their fits use a prefactor T_{θ}^{0} whose weak *T*-dependence disagrees with our continuum two-dimensional theory. We speculate that this can be explained by corrections to scaling: there is an appreciable density of fermionic carriers not bound into Cooper pairs, and this would reduce the effective bosonic superfluid density which enters our scaling forms with a *T*-dependent factor. Corson *et al.*⁴ motivated the scaling using the vortex theory¹⁸, but did not consider bound vortex pairs, whose contributions do not obey their scaling assumptions.

We have delineated the distinct dynamic regimes of thermally fluctuating superconductors: the low-*T* phase-only hydrodynamics, the classical vortex hydrodynamics in the vicinity of T_{KD} and the quantum-critical region where phase and vortex fluctuations strongly coupled—here we proposed a dynamical model in which non-linear 'mode-coupling' terms demanded by the Poisson bracket, equation (6), dominate the universal, low-frequency, dissipative dynamics. Our approach could be extended to other two-dimensional systems, such as the magnetic-field-tuned superconductor–insulator transition, where a coupling to the external field would be required in equation (4), and quantum Hall transitions.

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Optical microscopy using a single-molecule light source

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Rapid progress in science on nanoscopic scales has promoted increasing interest in techniques of ultrahigh-resolution optical microscopy. The diffraction limit can be surpassed by illuminating an object in the near field through a sub-wavelength aperture at the end of a sharp metallic probe^{1,2}. Proposed modifications^{3,4} of this technique involve replacing the physical aperture by a nanoscopic active light source. Advances in the spatial⁵ and spectral⁶ detection of individual fluorescent molecules, using near-field and far-field methods⁷, suggest the possibility of using a single molecule^{8,9} as the illumination source. Here we present optical images taken with a single molecule as a point-like source of illumination, by combining fluorescence excitation spectroscopy¹⁰ with shearforce microscopy¹¹. Our single-molecule probe has potential for achieving molecular resolution in optical microscopy; it should also facilitate controlled studies of nanometre-scale phenomena (such as resonant energy transfer) with improved lateral and axial spatial resolution.

Over nearly two decades the resolution in scanning near-field optical microscopy (SNOM) using aperture probes has levelled out at 3a value of \sim 50 nm. In addition to technological difficulties such as reproducibility of tip fabrication, the resolution in this mode of SNOM is fundamentally limited because the finite skin depth of real metals leads to a finite effective aperture size that one can achieve in the laboratory³. To overcome this problem, some researchers have pursued apertureless SNOM in which a sharp tip is used to scatter the near field of the sample¹², but in this configuration the resolution is limited by the radius of curvature of the probe used. The use of a nanoscopic active medium as a probe for SNOM has the advantage that its size can be reduced to that of a single atom or molecule. In this case, the resolution should only depend on the separation between the source and the sample.

Figure 1a shows the schematics of a combined scanning confocal microscope and scanning near-field optical microscope that constitutes the heart of our experimental setup¹³ operating at T = 1.4 K. A commercial piezo-electric scanner controls the position of the sample in front of a tapered optical-fibre probe. A microscope objective with a numerical aperture of 0.8 can be used to illuminate the sample or to collect the light from it. For adjusting the focus of the objective and for positioning the fibre probe, homemade piezo-driven translation stages are used¹⁴. We use the shearforce signal from a quartz tuning fork to monitor or to regulate the separation between the sample surface and the probe¹⁵. In the experiments described here, the end of the fibre probe contained a micron-sized p-terphenyl crystal (see Fig. 1b) that was doped with terrylene molecules at a nominal concentration of about 10^{-7} . To prepare such a probe, we first co-sublimated p-terphenyl and terrylene to obtain crystals of the order of a few micrometres in size on a cover slide. The spatial doping concentration of the

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microcrystals was then examined by collective excitation of the terrylene molecules at $\lambda = 514$ nm in a confocal arrangement under ambient conditions^{16,17}. Once a microcrystal of a desired size and doping was located, it was carefully glued to the end of an uncoated chemically etched single-mode fibre, and this probe was transferred into the cryogenic setup indicated in Fig. 1a.

Individual terrylene molecules in the micro-crystal were identified by fluorescence excitation spectroscopy^{10,18} at T = 1.4 K. The light of a ring-dye laser operating at $\lambda \approx 578$ nm was first intensity stabilized using an electro-optical modulator and then coupled into the optical fibre leading to the micro-crystal in the cryostat. Because the sharp zero-phonon lines of the terrylene molecules are inhomogeneously distributed in frequency space, individual molecules can become excited when scanning the laser frequency. The red-shifted fluorescence of a selectively excited molecule at $\lambda \approx 630 \,\text{nm}$ is then detected by an avalanche photodiode after passing through the microscope objective and spectral filters. In this manner, we could typically record resonance lines narrower than 50 MHz, count rates of 100 kHz and a signal-to-noise ratio of the order of 100:1 (Fig. 1c). Aside from the exquisitely narrow spectral lines and a high signal-to-noise ratio, our system also proves to be extremely photostable in contrast to the dye molecules that are typically studied at room temperature⁵. These features are the main reason for choosing this system for our present work.

Once the laser was tuned into resonance with a single terrylene molecule, the probe was moved nearer the sample until the onset of a shear-force signal was observed at a separation of about 20 nm. The sample was then retracted to a given fixed axial position from the tip and scanned laterally while monitoring the shear-force signal as a safety measure. The sample consisted of 25 nm high triangular aluminium islands arranged in a hexagonal lattice with a period of 1.7 μ m on a cover slide¹⁹. Figure 2a displays a topography image of the region that was studied in this experiment recorded with an atomic force microscope after the sample was taken out of the cryostat. The arrow indicates an area where a metallic island is largely missing, which results in two nanostructures. The image in Fig. 2b shows the raw fluorescence signal of a single molecule





positioned in front of this sample just before the onset of the shear-force signal. The photon counts were integrated for 20 ms per pixel during which period the laser frequency was scanned back and forth once over an interval of ± 250 MHz around the resonance to compensate for small laser drifts and occasional spectral diffusions²⁰. As expected, the signal drops substantially each time a metallic island blocks the transmission of the molecular fluorescence. Although the triangular form of the islands does not manifest itself strongly, the image shows a striking reproducibility among different lattice periods. For example, the set of islands labelled *i* can clearly be distinguished from that labelled *ii* corresponding to the two possible orientations of the triangular structures. Also, the bright regions display a systematic pattern, with the exception of the area around the defect.

In Fig. 3 we show this image (Fig. 3e) together with three other optical images taken with the same single molecule at the same lateral position but with different distances between the crystal extremity and the sample (350 nm, 80 nm, 50 nm and 20 nm in Fig. 3a, b, d and e, respectively). The scans were recorded in the chronological order of Fig. 3b, a, e and d. These images are very lightly filtered by averaging the signal from each pixel with a weighting factor of 8 and its 8 neighbouring ones each with a weight of 1. To account for variations in the excitation intensity among different runs, the count rates in all four images have been normalized and a common colour scale is used so that one can compare them directly. Individual metallic islands and their hexagonal arrangement can be clearly identified in all cases, but the tiny defect structures give rise to a significant signal only at the smallest molecule-sample separation achieved in Fig. 3e. The fact that the contrast and the resolution are considerably worse at a larger distance also becomes evident when comparing Fig. 3a with the other images.

Bringing the sample closer in steps as small as 30 nm from Fig. 3b to d to e results in a substantial but continuous evolution of the bright and dark contrast patterns although these remain very robust within each figure. This can be seen more quantitatively in Fig. 3f where two cross-sections α and β exactly 1.7 μ m apart are plotted. Note that the signal undergoes oscillations above and below the background indicated by a baseline. We attribute the origin of this effect and the rich images of individual metallic islands (for example, in Fig. 3e) to the polarization-sensitive interaction of the molecular dipole radiation with the aluminium triangles. Such interference-like phenomena have been reported in different



Figure 2 Topography and optical images of the sample. **a**, An atomic force microscope (AFM) topography image of the area of the sample used in this experiment. The arrow points to the remains of an island that is largely missing. We note that, although we have also recorded shear-force topography images of the sample taken at 1.4 K, because of our micrometre-size tip dimensions the resolution is not as good as that of an AFM. **b**, The optical raster-image recorded using the fluorescence of a single terrylene molecule for illumination. This scan contains 100×80 pixels with an integration time of 20 ms per pixel. The colour scale displays the raw experimental count rates in units of thousand counts per second (kc.p.s.).

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experimental^{21,22} and theoretical^{23,24} works on conventional SNOM with polarized illumination. These studies have concluded that image interpretation becomes non-trivial for certain combinations of the electric field orientation and sample morphology and that the image quality depends on the illumination and detection polarization as well as the detection angle²². In the case of a single-molecule emitter, one should also take into account the modification of its spectral properties owing to the influence of the sample surface^{25,26}. A detailed theoretical and experimental study of these properties will be the topic of a future publication.

Clearly, the complex contrast observed in the images complicates the assignment of a resolution. When dealing with finite-sized structures, one common method for evaluating the resolution is measuring the edge sharpness of the obtained signal²⁷. In our case, however, following this guideline is not straightforward because of the oscillations about the background. We therefore merely note that the signal in Fig. 3f drops below the baseline from its 10% to 90% value over a length of about 180 nm which is less than onethird of the emission wavelength.

When extending the principles of scanning optical near-field microscopy to the case of a point-like electric dipole probe, the resolution is expected to depend only on the probe-sample separation. From a practical point of view, addressing molecules



Figure 3 Images taken with the same molecule positioned at four different distances from the sample. These images are very lightly filtered (see text). The separation between the micro-crystal's extremity and the sample was 350 nm (**a**), 80 nm (**b**), 50 nm (**d**) and 20 nm (**e**). **e**, A more complete view of the run shown in Fig. 2b where in the last part the molecular spectrum became unstable and underwent a large jump of the order of hundred GHz (ref. 29) when the probe came into contact with the sample, so that a very small change in the shear-force signal was detected. **c**, The fluorescence signal along the cross-section shown in **b**. Note the sudden drop of the signal to ~200 counts per pixel (about 0.2 on the normalized scale) while our residual fluorescence corresponded to about 30 counts per pixel. Here the molecule has undergone a small spectral jump to the edge of the frequency scan range which was corrected for after a few scan lines by adjusting the laser frequency manually. **f**, the fluorescence signals along the two cross-sections of the four images taken over a period of over 2 h reveals the stability of the system and lack of any lateral drifts.

that lie very close to the probe end should lead to higher spatial resolution and in the extreme limit to a molecular resolution. To achieve this, we plan to use sub-micrometre crystals and implement various techniques for identifying single molecules at the end of the crystal, for example, through evanescent excitation. In addition to the detection of the fluorescence intensity, the internal quantum mechanical properties of the molecule such as fluorescence lifetime^{26,28} and energy level shifts²⁶ can be used to extract information about its immediate environment. Our single-molecule probe could be also used to study a variety of phenomena such as surface-enhanced Raman spectroscopy and resonant energy transfer with an unprecedented lateral and axial spatial resolution.

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Intercalation of alkylamines into an organic polymer crystal

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Organic solid-state synthesis allows formation of products that are difficult or impossible to produce by conventional methods. This feature, and the high degree of reaction selectivity that can be achieved, is a direct result of the control over the relative orientation of the reactants afforded by the solid state. But as the successful development of 'topochemical reactions' requires the careful design of suitable reactant crystals, the range of both reactions and products amenable to this approach has been limited^{1,2}. However, recent advances in organic crystal engineering, particularly the rational design of complex solid architectures through supramolecular preorganization³⁻⁹, have renewed interest in topochemical reactions. Previously, we have orientated muconate monomers-diene moieties with a carboxylate group on each end—using long-chain n-alkylammonium ions, such that the topochemical photopolymerization of the solid-state reactants produces layered crystals of stereoregular and highmolecular-mass polymers¹⁰⁻¹³. Here we show that these polymer crystals are capable of repeated, reversible intercalation¹⁴ by conversion to the analogous poly(carboxylic acid), followed by transformation into a number of poly(alkylammonium



Figure 1 Photoreactions and polymer transformation performed in the crystalline state. Depending on the carbon number of the *n*-alkylammonium group, alkylammonium (Z,Z)-muconate undergoes either topochemical polymerization or topotactic *EZ*-isomerization when exposed to photoirradiation in the crystalline state. The ammonium polymer crystals obtained by the topochemical polymerization are converted in the crystalline state to acid polymer crystals via polymer transformation. The reverse reaction from polyacid to the ammonium polymers also occurs.

muconate)s upon addition of the appropriate amine. Introduction of functional groups into these crystals may allow the design of organic solids for applications such as molecular recognition, separation and catalysis, thereby extending the range and practical utility of current intercalation compounds^{15–20}.

Alkylammonium (Z,Z)-muconates undergo polymerization in the crystalline state under photoirradiation. When a higher *n*-alkyl group (carbon number $m \ge 10$) is introduced in the N-substituent, the photoproduct is an insoluble polymer crystal (Fig. 1; see Supplementary Information for preparation and polymerization procedures). Derivatives with a lower *n*-alkyl substituent (m < 10) favour topotactic isomerization to the corresponding (E,E)-isomers. In Table 1, the results of the photopolymerization are given together with the values of 2θ and interplanar distance (d) determined by small-angle powder X-ray diffraction for both monomer and polymer crystals. No peak was observed in the small-angle region for the monomer crystals with m < 10, in contrast to the intense peaks at $2-3^{\circ}$ for the monomer and polymer crystals with m \geq 10 (Fig. 2a). They correspond to 30–45 Å of the *d* value, which increased in proportion to the number of carbons in the N-alkyl group. The topochemically polymerizable (Z,Z)-muconates have similar crystal lattice parameters to each other^{11,12}: monoclinic, $P2_1/c$, a = 10.2320 Å, b = 4.9310 Å, c = 11.4970 Å, $\beta = 107.146^\circ$, $V = 554.29 \text{ Å}^3$, Z = 2 for the ethyl ester; monoclinic, $P2_1/a$, a =10.98 Å, b = 4.862 Å, c = 17.72 Å, $\beta = 97.93^{\circ}$, V = 936 Å³, Z = 4 for the benzylammonium salt; and monoclinic, $P2_1/a$, a = 11.033 Å, b =4.9360 Å, c = 18.011 Å, $\beta = 95.055$, V = 977.0 Å³, Z = 4 for the 2chlorobenzylammonium salt. A lamellar structure of alkylammonium cation and muconate dianion layers is observed in the monomer crystals of the benzylammonium derivatives, and this ordered structure remains during the topochemical polymerization. Polymerization behaviour that depends on the length of the alkyl



Figure 2 Small-angle X-ray scattering. **a**, Diffraction profiles of poly(*n*-alkylammonium (Z,Z)-muconate) crystals as polymerized. **b**, Relationship between the carbon number (*m*) of the *n*-alkyl group and the *d* values for *n*-alkylammonium (Z,Z)-muconates. Circles, monomer crystals; squares, polymer crystals obtained as polymerized under photoirradiation in the crystalline state after removing unreacted monomers; triangles, polymer crystals obtained by the crystalline-state polymer transformation from poly(muconic acid) with corresponding *n*-alkylamines.