. After absorption (we do not specify whether the 108 absorption is real or virtual) of an excitation photon, the 109 excited electronic state^{31,37} may interact and exchange energy with the phonon bath or, in the case of metals, with the bath of thermally excited charge carriers around the Fermi level. After a number of interactions, the excited electronic state will reemit a photon, which can possess a lower or higher energy than that of the excitation photon. $^{40-42}$ For a nonresonant excitation, the probability of more than one interaction is 116 negligible, and the main contribution to secondary emission is Raman scattering.³⁴ This is the case, for example, of insulators excited well below their electronic absorption edge. For 119 resonant excitation, a relatively long-lived excited state is 120 prepared. It will have enough time to interact repeatedly with 121 thermal baths, particularly with phonons. This is the case of 122 organic dye molecules or semiconductors in which relaxed 123 fluorescence is observed. We also note that fluorescence always 124 presents hot bands on the anti-Stokes side of the excitation 125 laser. In most fluorescence detection schemes, however, these 126 hot bands are ignored, but they are far from negligible in 127 heavily doped samples.⁴³

Metal nanoparticles fall between those two extremes because the excited electronic state, an electron—hole pair, relaxes very rapidly by interacting with other charge carriers and with phonons. The photoluminescence lifetimes are on the order of tens of femtoseconds, and therefore, there is not enough time to obtain a fully relaxed luminescence. In other words, the photoluminescence is always "hot". It is worth noting that Raman scattering, corresponding to the lowest order of interaction with baths, will be an important contribution to photoluminescence. However, second and higher orders may also contribute significantly. Because all these processes obey a Boltzmann-type relationship between anti-Stokes and Stokes emission, they cannot be easily distinguished from each other on the basis of their temperature dependence.

The anti-Stokes emission is highly sensitive to temperature, 142 and thus, it can be used for thermometry. The In this Letter, we 143 present a simple procedure to extract the absolute temperature 144 from the anti-Stokes photoluminescence spectrum of individual gold nanorods without the need of any previous 146 temperature calibration. We show that we can determine the 147 particle temperature *in situ* with an accuracy of 6% by 148 recording a single anti-Stokes spectrum (with an acquisition 149 time of 3 min). Moreover, by performing this measurement at 150 different excitation powers we can obtain the temperature of 151 the surrounding medium with an accuracy better than 2%.

Phenomenological Model for the Luminescence 153 Emission. In a nutshell, we consider the luminescence 154 emission as radiative recombination of electron—hole pairs 155 created by the decay of the plasmon, after their interaction 156 with thermal baths. Before the recombination, carriers may 157 interact with the baths one or more times, leading to secondary 158 light emission with an energy different from the initial internal 159 energy of the pair. The anti-Stokes spectral contribution arises 160 from interactions that increase the energy of the pair, whereas 161 the Stokes emission corresponds to a decrease in energy. The 162 emission process will be enhanced by the surface plasmon; 163 therefore, the luminescence spectrum will be modulated with 164 the plasmon shape. A schematic representation of these ideas is 165 shown in the Supporting Information.

Exciting a gold nanorod with a monochromatic beam at its 167 resonance frequency, ω_{SPR} , generates a collective oscillation of 168 the gas of conduction electrons, also called a plasmon. The 169 plasmon decays by forming a pair of hot electron and hole with 170 an internal energy equal to the exciting photon energy, 46-48 171 i.e., $E_{a-b} = \hbar \omega_{L}[2]$ 172

i.e., $E_{\rm e-h}=\hbar\omega_{\rm L}$ [3] 172

This hot electron—hole pair has a small probability of 173 recombining radiatively, i.e., of re-emitting its high electronic 174 energy as a photoluminescence photon. If the electron and 175 hole have interacted only with static surfaces or defects, their 176 energies will be the same. Therefore, the emitted photon will 177 have the same energy as the incoming photon and will not 178 contribute to the measured photoluminescence. It will be 179 blocked by the notch filter used to remove the exciting laser 180 from detection. If, however, the electron and hole have 181 interacted with a phonon or a thermally excited electron or 182 hole, they may have lost or acquired energy. In both cases the 183 energy available upon recombination cannot much exceed $5\omega_{\rm L}$ 184 $+k_{\rm B}T$, where $k_{\rm B}$ represents Boltzmann's constant and T the 185 absolute temperature.

Radiative recombination gives rise to emission spectrally and 187 spatially distributed throughout the particle over a broad 188 frequency band with an exponential cutoff at $\hbar\omega_{\rm L}$ + $k_{\rm B}T$. The 189 weak recombination emission can be greatly enhanced by the 190 surface plasmon resonance, acting as an antenna. With this 191 model, the following predictions can be made. First, the 192 emission spectrum must follow the plasmon spectrum if the 193 excitation laser is well above the plasmon resonance as shown 194 in Figure 1, green line. If the excitation falls within the plasmon 195 fl resonance, the spectrum is expected to follow the plasmon 196 spectrum multiplied by a Bose-Einstein statistics factor arising 197 from phonon population (here, we assume that the coupling to 198 the phonons dominates the process, while refs 34 and 36 199 assume that carrier-carrier interactions dominate). Thus, 200 under our assumption, the emission should be proportional 201 to the phonon occupation number \overline{n} for anti-Stokes and \overline{n} + 1 202 for Stokes processes, with

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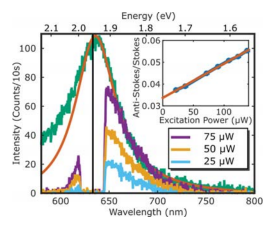


Figure 1. Luminescence emission spectra of a single gold nanorod. The green curve is the measured luminescence emission under 532 nm excitation, and the red curve shows the extracted $I_{\rm SPR}(\omega)$ from this spectra. The other curves are the emission of the same particle under 633 nm irradiation at three different powers indicated in the legend. The inset shows the anti-Stokes-to-Stokes ratio as a function of the excitation power, overlapped with a linear fit in red. The dip centered on the laser wavelength is caused by the notch filter used to prevent the excitation laser from reaching the detectors. For these spectra, the acquisition time was 10 s.

$$\overline{n} = \left(\exp\frac{\hbar\Omega}{k_{\rm B}T} - 1\right)^{-1} \tag{1}$$

where $\hbar\Omega$ is the phonon energy. If carrier—carrier interactions dominate, the occupation number world obey Fermi statistics,

7 i.e.,
$$n = \left(\exp\left(\frac{\hbar\Omega}{k_{\rm B}T}\right) + 1\right)^{-1}$$

With this model, we can also predict that the emission should be polarized. For the strong longitudinal plasmon of gold nanorods, this polarization coincides with the longitudinal axis of the particle. Moreover, the lifetime should be determined by the lifetime of hot electrons and holes and should be significantly shorter than the thermalization time of the carriers. Indeed, a few interactions would suffice to reduce the carriers' energy significantly, and therefore, the electron and hole would not have the energy required to produce an optical photon. One important assumption for this model is that the emission spectrum of radiative recombination is much plasmon resonance should excite the electron—hole pairs with leave the same efficiency as well above the plasmon resonance.

Application to Nanothermometry. According to the model just described, the anti-Stokes emission spectrum follows the form

$$I(\omega) = I_{\text{SPR}}(\omega) \left(\exp \frac{\hbar(\omega - \omega_{\text{L}})}{k_{\text{B}}T} - 1 \right)^{-1}$$
 (2)

227 where $I(\omega)$ is the emitted intensity, ω is the angular frequency 228 of the photons, $\omega_{\rm L}$ is the frequency of the exciting laser, and 229 $I_{\rm SPR}(\omega)$ is the surface plasmon resonance spectrum. The only 230 remaining free parameter is the temperature T (and a 231 normalization constant not included in eq 2).

The procedure we propose to obtain the absolute temperassauture T of gold nanorods from the anti-Stokes luminescence emission without the need of any previous temperature 234 calibration involves the following steps:

1. Obtain the surface plasmon resonance spectrum of the 236 particle. This is usually expressed as a Lorentzian 237 function, 11 i.e., 238

$$I_{SPR}(\omega) = \frac{(\Gamma/2)^2}{(\omega - \omega_{SPR})^2 + (\Gamma/2)^2}$$

where ω is the photon energy frequency, $\omega_{\rm SPR}$ is the $_{239}$ resonance frequency, and Γ is the width of the surface 240 plasmon resonance. In our case, we detect the spectrum 241 of photoluminescence excited at 532 nm to extract $\omega_{\rm SPR}$ 242 and Γ , with the procedure explained in the Supporting 243 Information.

- Excite near the longitudinal plasmonic resonance and 245 detect the blue-shifted anti-Stokes emission spectra. For 246 this we employed a 633 nm laser as a source.
- 3. Fit the high-energy part of the spectrum using eq 2 with 248 *T* as the *only* free parameter. 249

We emphasize that we cannot simply use the anti-Stokes to 250 Stokes intensity ratio to obtain the temperature of the particle, 251 as is commonly done with Raman lines of molecules, 50,51 due 252 to the presence of the strong plasmonic enhancement of the 253 emission that must be considered in addition to the Boltzmann 254 factor.

Experimental Methods. All the measurements in this 256 work were performed with a home-built confocal microscope 257 equipped with a spectrometer (Acton 500i) in the emission 258 path. We focused our lasers to a diffraction-limited spot using a 259 60×, NA 1.4 oil immersion objective (Olympus) 22d collected 260 the emitted photons through the same objective. This provided 261 high excitation and collection efficiency. We employed a 532 262 nm (CNI) laser for characterizing the nanorods' plasmon and a 263 633 nm HeNe (Thorlabs) to excite the nanorods in resonance. 264 The intensity of the laser was controlled via an acousto-optic 265 modulator. We provide a scheme of the setup and more details 266 in the Supporting Information.

Wet-chemically synthesized nanorods 52 with average dimen- 268 sions of 21 M \times 50 nm and a plasmon resonance around 650 59 nm were spin-coated onto clean coverslips, controlling the 270 superficial concentration to separate individual nanorods 271 within the diffraction-limited spot. 11 The characterization of 272 the gold nanorods sample with transmission electron 273 microscopy (TEM) and the ensemble extinction spectrum 274 are presented in the Supporting Information.

The samples were mounted in a flow cell that allowed us to 276 increase the temperature of the medium up to 60 °C and to 277 monitor it through a Pt100 resistance thermometer placed 1 278 mm away from the observation area. To compensate for the 279 drift of the setup while increasing the temperature, we 280 developed a computer program to continuously track a 281 reference particle. The same program was responsible for 282 recording the temperature and triggering the spectrometer. In 283 this way, complete data sets were acquired at different 284 temperatures, with excitation at 532 and 633 nm, at different 285 laser intensities. A spectrum with 532 nm laser excitation was 286 taken after every cycle to ensure that the particle under study 287 had not reshaped due to high excitation power.

When recording anti-Stokes spectra to extract the temper- 289 ature of the particles, accumulations of each spectrum were 290 recorded with an exposure time of 30 87 his not only allowed 291 us to lower the noise of the measurement because of a longer 292

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293 exposure time B min in total), but also allowed us to remove 294 bright pixels generated by cosmic rays. Having several 295 accumulations is also useful to monitor changes in the 296 intensity of the spectra during the acquisition itself. These 297 changes can be due to a drift of the setup while measuring or 298 to reshaping of the particle. If the reshaping was confirmed by 299 comparing the spectra acquired with the 532 nm laser, 53 the 300 measurements where rejected. If the changes in the observed 301 emission spectra were due to drift of the setup, the particular 302 data set was not taken into account. For the purposes of this 303 work, the excitation intensity is crucial for characterizing the 304 method; if the particle is not in focus, it would result in an 305 overestimation of the excitation power.

Results. The proposed model for the anti-Stokes emission requires the plasmon spectrum $(I_{\rm SPR}(\omega))$ in eq 2) in order to fit 308 the emission at shorter wavelengths and extract the particle 309 temperature. It has been shown that both scattering and 310 luminescence spectra roughly overlap over a broad range of 311 wavelengths. Therefore, exciting gold nanorods with 532 nm 312 allows us to record the longitudinal plasmon spectra, as shown 313 in the green solid curve of Figure 1. It has to be recalled that 314 the luminescence spectrum is not a perfect Lorentzian since 315 there is a broadband contribution also observed in bulk gold. The procedure to extract the SPR profile from such a 317 measurement is explained in the Supporting Information. We 318 show in Figure 1 the extracted surface plasmon profile in the 319 red solid curve.

The other curves in Figure 1 show the luminescence 211 emission of the same nanorod with irradiation at 633 nm at 322 different powers, ranging from 25 to 75 μ W at the back 323 aperture of the objective. The vertical black line shows the 324 wavelength of the laser. The Stokes part of the spectrum at 325 longer wavelengths than the excitation shows the same shape 326 as the plasmon emission observed under 532 nm excitation, 327 apart from a normalization factor. From the figure, it can 328 readily be seen that the shape of the anti-Stokes emission, at 329 shorter wavelengths than excitation, is exponential-like and 330 does not follow the Lorentzian shape of the Stokes emission. 331 The dip between Stokes and anti-Stokes is caused by the notch 332 filter that prevents direct excitation light from reaching the 333 detectors.

The inset of Figure 1 shows the anti-Stokes-to-Stokes ratio of the integrated luminescence for different laser excitation intensities. It is possible to see that even though the photoluminescence process is linear, the anti-Stokes intensity increases slightly more rapidly than the Stokes emission. We already exploited this phenomenon to image gold nanorods in high-background conditions. For more information on the power dependence of both the anti-Stokes and Stokes luminescence, please refer to the Supporting Information.

To further characterize the anti-Stokes emission in gold manorods, we measured the emission for 105 nanorods with different plasmon resonances under the same 633 nm 46 excitation and calculated the ratio of integrated anti-Stokes to Stokes emissions. Figure 2 shows the experimental ratios as blue circles versus the surface plasmon resonance (SPR) of the particle. The vertical black line marks the laser wavelength. The particles measured had resonances between 600 and 690 mm; the ones showing the maximum ratio of anti-Stokes to Stokes are those with a resonance to the blue of the laser. For these particles, the longitudinal plasmon is enhancing preferably the anti-Stokes emission. For particles with a resonance at the laser wavelength, the anti-Stokes and the

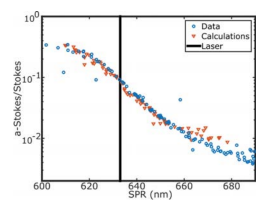


Figure 2. Characterization of anti-Stokes emission for different surface plasmon resonance. Ratio of the anti-Stokes to Stokes emission under 633 nm excitation as a function of the resonance wavelength of the particle. The blue circles are experimental results (105 different nanorods), while the red triangles are the results of numerical simulations with eq 1 (82 nanorods with different aspect ratio). There is a very good agreement between experiment and calculations. Particles with a resonance to the blue of the laser (indicated by the vertical black line) have an increased anti-Stokes emission.

Stokes emission have similar enhancement and show a ratio 356 close to 10%.

Figure 2 also shows as red triangles the results of numerical 358 calculations showing an excellent overlap between the 359 measured and the calculated data. To obtain these results, 360 we numerically calculated the absorption cross section of 82 361 particles with the ADDA package using a fixed width of 21 362 nm and different lengths to achieve different SPR wavelengths. 363 Each calculated absorption spectrum was fitted by a Lorentzian 364 and used as $I_{\rm SPR}(\omega)$ in eq 2. Assuming a diffraction-limited 365 laser spot and using the calculated absorption cross section, we 366 calculated the temperature of the particle. This value was used 367 in eq 2 to compute the anti-Stokes emission spectrum. The 368 Stokes emission was set proportional to the excitation power 369 with a shape given by the calculated absorption spectrum. 370 Since both anti-Stokes and Stokes emissions are proportional 371 to the excitation power, this term cancels out when computing 372 the ratio. The laser power therefore only enters into the 373 equation when calculating the temperature of the particles. It is 374 remarkable that the agreement between data and calculations 375 was achieved without free parameters, solely taking into 376 account the transmission spectra of the filters.

We then use our protocol to extract the temperature of an 378 individual nanorod. The anti-Stokes spectra shown in Figure 1—22 were recorded at four different excitation intensities, while the 380 full lines are the fits using our model from eq 2; again, there is 381 an excellent agreement between data and model. For every 382 anti-Stokes measurement we have also acquired the full 383 plasmon spectrum exciting with a 532 nm laser before and 384 after the temperature extraction, to verify that the particle did 385 not reshape while being excited at resonance.

The inset in Figure 3 shows the temperatures resulting from 387 fs the fits at different irradiation intensities (blue dots). Note that 388 the absolute temperatures of the particle at each excitation 389 power were calculated without any calibration. As expected, 390 the temperature of the nanorod varied linearly with excitation 391 intensity or equivalently with the absorbed energy. Thus, this 392 method provides an *in situ* way to measure the temperature 393 reached by nanoparticles when they are excited with resonant 394

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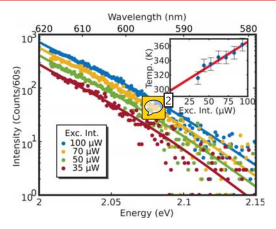


Figure 3. Anti-Stokes emission of a single nanorod at different irradiation powers. We used the model from eq 2 to fit the experimental data. There is an excellent agreement between data and model. The inset shows the extracted temperature at each power (blue dots) and a linear extrapolation of the data to 0 μ W excitation power. The value obtained for room temperature was 293 K, while the measured value was 296 K.

395 monochromatic light. Additionally, from these data sets it is 396 also possible to calculate the temperature at 0 μW excitation 397 power, i.e., room temperature, by extrapolating the results with 398 a linear fit. The value we obtained in this case is 293 ± 6 K, 399 while room temperature was 296 K, a 2% accuracy.

The accuracy of the obtained temperature depends on the 401 signal-to-noise ratio of the anti-Stokes spectra and on the 402 accurate modeling of the photoluminescence. The first step in 403 the protocol is the determination of the surface plasmon 404 spectral profile, $I_{SPR}(\omega)$ in eq 2. In this Letter, we obtained this 405 term by fitting an exponential background plus a Lorentzian to 406 the spectra obtained at 532 nm excitation. We note that this 407 choice was made for experimental convenience in our setup, 408 but other options to obtain the SPR profile are suited for the 409 procedure as well. The error bars in the inset of Figure 3 and in 410 the following figures are the result of the estimated variance in 411 the fit parameters for the anti-Stokes spectra using eq 2, step 3 412 in our protocol. More details about the error determination for 413 the final temperature extracted are given in the Supporting 414 Information.

As expected from the model, the anti-Stokes emission 416 depends not only on the particle's intrinsic properties but also 417 on the temperature of the surrounding medium.⁵⁴ In order to 418 further test this point, we changed the temperature of the sample in a controlled manner and recorded the luminescence emitted by a single nanorod.

For this set of experiments, we employed an air objective (60×, NA 0.9, Olympus) to avoid the presence of a heat sink 423 directly in contact with the observed area. We employed higher laser powers compensate for the lower excitation efficiency. 425 At each temperature, seven 6 ectra (with 30 s integration time 426 each) were acquired at different 633 nm excitation powers and 427 also a spectrum of the plasmon before and after each 428 measurement in order to monitor any possible reshaping of 429 the particles during the experiment.

Figure 4 shows the extracted temperature of a particle at 431 varying excitation powers and at different water temperatures. 432 The blue squares are the results of the measurement at 60 $^{\circ}$ C, 433 while the green crosses are measured at 40 °C and the yellow 434 circles at 20 °C. The full lines are fits with the same slope for

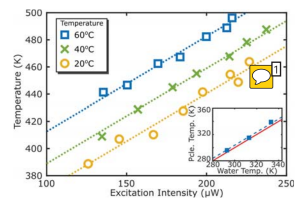


Figure 4. Calibration-free temperature measurement. Extracted temperatures from the anti-Stokes-luminescence emission of an individual nanorod at different excitation powers and at different sample temperatures. The dashed lines are fits with the same slope for the three temperatures. The squares in the inset plot show the local temperature of the sample obtained by extrapolating the temperature at zero excitation power as a function of the water temperature. The red line represents the expected curve if both temperatures are identical (equal). The dashed blue line is a fit to the data points with unit slope that shows a systematic offset of 3.8 K, a 1.2% difference.

the three sets of data. Using these fits we can extrapolate the 435 temperature value at zero excitation power to obtain the water 436 temperature in each case. The values obtained with the 437 extrapolation to 0 μ W excitation power were 296 \pm 4, 315 \pm 4, 438 and 339 \pm 4 K for water temperatures of 293, 313, and 333 K, 439 respectively. The inset plot in Figure 4 presents these points 440 and a red solid line with the expected curve if both 441 temperatures are identical. The dashed line shows a fit of the 442 data that evidence a small systematic offset of 3.8 K. This 443 represents an inaccuracy of 1.2%, which is a good result for a 444 calibration-free method.

Figure 4 clearly shows that the extracted temperature varies 446 with the temperature of the surrounding medium. More 447 strikingly, the method does not require any previous 448 calibration nor adjustment. Notably, the presented calibra- 449 tion-free procedure would allow us to perform the same 450 measurements in any other setup and could act as a reference 451 for calibration of other nanothermometers.

Being able to control and monitor temperature at the 453 nanoscale is of utmost importance in different fields ranging 454 from photothermal therapy⁵ to nanofabrication. 55 In this work, 455 we have shown a simple procedure that allows us to measure 456 the temperature of single gold nanorods irradiated by a 457 monochromatic continuous laser and without any previous 458 calibration. The level of accuracy of the temperature 459 measurement depends on several factors, but for a single 460 nanorod, it can be estimated to be better than 6-K3ith an 461 integration time of 3 14 n without any previous calibration.

The model employed for describing the anti-Stokes emission 463 takes into account the surface plasmon resonance of the 464 particles under study, which is responsible for enhancing the 465 emission, as well the electron-hole pair interaction with the 466 thermal baths. Particles with a resonance to the red of the 467 excitation wavelength would be more reliable in the temper- 468 ature extraction procedure but would also exhibit a lower 469 emission toward shorter wavelengths. The trade-off between 470 both effects and the possibility to fully characterize the 471 plasmon resonance will determine the specific particles that are 472 better suited for each application.

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A possible improvement of this technique would be the use 475 of nanostructures with a narrow shape distribution such as gold 476 bipyramids. 56 Such structures would be ideal candidates for temperature extraction since they present negligible size 478 dispersion, and thus, their plasmon can be measured in bulk 479 or determined from theory, avoiding the need of a second 480 excitation source.

The proposed method does not require any temperature 482 calibration since the only free parameter of the model is the 483 absolute temperature of the nanoparticle under study. 484 Moreover, the recording of the anti-Stokes spectrum is readily 485 achievable in any confocal microscope with a coupled 486 spectrometer. A 6 K accuracy may suffice for several 487 applications; it is important to point out that this value can 488 be improved in different ways: by carefully selecting the 489 particles that show the most favorable plasmon resonance; by 490 determining the plasmon resonance through white-light 491 scattering, reducing the uncertainty in the fit; and by increasing 492 the exposure times to increase the signal-to-noise ratio. A 493 cheaper alternative would have two detectors and band-pass 494 filters to detect the anti-Stokes and the Stokes emission 495 independently. However, this approach would require a 496 temperature calibration.

ASSOCIATED CONTENT

498 Supporting Information

499 The Supporting Information is available free of charge on the 500 ACS Publications website at DOI: 10.1021/acs.nano-

> -Stokes emission form gold nanorods, experimental Fup, gold nanorods sample characterization, determination of the error in the temperature extraction, and gold nanorod temperature numerical calculations (7)DF)

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512 Notes

513 The authors declare no competing financial interest.

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Anti-Stokes emission from gold nanorods, determination of surface plasmon resonance, experimental setup, gold nanorods sample characterization, luminescence power dependence, determination of the error in the temperature extraction, gold nanorod temperature numerical calculations.

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