nature electronics

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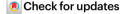
https://doi.org/10.1038/s41928-023-01047-2

Complementary carbon nanotube metaloxide-semiconductor field-effect transistors with localized solid-state extension doping

Received: 5 November 2022

Accepted: 12 September 2023

Published online: 19 October 2023



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Low-dimensional semiconductors such as one-dimensional carbon nanotubes could be used to shrink the gate length of metal-oxidesemiconductor field-effect transistors (MOSFETs) below the limits of silicon-based transistors. However, the development of industry-compatible doping strategies and polarity-control methods for such systems is challenging. Here we report top-gate complementary carbon nanotube MOSFETs in which localized conformal solid-state extension doping is used to set the device polarity and achieve performance matching. The channel of the transistors remains undoped, providing complementary metal-oxidesemiconductor-compatible n- and p-MOSFET threshold voltages of +0.29 V and -0.25 V, respectively. The foundry-compatible fabrication process implements localized charge transfer in the extensions from either defect levels in silicon nitride (SiN_x) for n-type devices or an electrostatic dipole at the SiN_x/aluminium oxide (Al₂O₃) interface for p-type devices. We observe SiN_x donor defect densities approaching 5×10^{19} cm⁻³, which could sustain carbon nanotube carrier densities of 0.4 nm⁻¹ in the extensions of scaled nanotube devices. Our technique is potentially applicable to other advanced field-effect transistor channel materials, including two-dimensional semiconductors.

Semiconducting single-walled (SW) carbon nanotubes (CNTs) are an ideal channel material for scaled, high-performance transistors because of their charge carrier mobility, injection velocity and 1 nm diameter $^{1-7}$. Recent advances in CNT transistor components have included contact resistances approaching the theoretical limit at 10 nm contact length 8 , thin high-capacitance top-gate dielectrics 9 and scaled devices with

record d.c. performance $^{10-14}$. However, from a foundry perspective, record d.c. performance observed in a scaled individual device does not, in itself, indicate complementary metal–oxide–semiconductor (CMOS) relevance. A focus on modules and techniques, which can potentially leverage demonstrated d.c. performance in the complex parameter space of CMOS manufacturing, is now needed. Key aspects include device

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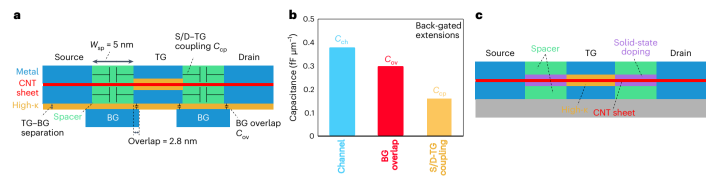


Fig. 1 | **CNT device architecture comparison. a**, Device architecture with local back gates (BG) to induce extension doping. **b**, Comparison of channel capacitance $C_{\rm ch}$ (0.37 fF μ m⁻¹) with parasitic $C_{\rm ov}$ (0.29 fF μ m⁻¹) between extension BG and top gate (TG) and parasitic coupling capacitance between gate and

S/D electrodes $C_{\rm cp}$ (0.16 fF μm^{-1}) of architecture shown in **a. c.**, MOSFET device architecture using localized extension all-around conformal solid-state extension doping. $W_{\rm sp}$ and κ are the spacer width and the relative permittivity, respectively.

architectures that minimize parasitic capacitance, device uniformity and the enablement of established CMOS metrology techniques, all of which need to be achieved while simultaneously promoting design simplicity and using standard CMOS design and process steps.

To meet the d.c. targets of the International Roadmap for Devices and Systems (IRDS)¹⁵ for beyond 2030, including their on-current ($I_{\rm on}$) and gate-length ($I_{\rm g}$) targets, gate-all-around-like architectures will be required². Global gate architectures (gate-overlapping contacts, extensions and channel) are not d.c. competitive because of a degraded above-threshold transition, leading to an $I_{\rm on}$ that is reduced up to three times for a given off-current $I_{\rm off}$ (ref. 16). Parasitic capacitance optimization and design simplicity are equally important (Fig. 1) to meet CMOS power–speed requirements. Extension doping has been identified as a key obstacle to power–speed performance because of lack of CNT-compatible, localized and foundry-friendly extension doping techniques.

However, architectures using local gates to induce extension doping (extension gates; Fig. 1a) are prohibitive because of parasitic capacitances and design complexity. Using parameters from 2031 IRDS targets including 3σ critical dimension control and 3σ overlay (see Supplementary Fig. 1 for details), the overlap capacitance (C_{ov}) between extension back gates and the top gate is estimated to introduce a parasitic capacitance exceeding 78% of the intrinsic device channel capacitance (Fig. 1b). This analysis does not include parasitic capacitance from further routing for extension gates and increased design complexity. Additionally, because of geometrical constraints, the planar nature of extension gates and their position from the CNT sheet provides inefficient extension doping and may require non-standard CMOS voltage.

In contrast, a metal–oxide–semiconductor field-effect transistor (MOSFET) device architecture using localized extension all-around conformal solid-state extension doping (Fig. 1c) could accomplish the necessary extension doping using charge transfer from defect levels in dielectrics or a broken-gap band alignment between the dielectrics and the CNT in the extension regions only. This approach can minimize parasitic capacitance, optimize doping efficiency with an undoped channel and provide design simplicity and CMOS foundry compatibility. In particular, predictive technology computer-aided design (TCAD) modelling has shown that this MOSFET architecture could meet IRDS requirements in terms of threshold voltage $V_{\rm T}$, $I_{\rm on}$, $I_{\rm off}$ and subthreshold swing 19. (A summary of key CNT device architectures previously used 16,17,20–32 is provided in Supplementary Fig. 2.)

In this article, we report top-gate complementary CNT MOSFETs on the basis of the architecture in Fig. 1c, in which device polarity is set and performance matching is achieved, using localized conformal solid-state extension doping, a module that is compatible with CMOS foundry resources. Localization of doping to the extensions only—the channel remains undoped, as in advanced silicon CMOS—is

used to reduce device variability in short-channel devices. We report impedance-voltage measurements and the extraction of interface state density $D_{\rm it}$ as a function of gap energy. The location of interface states in energy space is essential to support identification of their structural origin and develop feasible passivation schemes. Furthermore, we use TCAD to understand the doping mechanisms and extract the density of ${\rm SiN}_{\rm x}$ defects responsible for charge transfer doping. We also use TCAD to compare our experimental results with the future requirements of highly scaled devices.

CNT MOSFET fabrication, device structure and d.c. characteristics

Complementary top-gate CNT MOSFETs are fabricated on high-resistivity ($2 \times 10^4 \,\Omega$ cm) Si wafers coated by ALD with 20 nm HfO₂. Subsequently, a high-density network of SW CNTs is deposited on the HfO₂ surface. The CNT parameters include a mean value for diameter d and length of 1.42 nm (corresponding to a bandgap of 0.55 eV) and 1 μm, respectively. The linear density is about 30 μm⁻¹ (Supplementary Fig. 3). Source and drain contacts are patterned by lift-off, and palladium (Pd) contact metal is used for both n- and p-type MOSFETs³³. The gate stack consists of low-temperature (LT) AlO_x (1.5 nm) deposited directly on the CNT network, followed by Al₂O₃ (4 nm) ALD at 250 °C and Pd gate metal (30 nm) deposition. The LT AlO, process was previously shown to achieve conformal gate oxide coverage on CNTs while being compatible with subsequent ALD growth of high-κ dielectrics³⁴ and providing bilayer gate stacks with equivalent oxide thickness as low as 0.6 nm (refs. 9,34). N-type extensions are realized by electron transfer from defect levels in plasma-enhanced chemical vapor-deposited SiN_x (ref. 29) (deposition rate is about 0.13 nm s⁻¹), whereas p-type extensions rely on an electrostatic dipole at the interface of a SiN_x-Al₂O₃ bilayer. Extension doping also defines the contact polarity, avoiding the use of air exposure or process-sensitive low-working-function contact metals³⁵⁻³⁷. Fabricated top-gate n- and p-type MOSFETs have a gate length L_g , gate width W_g and source-drain distance L_{sd} of 1 μ m, $150 \, \mu m$ and $3 \, \mu m$, respectively. The detailed device fabrication flow is provided in Methods.

Figure 2 summarizes the structural and electrical device properties. Cross-section schematics illustrate the device structure (Fig. 2a,b), which is verified by cross-section transmission electron microscopy (TEM) (Fig. 2c,d) for both n- and p-type MOSFETs. The devices' polarity is set by the extension regions by using either a 100 nm SiN_x layer in direct contact with the CNT network (n-type MOSFET) or a 5 nm SiN_x layer situated on top of the Al₂O₃ gate dielectric stack (p-type MOSFET). Electrical n- and p-type CNT MOSFET properties (I-V) include transfer characteristics (drain current I_d versus gate-source voltage V_{ds} with source-drain voltage V_{ds} as parameter) on a logarithmic

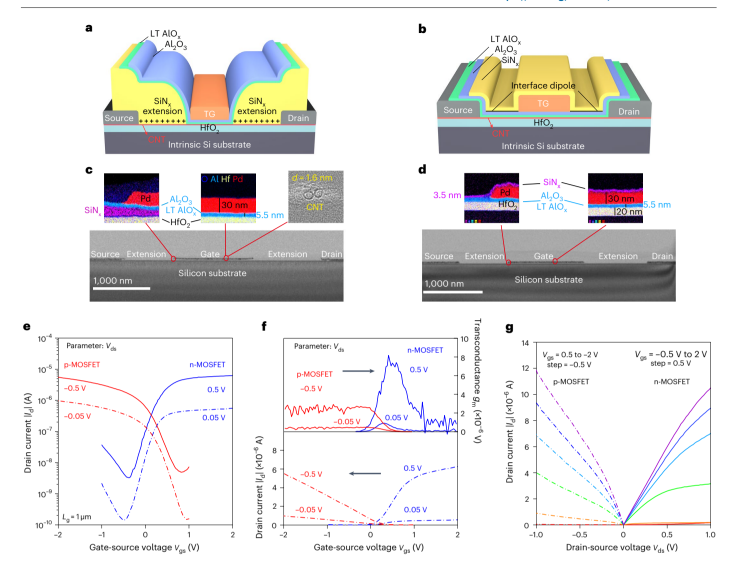


Fig. 2 | Top-gate complementary CNT MOSFETs with the devices' polarity set by localized conformal solid-state extension doping. a, Schematic of a top-gate n-type CNT MOSFET. Defects in the SiN_x at energy levels above the CNT conduction band edge E_c donate electrons to the CNT, rendering the extensions and contacts n-type, which sets the device's polarity to n-type. b, Schematic of a top-gate p-type CNT MOSFET. A dipole layer at the $\mathrm{SiN}_x/\mathrm{Al}_2\mathrm{O}_3$ interface turns extensions and contacts p-type, which sets the device's polarity

to p-type. **c**, TEM cross-section of a completed n-type CNT MOSFET. **d**, TEM cross-section of a completed p-type CNT MOSFET. **e**, Transfer characteristics of the complementary CNT MOSFETs at $|V_{\rm ds}| = 0.05$ V and 0.5 V at logarithmic scale. **f**, $I_{\rm d}$ – $V_{\rm gs}$ and $g_{\rm m}$ – $V_{\rm gs}$ characteristics of the complementary CNT MOSFETs at $|V_{\rm ds}| = 0.05$ V and 0.5 V at linear scale. **g**, Output characteristics $I_{\rm d}$ – $V_{\rm ds}$ of the complementary CNT MOSFETs with $V_{\rm gs}$ as parameter. The gate oxide (Al $_{\rm 2}$ O $_{\rm 3}$) thickness for both n- and p-type MOSFETs is 5.5 nm.

(Fig. 2e) or linear scale with transconductance g_m versus $V_{\rm gs}$ (Fig. 2f) and output characteristics $I_{\rm d}$ versus $V_{\rm ds}$ with $V_{\rm gs}$ as parameter (Fig. 2g). Good performance matching between n- and p-type devices is demonstrated with $V_{\rm T}$ (0.29 V and -0.25 V), on-current $I_{\rm on}$ (10.5 μ A, 12 μ A), on-resistance $R_{\rm on}$ (84 k Ω , 57 k Ω) and an $I_{\rm on}/I_{\rm off}$ ratio of 10^4 . Here, $V_{\rm T} = V_{\rm gs}$ at maximum transconductance g_m ($V_{\rm ds} = 0.05$ V), $I_{\rm on} = |I_{\rm d}|$ at $|V_{\rm ds}| = 1$ V and $|V_{\rm gs}| = 2$ V, $R_{\rm on} = V_{\rm ds}/I_{\rm d}$ at $|V_{\rm ds}| = 0.05$ V and $|V_{\rm gs}| = 2$ V and $I_{\rm on}/I_{\rm off} = I_{\rm dmax}/I_{\rm dmin}$ at $|V_{\rm ds}| = 0.05$ V. Note that performance matching is accomplished with only one structural modification—the extension doping—whereas all other components including gate stack, contacts, channel and substrate are identical. The extension doping is localized, and the channel remains undoped: a key feature enabling CMOS-compatible $V_{\rm T}$ for both n- and p-type devices.

MOS impedance of one-dimensional CNT channels

Quantifying carrier density and mobility in a FET channel is key to understanding and optimizing the device's performance metrics. However, no previous experiment has directly measured the channel charge in a CNT

because of the difficulty of obtaining the gate-to-channel capacitance in an environment overwhelmed by large parasitic gate-to-contacts and gate-to-substrate capacitance inherent to previously reported device architectures (Fig. 1). The carrier density has typically been estimated from electrostatic models using carrier mobility as a parameter to fit measured d.c. I-V characteristics^{3,4,25}. In this study, direct measurement of the channel charge is enabled by a combination of developments that manage parasitic capacitance and minimize its effect on the measured small-signal gate capacitance $C_{\rm g}$. These factors include a top-gate MOSFET architecture with localized solid-state extension doping, eliminating overlap of metal electrodes and associated parasitic capacitance (Fig. 2a,b); use of a high-resistivity Si substrate to minimize parasitic substrate capacitance; and on-chip zero-open and zero-short calibration structures for de-embedding measured $C_g - V_g$ characteristics from any remaining parasitic capacitance components. With all factors above combined, $C_g - V_g$ data are obtained with high confidence over a frequency range f of 100 kHz to 1 MHz.

Figure 3a shows scanning electron microscope images of layouts for CNT capacitor and on-chip zero-open and zero-short calibration

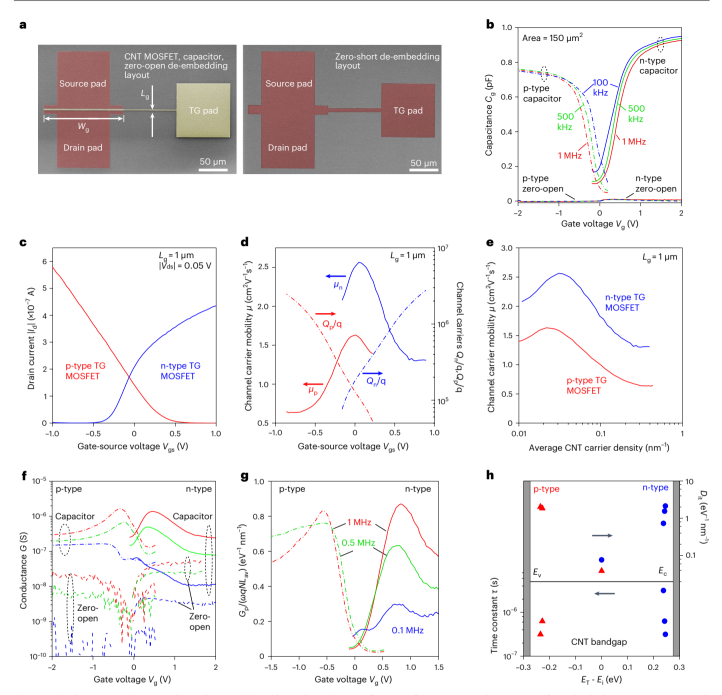


Fig. 3 | Device layout, capacitance-voltage characteristics, channel-carrier mobility and density. a, False-colour scanning electron microscope pictures of device layouts. The layout of the TG CNT MOSFET and the CNT capacitor are identical; the zero-open de-embedding structure shares the same visible layout elements (left). For MOSFET I_d – V_g , measurements, $|V_{ds}|$ = 0.05 V. Both source and drain terminals are grounded during C_g – V_g measurements. L_g and W_g are 1 µm and 150 µm, respectively. Zero-short de-embedding structure (right). b, Multifrequency C_g – V_g characteristics of CNT capacitors and of zero-open de-embedding structures after zero-open and zero-short calibration. c, CNT MOSFET I_d versus V_g s transfer curves at $|V_{ds}|$ = 0.05 V. d, Channel-carrier mobility μ and channel-carrier density Q as a function of V_g s, e, Channel-carrier mobility μ as

function of average carrier density n or p of an individual CNT. Maximum network channel mobility μ_n of 2.6 cm² V⁻¹s⁻¹ and μ_p of 1.6 cm² V⁻¹s⁻¹ is observed at an average carrier density of an individual CNT n = 0.031 nm⁻¹ and p = 0.022 nm⁻¹ for n-type and p-type MOSFETs, respectively. **f**, Multifrequency G- V_g characteristics of CNT capacitors and zero-open de-embedding structures after zero-open and zero-short calibration. The measurement frequencies are 1, 0.5 and 0.1 MHz, indicated with red, green and blue colour, respectively. **g**, Parallel conductance of interface states of CNT capacitors as a function of V_g . **h**, Interface state density D_{it} and corresponding characteristic time constant τ as a function of E_{τ} - E_{i} in the CNT bandgap.

structures. Note that the capacitor layout differs from the zero-open calibration structure only by the presence of CNTs in the active area of the capacitor; visible layout elements are identical. The capacitor layout also matches that of the MOSFET; the difference here is in the measurement condition. After completion of zero-open and zero-short

calibration using the on-chip calibration structures, multifrequency $(0.1 \le f \le 1 \, \text{MHz}) \, C_g - V_g$ characteristics of CNT capacitors and zero-open calibration structures are measured (Fig. 3b). N- and p-type zero-open structures measure a capacitance not exceeding 10 fF (Fig. 3b), keeping parasitic capacitance at 1–10% of CNT gate capacitance over the studied

gate-voltage and frequency range. To obtain carrier mobility, further data required are MOSFET I_d – $V_{\rm gs}$ curves at low $V_{\rm ds}$ (Fig. 3c) and parasitic device resistance $R_{\rm p}$, which is derived from resistor I–V measurements (Supplementary Fig. 4).

The total measured carrier density under the gate $Q = \int C_{\sigma} dV_{\sigma}$ where 1 MHz C_g - V_g curves are used to minimize the effect of trapped charge on Q. The channel-carrier mobility is obtained using $\mu = I_d L_g^2 / Q V_{ds,int}$ (ref. 38) where the intrinsic voltage $V_{ds,int} = V_{ds} - I_d R_p$. Figure 3d shows the channel-carrier mobility μ_n , μ_n and the channel charge $Q_{\rm n}$, $Q_{\rm n}$ of the CNT network as a function of $V_{\rm gs}$ for n- and p-type CNT MOSFETs, respectively. Channel-carrier mobility μ_n , μ_n versus average carrier density *n* and *p* of individual CNTs for n- and p-type MOSFETs is shown in Fig. 3e. The carrier densities observed in an individual CNT are obtained using $n = Q_n/(N_n L_{av} q)$ and $p = Q_p/(N_p L_{av} q)$, where N_n and N_n are the number of CNTs, L_{av} is the average CNT length under the gate and q is the unit charge. N_n and N_n are estimated using the ratio between total measured accumulation capacitance (9.2×10^{-13}) and 7.5×10^{-13} F for the n- and p-type MOSFET, respectively) and the average CNT accumulation capacitance per unit length $C'_{\rm mod}$ = 1.31 × 10⁻¹⁰ F m⁻¹ for the gate stack in Fig. 2, as calculated from electrostatic models³⁹ and the CNT density of states⁴⁰ (Methods). The values obtained are $N_n = 4,966$ and $N_p = 4,048$ for the n- and p-type MOSFET, respectively. The corresponding linear CNT densities are $N_p/W_g = 33$ and $N_p/W_g = 27 \,\mu\text{m}^{-1}$, which are in good agreement with the CNT density obtained from atomic force microscopy measurement (Supplementary Fig. 3). Note that for a CNT network, the carrier mobility including its peak position is determined by the conductivity of junctions connecting the CNTs and is not representative of the intrinsic CNT carrier transport. However, the demonstrated technique can be readily applied to high-purity, high-density, well-aligned CNTs when such substrates become available to experimentally obtain intrinsic CNT carrier mobility versus carrier density.

The ability to measure gate-to-channel impedance including C-Vand conductance-voltage (G-V) characteristics further enables the extraction of density and characteristic time constants of electronic states located at the interface between a CNT and the gate oxide. Figure 3f shows multifrequency $G-V_g$ characteristics of CNT capacitors and zero-open de-embedding structures. Note that the parasitic conductance signal obtained from zero-open structures is typically two to three orders of magnitude below the signal observed on the CNT capacitors. Conductance peaks, the hallmark of interface states, are only found on CNT capacitors, which provides further evidence that the conductance signal originates from the interface between the CNTs and the gate oxide. Interface state parallel conductance $G_{\rm p}$ is obtained from measured capacitance $C_{\rm g}$ and conductance G^{41} . Figure 3g shows the normalized interface state parallel conductance $G_{\rm p}/(\omega q N L_{\rm av})$ of an individual CNT, where ω and N are the angular frequency and number of CNTs under the gate, respectively. Density D_{it} and characteristic time constant τ of interface states are obtained from the peak value of each curve⁴¹ and are summarized in Fig. 3h as a function of the energy level E_T of an interface state relative to the intrinsic energy level E_i of the CNT. Interface states densities of 1–3 eV⁻¹ nm⁻¹ are observed in the vicinity of the band edges of an individual CNT with time constants around 1 µs.

The frequency dispersion observed in the measured $C_{\rm g}$ – $V_{\rm g}$ curves (Fig. 3b) is further used to obtain a first $D_{\rm it}$ estimate in the vicinity of $E_{\rm i}$ using the measured frequency dispersion at the capacitance minimum by applying the combined high-low frequency capacitance method ⁴¹. For a single CNT, $D_{\rm it}$ values of 0.076 and 0.039 eV⁻¹ nm⁻¹ are obtained for n- and p-type capacitors, respectively. Note that the derived density is a lower limit because further frequency dispersion is to be expected below 100 kHz.

Localized and tunable doping polarity for n- and p-type MOSFET extensions

Localized and tunable doping providing both n- and p-type polarity is a key ingredient of the MOSFET architecture presented in Fig. 2.

Experimental evaluation of doping efficiency is on the basis of global back-gate FETs, and doping quantification leverages electrostatic models. Cross-section schematics of CNT FETs used to develop n- and p-type solid-state doping are shown in Fig. 4a,b, respectively. All devices consist of a p $^{+}$ silicon back gate separated from the CNT network channel by a 15 nm HfO $_2$ film. The separation of the Pd contacts defines the gate length (1 µm); the device width is 50 µm. The FET polarity is determined by dielectric films subsequently deposited on top of the CNT network and simultaneously on the entire device, providing global doping.

For n-type doping, single SiN_x films are deposited under different conditions including deposition time, optimized NH₂ and SiH₄ flow rates and NH_3/SiH_4 flow ratio R = 3 (set A), whereas p-type doping is observed by depositing a SiN_x-LT AlO_x bilayer using R = 3 and 100 (set B). In Fig. 4c, selected I_d versus V_{gs} curves ($V_{ds} = 50$ mV) are shown for both sets A and B. Typically, flatband voltage shift $\Delta V_{\rm T}$ has been used to quantify doping. In an ideal device, all points along the I_d - V_{gs} curve are shifted by the same amount of $V_{\sigma s}$, and any point can be chosen to quantify doping. We found $\Delta V_{\rm T}$ to be less suitable for doping quantification because of strong charge trapping at V_T (the Fermi level E_F is located close to the CNT band edge, where a large trap density resides) and a notable effect from Schottky barriers at contacts for high current levels. Both parameters above introduce further doping-dependent shifts and stretch out of the I_d - V_{gs} , making quantitative extraction of doping density difficult. Here, we introduce the metric V_{\min} , with $V_{\min} = V_{gs}$ at minimum I_d . V_{min} is a distinct bias point with underlying physical meaning where the ambipolar device switches from one polarity to the other and the channel Fermi level resides at the intrinsic level E_i . We found V_{\min} to be not only an important quantity to identify the type of doping, the relative doping strength and quantitative doping parameters (discussed further below), but also a highly relevant metric for de-embedding doping effects from other mechanisms such as trapped charge (which is minimal at E_i because of either emission or neutralization (by means of recombination) of charge previously captured at the CNT band edge) and the effect of Schottky barriers at contacts, which is minimized at low current levels 42,43 . The position of V_{\min} is indicated on the I_d – V_{gs} curves in Fig. 4c. When a single SiN_x film is deposited (set A, n-type doping), V_{\min} shifts to more negative values with increasing SiN_x deposition time; this is accompanied by formation of a stronger n-type branch, whereas the p-type branch gradually disappears. However, when a SiN_x layer is deposited on LT AlO_x, forming a bilayer (set B), a positive V_{\min} shift is obtained, resulting in p-type doping with reinforced p-type and suppressed n-type branch.

Next, the metric ΔV_{\min} is introduced with $\Delta V_{\min} = V_{\min} - V_{\text{ref}}$, where $V_{\text{ref}} = V_{\text{min}}$ of the undoped reference case. The voltage of the undoped reference case V_{ref} is set by the intrinsic energy level E_i of the CNT with $V_{\rm ref} = E_{\rm F} - E_{\rm i} = (5.17 \, \text{eV} - 4.5 \, \text{eV})/q = 0.67 \, \text{V}$, where $E_{\rm F}$ is the Fermi energy of the back-gate p⁺Si substrate. On the basis of the definition of V_{ref} , n-type and p-type doping is observed for $\Delta V_{\min} < 0$ and $\Delta V_{\min} > 0$, respectively. In Fig. 4d, a summary of experimentally obtained ΔV_{min} data is shown as a function of SiN_x deposition time for both single SiN_x films and SiN_x -LT AlO_x bilayers. For all samples, ΔV_{min} continues to decrease with increasing SiN_x deposition time over the entire range experimentally investigated. Because only SiN_x defect levels in the vicinity of the CNT contribute to doping of the CNT (further discussed below), we consider that continuing changes of composition and atomic-bonding structure in the initially deposited SiN_x layer are likely facilitated by diffusion of species towards the interface, leading to increased SiN_x defect density in the immediate vicinity of the CNT and the SiN_x-HfO₂ interface as SiN_x deposition continues. For deposition of a SiN_x-LT AlO_x bilayer, the ΔV_{\min} versus SiN deposition time dependence is shifted along the ΔV_{\min} axis by +1.5 V, indicating the presence of an electrostatic dipole at the SiN_x-LT AlO_x interface. Note that similar dipole layers were reported earlier at SiO₂-Al₂O₃ interfaces⁴⁴.

To explain the experimentally observed dipole, models including an oxygen-density-difference model⁴⁴ and oriented Al-O-Si bonds⁴⁵

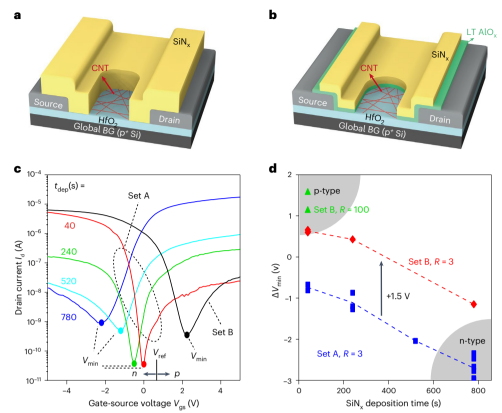


Fig. 4 | **Development of solid-state extension doping. a**, Cross-section schematic of a back-gate CNT FET used to develop n-type solid-state doping. **b**, Cross-section schematic of a BG CNT FET used to develop p-type solid-state doping. In both **a** and **b**, the opening in the dielectric layer is for illustration purposes only to reveal the CNT network. **c**, I_d versus V_{gs} curves ($V_{ds} = 0.05$ V) for single SiN_x films of different deposition time t_{dep} and R = 3 for n-type doping (set A, SiN_x t_{dep} as parameter) and for a SiN_x/LT AlO_x bilayer (set B) with R = 100 for SiN_x

for p-type doping. **d**, Summary of experimentally obtained ΔV_{\min} as a function of SiN_x deposition time t_{dep} for both single SiN_x films (set A) and SiN_x-LT AlO_x bilayers (set B, R=3,100). The ΔV_{\min} curve for bilayers (R=3) is shifted against the single layer SiN_x by +1.5 V along the ΔV_{\min} axis, indicating the presence of an electrostatic dipole at the SiN_x-LT AlO_x interface. The preferred corners for n-and p-type doping are indicated. The lines are a guide to the eye only.

were proposed. Because Al–O–Si bonds have similar polarity at $\mathrm{SiN_x-Al_2O_3}$ and $\mathrm{SiO_2-Al_2O_3}$ interfaces, the latter model can explain our observation of a dipole at the $\mathrm{SiN_x-Al_2O_3}$ interface. Although dipole layers are also reported at $\mathrm{HfO_2-Al_2O_3}$ interfaces 46 , the doping efficiency of such a dipole is less because of this interface and its corresponding dipole being located between the CNTs; this is in contrast to the $\mathrm{SiN_x-Al_2O_3}$ interface, where the dipole is situated perpendicular to the CNT surface and surrounds the CNT to a large extent. The observation of n- and p-type MOSFET V_T being essentially symmetric to zero also excludes any effect due to a dipole at the $\mathrm{HfO_2-Al_2O_3}$ interface and the assumption of a noticeable charge in the LT $\mathrm{AlO_x}$ layer because both polarity devices share the $\mathrm{HfO_2-Al_2O_3}$ interface and the LT $\mathrm{AlO_x}$ layer in the gate region.

Thus, doping is found to be tunable with the presented techniques. The preferred corners for n- and p-type doping are located in the lower-right and upper-left areas, as indicated in Fig. 4d. Consequently, single SiN_x films with deposition time exceeding 650 s (R=3) and SiN_x –LT AlO_x (SiN_x : 5 nm, R=100; AlO_x : 1.5 nm) bilayers are selected for localized n- and p-type extension doping of our top-gate and complementary CNT MOSFETs, respectively.

Beyond the experimental observation of n- and p-type doping tunability, a deeper understanding of the physical mechanism behind doping by means of spatial charge transfer from localized defect levels in dielectrics is required to extract physical parameters, optimize such techniques and understand their limitations. In the past, the effect of spatial charge transfer from dielectrics has often been ascribed to a fixed charge uniformly distributed throughout the depth of the dielectric layer. Such models are simplistic and do not consider that

each charge is associated with an energy level in the energy band structure of a solid. In the case of SiN_v, the defect structure is complex^{47,48}.

For illustration purposes, we use here the well-known Si dangling bond centre =Si⁰ termed K⁰ (ref. 48). The K⁰ centre is amphoteric, having a lower donor type transition level +/0 and a higher acceptor type transition level 0/-. A tight-binding recursion method⁴⁹ placed the +/0 transition level at -4.3 eV (ref. 50), which is essentially identical to the conduction band minimum E_c of the CNTs considered here. Calculated energy-band parameters have often been afflicted with substantial uncertainty; for the purpose of this treatise, we assume that the energy E_D of the +/0 transition level is situated 0.6 eV above the E_c of the CNT ($\Delta E = 0.6$ eV). Note that for electron donation, the energy E_D of the SiN_x donor level must be located above E_c of the CNT channel. The density N_K of ionized K⁰ centres follows Fermi–Dirac statistics

$$N_{\rm K}^{+} = N_{\rm K}^{0} \left[1 - \frac{1}{1 + e^{\frac{E_{\rm D} - E_{\rm F}}{kT}}} \right]$$

where N_K^0 , k, T are the density of K^0 defect centres, the Boltzmann constant and the absolute temperature, respectively. Solving Poisson's equation

$$\nabla (\kappa \nabla \varphi) = -\frac{\rho(\varphi)}{\varepsilon_0} = -\frac{q}{\varepsilon_0} \left\{ N_{K}^{+}(\varphi) + p(\varphi) - n(\varphi) \right\}$$

using TCAD^{40,51}, where φ , ρ , and ε_0 are the electrostatic potential, the volume charge density, and the permittivity of vacuum, respectively,

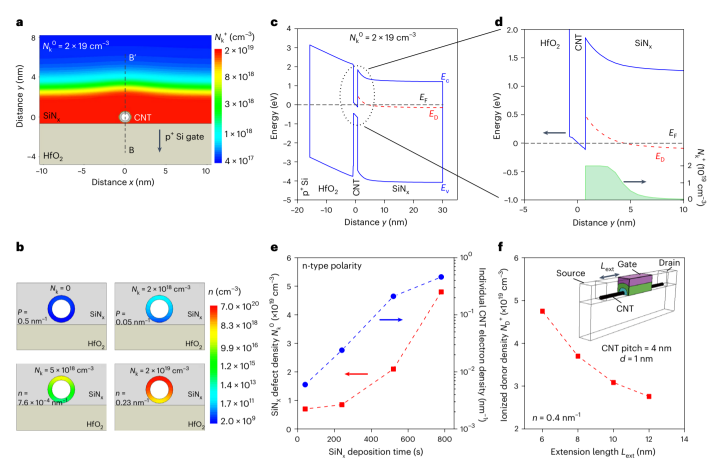


Fig. 5 | **Modelling of solid-state extension doping. a**, 2D density distribution of ionized K^0 defect centres $N_{\rm k}^+$ for a cross-section perpendicular to the sourcedrain path in Fig. 4a. The SiN_x N_{K^0} centre density and energy position ΔE of the +/0 level used in the simulations are uniform throughout the SiN_x film and equal to $N_{K^0} = 2 \times 10^{19} \, {\rm cm}^{-3}$ and $\Delta E = 0.6 \, {\rm eV}$, respectively. The ionized charge $N_{\rm k}^+$ originating from the SiN_x K^0 centres is confined to the vicinity of the CNT and the first few nanometres of the SiN_x layer. **b**, CNT carrier concentration distributions and electron and hole concentrations per unit length for SiN_x K^0 centre density ranging from 0 to 2 × 10¹⁹ cm⁻³. **c**, Energy-band diagram for identical parameters

along the line B–B' shown in **a**. The work function of the p* Si substrate and the CNT diameter are 5.17 eV and 1.45 nm, respectively. **d**, Detail of energy-band diagram in **c** including ionized defect centre density $N_{\rm K}$ *. **e**, SiN_x defect density $N_{\rm K}$ 0 and electron density n of individual CNTs as function of SiN_x deposition time. **f**, Calculated ionized donor density $N_{\rm D}$ * versus extension length $L_{\rm ext}$ required to maintain a CNT electron density n in the extension of 0.4 nm⁻¹ for a potential future scaled planar device architecture, as shown in the inset. Here, ionized donors of concentration $N_{\rm D}$ * are assumed to be uniformly distributed in the extension volume with length $L_{\rm ext}$ and thickness of 5 nm.

we obtain the two-dimensional (2D) density distribution of ionized K^0 centres (N_K^+) shown in Fig. 5a for a cross-section perpendicular to the source-drain path of the device shown in Fig. 4a. The depth of the space charge region along the y axis can be estimated as

$$d \cong \sqrt{2\kappa\varepsilon_0 \Delta E/qN_{\rm K}^0} = 4.5 \,\mathrm{nm}$$

using the parameters given in Fig. 5 and a κ value of 6 for SiN_{κ}.

In the vicinity of the CNT, electron transfer from K^0 centres to the CNT is energetically favoured, leading to simultaneous n-type doping of the CNT (Fig. 5b) and K^0 centre ionization (Fig. 5a). For $N_{K^0}=0$, CNT polarity is p-type because of the work function (5.17 eV) of the p^+ Si back gate; however, CNT polarity is increasingly converted to n-type with higher N_{K^0} density (Fig. 5b). Figure 5c shows the energy-band diagram along the line B–B' in Fig. 5a. SiN $_x$ energy bands are flat, and space charge is essentially zero beyond a SiN $_x$ depth d of 5 nm. This is further illustrated in Fig. 5d, which shows a detail of the energy-band diagram in Fig. 5c including ionized defect centre density N_K^+ . To finally close the loop between the experimentally extracted values of V_{\min} for the investigated SiN $_x$ deposition conditions (Fig. 4c) and the models developed above, the SiN $_x$ defect density N_{K^0} and the CNT carrier density are

extracted for different SiN_x deposition times (Fig. 5e). This is accomplished by finding the density N_{K^0} for each V_{\min} shown in Fig. 4c and then calculating the hole and electron density (Fig. 5b) at zero bias using the identified N_{K^0} values. Thus the SiN_x defect parameters are determined from electrical measurements. Figure 5f shows the calculated ionized donor density N_{D}^+ versus extension length L_{ext} required to maintain a CNT electron density n of 0.4 nm $^{-1}$ for a potential future scaled device architecture, as shown in the inset, as calculated from electrostatic models 39 and the CNT density of states 40 . Our experimentally obtained N_{K^0} values approach required values for extension doping of such scaled devices. To compromise between performance and leakage, we use 0.4 nm $^{-1}$ as a doping strength. This target allows us to meet contact resistance requirements, especially for short contact length 2 ; source-to-drain direct tunnelling, which increases with higher extension doping, meets targets as well.

Doping due to spatial charge transfer also has limitations, as implied in Fig. 5f, because the electrostatic potential generated from ionized dopants is relatively inefficient to induce charge carriers in a CNT near contact and gate-metal interfaces because of charge screening. Thus, doping CNTs at short extension length and CNT pitch is even more challenging. Limitations due to metal screening apply to all spatial charge-transfer techniques; for example, using extension gates

to induce electrostatic potential throughout a short region between source/drain (S/D) and gate metal is even less efficient and, further, not applicable to stacked nanosheets such as reported for 2D stacked nanosheet MOSFETs⁵². Another approach is substitutional doping; however, even assuming the absence of physical damage to the CNT, theory shows stronger Coulomb scattering for substitutional doping compared to spatial charge-transfer doping leading to more severe performance degradation of CNFETs⁵³, and strong device variations may result from random substitutional doping effects in a small volume. Spatial charge transfer from dielectrics for extension doping is also a critical component for future stacked 2D nanosheet devices⁵². 2D nanosheet MOSFETs require a vertically stacked architecture to meet IRDS requirements¹⁵. A promising approach is to provide extension doping by inner spacer dielectrics.

Conclusions

We have reported process modules for CNT MOSFET architecture fabrication with 2031 IRDS targets in mind. These modules could potentially be used to build a future, foundry-relevant CNT CMOS technology. We fabricated top-gate complementary CNT MOSFETs for which the devices' polarity is set and performance matching is accomplished exclusively by localized conformal solid-state extension doping, a module that is compatible with CMOS foundry resources. Localization of doping to extensions only is an important aspect of this approach and reduces device-to-device variability in a short-channel device as dopant fluctuations are a substantial source of variability. Enabled by the extension doping module, we provided impedance-voltage measurements and the extraction of interface state density $D_{\rm tt}$ as a function of gap energy. The location of interface states in energy space is essential to support identification of their structural origin and develop feasible future passivation schemes.

Methods

Purification of semiconducting CNTs

Semiconducting CNTs are purified using a 1:1 ratio by weight of arc-discharge CNT powder (Carbon Solution, Inc., AP-CNT) and conjugated polymer PCz (poly[9-(1-octylonoyl)-9H-carbazole-2,7-diyl)]) (American Dye Source, Inc.), which are dispersed at a concentration of 2 mg per ml in toluene. This mixture is sonicated with a tip sonicator (Sonic, VCX-750, 7 mm probe tip) for 30 min at 600 W in a 20 °C cool flowing water bath to dissipate excess heat. The dispersed solution is centrifuged in a swing bucket rotor (Beckman, Optima XPN-90, SW 32 Ti Swinging Bucket Rotor) at 50,000g for 20 min to remove the undispersed material. The top 90% of the supernatant is collected. Then the centrifugation process is repeated at 50,000g for 2 hours to further remove the metallic CNT. Finally, the top 90% supernatant is collected for the CNT network deposition by the immersion method.

CNT network deposition

CNT networks were deposited by the immersion method. The target substrates were immersed in the diluted CNT in toluene solution for 18 hours to form randomly orientated, uniform high-density CNT networks. After the deposition, the target substrate was soaked in toluene for 20 min and baking in ambient at 180 °C for 30 min.

Fabrication process of network CNT-based transistors

A 500 μ m thick p⁺-doped Si wafer (University Wafer no. 785) was used as the substrate for the back-gate FET devices for doping development, and a 500 μ m high-resistivity Si wafer (University Wafer no. 3328) was used as the substrate for the complementary MOSFET devices. To remove unwanted impurities, Radio Corporation of America treatment was done, followed by a spin–rinse–dry cycle. A 20 nm HfO₂ film was deposited on the wafer at 200 °C using an ALD system (Fiji F202 Cambridge Nanotech) followed by CNT network deposition. The uniformity of the network CNTs across the entire wafer was confirmed by SEM

(FEI Apreo). The density of the network CNTs and length distribution were verified by atomic force microscopy (VEECO Dimension 3100). The diameter distribution of the CNTs was extracted using TEM (Supplementary Fig. 3). The source and drain metal contact were defined by photolithography (Heidelberg MLA150), and 30 nm Pd was deposited by e-beam evaporation (Temescal BJD 1800). The channel regions were protected by photoresist that was defined by photolithography, after which the excess CNTs (CNTs outside the FET structure) were removed by oxygen plasma etch (Oxford PlasmaLab 80). Subsequently, the residual photoresist on the channel region was removed using AZ NMP. The wafer was then diced, and the next steps were done on small chips.

Back-gate CNT FET for doping study

After the residual photoresist was removed with AZ NMP, SiN_x thin films were deposited using plasma-enhanced chemical vapor deposition (PECVD; Oxford PlasmaLab 80plus). Various thicknesses of SiN_x were deposited ontop of the substrate by changing the deposition time. During all runs with different SiN_x deposition times, the system was kept at a temperature of 270 °C, and plasma was run at a frequency of 13.56 MHz at 20 W. The pressure of the chamber was kept at 900 mTorr with, for example, 5% SiH_4 (dilute in He) at a flow rate of 200 sccm and a NH_3 flow rate of 30 sccm (R=3). To open the contacts, openings were defined by photolithography and SiN_x etched using buffered oxide etchant (BOE). The detailed fabrication flow of the back-gate FETs is shown in Supplementary Fig. 5.

Top-gate CNT n-MOSFET

The chip with predefined source and drain contacts was first baked at 180 °C for 5 mins to remove moisture. Afterwards, 100 nm SiN $_{\rm x}$ (R=3) was deposited using the PECVD tool. To deposit the gate oxide for the gate region, the SiN $_{\rm x}$ in the gate region was removed using wet-etch methods. The gate region was defined by photolithography and the SiN $_{\rm x}$ etched with BOE. Subsequently, the entire chip was immersed in AZ NMP overnight to remove all photoresist. A LT AlO $_{\rm x}$ film with a thickness of 1.5 nm was deposited by ALD at 50 °C followed by an additional 4 nm high-temperature Al $_{\rm 2}O_{\rm 3}$ that was also deposited by ALD but at 250 °C. To deposit the top gate, the pattern was defined by photolithography and 30 nm Pd was deposited by e-beam evaporation. Vias over contact pads were patterned by lithography, and then SiN $_{\rm x}$ and gate dielectrics were etched with BOE to enable electrical measurements. The detailed MOSFET process flow is shown in Supplementary Fig. 6.

Top-gate CNT p-MOSFET

The chip with predefined source and drain contacts was first baked at 180 °C for 5 mins to avoid any moisture from the previous process. A 1.5 nm LT AlO $_{\rm x}$ film was deposited on top of the chip by ALD at 50 °C. Then an additional 4 nm Al $_{\rm 2}O_{\rm 3}$ were deposited by ALD at 250 °C. Similar to the n-MOSFET top-gate process, the gate region was defined by photolithography and the 30 nm Pd gate metal was deposited by e-beam evaporation. Next, 5 nm of SiN $_{\rm x}$ (R = 100) was deposited using the PECVD tool. To perform the electrical measurement, openings on contact pads were made by photolithography and BOE etch.

I-V measurement

All the transistor characterization measurements were performed in a common-source configuration. Electrical I-V data was measured using a shielded probe station and an Agilent B1500A semiconductor device parameter analyzer equipped with high-resolution source monitor units. The transfer characteristics ($I_{\rm d}-V_{\rm gs}$) were performed at a bias sweep rate of 120 mV s⁻¹ with steps of 30 mV. For the output characteristics ($I_{\rm d}-V_{\rm ds}$), the drain voltage ($V_{\rm ds}$) was swept from 0 to +1 V or -1 V for n-MOSFET and p-MOSFET, respectively, in steps of 20 mV and at a sweep rate of 100 mV s⁻¹. The gate-source voltage $V_{\rm gs}$ was swept from the off state of the device ($V_{\rm gs}=-0.5$ V to 2 V for n-MOSFET and $V_{\rm gs}=0.5$ V to -2 V for p-MOSFET). The back-gate FET transfer characteristics were combined from two different $V_{\rm gs}$ sweeps. In each sweep, $V_{\rm gs}$ was varied

either from 0 to +5 V or from 0 to -5 V in steps of 100 mV at a sweep rate of 400 mV/s. This technique was used to minimize the effect of charge trapping at a large-gate bias of up to 5 V, which can lead to shift of the $I_{\rm d}$ - $V_{\rm gs}$ curve along the $V_{\rm gs}$ axis.

C-V measurement

Multifrequency C-V and G-V measurements were carried out using a precision LCR metre (Agilent 4284A) connected to a Keithley 2420. The 'high' terminal was connected to both source and drain and the 'low' terminal to the gate. A small-signal ac excitation with amplitude of 25 mV was applied to the device, and the d.c. bias was swept with a rate of 12 mV s⁻¹ in steps of 30 mV. An external bias (Keithley 2420) was connected to the LCR metre using an adaptor (Keysight 16065C) to control the bias applied to the sample. Simultaneous multifrequency sweeps were performed by sweeping the bias and obtaining the capacitance at three different frequencies at each bias point. The device was encapsulated in a Faraday cage to cancel out any noise from the environment. Before any measurements, the parasitic capacitances were cancelled out by performing an open-and short calibration on fabricated on-chip open-and-short de-embedding structures. All devices were measured at three different frequencies: 1 MHz, 500 kHz and 100 kHz.

TCAD simulations

Electrostatic models are implemented in either Nano Device Simulator 39 or Sentaurus Device framework 51 . An analytical CNT density-of-state model depending on CNT diameter is used to compute the carrier density in the CNT 40 .

For the simulations supporting Fig. 3, the number of CNTs under the gate *N* is estimated from the accumulation capacitance as follows:

$$N = \frac{C_{\text{exp}}}{C'_{\text{mod}} L_{\text{av}}} = \frac{C_{\text{exp}}}{C'_{\text{mod}} (L_g/\cos\varphi)}$$

where $C_{\rm exp}$, $C_{\rm mod}$, $L_{\rm av}$ and φ are the total measured accumulation capacitance, the calculated average accumulation capacitance per CNT unit length and the average length and average orientation of CNTs under the gate, respectively, with $L_{\rm av} = L_{\rm g}/\cos\varphi$. $C_{\rm mod}' = 1.31 \times 10^{-10}~{\rm F~m^{-1}}$ is obtained using the Poisson solver in the Nano Device Simulator ³⁹ and the CNT density-of-state model ⁴⁰ for the geometry of the gate-stack cross-section in Fig. 2 (5.5 nm Al₂O₃ with k = 7.8 (ref. 9)). The corresponding oxide capacitance per unit length is $1.65 \times 10^{-10}~{\rm F~m^{-1}}$. Assuming an average orientation of 45°, $N_{\rm n}$ = 4,966 and $N_{\rm p}$ = 4,048 are obtained. Note that the quantities extracted for individual CNTs including carrier density n or p (= $Q/(NL_{\rm av}q)$, normalized interface state parallel conductance $G_{\rm p}/(\omega qNL_{\rm av})$ and interface state density $D_{\rm it} = 2.4G_{\rm p,peak}/(\omega qNL_{\rm av})$ do not depend on the assumption of average CNT orientation φ because $NL_{\rm av} = C_{\rm exp}/C_{\rm mod}$, which is independent of φ . Here, $G_{\rm p,peak}$ is the peak value of the $G_{\rm p}$ versus $V_{\rm g}$ dependence.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Acknowledgements

We acknowledge the use of facilities and instrumentation supported by the National Science Foundation through the University of California San Diego Materials Research Science and Engineering Center DMR-2011924.

Author contributions

Z.Z. and M.P. contributed equally. Z.Z. conducted the device fabrication and measurement. M.P. developed the impedance and doping models, device layout and data analysis. G.P., W.E.S., N.S. and T-E.L. contributed to device fabrication, and S.N. supported impedance measurements. S.-K.S. and G.D. contributed to device modelling. T.-A.C. provided the CNT substrates. S.L.L., V.D.-H.H. and C.-F.H. provided TEM analysis. I.R., A.C.K., P.B. and H.-S.P.W. guided the project. Z.Z. and M.P. prepared the paper draft, and all the authors commented on the final version.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41928-023-01047-2.

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Peer review information *Nature Electronics* thanks the anonymous reviewers for their contribution to the peer review of this work.

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