SOLAR ENERGY
Thermoelectrics on the roof

ORGANIC SPINTRONICS
Supramolecular spin valves

NANOMEDICINE
Communication-enhanced targeting

Nanowires without end
Arrays of indefinitely long uniform nanowires and nanotubes

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Nanowires are arguably the most studied nanomaterial model to make functional devices and arrays1–4. Although there is remarkable maturity in the chemical synthesis of complex nanowire structures, their integration and interfacing to macro systems with high yields and repeatability5–7 still require elaborate aligning, positioning and interfacing and post-synthesis techniques8,9. Top-down fabrication methods for nanowire production, such as lithography and electrospinning, have not enjoyed comparable growth. Here we report a new thermal size-reduction process to produce well-ordered, globally oriented, indefinitely long nanowire and nanotube arrays with different materials. The new technique involves iterative co-drawing of hermetically sealed multimaterials in compatible polymer matrices similar to fibre drawing. Globally oriented, endlessly parallel, axially and radially uniform semiconducting and piezoelectric nanowire and nanotube arrays hundreds of metres long, with nanowire diameters less than 15 nm, are obtained. The resulting nanostructures are sealed inside a flexible substrate, facilitating the handling of and electrical contacting to the nanowires. Inexpensive, high-throughput, multimaterial nanowire arrays pave the way for applications including nanowire-based large-area flexible sensor platforms, phase-change memory, nanostructure-enhanced photovoltaics, semiconductor nanophotonics, dielectric metamaterials, linear and nonlinear photonics and nanowire-enabled high-performance composites.

One-dimensional nanostructures such as nanowires, nanotubes and nanoribbons are continuing to be at the forefront of nanoscience and nanotechnology. Two distinct approaches for the fabrication of these structures are the top-down and bottom-up philosophies10. During the past decade, chemical synthesis demonstrated the impressive success of the bottom-up approach in achieving controllable composition and morphology11 and prototype functional devices12,13, but less so for the integration of high-density device assemblies. Post-synthesis assembly techniques, designed for this end, such as electric14, magnetic-field-assisted alignment and dielectrophoresis5, optical and optoelectronic tweezers15, laminar flow in microfluidic channels13 and micro contact printing16 still do not achieve low cost, high throughput, and good reproducibility with high-precision addressability. For some applications chemical synthesis is simply not a fitting choice; for example, it is still difficult to produce very long aligned nanowires12,17, and to obtain nanowires on large area substrates18, or polymer substrate nanowire integration19, requires separate high-temperature nanowire synthesis and subsequent transformation.

On the top-down side, patterning by lithography is a powerful but costly micro- and nanofabrication tool perfected by the semiconductor industry. However, for one-dimensional nanostructure fabrication, it is not much favoured owing to scientific challenges such as resolution, surface roughness, limited chemical composition and its low throughput10. Various hybrid techniques are developed for nanowire fabrication to expand the toolkit, such as using photonic crystal fibres20 as microfluidic reactors21 to synthesize nanowires (templating) and meniscus-controlled solution evaporation17 (growth and winding process) to obtain very long nanowires. A few other successfully fabricated top-to-bottom micro- or nanostructures are the photonic crystal fibre, which consists of microtubular enclaves inside a silica fibre22 for optical guiding, thermally drawn composite fibres integrating semiconductors, metals and insulators featuring micro- and nanosizes for advanced functionality11,20, and silica nanofibres obtained by tapering25. Direct attempts to produce very long metallic26 and semiconducting27,28 micro- and nanowires inside viscoelastic matrices using thermal size reduction similar to the Taylor wire process29 could not produce ordered structures. Arbitrarily distributed filaments are obtained, making it impossible to identify/address each filament as single nanostructures.

Here we report the first successful fabrication of arrays of millions of ordered indefinitely long nanowires and nanotubes in a flexible polymer fibre. Using a new iterative size-reduction process, semiconducting nanowires and hollow-core piezoelectric polymer nanowires, with diameters below 15 nm, are obtained as well-ordered high-density arrays. The nanostructure sizes are uniform for hundreds of metres along the fibre and radially homogeneous in the cross-section. Unique multimaterial core–shell nanowire arrays were obtained and nanowires and nanotubes can be extracted to obtain extremely long free-standing slivers.

To fabricate nanowires that bridge the macroscopic and the nanoscopic world, we designed a multistep iterative thermal size-reduction process inspired from composite-fibre drawing from polymer reels. We start with a macroscopic polymer rod with an annular hole which is tightly fitted with a thermoelastically compatible material that is to become nanowires on multiple layers. Post-synthesis assembly techniques, designed for this end, such as electric14, magnetic-field-assisted alignment and dielectrophoresis5, optical and optoelectronic tweezers15, laminar flow in microfluidic channels13 and micro contact printing16 still do not achieve low cost, high throughput, and good reproducibility with high-precision addressability. For some applications chemical synthesis is simply not a fitting choice; for example, it is still difficult to produce very long aligned nanowires12,17, and to obtain nanowires on large area substrates18, or polymer substrate nanowire integration19, requires separate high-temperature nanowire synthesis and subsequent transformation.

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A new nanofabrication technique, based on iterative size reduction, to produce ordered, indefinitely long nanowire and nanotube arrays. A macroscopic multimaterial rod is reduced to ordered arrays of nanowires by thermal size reduction in a protective polymer matrix in successive steps. Each step starts with structures obtained from a previous step, resulting in geometrical size reduction and increment in wire number and length. Using the technique, we produced millions of kilometre-long nanowires with sub-10 nm diameter and an aspect ratio of $10^{11}$. As an example, a 10 mm amorphous semiconducting rod (Ge–As–Se–Te) is reduced to hundreds of metres of single 200 µm wire (reduction factor $\times 50$), $\sim 30$ wires of 5 µm diameter (reduction factor $\times 50$) and $\sim 1,000$ wires of 250 nm (reduction factor $\times 50$). Nanowire arrays are obtained as embedded in a dielectric polyetherimide encapsulation. Wire-array cross-sections with well-ordered wires and extracted, globally oriented slivers are shown after each step. Inset: Transmission electron microscopy image of a single 32-nm-thick nanowire obtained by further scaling down the third-step nanowires.
Figure 2 | Globally ordered, multimaterial nanowire, nanotube and cylindrical core–shell arrays. Nanowire and nanotube arrays are extracted from polymer matrix by chemical etching, retaining their global alignment. a, As$_2$Se$_3$ semiconducting nanowires. b, As$_2$Se$_3$ nanowire core with PVDF encapsulation forming a glass–polymer cylindrical core–shell structure. c, High-refractive-index low-bandgap semiconducting Ge$_{15}$As$_{25}$Se$_{15}$Te$_{45}$ nanowire slivers. d, Hollow-core piezoelectric polymer (PVDF) nanotube slivers with ~20 nm wall thickness. Hollow cores of the tubes are evident from the creases in the slivers after extraction by chemical etching.

wires. The total number of nanowires and their ultimate size distribution is determined by the number of packed fibres after each step, the total number of iterative steps and the reduction factor as shown in Fig. 1.

We demonstrate nanowire production using chalcogenide semiconducting glasses in Fig. 1. Stable glass-making non-oxide chalcogen compounds have low softening points, making them amenable for thermal co-drawing inside polymer matrices owing to their excellent match of thermomechanical properties (Supplementary Fig. S2). We synthesized a 10-mm-diameter glass-making chalcogenide Ge$_{15}$As$_{25}$Se$_{15}$Te$_{45}$ (GAST, $T_g$ $\sim$ 190 $^\circ$C) from high-purity elemental constituents using the conventional melt-quenching technique$^{30}$ (see the Methods section). Around the glass rod, a high-temperature engineering polymer (polyetherimide, $T_g$ $\sim$ 220 $^\circ$C) is rolled and thermally consolidated in vacuum. The composite structure is then thermally drawn to reduce radial size (Supplementary Fig. S3). Typically, the process is carried out at 50 cm min$^{-1}$ drawing speed while feeding the macroscopic rod at 1.5 mm min$^{-1}$ into a furnace at 275 $^\circ$C. The ‘fibre wires’ obtained from the first drawing are then packed and vacuum consolidated to prepare a second-step macroscopic rod. Vacuum consolidation ensures structural homogeneity and purity from defects that may adversely affect the subsequent thermal drawing process (see the Methods section). Beginning with a 10-mm-diameter GAST rod, the diameter is successively reduced to 200 nm, 50 nm and 250 nm within each drawing step. Further reduction in the third step results in sub-50 nm nanowire arrays. In Fig. 1, we show a perfectly ordered array of GAST micro- and nanowires in the polyetherimide polymer cross-section. It is remarkable that the macroscopic structure is undistorted after the drawing steps. We have also demonstrated that the micro- and nanowires can be extracted from the polymer matrix retaining their global alignment simply by exposing the polymer section to organic etching agents (dichloromethane (DCM) or dimethylacetamide; see the Methods section). Resulting free-standing micro- and nanowire slivers are shown in Fig. 1.

We fabricated a variety of multimaterial nanowire and nanotube structures by using the same technique (Fig. 2; Supplementary Figs S4–S7). Regular nanowire arrays are fabricated from the highly nonlinear glass As$_2$Se$_3$ (Fig. 2a), ordered core–shell arrays of As$_2$Se$_3$–PVDF (polyvinylidene fluoride; Fig. 2b), high-refractive-index GAST glass (Fig. 2c), and photoconductive and phase-change Se glass (Supplementary Figs S4–S6). Separately, nanotubes of piezoelectric PVDF (melting temperature $T_m$ = 171 $^\circ$C) are obtained using the same process by using PVDF sheets as the filling material and polysulphone (PSU, $T_g$ $\sim$ 190 $^\circ$C) as the jacket layer (Fig. 2d). PVDF sheets take their tubular shape through melting at the drawing temperature ($T_{draw}$ = 240 $^\circ$C) followed by self-organization due to high surface energy between the fluoride polymer and the protective PSU jacket. Macroscopic PVDF tubes can be seen after the first and second step drawings, and more interestingly tubular shapes are conserved in the
We produced kilometre-long nanowires and nanotubes with any diameters within the 100 µm–10 nm range by preserving ordered geometry. **a**, As$_2$Se$_3$–PVDF core–shell nanowires scaled regularly from 200 µm to 14 nm. **b**, Hollow-core PVDF nanotubes scaled regularly from 200 µm to 20 nm without losing structural integrity. Smearing around the nanowires is caused during sample preparation with ultramicrotomy.

Nanoscale, resulting in very long arrays of nanofluidic channels in a flexible polymer fibre. Thermal melt-drawing of globally aligned PVDF nanotubes is an interesting alternative to existing templating and sol–gel methods for producing high-density nanotube channels.

Nanowire fabrication with low-melting-temperature metals and metal alloys (Sn, SnPb, SnAg) was also explored using the same process. Fibre drawing is ideally designed for glassy materials; however, if it is possible to melt the crystalline substance without oxidizing inside a protective glassy matrix during thermal draw, crystalline rods will melt and recrystallize as elongated wires conserving the structural integrity. Recently, it was shown that other phases (such as metals and semimetals) can be drawn in an encapsulating glassy matrix. Here, drawing occurs in a regime dominated by viscous forces, allowing for internal low-viscosity metallic domains to be arranged in cross-sections confined by viscous glassy boundary layers. However, this latter type of composite drawing becomes especially challenging with submicrometre filament sizes. The Taylor process, other attempts and the authors’ own previous work suggests that crystalline materials can be co-drawn in glassy matrices when the drawing conditions are just right. To demonstrate that it is indeed feasible to fabricate ordered arrays of micrometre- and sub-micrometre-scale metallic wires in polymer fibres, we selected tin and tin alloys owing to their low melting temperature ($T_m = 223$ °C). The drawing process is carried out with similar parameters to those of chalcogenide glasses. We started with a 1.6-mm-diameter soldering alloy Sn$_{96.5}$Ag$_{3.5}$ tightly fitted inside a hollow-core polyethersulphone rod, and scaled successively to 60 and 4 µm (Supplementary Fig. S7).

We demonstrate the utility of the top-down size-reduction technique by producing wires with sizes spanning the whole micro- and nanoscale (Fig. 3). Beginning with the same macroscopic rod, structures of six orders of magnitude in size can be obtained. We demonstrate this facility with core–shell As$_2$Se$_3$–PVDF nanowires and hollow-core PVDF tubes. As$_2$Se$_3$–PVDF core–shell nanowires were regularly scaled down to 61 µm, 1 µm, 220, 122, 66, 41 and 14 nm (Fig. 3a). Hollow-core PVDF nanotubes were scaled regularly, conserving their structural integrity, from 100 µm down to 20 nm (Fig. 3b). It is remarkable that we produce extended lengths of micro- and nanotubes that are highly sought materials in micro- and nanofluidics research. The wires and tubes are obtained as ordered arrays in the surrounding polymer matrix. In the micrographs, smearing around the nanowires is caused during
10-m-long nanowire array
2 µm
20 µm
1 µm 170 180 190 200 210 220 230 240 250
Nanowire diameter (nm)

Figure 4 | Radial and axial uniformity of the nanowire arrays. a, A polymer-embedded nanowire array rolled around a pencil truly spans macroscopic and nanoscale worlds. Cross-sectional SEM micrographs from both sides of a 10-m-long polymer fibre that contains hundreds of As$_2$S$_3$–PVDF core–shell nanowires prove that nanowire arrays are axially uniform to less than 1% for macroscopic distances. b, High-precision hexagonal packing of core–shell nanowires in the polymer matrix. c, Radial size distribution of the nanowires, shown as a histogram, is uniform with a standard deviation of 6.5%.

sample preparation with microtomy, making it challenging to image embedded nanowires in the polymer matrix. For glassy filling materials, the ultimate size reduction limit depends on the drawing dynamics and viscoelastic properties of the matrix. Amorphous glasses and polymers that soften during drawing can be easily reduced to 10 nm radial size without axial breakdown. We infer that, with feed-in and draw speeds tuned and controlled carefully, the drawing process can yield molecular wires; the ultimate size reduction limit is still an enticing question.

We demonstrated radial and axial uniformity of the As$_2$S$_3$–PVDF core–shell nanowires by taking cross-sectional scanning electron microscopy (SEM) images (FEI, Nova NanoSEM 600) from both sides of a 10-m-long nanowire-embedded fibre, rolled conveniently on a pencil (Fig. 4). It is extraordinary that the conformal surface coverage of a macroscopic object with nanowires can be achieved trivially. Roughly 400 of the As$_2$S$_3$–PVDF core–shell nanowires with 200 nm diameter are obtained with two-step iterative drawing. The two-dimensional array structure and integrity of the embedded nanowire arrays are retained for macroscopic lengths. The size distribution along the fibre length is uniform within 1% (Fig. 4a). High-precision hexagonal packing of core–shell nanowires in the matrix is shown in Fig. 4b. The size distribution of the nanowires from the second step is shown as a histogram and found to be uniform within 6.5% (Fig. 4c).

Finally, we investigated the size-dependent photoconductivity of selenium micro- and nanowire arrays fabricated using the described
technique. Elemental selenium has phase-dependent electrical conductivity and photoconductivity properties, making it suitable for phase-change memory, optoelectronics and photodetection applications. In a three-step drawing process, we obtained a 56 μm wire, an array of 100 × 8 μm wires and an array of 10,000 × 190 nm nanowires. The structure of the nanowires inside the polyethersulphone polymer matrix is conserved after the drawing steps (Fig. 5). The uniformities of the selenium nanowires from the second and third steps are found to be ±3% and ±11% respectively (Supplementary Fig. S5). Factors contributing to the overall size distribution are slight variations in the fibres that are used to make the macroscopic composite, and the total accumulated size variation from each previous step. Amorphous selenium wires are obtained after drawing, as revealed by X-ray diffraction and transmission electron microscope diffraction (FEI, Tecnai G2 F30). The wires are crystallized through heat treatment, by annealing above the crystallization ($T_c = 110^\circ$C) but below the melting temperature$^{[27,35]}$ and by exposing to the organic reagent pyridine (Fig. 5a, see Methods section). After crystallization we observe that the conductivity increases to $10^{-4}$ S cm$^{-1}$ consistent with bulk selenium conductivities$^{[34]}$. Electrical measurements were carried out with nanowire arrays by cleaving fibre ends with a razor blade and electrical contacting was achieved using silver paint on the facets of the fibre. Step 3 nanowires were chemically crystallized after extraction from the polymer matrix. Electrical gain due to photoconductivity is measured by shining white light (50 W) while applying voltage bias (Fig. 5b). Step-3 wires make Ohmic contact and feature photoconductivity after crystallization (Fig. 5c). The photoconductance of the nanowires is observed to be a function of the nanowire number and size in the array. We observed that the $I/I_0$ gain (photocurrent versus dark current) was an order of magnitude larger for step-2 wires and two orders of magnitude larger for step-3 wires with respect to the single selenium microwire from the first drawing (Fig. 5d). The switching between on/off states is also faster with the smaller nanowire sizes. We attribute the logarithmic increase in photoconductance to the high absorption of selenium, increased surface-area-to-volume ratio and enhanced scattering of the incoming beam.

We have demonstrated a unique, high-throughput, lithography-free nanofabrication method to produce macroscopically large nanowire and nanotube arrays. The nanostructures are indefinitely long, ordered, uniform, high-density arrays embedded in a flexible polymer fibre matrix. Diverse multimaterial sets and disparate combinations are used to obtain crystalline semiconducting nanowire arrays, chalcogenide nanowire array waveguides, piezoelectric polymer nanotubes and cylindrical core–shell structures. It is an open question if molecular wires can be obtained using the method. The ultimate size reduction for the nanowires needs to be studied by optimizing drawing parameters for thermoviscous materials and materials that melt during drawing separately. On the other hand, the material set can also be widened to include semiconductors, such as Si and Ge, that soften/melt at higher temperatures using suitable glassy matrices.

We expect that a whole new family of radically novel nanowire applications will be enabled owing to the unique

Figure 5 | Photoconductivity of selenium microwire and nanowire arrays. a, Amorphous selenium wires are crystallized through heat treatment or by exposing to the organic reagent pyridine. SEM images of regularly scaled-down individual selenium wires. The electrically conductive hexagonal crystallographic phase is obtained by X-ray and electron diffraction. b, A schematic representation of the photoconductance measurement geometry. c, The photoconductance from a selenium nanowire in the dark and on white-light illumination. d, A logarithmic increase in the ratio of photocurrent versus dark current ($I/I_0$) and a reduction in switching time ($\tau$) with reducing nanowire diameter are observed. The performance increase is attributed to the high optical density of selenium, increased surface-area-to-volume ratio and enhanced scattering in the polymer matrix.
Methods

Synthesis of chalcogenide-glass rods. As$_x$Se$_y$ and Se rods, 10 mm in diameter and 15 cm in length, are prepared by melting commercially bought glasses (Amorphous Materials) and pellets (Alfa Aesar) in a vacuum-sealed quartz tube followed by water quenching. Ge$_x$As$_y$Se$_z$Te$_{4-x}$ glass rods (the same diameter) are prepared from high-purity elements (Alfa Aesar) using the sealed-ampoule melt-quenching technique. The weighted materials are placed in a quartz tube under nitrogen atmosphere in a glove box (H$_2$O, O$_2$ below 0.1 ppm). The quartz tube is kept at 330 °C for an hour under vacuum to remove surface oxides and sealed (10$^{-4}$ torr). The sealed ampoule is then heated to 950 °C, rocked for 18 h and then quenched in iced water.

Macroscopic-composite preparation, consolidation and thermal size reduction. Macroscopic rods that will undergo axial elongation and radial size reduction are structures approximately 30 mm in diameter. These structures are prepared in a clean pressure flow room, by tightly rolling polyethersulphone or polyetherimide films around chalcogenide glass rods (10 mm diameter) until the diameter is 30 mm (see Supplementary Fig. S1). Then, the macroscopic structure is thermally consolidated under vacuum (10$^{-7}$ torr), above the glass transition temperature of the polymer and glasses to fuse (260 °C for 30 min). Step-2 and higher-step structures are prepared with previously obtained polymer–wire fibres and a hollow-core polymer rod fabricated by rolling polymer sheets on a Teflon rod, consolidation and removing the Teflon rod. About 100 fibres with 0.5 mm diameters are cut as 10 cm fibres, tightly packed and placed inside the hollow-core polymer rod. The structure is consolidated once more to fuse packed fibres to the outer polymer jacket (250 °C for 15 min). The second consolidation step ensures structural integrity during drawing. The macroscopic structure is heated and drawn into hundreds of metres of micro- and nanowire in a vertical two-zone furnace, with a top-zone temperature of 275 °C and bottom zone 200 °C (see Supplementary Fig. S3). The nanowire diameter is controlled using a laser micrometer by monitoring the fibre diameter.

Nanowire and nanotube extraction. The polymer encapsulating the nanowires and nanotubes was extracted for electron microscopy imaging in the longitudinal dimension by DCM (Carlo Erba). Nanowire or nanotube arrays embedded in polymer were fixed on a glass substrate by means of aluminium foil, and then half soaked in bottles containing DCM. The extracted nanostructures were further rinsed gently with DCM to remove residual polymer. A final 0.5 °C plasma treatment (Nanoplas DSB6000) for several hours was applied for chalcogenide and metal nanowires to remove organic residues.

Nanowire crystallization by thermal and chemical methods. Step-1 selenium nanowires were crystallized by annealing at 150 °C for 1 h. Step-2 nanowire crystallization occurred after keeping nanowire arrays at 220 °C for 15 min and cooling to room temperature in 1.5 h. For crystallization of step-3 selenium nanowire arrays, extracted nanowire arrays were soaked overnight in a 10% pyridine (Riedel-de Haen) solution in isopropanol.

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References


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**Author contributions**


**Additional information**

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Arrays of indefinitely long uniform nanowires and nanotubes

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In the version of this Letter previously published, the key for Fig. 5c was incorrect. This error has now been corrected in the HTML and PDF versions.
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Nanowire array production by iterative thermal size reduction technique

Figure S1 | Nanowire array production from a macroscopic rod by iterative thermal size reduction. Step 1: A macroscopic cylindrical rod (diameter 10 mm, length 200 mm) is fabricated from the material that is to become nanowires upon axial elongation. An thermomechanically suitable polymer sheet (PES, PEI, PSU) is tightly rolled around the rod in a clean room until the final thickness is 30 mm. The composite is then thermally consolidated under vacuum, above the glass transition temperature, in order to fuse the polymer and the cylindrical rod. Finally the composite is drawn in a furnace to obtain hundreds of meters of microwires embedded in a polymer fiber matrix. Step 2: About 100 fibres with 0.5 mm diameters are cut into 200 mm fibres, tightly packed and a polymer cladding is rolled around the fibers, and consolidated; or alternatively cut fibres are inserted inside a pre-consolidated hollow polymer rod and consolidated. Second step drawing results in a submicron wire arrays in the cross section of the fibre. Step 3: The same procedure, followed in the second step is iterated, to obtain hierarchically positioned arrays of arrays of nanowires. The nanowire diameters at each step is geometrically reduced and at each step controlled by monitoring the enwrapping fibre diameter.
Dynamical scanning calorimetry (DSC) of nanowire materials and encapsulating polymers.

Nanowire and nanotube materials, and the encapsulating polymer matrices that undergo controlled thermal size reduction are thermally characterized using dynamical scanning calorimetry (DSC). Encapsulating high temperature polymers, polyethersulphone (PES) and polyetherimide (PEI), have glass transition temperatures at around 220 °C, above which they become amenable for drawing. Chalcogenide glasses, Ge_{15}As_{25}Se_{15}Te_{45} and As_{2}Se_{3} have glass transition temperatures around 190 °C. No crystallization peak is observed for these glasses in this temperature range. Semimetallic selenium glass melts at 221 °C and has a crystallization temperature of 110 °C. Metallic alloy Sn_{95}-Ag_{5} has a melting temperature of 223 °C. Polysulphone (PSU) has glass transition temperature at 190 °C and polyvinylidene fluoride (PVDF) has a melting temperature at 170 °C.

Figure S2 | Dynamical scanning calorimetry (DSC) of nanowire materials and encapsulating polymers.
Figure S3 | Low temperature, multimaterial fibre drawing tower used for the iterative size reduction of composite macroscopic rods down to nanowires and nanotubes. The 2.5 m tower consists of a thermally isolated double zone furnace, feeding, capstan, thickness, and tension meters units. Thermal size reduction takes place at a 2.5 cm hot region of the furnace under applied axial stress ($T=50$–$300$ g). Nanowire diameter is controlled using the polymer fibre diameter and tension as a feedback to the down-feed, capstan and furnace temperature controllers. Polymer fibre thickness is measured with a laser micrometer and fibre tension by a transverse tensiometer. Smaller reduction factors (10–20×) are obtained at relatively lower temperatures (260–270 °C) and smaller drawing speeds (0.05 m/min). Higher reduction factors (50–200×) are obtained with higher temperatures (280–300 °C) and higher drawing speeds (1–5 m/min). 10 kilometer-long continuous nanowire array is produced from 30-cm long macroscopic rod with a reduction factor 200.
Selenium micro and nanowire array

**Figure S4 | Ordered selenium nanowire arrays obtained by three step iterative size reduction.** Selenium is a glass making chalcogenide element with a crystallization temperature of 110 °C and melting temperature of 230 °C. Therefore it is molten at the drawing temperature of 270 °C. XRD diffraction studies indicate that Se is amorphous after each thermal drawing/size reduction. **a-b,** In the first step a 10 mm Selenium rod is reduced to 250-50 µm single wire (reduction factor 40-200x). Extracted microwires retain their global alignment. **c-d,** For the second step, ~100 selenium wires are cut and tightly packed and redrawn to 7 µm (reduction factor 34×) to obtain hundreds of meters long ordered selenium microwires. **e-f,** In the final step, previously obtained selenium microwire arrays are cut, packed and redrawn to 250 nm hierarchically ordered nanowire arrays (reduction factor 28×, total reduction factor is 40,000×).
Selenium micro- and nanowire diameter distributions after each size reduction step

Figure S5 | Selenium wire size distribution histograms after each iterative reduction step. 

a, After the second size reduction step, the size distribution of a total of 100 microwires are found to be 8.6±0.3 μm, corresponding to ±3% variation in diameter. 

b, After the third step size distribution of about 10,000 selenium microwires are found to be 262±30 nm, corresponding to a ±11% variation. Factors contributing to overall size distribution are slight variations in the fibers that are used to make the composite, and total accumulated size variation from each previous step.
Crystal structure of Selenium nanowires

Figure S6 | High-resolution transmission electron microscopy (HR-TEM) images of Selenium nanowires. a, HR-TEM image of a single crystalline selenium nanowire taken from nanowire cross section. b, HR-TEM image showing polycrystalline domains of a selenium nanowire. The insets show the electron diffraction patterns. Heat or solvent treatment of the amorphous selenium nanowires above the crystallization temperature produce nanowires that are single crystalline and/or polycrystalline.
Metal (Sn-Ag) microwire array

Figure S7 | Ordered arrays of metallic (Sn$_{95}$-Ag$_{5}$) microwires embedded in a dielectric polymer. Unlike glassy substances, metals have a sharp thermal phase change behaviour. Their viscosity change abruptly at the melting temperature. Therefore thermal size reduction of metallic filling materials is more challenging. We used a low melting temperature metal alloy, Sn$_{95}$-Ag$_{5}$ ($T_m=231$ °C). a-b, In the first step, a 1.6 mm Sn-Ag rod is reduced to 61 μm single wire (reduction factor 26×). Polymer embedded single metallic microwire and extracted microwire is shown. c-d, In the second step, ~70 fibres with metallic wires are cut and tightly packed and redrawn to 4 μm (reduction factor 15×) to obtain hundreds of meters long, ordered metallic microwires. e, Close-up SEM micrograph shows hexagonal structure of microwire array. f, Size distribution of the microwires are determined to be 3.6±0.1 μm. Due to high surface energy of metals, and melt not being able sustain shear stresses, micro wires could not be drawn further continously; beads, discontinuities and structural deformation were observed upon third drawing step (not shown).