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Article:

Palaferri, D, Todorov, Y, Bigioli, A et al. (11 more authors) (2018) Room-temperature nine- μm -wavelength photodetectors and GHz-frequency heterodyne receivers. *Nature*, 556. pp. 85-88. ISSN 0028-0836

<https://doi.org/10.1038/nature25790>

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Room temperature 9 μm photodetectors and GHz heterodyne receivers

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Room temperature operation is mandatory for any optoelectronics technology which aims to provide low-cost compact systems for widespread applications. In recent years, an important technological effort in this direction has been made in bolometric detection for thermal imaging¹, which has delivered relatively high sensitivity and video rate performance (~ 60 Hz). However, room temperature operation is still a major challenge for semiconductor photodetectors in the 8–12 μm wavelength band², and all developments for applications such as imaging, environmental remote sensing and laser-based free-space communication³⁻⁵ have therefore had to be realised at low temperatures. For these devices, high sensitivity and high speed have never been compatible with high temperature operation^{6,7}. Here, we show that a 9 μm quantum well infrared photodetector⁸, implemented in a metamaterial made of subwavelength metallic resonators⁹⁻¹², has strongly enhanced performances up to room temperature. This occurs because the photonic collection area is increased with respect to the electrical area for each resonator, thus significantly reducing the dark current of the device¹³. Furthermore, we show that our photonic architecture overcomes intrinsic limitations of the material, such as the drop of the electronic drift velocity with temperature^{14,15}, which constrains conventional geometries at cryogenic operation⁶. Finally, the reduced physical area of the device and its increased responsivity allows us to take advantage of the intrinsic high frequency response of the quantum detector⁷ at room temperature. By beating two quantum cascade lasers¹⁶ we have measured the heterodyne signal at high frequencies, above 4 GHz. These wide band uncooled detectors shall have therefore a significant impact on technologies such as multichannel coherent Gigabit/s data transfer¹⁷ and high precision molecular spectroscopy¹⁸.

37 An important intrinsic property of inter-subband (ISB) quantum well infrared photodetectors
38 (QWIPs) based on III-V semiconductor materials that has not yet been exploited is the very short
39 lifetime of the excited carriers. The typical lifetime is of the order of few picoseconds⁷, which
40 leads to two important consequences: the detector frequency response can reach up to 100
41 GHz, and the saturation intensity is extremely high (10^7 W/cm²)¹⁹. These figures are ideal for a
42 heterodyne detection scheme where a powerful local oscillator (LO) can drive a strong
43 photocurrent, higher than the detector dark current, that can coherently mix with a signal
44 shifted in frequency with respect to the LO. Notably, these unique properties are unobtainable
45 in infrared inter-band detectors based on mercury-cadmium-telluride (MCT) alloys, which have
46 a much longer carrier lifetime and therefore an intrinsic lower speed response^{2,20,21}. Yet, the
47 performance of all photonic detectors is limited by the high dark current that originates from
48 thermal emission of electrons from the wells, and rises exponentially with temperature, thus
49 imposing cryogenic operation (~ 80 K) for high sensitivity measurements. Previously, highly
50 doped ($\sim 1 \times 10^{12}$ cm⁻²)²², photovoltaic²³ 10 μ m QWIPs and QCDs²⁴ with large number of quantum
51 wells have been observed to operate up to room temperature, but only when illuminated with
52 powerful sources as CO₂ or free electron lasers.

53 In the present work, we show that this intrinsic limitation in QWIP detectors can be overcome
54 through use of a photonic metamaterial. We are able to calibrate our detector at room
55 temperature using a black body emitting only hundreds of nW, orders of magnitude smaller
56 than that required previously. To date, room temperature performance with values comparable
57 to those that we report here has only been demonstrated in the 3–5 μ m wavelength range,
58 using quantum cascade detectors (QCDs)²⁴⁻²⁶ and MCT standard detectors²⁷.

59 The photonic metamaterial structure is shown in Fig. 1a. The GaAs/AlGaAs QWIP⁸ contains $N_{qw} =$
60 5 quantum wells absorbing at 8.9 μ m wavelength (139 meV) that has been designed according
61 to an optimized bound-to-continuum structure from ref. 7. The absorbing region is inserted in
62 an array of double-metal patch resonators⁹⁻¹², which provides sub-wavelength electric field
63 confinement and act as antennas. The resonant wavelength is fixed by the patch size s through
64 the expression $\lambda = 2sn_{eff}$, where $n_{eff} = 3.3$ is the effective index⁹. The structures with $s = 1.3$ μ m
65 are thus in close resonance with the peak responsivity of the detector.

66 In our structure, the microcavity increases the device responsivity by a local field enhancement
67 in the thin semiconductor absorber¹⁰, while the antenna effect extends the photon collection
68 area of the detector, A_{coll} , making it much larger than the electrical area $\sigma = s^2$ of the device¹³.
69 As the detector photocurrent is proportional to A_{coll} , while the dark current is proportional to σ ,
70 for the same number of collected photons there is therefore a substantial reduction of the dark
71 current that results in a net increase of the detector operating temperature.

72 Besides the collection area A_{coll} , which defines the absorption cross section per patch resonator,
73 another crucial parameter is the contrast C of the reflectivity resonance shown in Fig. **1b**. This
74 parameter quantifies the fraction of the incident photon flux absorbed collectively by the array.
75 As shown in Fig. **1c**, the contrast can be adjusted by changing the array periodicity p ¹⁰. Optimal
76 detector responsivity is obtained at the *critical coupling point*, $C = 1$, where all incident radiation
77 is coupled into the array. The collection area per patch is related to the contrast according to
78 the expression $A_{\text{coll}} = Cp^2\xi$, where the factor $\xi = 0.7$ takes into account the polarizing effect of
79 the connecting wires (Methods)¹³. From the data in Fig. **1c**, the critical coupling is obtained with
80 a period $p = 3.3 \mu\text{m}$, which corresponds to a collection area $A_{\text{coll}} = 7.5 \mu\text{m}^2$, four times larger than
81 the electrical area $\sigma = 1.7 \mu\text{m}^2$ of the patch.

82 The device processing has been optimized in order to generate current solely under the metallic
83 square patches and not below the 150 nm wide leads connecting them. To this end we have
84 realised ohmic contacts between the patches and the underlying semiconductor layers using
85 PdGeTiAu annealed alloy, while a Schottky barrier, made by depositing TiAu, prevents vertical
86 current between the metallic wire and the semiconductor. Moreover, all cavities are connected
87 to an external wire-bonding pad insulated by an 800-nm-thick Si_3N_4 layer (Methods). Thanks to
88 all these precautions the conductive area is reduced to the sum of the areas of all the patch
89 resonators, which prevents additional dark current from flowing across the device.

90 In order to quantify the detector performance, we have compared the detector array with a
91 reference device, here referred to as “mesa”, where the same absorbing region is processed
92 into 200 μm diameter circular mesa and light is coupled in through the 45°-polished substrate
93 edge⁷. The mesa reference provides the intrinsic photo-response of the detector (Methods). In
94 Fig. **2a** we compare the peak responsivities for the two configurations, obtained with a
95 calibrated black body source at 1000°C (Methods). The mesa device could be characterized only
96 up to 150 K, as the photo-current becomes undetectable at higher temperatures. The array
97 detectors show a seven-fold enhancement of the responsivity at low temperatures. Most
98 remarkably, the responsivity could be characterized up to room temperature, where the
99 measured responsivity (0.2 A/W) is comparable with the best responsivity for the mesa device
100 measured at around 50 K. We were thus able to record photo-current spectra up to room
101 temperature, Fig. **2b**, which is, to our knowledge, the first type of such measurement with a
102 QWIP operating in the 9 μm band using a thermal source.

103 By quantifying carefully the number of photons absorbed in each geometry (Methods), we were
104 also able to extract the photoconductive gain g for each structure (Fig. **2c**). We recall that the
105 gain provides the number of electrons circulating per photon absorbed in the QWs^{7,28}, and is an
106 intrinsic property of the absorbing region. All our devices show the same values of the gain as a
107 function of temperature, irrespective of their fabrication geometry, which proves that the

108 material properties are identical for the two structures. Following Ref.7, the photoconductive
 109 gain is proportional to the electron drift velocity in the AlGaAs barriers and its temperature
 110 dependence is linked to microscopic scattering processes in polar materials^{14,15}. Our results fit
 111 well the temperature dependence of the drift velocity described on ref. 14. The derived low
 112 temperature value of the drift velocity is of the order of 6×10^6 cm/s as expected at an electric
 113 field of 20 kV/cm for an Al concentration in the range 20–30%²⁹. These results account for the
 114 temperature drop of the responsivity observed in Fig. 2a. Above 200 K, the gain acquires an
 115 almost constant value $g = 0.25 - 0.2$, of the order of $1/N_{qw}$. This implies that photoexcited
 116 electrons can only *travel* from one well to the next adjacent well, as the mean free path of the
 117 electrons is now shorter than the distance between two wells. Very interestingly, in this limit, it
 118 clearly appears that a detector based on a single quantum well would be advantageous at high
 119 temperatures. These results illustrate how our devices give access to the high temperature
 120 physics of quantum detectors, a unique regime unexplored so far.

121 The best assessment of detector performance is the specific detectivity⁷ $D^* = \frac{R\sqrt{A_{det}}}{\sqrt{A_{eg}I}}$ plotted in
 122 Fig. 3a for the mesa reference and for the patch devices. The experimental results are compared
 123 with our model that describes the impact of the photonic design on the detectivity as a function
 124 of the temperature¹³. For clarity, in Fig. 3b we provide the ratio between the detectivities. At
 125 low temperature, we observe an enhancement of only a factor of two. Here, the dark current is
 126 negligible and the main source of noise is the background photocurrent induced by the 300 K
 127 black body of the environment. In this regime higher responsivity means also higher background
 128 noise, and the detectivity enhancement scales with the square root of the responsivities ratio
 129 i.e. $(R_{array}/R_{mesa})^{1/2} = 2.6$. The situation is totally different at high temperature, where the dark
 130 current is the dominant contribution to the noise. In this case the detectivity enhancement is

$$131 \quad R_{array}/R_{mesa} (A_{coll}/\sigma)^{1/2} \sim 14, \quad (1)$$

132 and the actual performance of the arrays at 300 K is equivalent to the performance of the mesa
 133 reference at 150 K, doubling the temperature of operation. This is a significant improvement,
 134 well beyond that is predictable from the low temperature operation. Our device concept
 135 therefore takes advantage of both the responsivity enhancement and the strong suppression of
 136 the dark current owing to the antenna effect, as expressed by the factor $(A_{coll}/\sigma)^{1/2}$. As explained
 137 in Ref. 13, the combination of the microcavity and the antenna effect thus slows down the
 138 decrease of the detectivity with temperature, pushing the detector operation to much higher
 139 temperatures than expected.

140 By exploiting our photonic concepts we have achieved high temperature operation with relative
 141 high sensitivities. We now seek to benefit from the inherent very high frequency response
 142 together with the reduced electrical capacitance of our devices in order to use them as

143 heterodyne receivers. In this case, by increasing the power of the local oscillator one may
144 achieve the ultimate heterodyne sensitivity set only by the detector absorption coefficient.

145 This realization is depicted in Fig. **4a**, where we show schematically the heterodyne
146 arrangement that we used to probe our detector at room temperature. It consists of two single
147 mode distributed feedback (DFB) quantum cascade lasers (QCLs)¹⁶ operating at $\lambda = 8.36 \mu\text{m}$.
148 The lasers, used respectively as signal and local oscillator are made collinear by a beam splitter
149 (BS) before they impinge on the detector. The latter is connected via wire bonding to a high
150 frequency coaxial cable that is connected to a spectrum analyser. Each laser has a linewidth of
151 the order of one MHz when current and temperature are stabilised. By adjusting the
152 temperature of each laser, their frequencies are tuned within few GHz (Methods).

153 When the detector is illuminated by both lasers a clear heterodyne signal appears on the
154 spectrum analyser. In Fig. **4a** we show a measurement at 1.06 GHz, with a 40 dB signal-to-noise
155 ratio. We have measured heterodyne signals up to 4.2 GHz as it is illustrated in Fig. **4b**. Our
156 bandwidth is presently limited by a strong impedance mismatch between the detector and the
157 external circuit. In Fig. **4c** we report the characterisation of the sensitivity of the heterodyne
158 receiver at room temperature. The blue dots correspond to the direct current (DC) saturation
159 curve for the LO, while the red curve is the heterodyne signal at 1 GHz as a function of the signal
160 power. The straight line is a linear fit for the LO saturation curve. The saturation experiment
161 shows that the detector responds linearly up to 78 mW ($\sim 3.1 \text{ kW/cm}^2$) of incident power.
162 Moreover, the linear fit intercepts the 1 Hz integration band for a power of $\sim 0.5 \text{ nW}$, in very
163 good agreement with the measured room temperature detectivity from Fig. **3a**. As can be
164 observed from Fig. **4c**, the heterodyne data are very well fitted with a square root dependence
165 (dashed line) and can reach a signal-to-noise ratio of unity for an incident power of a few pW
166 and an integration time of the order of 10 ms. This clearly shows the strength of the heterodyne
167 technique that let us envision sensitivity in the thermal region at $\lambda = 9 \mu\text{m}$ which is unreachable
168 with any other technique at room temperature. Note that in our experiment the photocurrent
169 induced by the LO, $I_{\text{LO}} \sim 0.5 \text{ mA}$ is still dominated by the detector dark current, $I_{\text{dark}} \sim 3.5 \text{ mA}$. By
170 increasing the LO power and/or decreasing the temperature of the detector by few tens of
171 degrees using thermo-cooled elements, these detectors could reach the ultimate heterodyne
172 detection limit, set by their absorption efficiency^{7,13} and the relative intensity noise of the local
173 oscillator³⁰.

174 In conclusion, we have demonstrated metamaterial photonic detectors operating room
175 temperature with high sensitivity in the second atmospheric window at $\lambda \sim 9 \mu\text{m}$. While our
176 detectors show lower DC detectivity than microbolometers, they have an extremely fast
177 frequency response of tens of GHz. Using a quantum cascade laser as a local oscillator, we
178 have implemented a heterodyne detection setup, and validated that these uncooled detectors

179 can operate as coherent heterodyne receivers up to 4.2 GHz. The heterodyne scheme has,
180 indeed, a tremendous potential for sensitive detection in the mid-(far-) infrared that may
181 outperform all others competing technologies. The combination of high sensitivity with high
182 frequency response (tens of GHz) is the essence of this new class of metamaterial detectors.
183 Nonetheless, we recall that when installed on Peltier elements, the DC detectivity of our
184 devices is comparable to that of uncooled microbolometers.

185 Our devices will be of extreme relevance for the detection of coherent signals (lasers), in
186 particular for free space high-data-rate transfer¹⁷ and dual comb spectroscopy³¹, which is an
187 emerging high resolution spectroscopic technique, for which high speed detectors are
188 essential. In general, well-established applications such as optical free space communications,
189 thermal imaging and environmental remote sensing will greatly benefit from our coherent
190 sensitive detection. Moreover, our estimates show that the heterodyne scheme could also
191 serve for the generation and synthesis of microwaves (up to few hundreds GHz) with quite
192 good efficiency of the order of few percent. Finally, we point out that these coherent
193 detectors are ideally suited to be implemented into photonic integrated circuits (PIC's) where
194 the local oscillator is combined with the heterodyne receiver.

195

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275 **Acknowledgements**

276 We acknowledge financial support from the FP7 ITN NOTEDEV project (Grant. No. 607521), the
277 ERC grant “ADEQUATE”, the French National Research Agency (ANR-16-CE24-0020 Project
278 “hoUDINi”), and the EPSRC (UK) projects “COTS” and “HYPERTERAHERTZ” (EP/J017671/1,
279 EP/P021859/1). EHL and AGD acknowledge the Royal Society and the Wolfson Foundation, and
280 thank Dr L. Chen for skilled support with the device processing.

281 **Author contributions**

282 D.P., Y.T. and C.S. conceived the experiments, designed the QWIP structure, analysed the data
283 and wrote the manuscript. D.P. fabricated the QWIP devices and performed measurements and
284 data analysis together with A.B. A.M. and D.G. helped with the heterodyne measurements. A.C.
285 calibrated the blackbody for the responsivity measurements and helped with the
286 characterization of the mesa device. A.V. helped with data analysis. L.L., A.G.D. and E.H.L. grew
287 the QWIP structure and provided the wafer-bonding for the double-metal processing. F.K., M.B.
288 and J.F. provided the DFB QCLs for the heterodyne experiment. All the work has been realised
289 under the supervision of C.S.

290 **Author information**

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292 The authors declare no competing financial interests.

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298 **Figure 1|Device concept.** Double-metal patch antenna, with the various metallic layers employed for
299 electrical contacts (Methods). The absorbing region contains a QWIP structure (386 nm) with five QWs
300 Si-doped at $n=7\times 10^{11} \text{ cm}^{-2}$. For this metamaterial structure the photon collection area, A_{coll} , is much larger
301 than the electrical area σ . The scale bar on the image is 500 nm. **b**, Reflectivity spectrum (blue curve) of
302 a patch antenna array with $s=1.30 \mu\text{m}$ and a period $p=3.30 \mu\text{m}$. The dashed line is a Lorentzian fit
303 providing the absorption contrast C . **c**, Contrast C and collection area A_{coll} as a function of the array unit
304 cell area $\Sigma=p^2$. The observed saturation of A_{coll} is in agreement with theoretical predictions¹³.

305 **Figure 2|Detector characterizations.** **a**, Peak responsivity, measured with a calibrated 1000° C blackbody
306 source, of QWIP devices fabricated in 200 μm diameter mesa (circles), and into patch resonator arrays
307 with $s=1.35 \mu\text{m}$ (squares) and $s=1.30 \mu\text{m}$ (triangles). **b**, Normalized photocurrent spectra of the $s=1.30$
308 μm array at 78 K, 200 K and 295 K. **c**, Photoconductive gain and electronic drift velocity of the three
309 devices presented in **2a** as a function of temperature, for 0.5 V bias voltage (21 kV/cm electric field).
310 The drift velocity is obtained using a QW capture time of 5 ps (see ref. 7 and Methods).

311 **Figure 3| Detectivity as a function of the temperature.** **a**, Specific detectivity (2π field of view) as a
312 function of the temperature and at a bias of 0.5 V, for the reference mesa (circles) and two arrays
313 structures: $s=1.30 \mu\text{m}$ (triangles) and $s=1.35 \mu\text{m}$ (squares). The red line is a fit of the reference using
314 $d(T)=d_0/[1+d_1T\exp(-E_{\text{act}}/k_B T)]^{1/2}$ where d_0 and d_1 are fit parameters, $E_{\text{act}} = 120 \text{ meV}$ is the activation energy
315 and k_B is the Boltzmann constant. The blue curve is the model of quantum detectors embedded in patch
316 resonators described in ref. 13. **b**, Ratio between the detectivities in the two different detector
317 geometries. Dots show the corresponding BLIP temperatures: $T_{\text{BLIP}}^{\text{mesa}} = 70 \text{ K}$ (mesa) and $T_{\text{BLIP}}^{\text{cavity}} = 83 \text{ K}$
318 (patch cavity arrays).

319 **Figure 4|Tunable heterodyne experiment and results.** **a**, Heterodyne arrangement involving DFB
320 QCLs and a cavity array QWIP at room temperature . A 40 dB heterodyne power spectrum is shown,
321 acquired using a spectrum analyser with 1 MHz resolution bandwidth. **b**, Normalized heterodyne power
322 signal (in linear scale).. **c**, Log-log plot of the signal-to-noise ratio as a function of the signal QCL power,
323 for LO power of 40 mW. The noise of the QWIP is calculated using the measured gain and dark current
324 values at room temperature.

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333 **Methods**

334 **QWIP fabrication.** The QWIP structure is grown by MBE (molecular beam epitaxy). It consists of
335 five GaAs quantum wells (QWs), each with a thickness $L_{\text{QW}} = 5.2$ nm and each n -doped across
336 the central 4 nm region with Si at a density of $N_{\text{d}} = 1.75 \times 10^{18} \text{ cm}^{-3}$, providing a sheet density of
337 $n = 7 \times 10^{11} \text{ cm}^{-2}$. The QWs are separated by $\text{Al}_{25}\text{Ga}_{75}\text{As}$ barriers of thickness $L_{\text{b}} = 35$ nm. At the top
338 and bottom of this periodic structure GaAs contact layers are grown, with thicknesses $L_{\text{c,top}} =$
339 100.0 nm and $L_{\text{c,bottom}} = 50.0$ nm and doping $N_{\text{d,top}} = 4.0 \times 10^{18} \text{ cm}^{-3}$ and $N_{\text{d,bottom}} = 3.0 \times 10^{18} \text{ cm}^{-3}$,
340 respectively. The double-metal structures are obtained through wafer-bonding on a GaAs host
341 substrate using 500 nm gold layers, and by selectively etching down to an etch-stop $\text{Al}_{65}\text{Ga}_{35}\text{As}$
342 layer grown before the bottom contact. As shown in Fig. **1a**, the patch-antennae are connected
343 by 150 nm thin metallic wires which are realized using electron-beam lithography (consecutive
344 alignments allow different metallic alloy contacts). The final structure is obtained by ICP etching
345 of the semiconductor region between the antennae. The Schottky barrier under the thin
346 metallic wires prevents vertical dark current flow between the metal and the semiconductor³².
347 The 45° facet substrate-coupled geometry consists of a 200 μm diameter circular mesa, with
348 annealed Pd/Ge/Ti/Au as a top contact and annealed Ni/Ge/Au/Ni/Au as a diffused bottom
349 contact.

350 Extended Data Fig. 1 shows a scanning electron microscope (SEM) image of the quantum
351 detector device made of our metamaterial photonic concept. The pixel of the device is 50×50
352 μm^2 . The external pad is connected to the array by the 150 nm wires and is insulated from the
353 bottom ground plane by 800 nm thick Si_3N_4 layer. The TiAu pad connects the device to the
354 external circuit by wire bonding.

355 **Reflectivity and photocurrent analysis.** Reflectivity spectra and photocurrent spectra were
356 obtained using a Bruker Vertex interferometer. Reflectivity measurements were performed at a
357 15° incident angle and at room temperature, and the incident light was polarized perpendicular
358 to the 150 nm thin connecting wires. For the photocurrent spectra, QWIP devices were mounted
359 in a cryostat with an internal cooled metallic shield and a ZnSe optical window. Photocurrent
360 and responsivity were measured using a blackbody source at 1000 °C, which was calibrated with
361 an MCT detector. The source is focused onto the detector by two gold parabolic mirrors ($f/1$ and
362 $f/3$), providing typical field of view of 60°. The photocurrent is measured with a lock-in
363 technique using an optical chopper at 1059 Hz and a shunt resistance connected to the voltage
364 input of a lock-in amplifier Stanford Research SR1830, without using pre-amplifiers.

365 **Light polarization dependence.** Our structures support two fundamental modes, TM_{100} and
366 TM_{010} , which are represented in Extended Data Fig. 2a. This figure shows the vertical electric
367 field E_z in the plane of the resonator, obtained through finite elements simulations. The electric

368 field distribution follows a standing wave pattern, with a node in the center of the square and
 369 maxima at the edges. The connecting wires perturb the TM_{010} mode slightly, which results in a
 370 lower coupling efficiency for this mode. As a result, the total photoresponse of the antenna-
 371 coupled device has a co-sinusoidal dependence with the light polarization of the normally
 372 incident wave.

373
 374 In Extended Data Fig. 2b, we plot the peak value of the photocurrent for a $s = 1.30 \mu\text{m}$ structure
 375 as a function of the polarization of a plane wave incident on the array (open circles), with the
 376 90° direction corresponding to the direction of the connecting wires. The angular integral of the
 377 cavity photocurrent peak $I_{\text{photo}}(\theta)$ plotted in Extended Data Fig. 2b gives a polarization coupling
 378 coefficient $\xi_{\text{array}} = \int_0^{2\pi} I_{\text{photo}}(\theta) d\theta = 71\%$. The contrast value C of the TM_{100} polarized light is
 379 obtained from the measurement of Fig. 1b. For comparison, in the same graph we also plot the
 380 polarization dependence of the photoresponse measured for the mesa geometry (open
 381 squares). Here the 0° direction corresponds to the growth direction of the QWs, and the
 382 incident wave propagates normally to the 45° polished facet. This polar plot therefore recovers
 383 the inter-subband selection rule, as expected⁷.

384 **Definition of the collection area A_{coll} .** As all incident radiation that is not absorbed is reflected,
 385 the contrast C provides directly the fraction between the incident P_i and absorbed flux P_a for
 386 each patch, $C = P_a/P_i$. If we note by the incident photon flux Φ_i , then the power received by each
 387 antenna is $P_i = \Phi_i p^2$, and the power absorbed is by definition $P_a = \Phi_i A_{\text{coll}}$. Then using $C = P_a/P_i$ we
 388 obtain $A_{\text{coll}} = Cp^2$; in the main text we also add a corrective factor of $\xi_{\text{array}} = 0.7$ owe to the
 389 polarizing effect of the wires, as described in the previous paragraph.

390 **Responsivity, gain and specific detectivity** In Extended Data Fig. 3a we show the responsivity
 391 curves as function of voltage for both the mesa and the patch cavity with $s = 1.35 \mu\text{m}$. The
 392 decrease of the responsivity with temperature is attributed to the thermal dependence of the
 393 charge carrier drift velocity and to an increased phonon-electron interaction^{14,15} (see Fig. 2c).
 394 Note that QWIP devices show the typical negative differential photoconductivity, identified as
 395 the Gunn effect, which consists of a photocurrent decrease as function of voltage at specific
 396 critical fields, at which inter-valley electron scattering is induced in GaAs⁷.

397 The responsivities of the mesa can be expressed by considering the voltage dependent
 398 photoconductive gain $g(T, V)$ of the detector active region and the peak inter-subband energy
 399 $E_{21} = 143 \text{ meV}$ (taking into account many-body effects) :

$$400 \quad R_{\text{mesa}}(E_{21}, T, V) = \eta_{\text{isb}}(E_{21}) eg(T, V) t_{\text{GaAs}} \xi_{\text{mesa}} / E_{21} \quad (2)$$

401 where $\eta_{isb} = 5.0\%$ is the absorption coefficient for the five QW system in the 45° facet geometry,
 402 e is the electron charge, $t_{GaAs} = 0.67$ is the substrate transmission coefficient at $8.6 \mu\text{m}$ and
 403 $\xi_{mesa} = 0.5$ is the polarization factor (only one polarization of the incident light is coupled with
 404 the 45° facet). Analogously to Eq. (2), we can define¹³:

$$405 \quad R_{array}(E_{21}, T, V) = \frac{B_{isb}(E_{21})}{B_{isb}(E_{21}) + Q_{ohm}^{-1} + Q_{rad}^{-1}} eg(T, V) C \xi_{array} / E_{21} \quad (3)$$

406 where $Q_{ohm} = 4$ and $Q_{rad} = 22$ represent the ohmic and radiative dissipation of the double metal
 407 cavity, respectively, obtained by reflectivity measurements. Indeed, the Lorentzian fit of the
 408 reflectivity resonance from Fig. 1b in the main text provides the FWHM and the sum
 409 $1/Q_{ohm} + 1/Q_{rad}$, and Q_{rad} is calculated from the analytical expression provided in Ref.13.

410 The dimensionless parameter B_{isb} quantifies the energy dissipation through inter-subband
 411 absorption and is expressed by a lorentzian lineshape:

$$412 \quad B_{isb}(E) = f_w \frac{E_p^2}{4E_{21}} \frac{\hbar\Gamma}{(E - E_{21})^2 + \frac{(\hbar\Gamma)^2}{4}} \quad (4)$$

413 where $f_w = N_{QW}L_{QW}/L = 0.067$ is the filling factor of the absorbing QWs on the overall thickness, E_p
 414 $= 47.2 \text{ meV}$ is the inter-subband plasma energy, and $\Gamma = 15.0 \text{ meV}$ is the full-width-at-half-
 415 maximum of the mesa photo-response, obtained by a fit to the experimental data. We obtain a
 416 similar value $B_{isb} = 0.07$ for the two resonant cavities $s = 1.30 \mu\text{m}$ and $s = 1.35 \mu\text{m}$. The absorption
 417 coefficient in the antenna-coupled QWIPs is described by the branching ratio $\eta_{array} =$
 418 $\frac{B_{isb}}{B_{isb} + Q_{ohm}^{-1} + Q_{rad}^{-1}} = 18.9\%$. Using Eq. (2) and Eq. (3) with the measurement data in Fig. 2a, we
 419 obtain very similar values for the photoconductive gain for the mesa and the array, as shown for
 420 the data at 0.5 V (21 kV/cm) in Fig. 2a. This confirms that the absorbing regions for the two
 421 geometries are identical. Furthermore, the data shows an exponential decrease of the gain as a
 422 function of temperature. Following Ref. 7 the photoconductive gain can be defined as:

$$423 \quad g = \frac{\tau_{capt} v_d}{N_{QW} L_p} \quad (5)$$

424 where $\tau_{capt} = 5 \text{ ps}$ is the capture time, v_d is the drift velocity, $N_{QW} = 5$ is the number of quantum
 425 wells and $L_p = 40.2 \text{ nm}$ is the length of a period in the structure. The thermal dependence of the
 426 gain is related directly to the drift velocity and therefore to the electron mobility. Following Ref.
 427 14 we can express the temperature dependence as:

$$428 \quad g(T) = \frac{1}{\frac{1}{g_0} + \frac{B}{\exp\left(\frac{E_{LQ}}{k_B T}\right)} + \left(\frac{E_{AC}}{k_B T}\right)^{3/2}} \quad (6)$$

429 Here $E_{LO}=36$ meV is the longitudinal optical phonon energy in GaAs, and the fit parameter
430 $g_0=1.25\pm 0.03$ expresses the value of the gain at equilibrium (without thermal scattering
431 dependence). The second term in the denominator represents the polar optical scattering (see
432 Ref. 15) where the parameter $B=24.4\pm 1.6$ is a dimensionless polar constant and the third term
433 represents the deformation potential scattering caused by interaction of carriers with acoustic
434 phonons, with a corresponding parameter $E_{AC}=0.07\pm 0.01$ meV which characterizes the acoustic
435 deformation potential. Eq. (6) provides very good fits of the experimental data, confirming the
436 model.

437 The values of photoconductive gain obtained in this way are used to calculate the detectivity as
438 function of applied voltage, at different temperatures, as illustrated in Extended Data Fig. 3b.

439 **Heterodyne measurement.** The two beams from the QCLs are made collinear using f/0.5
440 germanium lenses and a beam splitter, and then focused onto the detector by a f/1.5 lens and a
441 $\lambda/4$ waveplate to avoid optical feedback (Fig. 3a). The two lasers are DC biased with a voltage
442 supply and are mounted in two Janis cryostats to stabilize their temperatures using liquid
443 nitrogen flow. The QWIP is polarized by a Keithley 2450 sourcemeter and the heterodyne
444 signal is sent to a spectrum analyser Agilent E4407B using a bias tee. In this arrangement the
445 QWIP detector is at room temperature, without using any cooling system. The QC laser used as
446 the LO is kept at a temperature 254 K while the QC laser used for the signal is kept at 293 K.
447 With the temperature stabilized, it is possible to tune the spectral position of the two DFBs by
448 slightly changing the applied DC current, according to the tuning coefficients $\beta_{LO}=378$ MHz/mA
449 and $\beta_S=413$ MHz/mA (extracted from a linear fit to the emission frequency of the lasers as a
450 function of temperature and bias).

451 In the case of a high power LO, the NEP of the heterodyne can be written⁷ $NEP_{het}=E_{21}/(\eta\tau)$
452 where η is the absorption coefficient of the QWIP and τ is the integration time (set by the
453 integration bandwidth Δf as $\tau=1/\Delta f$). For our device in the microcavity array we have a
454 theoretical limit of NEP_{het} of less than 1 aW for an integration time $\tau=1$ s at 300 K. In the
455 experiment shown in Fig. 4, the signal-to-noise ratio is still mainly limited by the dark current.
456 The square root fit of the signal-to-noise ratio can be extrapolated to 1, which provides NEP_{het}
457 ~ 10 fW for an integration time of 1 s ($NEP_{het} \sim 1$ pW for an integration time of 10 ms), that is
458 still four orders of magnitude higher than the theoretical limit. These estimations indicate that a
459 high power LO could achieve sensitivities at the single photon level at room temperature.

460 **Linearity and Heterodyne Measurement** In Extended Data Fig. 4 we show the spectra of the
461 two QCLs compared to the room temperature response of the QWIP in the microcavity array
462 geometry. We notice that the lasers are detuned from the maximum intersubband absorption,
463 resulting in a detector photoresponse that is half of the maximum achievable. This is an
464 important remark because the responsivity and detectivity values we report in Figs. 2 and 3

465 correspond to the peak values of detector photoresponse. The background-limited NEP (noise
466 equivalent power) is defined as $NEP = \sqrt{A_{det}}/D^*$. The detector area A_{det} corresponds to the
467 $50 \times 50 \mu\text{m}^2$ area of the whole array, which is equal to the number of patches N_{patch} multiplied by
468 the array unit cell area $\Sigma = p^2$. Indeed, in the critical coupling point, all incident radiation is
469 absorbed by the array, and therefore the collection area for each patch A_{coll} coincides with the
470 array unit cell $\Sigma = p^2$. Using our measured value of detectivity at 295 K for the cavity with $s = 1.30$
471 μm at 0.5 V (Fig. 3) we have $D^* = 2.8 \times 10^7 \text{ cmHz}^{0.5}/\text{W}$ and $NEP = 0.2 \text{ nW/Hz}^{0.5}$. Taking into account
472 the 50% spectral overlap, we obtain $NEP = 0.4 \text{ nW/Hz}^{0.5}$, which agrees with that observed from
473 the linearity measurement in Fig. 4c. Therefore the data presented in the main text are perfectly
474 consistent.

475 **References:**

476 **32.** Sze, S.M. and Kwok, Ng. *Physics of semiconductor devices*, Wiley, New Delhi India (2011)

477 **Data availability statement**

478 The authors declare that all data supporting the findings of this study are available within the
479 paper and its supplementary information files.

480 **Extended Data Figure 1 | Global view of the device**

481 Scanning electron microscope (SEM) picture of mid-infrared QWIP structure embedded into
482 $50 \times 50 \mu\text{m}^2$ array of patch resonators. We have indicated the top TiAu contact evaporated onto a
483 800 nm thick Si_3N_4 insulating layer.

484
485 **Extended Data Figure 2 | Polarization dependence of the photo-response**
486 **a**, Finite element simulation of the E_z field component coupled with the patch cavity QWIP, for
487 the TM_{100} and the TM_{010} modes. **b**, Polar graph of the cavity photocurrent peak as function of
488 the wire grid polarization angle. The photocurrent is normalized at its maximum at 0° . The open
489 circles are the results for the cavity array, where the 90° direction corresponds to the
490 connecting wires. The open squares are the results for the mesa geometry, where the 0°
491 direction corresponds to the growth direction of the QWs.

492 493 **Extended Data Figure 3 | Mesa and cavity array detector characteristics**

494 **a**, Responsivity of the mesa and the $s = 1.35 \mu\text{m}$ antenna-coupled devices as function of applied
495 voltage. The temperature in K of the QWIP is indicated for each measured curve. **b**, Specific
496 detectivity for the mesa and the microcavity devices as a function of the applied bias at
497 different temperatures.

498 499 **Extended Data Figure 4 | Spectral characteristics of the two lasers and the QWIP detector**

500 **a**, Emission spectra of the QC lasers (QCL_O and QCL_S) compared to the room temperature
501 response of the microcavity QWIP. **b**, Blown up version of the spectrum showing the two
502 distinct QCL emission lines. The QCL_O was operated at 330 mA with temperature stabilized at
503 293 K, and the QCL_S was operated at 280 mA with temperature stabilized at 254 K.







