High-density magnetoresistive random access memory operating at ultralow voltage at room temperature

Jia-Mian Hu\textsuperscript{1,2}, Zheng Li\textsuperscript{1}, Long-Qing Chen\textsuperscript{2} & Ce-Wen Nan\textsuperscript{1}

The main bottlenecks limiting the practical applications of current magnetoresistive random access memory (MRAM) technology are its low storage density and high writing energy consumption. Although a number of proposals have been reported for voltage-controlled memory device in recent years, none of them simultaneously satisfy the important device attributes: high storage capacity, low power consumption and room temperature operation. Here we present, using phase-field simulations, a simple and new pathway towards high-performance MRAMs that display significant improvements over existing MRAM technologies or proposed concepts. The proposed nanoscale MRAM device simultaneously exhibits ultrahigh storage capacity of up to 88 Gb inch\textsuperscript{-2}, ultralow power dissipation as low as 0.16 fJ per bit and room temperature high-speed operation below 10 ns.
Magnetoresistive random access memory (MRAM), because of its moderately fast access time and almost unlimited endurance, has offered a tantalizing application potential as next-generation non-volatile integrated memories. However, mass production of MRAMs has long been hampered by their high writing energy from Ampère-current-induced magnetic field. This would also cause severe cross-talk among neighbouring cells when miniaturizing the device size for higher storage capacity. One promising solution is to develop voltage-write MRAM devices, that is, manipulating magnetization directly using electric voltage rather than current. Previous proposals include integrating a spin valve or magnetic tunnel junction (MTJ) unit onto a multiferroic layer (for example, CrO₂ (ref. 7) or BiFeO₃(ref. 8)), and more recently using ferroelectric (FE) oxide as the tunnel barrier as in a Fe/BaTiO₃/(La,Sr)MnO₃ junction where significant voltage control of tunnel magnetoresistance was realized at rather low temperatures. Nonetheless, no existing proposals have so far achieved low voltage, high storage density and room temperature operation all at the same time.

In this study, we illustrate a simple and new approach towards ultralow voltage-controlled MRAMs using phase-field simulations. We show that the proposed device is capable of achieving ultralow writing energy (less than 0.16J per bit), room temperature (or even wider range) operation and high storage density (up to 88 Gb inch⁻²) all at the same time, associated with reliable thermal stability (quality factor over 40 at T=300K) and comparable writing speed (sub-10ns).

Results
Ultralow voltage-controlled MRAM device structure. Figure 1a illustrates one possible layout of a memory cell array, which builds on planar complementary metal-oxide-semiconductor (CMOS) platforms. As in normal MRAMs, it has a typical 1-T(transistor)/1-magnetoresistive(MR) element cell architecture (Fig. 1b), where the readout is accomplished by sensing the resistance change of the cell. However, the most remarkable feature is its ideal zero current, gate-controlled-write process (see Methods section for the operation time sequence), which is fast and extremely power-efficient. There are virtually no cross-talks among neighbouring cells. It has a simple structure, which is easier to manufacture than conventional MRAMs, due to the absence of extra current-carrying lines. The simplicity offers a great potential for realizing high storage capacity. The basic building block, incorporating a spin valve or MTJ unit onto a (001)-oriented perovskite FE (for example, lead magnesium niobate-lead titanate, PMN-PT) layer (or maybe (001)-oriented rhombohedral FE layer with spontaneous polarization along (111) directions), is shown in Figure 1c. An external electric voltage is perpendicularly applied to the PMN-PT layer triggered by the connecting transistor. This produces large anisotropic in-plane piezostrains, which would be further imparted to its upper magnetic-free layer and induces a 90° in-plane magnetization rotation in the latter via the strain-mediated inverse magnetoelastic coupling. By operating the (011)-oriented PMN-PT layer below its FE coercive field, bistable in-plane piezostrains can be obtained, that is, the strain states can be maintained even when the applied voltages are switched off, as demonstrated by a permanent magnetization switching observed recently in polycrystalline Ni thin film grown on (011) PMN-PT. This interesting bistable magnetization switching should bring greater design flexibility to the present voltage-controlled MRAM devices, that is, the non-volatility can be achieved not only by virtue of high-quality epitaxial magnetic thin films whereby the magnetocrystalline anisotropy serves as the potential barrier, but also in polycrystalline and/or amorphous thin films, despite a little lower magnetoresistance ratio due to structural imperfections in the latter.

Compared with previous MRAM proposals, the present voltage-write MRAM device is actuated at an ultralow voltage, as discussed below. Second, the strain-mediated magnetoelastic coupling offers a straightforward way to electrically manipulate the magnetization at room temperature, whereas the promising Fe/BaTiO₃/(La,Sr)MnO₃ junction is limited to low temperature. Moreover, our proposed voltage-write MRAM device has a much wider operation temperature range than the voltage-controlled MRAM that uses CrO₂ with its Neel temperature (T₆=307K) close to room temperature. Third, the perpendicular writing voltage bias makes it easy to integrate it with existing CMOS platforms to fabricate nanoscale high-density memory devices. In contrast, multiferroic BiFeO₃-based MRAMs demonstrated so far require an in-plane actuation electric field. Such a transverse actuation voltage along the in-plane direction usually resorts to an inter-digitized transducer patterning, which is typically of a micron size, and intrinsically prohibits high storage capacity. This is also the case for the voltage-controlled MRAM using (001)-oriented FE layer.

Working and design principles of the device. For illustration, we perform phase-field simulations of this electric-field-controlled MRAM device by choosing a (001) Ni film as the free layer. A spin valve comprising a synthetic antiferromagnetic layer (Fig. 1c) is considered as the upper MR element while an MTJ is also available. An initial uniform (010) magnetization is assumed for the Ni-free layer, which is exchange coupled to its upper reference layer. This direction can be set by annealing the device in a magnetic field. As described above, a perpendicular electric field E is applied to the
magnetization states are exhibited at $E = 0$, for example, $m_x = 0.9538$ and 0.0364 for the $64 \times 64 \times 35$ nm$^3$ Ni-free layer, corresponding to a magnetization rotation of about 70°. This is also directly illustrated by their magnetic vector diagrams shown in Figure 2b. Moreover, the magnetization–$E$ curves become gradually flatter with the film thickness decreasing from 35 to 5 nm, along with a small increase in the maximum magnetization rotation around 5°. This can be attributed to the suppression of the out-of-plane magnetization component due to enhanced demagnetization in thinner magnetic thin films, which would in turn facilitate the in-plane magnetization rotation. The larger magnetization rotation further leads to a broader resistance change of the spin valve, as the device resistance is determined by the relative magnetization alignment, analytically expressed as\(^{(1)}\)

\[
R(m_x) = \frac{2(1 + R_{\text{MR}}) R_s}{2 + R_{\text{MR}} (1 + \sqrt{1 - m_x^2})}.
\]

Here $R_s$ denotes the device resistance with parallel magnetization configuration in the reference and free layer, and $R_{\text{MR}}$ the maximum magnetoresistance ratio, that is, the relative device resistance change between the parallel and antiparallel magnetization alignments. For illustration, $R_{\text{MR}}$ is taken as 7% for a normal spin valve\(^{(1)}\) with giant magnetoresistance effect. Corresponding to the greater magnetization switching, increased maximum resistance changes are exhibited as indicated in Figure 2c. This may suggest the first design principle of such voltage-write MRAM device: the thinner, the better. However, growing ultrathin films requires more sophisticated processing control. Besides, the interface magnetic anisotropy contribution may lead to additional effect on the magnetization switching\(^{(11)}\), especially in ultrathin (several monolayers) films\(^{(19)}\), but is not considered herein regarding the robust magnetoelastic coupling of Ni as well as the large piezoelectric response of PMN-PT, which should in principle produce a large strain-mediated magnetoelastic response\(^{(19)}\). Nonetheless, based on the results in Figure 2, 5 nm could be a well-compromised thickness for metallic Ni-free layer deposited on the PMN-PT. In particular, the actuation electric field required for the MRAM device is ultralow, that is, $|E| < 0.2$ V $\mu$m$^{-1}$ (Fig. 2c).

As small lateral size of the spin valve is highly desirable for pursuing high storage density, we examine the lateral size effect of the free layer on the $E$-induced magnetization switching while fixing the film thickness at 5 nm. As seen from Figure 3a, a square-shaped magnetization–$E$ loop is exhibited even when the lateral size reduces to 32 nm, and hence it can be used for bistable memory implementation. This naturally leads to the second design principle of this $E$-controlled MRAM device: the smaller, the better. However, it cannot be considered as a generalized conclusion, as the complex magnetization distribution near the lateral edges would, to some extent, hamper the in-plane magnetization rotation when the lateral sizes are shrunken. This is evidenced by a moderate reduction of the maximal magnetization rotation when the lateral size decreases from 64 to 32 nm (Fig. 3a). One may refer to the magnetic vector diagrams in Figure 3b to compare the different magnetization distributions near the square edges. In addition, the deterioration of thermal stability with reduced free layer volume is another issue that will be discussed below. On the other hand, for the free layers with lateral size larger than 128 nm, the $E$-induced in-plane magnetization rotation would be significantly cancelled out by the presence of multi-domains (see the corresponding magnetic vector diagram in Fig. 3b) driven by the demagnetization field in the lateral direction, and thus leads to much smaller relative resistance changes, as indicated in Figure 3c. It should be noted that the magnetization–resistance relationship in multi-domain-free layers might not exactly be described by equation (1), as different regions of magnetizations (namely, the magnetic domains) could separately contribute to the resistance change. Nevertheless, it is still rational to believe...
The thermal stability factor is defined as $F = f_{\text{struc}} V_{\text{i}}$, where $F$ is the energy barrier that separates the two stable magnetization states ($V_{\text{i}}$ denotes the volume of the free layer, $k_B$ the Boltzmann constant and $T$ the temperature set as 300 K). To ensure a reliable non-volatility, a thermal stability factor of higher than 40 is normally required. While common MRAMs rely on the intrinsic magnetic anisotropy energy (for example, the magneto-crystalline anisotropy) to retain the magnetization state, the energy barrier mostly results from the extrinsic elastic strain stimuli in the present proposal, which eliminates the specific crystal structure requirements for the free layer and offers more design flexibility. The potential barrier energy $f_{\text{struc}}$ can be estimated by calculating the free energy difference between the two stable magnetization states at $E = 0$. Similarly, fixing the film thickness to 5 nm, the lateral size dependence of the thermal stability factor is given in Figure 4, with their corresponding relative resistance changes shown for comparison. From the plots, it can be clearly seen that the free layer with lateral sizes from 64 to 160 nm exhibit both sufficient thermal stability and comparable resistance change, although the latter two attributes appear not possible to be simultaneously optimized. Nevertheless, the results indicate that Ni films with lateral sizes of between 64 and 160 nm, and thickness of about 5 nm, should be suitable for application as the free layers of such novel MRAM devices. If further reducing the lateral size for higher storage density, then efforts should be made to compensate the concomitant loss of the thermal stability, as well as the reduction of the maximum resistance change due to the hampered in-plane magnetization rotation.

As mentioned at the beginning, the ultralow power consumption, resulting from both the inherent energy efficiency of electrical switching and its ultralow actuation voltage, is the most advantageous quality of the present ultralow electric-voltage-controlled MRAM, far superior to any other existing or emerging MRAM technologies. Figure 2c shows such a small $E$ writing window from about −0.13 to 0.1 V $\mu$m$^{-1}$. That means, if using a 2-μm PMN-PT thick layer, an ultralow voltage of about 0.26 V would be sufficient for the device operation. This operation voltage is much lower than that reported previously in the proof-of-concept experiments. For further illustration, we estimate the writing energy as $\frac{1}{2} P_{\text{FE}} V_{\text{i}}$ in terms of charging the FE capacitor, where $P_{\text{FE}}$, $S$ and $V$ denote...
the remnant polarization, electrode area and the operation voltage, respectively. In this case, an extremely low writing energy of around 0.16 fJ per bit can in principle be obtained by taking \( P_e = 30 \mu \text{C cm}^{-2} \) (ref. 13), \( S = 0.0041 \mu \text{m}^2 \) (for a \( 64 \times 64 \text{ nm}^2 \) free layer electrode) and \( V = 0.26 \text{ V} \). This energy consumption is drastically lower than 70 pJ per bit for conventional MRAMs\(^*\), and even orders of magnitude lower than 0.1 pJ in the state-of-the-art spin-transfer-torque (STT) MRAM technology\(\dagger\), which is deemed as the most reachable solution for next-generation MRAMs, and is currently under intense industrial interests.

As this low actuation electric field remains below the FE coercive field of the PMN-PT (approximately ±0.2 V \( \mu \text{m}^{-1} \) (ref. 13)), the fatigue problem that FEs (including multiferroic BiFeO\(_3\)) typically suffer from repetitive polarization reversal can also be greatly relieved. This should allow an effective alleviation of the energy loss and improvement of device reliability.

Furthermore, the dielectric breakdown vulnerability of the present device can be avoided as the write operating voltage, far below the dielectric breakdown threshold of the PMN-PT (>10 V \( \mu \text{m}^{-1} \) (ref. 24)), is applied only on the PMN-PT layer. This structural feature of the present device is different from that in STT-RAMs, where the write and read current flow across the whole device and share the same tunnel.

**Writing speed and storage capacity.** Another important performance attribute of the voltage-controlled MRAM device is the writing speed, which typically takes less than 10 nanoseconds (ns) or with a possibility of sub-1 ns (see the write-time estimation and corresponding electric-field-induced dynamic magnetization switching in the Methods section). This operation speed is significantly higher than conventional magnetic-write MRAM (around 20 ns (ref. 1)), while comparable with the STT-MRAM devices (~3 to 10 ns (ref. 25)). Finally, the fully gate voltage-controlled operation allows a good compatibility of the present device with current CMOS platforms, opening up possibilities for ultrahigh densities. For instance, assuming a typical channel length of 45 nm for the bottom transistor (Fig. 1a) and the lateral size of 64 nm for the upper ME spin valve (or MTJ)/PMN-PT heterostructure, a storage density of around 88 Gbit/inch\(^2\) can in principle be realized, challenging traditional NAND Flash memory on mass data storage applications\(\ddagger\). For comparison, the properties obtained in the present device prototype are summarized in Table 1, including several other existing room temperature non-volatile memory technologies.

### Discussion

We have demonstrated a simple and new approach towards ultralow voltage-controlled MRAMs. The present voltage-controlled MRAM device can be fabricated on CMOS platforms, wherein the basic building block comprises of a spin valve or MTJ MR element grown on a (011) PMN-PT layer. Without necessarily requiring epitaxial growth of the magnetically free layer, simply utilizing a specifically engineered FE layer with voltage-induced bistable anisotropic in-plane piezostrains, one can realize most important attributes required by modern MRAM technology at the same time. Therefore, we expect our finding will stimulate future experimental and engineering efforts on developing such an electric-voltage-controlled device, which has superiorities over the previous device paradigms\(^2\)-\(^9\),\(^14\),\(^22\). The present device proposal represents a simple pathway to realizing high-density, low voltage-controlled MRAM applications at room temperature, either as standalone/embedded memories or logic circuits.

**Figure 5 | Device operation illustration.** (a) Cell array image of the voltage-controlled MRAM device. BL, PL and WL stands for bitline, plateline and wordline, respectively. (b) Illustration of writing time sequence. The state ‘1’ is written at the time stage \( t_3 \), where the voltages in BL2 and PL are in high and low level, respectively. Writing a ‘0’ (corresponding to the dash dot line in the row BL2) is accomplished at the time stage \( t_5 \) in a similar manner, but the BL2 and PL are in reverse polarities.

### Table 1 | Comparison of room temperature non-volatile memory technologies.

| Technology       | Storage capacity | Write time | Read time | Write energy (pJ per bit) | Write energy (pJ per bit) |
|------------------|------------------|------------|-----------|---------------------------|---------------------------|
| Flash-NAND       | >1 Gb            | >100 Mb    | 10 ns     | >0.01                      | >0.03                     |
| FeRAM            | >1 Gb\(\ddagger\) | >10 ns     | 10 ns     | >1 Gb\(\ddagger\)         | >1 Gb                     |
| MRAM             | 16 Mb\(\ast\)    | 20 ns      | 10 ns     | 70                         | 0.1                       |
| STT-RAM          | 1 Gb\(\ast\)     | 3–10 ns    | 10 ns     | 1.6 \times 10^4           |                           |
| SME-RAM          | 1 Gb\(\ast\)     | <10 ns     | 10 ns     | 1 Gb\(\ast\)              |                           |

\(\ast\)The proposed memory device prototype is denoted as the strain-mediated magnetoelectric random access memory (SME-RAM), where the estimated storage capacity, write time and write energy are summarized below. The read time of the present SME-RAM is assumed to be the same as conventional MRAM and STT-RAM devices, due to their similar magnetoresistive readout processes.

\(\dagger\)FeRAM indicates the ferroelectric random access memory. Performance parameters of Flash-NAND, FeRAM, MRAM and STT-RAM are taken from commercial products unless stated otherwise.

\(\ddagger\)Can be further scaled down and would possibly reach 64 MB or 128 MB (ref. 1).

\(\ddagger\)The 1-Gb STT-RAM has been demonstrated in research laboratories and could be industrialized in the near future (ref. 46).
Methods

Operation time sequence of the device. As seen in Figure 5a, the write voltage bias is applied between the Bitline (BL2) and the plateline (PL), which is controlled by the Wordline (WL) connecting to the gate of transistor. Corresponding writing time sequence is shown in Figure 5b. While to read, the sensing current (I_{sen}) travels through the transistor and the MR element from BL2 to BL1, to generate the high and low output voltages, that is,

\[
V_{\text{high}} = I_{\text{sen}} (R_{\text{mos}} + R_{\text{high}}), \\
V_{\text{low}} = I_{\text{sen}} (R_{\text{mos}} + R_{\text{low}}),
\]

where \( R_{\text{mos}} \) is the resistance of the transistor; \( R_{\text{high}} \) and \( R_{\text{low}} \) denote the bistable resistance states of the MR element resulting from the voltage-induced magnetization switching in the magnetic-free layer. The reference current (I_{ref}) is set to a specific value so that the corresponding reference voltage (V_r) is ideally between \( V_{\text{high}} \) and \( V_{\text{low}} \).

Introduction to the phase-field method. In the phase-field approach, the magnetic domain structure is described by the spatial distribution of the local magnetization vectors \( \mathbf{m} = \mathbf{M} \mathbf{m} = M_i (m_x, m_y, m_z) \), where \( \mathbf{M} \) and \( m_i \) (i = x, y, z) represent the saturation magnetization and the direction cosine, respectively. The temporal evolution of the magnetization configuration and thus the domain structure is governed by the LLG equation, that is,

\[
(1 + \gamma^2) \frac{d \mathbf{M}}{dt} = -\gamma_0 \mathbf{M} \times \mathbf{H}_{\text{eff}} - \frac{\gamma \sigma}{M_s} \mathbf{M} \times (\nabla \cdot \mathbf{H}_{\text{eff}}),
\]

where \( \gamma \) and \( \sigma \) are the gyromagnetic ratio and the damping constant, respectively, and \( \mathbf{H}_{\text{eff}} \) is the effective magnetic field, given as

\[
\mathbf{H}_{\text{eff}} = -\frac{1}{M_s} \frac{1}{M_s} \nabla \cdot \mathbf{M} = -\frac{1}{M_s} \nabla \cdot \frac{1}{M_s} \mathbf{M}.
\]

Here \( \mu_0 \) denotes the vacuum permeability, and \( F_{\text{tot}} \) is the total free energy of the magnetic layer,

\[
F_{\text{tot}} = F_{\text{mc}} + F_{\text{ms}} + F_{\text{ex}} + F_{\text{elas}} + F_{\text{dp}},
\]

where \( F_{\text{mc}}, F_{\text{ms}}, F_{\text{ex}}, F_{\text{elas}} \) and \( F_{\text{dp}} \) are the magneto crystalline anisotropy, magnetostatic, magnetic exchange and elastic energy, respectively. \( F_{\text{mc}} \) describes the ferromagnetic-type dipole coupling between the free layer and its upper reference layer, given as

\[
F_{\text{mc}} = -\mu_0 M_f H_{\text{dp}} m_i
\]

for an initial [010] magnetization; \( H_{\text{dp}} \) is the effective inter-layer dipole coupling field taken as 40 Oe for illustration (see more discussions on the influence of \( H_{\text{dp}} \) in Supplementary Fig. S2 and Supplementary Note 2). The magnetostatic energy \( F_{\text{ms}} \) can be written as,

\[
F_{\text{ms}} = -\frac{1}{2} \mu_0 M_f \int \mathbf{H}_{\text{d}} \cdot \mathbf{m} \, dV,
\]

where \( \mathbf{H}_{\text{d}} \) denotes the stray field arising from the long-range magnetization interactions in the system, given as

\[
\mathbf{H}_{\text{d}} \left( \begin{array}{c} \mathbf{M} \\ \mathbf{M} \end{array} \right) = N \begin{array}{c} \mathbf{M} \\ \mathbf{M} \end{array} \text{ being the average magnetization field}. \]

To incorporate the influence of the specific sample geometric size, a finite-size magnetostatic boundary condition is used for obtaining \( F_{\text{ms}} \) with the demagnetization factors \( N \) in the magnetostatic energy calculated numerically \( \gamma^2 \).

The mathematical expressions of \( F_{\text{elas}} \) and \( F_{\text{ex}} \) for cubic ferromagnets are the same as those given in refs 11 and 27.

The elastic energy \( F_{\text{elas}} \) can be written as

\[
F_{\text{elas}} = \frac{1}{2} \int c_{ijkl} \varepsilon_j \varepsilon_k \varepsilon_l \varepsilon_m \, dV = \frac{1}{2} \int c_{ijkl} \left( \varepsilon_j - \varepsilon_l^0 \right) \left( \varepsilon_j - \varepsilon_l^0 \right) \, dV,
\]

where \( c_{ijkl} \) is elastic constant, \( c_{ijkl} \) the elastic stiffness tensor and \( \varepsilon_i \) the total strain that can be represented as the sum of homogenous and heterogeneous strains following Khachaturyan’s mesoscopic elastic theory \( \gamma^2 \),

\[
\varepsilon_i = \varepsilon_i^0 + \eta_i.
\]
The heterogeneous strain $\eta$ is defined in such a way that $\int \eta dV = 0$, and the homogeneous strain $\varepsilon_{0}$ describes the macroscopic shape change of the magnetic thin film. Specifically, $\varepsilon_{11}$ and $\varepsilon_{22}$, namely, the homogeneous strains along the two in-plane crystal axes, can be expressed as,

$$\varepsilon_{11} = 0 + \varepsilon_{1}$$

(8)

$$\varepsilon_{22} = 0 + \varepsilon_{2}. $$

Here $\varepsilon_{1}$ and $\varepsilon_{2}$ denote the contribution from the piezoelectric transformation of the bottom PMN-PT layer, which we directly use the experimental data reported in ref. 13 as the input. We also assume that the PMN-PT layer is much thicker than the upper magnetic film to ensure a full strain transfer22. $\varepsilon_{i}$ is the biaxial residual strain resulting from the lattice and/or thermal mismatch. For a polycrystalline magnetic film deposited at room temperature, $\varepsilon_{i}$ almost equals zero; while for an epitaxial film, it depends on the specific substrate as well as the growth condition12. Note that $\varepsilon_{i}$ may have no influence on the in-plane magnetization rotation (Supplementary Fig. S3 and Supplementary Note 3), although it does have effects on the specific magnetic domain structure in the free layer11. Here it is taken as $-0.5$ for illustration. $\varepsilon_{ij}$ represents the stress-free strain describing the stress-free deformation of a cubic ferromagnet associated with the local magnetization change,

$$\varepsilon_{ij} = \frac{3}{2} \lambda_{ij0} (m_{ij} - \frac{1}{2}) (i \neq j).$$

(9)

$$\lambda_{ij0} = \frac{3}{2} \lambda_{ij} m_{ij} (i \neq j).$$

where $\lambda_{ii0}$ and $\lambda_{ij0}$ are the magnetostriuctive constants. By further considering a mixed boundary condition for the film-substrate system22, $\varepsilon_{ij}$ can be calculated by combining Khachatryan’s theory34 with the Stroh’s formalism of anisotropic elasticity35.

The temporal evolution of the local magnetization and thus the domain structures are obtained by numerically solving the LLG equation using the semi-implicit Fourier spectral method36. To illustrate the Ni-free layer with a size of 64 × 64 × 35 nm², the magnetic film is discretized into a three-dimensional array of cubic cells of 64Å × 64Å × 35Å, with the grid size $\Delta x = 0.8 \Delta y = \Delta z = 1$ nm in real space accordingly. A three-dimensional finite-size boundary condition is considered to calculate the geometric-size-related magnetostatic anisotropy energy as mentioned above. By varying the simulation zone size and/or the grid size, magnetic-free layers with different geometric sizes (for example, thickness and lateral size, see Figs 2 and 3) can then be treated. Each simulation starts with an initial (010) magnetization distribution accordingly and proceeds for long enough time to ensure a stabilized magnetization distribution, with a normalized time step $\Delta t = 0.01$. The Landau, elastic and magnetic parameters used for simulation are described in the literature16,17.

Write-time estimation. Such electric-field-controlled MRAM device can at least allow a sub-10 ns writing, which is much faster than conventional magnetic-field-switched MRAM (about 20 ns (ref. 1)) and comparable with practical STT-MRAM devices (about 3–10 ns (ref. 25). First, based on LLG equation, the average magnetization switching time is estimated as about 0.95 ns for magnetic-free layers with various lateral sizes (the thickness is set as 5 nm likewise), typically allows a GHz writing-time estimation with different geometric sizes (for example, thickness and lateral size, see Figs 2 and 3). Moreover, it is worth noting that there is no full polarization effect due to the thin film, see related discussion in ref. 40). It is rational to consider the stress to be applied almost instantaneously to the film51. Thus in total, the theoretical write time of such electric-field-controlled MRAM device can be estimated to be about 1.62 ns (0.95 + 0.67), well within a 10 ns range, required by future applicable MRAM technologies16. This result is comparable with the results reported for other related device proposals with similar strain-mediating electric-write procedure, such as multiferroic logic44 or nanowire memory devices45, which use either Terfenol-D(Ni (ref. 42)) particles, or FeGa (ref. 43) nanowires attached to a commercial Pb(Zr,Ti)O$_3$, layer, all with an estimated operation time less than 10 ns.

Electric-field-induced dynamic magnetization switching. The combination of phase-field model with micromagnetic simulation here in this paper allows us to go beyond the previous macro-spin model used for single-domain magnets51. Indeed, the free layer does not always behave as a single macro-spin when the thickness is large enough, which means the strain transfer time could be even shorter. Overall, the magnetic dynamic magnetization switching process could be far more complex than the simple model described above (such as the influences of the mechanical resonance frequency, and the Resistor-Capacitor time constant associated with charging the FE capacitor, see related discussion in ref. 40), it is rational to consider the stress to be applied almost instantaneously to the film51.

As for the piezoelectric transfer time $\tau$, we give a rough estimation on this short time span54,

$$\tau = \frac{d_1}{v_1} + \frac{d_2}{v_2}.$$ 

(10)

where $d_1$ and $d_2$ are the thicknesses; and $v_1$ and $v_2$ the velocity of sound in the FE and FM layer, respectively. For illustration, assuming $d_1 = 5$ nm (the Ni film-thickness), $d_2 = 3,000$ nm (it is well known that the speed of sound varies from substance to substance, but generally it travels very fast in solids, say, about 5,120 m s$^{-1}$ in iron at 293 K), then we have $\tau = 0.67$ ns. Note that we have considered the stress wave that covered the whole device system, but the effective mechanical coupling might be more likely to be confined in a closer vicinity of the interface, which means the stress transfer time could be even shorter. Overall, although the practical strain mediation situation could be far more complex than the simple model described above (such as the influences of the mechanical resonance frequency, and the Resistor-Capacitor time constant associated with charging the FE capacitor, see related discussion in ref. 40), it is rational to consider the stress to be applied almost instantaneously to the FM film51.

References

1. International Technology Roadmap for Semiconductors, 2007 Edition Emerging Research Devices, http://www.itrs.net (accessed October 2011).
2. Chatzistamatiou, X. & Lent, M. The emergence of spintronics in data storage. Nat. Mater. 8, 831–832 (2009).
3. Wu, K.-F. et al. Magnetotransport properties of magnetically soft spin-valve heterostructures. J. Appl. Phys. 97, 033903 (2005).
4. Binek, C. & Doudin, B. Magnetoelectronics with magnetoelectrics. J. Phys: Condens. Mater. 17, L39–L44 (2005).
5. Diemy, B. et al. Magnetotransport properties of magnetically soft spin-valve structures. J. Appl. Phys. 69, 4774 (1991).
6. Moodera, J. S., Kinder, L. R., Wong, T. M. & Meservey, R. Large magnetoresistance at room temperature in ferromagnetic thin film tunnel junctions. Phys. Rev. Lett. 74, 3273 (1995).
7. Chen, X., Hochstrasser, M. & Borisenko, P. & Kleemann, W. Magnetic exchange bias systems in spintronics. Appl. Phys. Lett. 89, 205208 (2006).
8. Behren, M. & Barthelme, H. Multiferroics: towards a magnetoelectric memory. Nat. Mater. 7, 425–426 (2008).
9. Garcia, V. et al. Ferroelectric control of spin polarization. Science 327, 1106–1110 (2010).
10. Liu, M. et al. Giant electric field tuning of magnetic properties in multiferroic ferrite/ferroelectric heterostructures. Adv. Funct. Mater. 19, 1826–1831 (2009).
11. Hu, J.-M. & Nan, C. W. Electric-field-induced magnetic easy-axis reorientation in ferromagnetic/ferroelectric heterostructures. Phys. Rev. B 80, 224416 (2009).
12. Ma, J., Lu, J.-M., Li, Z. & Nan, C.-W. Recent progress in multiferroic magnetic nanoelectronic composites: from bulk to thin films. Adv. Mater. 23, 1062–1087 (2011).
13. Wu, T. et al. Giant electric-field-induced reversible and permanent magnetization reorientation on magnetoelectric Ni(011) [Pb(Mg$_{1/3}$Nb$_{2/3}$O$_3$)$_{0.99}$PbTiO$_3$] heterostructure. Appl. Phys. Lett. 98, 012504 (2011).
14. Chu, Y. H. et al. Electric-field control of local ferromagnetism using a magnetoelectric multiferroic. Nat. Mater. 7, 478–482 (2008).
15. Hu, J.-M., Li, Z., Wang, J. & Nan, C. W. Electric-field control of strain-mediated magnetoelectric random access memory. J. Appl. Phys. 107, 093912 (2010).
16. Pertsev, N. A. & Kohlstedt, H. Resistive switching via the converse magnetoelectric effect in ferromagnetic multilayers on ferroelectric substrates. Nanotechnology 21, 245002 (2010).
17. Cavaco, C., van Kampen, M., Lagae, L. & Borghs, G. A room-temperature electrical field-controlled magnetic memory cell. J. Mater. Res. 22, 2111–2115 (2007).
18. Wolf, S. A. et al. Spintronics: a spin-based electronics vision for the future. Science 294, 1488–1495 (2001).
19. Hu, J.-M., Nan, C.-W. & Chen, L.-Q. Size-dependent electric voltage controlled magnetic anisotropy in multiferroic heterostructures: interface-charge and strain comodiated magnetoelectric coupling. Phys. Rev. B 83, 134408 (2011).
20. Ikeda, S. et al. A perpendicular-anisotropy CoFeB–MgO magnetic tunnel junction. Nat. Mater. 9, 721–724 (2010).
21. Ohno, H. A window on the future of spintronics. Nat. Mater. 9, 952–954 (2010).
22. Maruyama, T. et al. Large voltage-induced magnetic anisotropy change in a few atomic layers of iron. Nat. Nanotechnol. 4, 158–161 (2009).
23. Bedau, D. et al. Ultrafast spin-transfer switching in spin valve nanopillars with perpendicular anisotropy. Appl. Phys. Lett. 96, 022514 (2010).
24. Park, S.-E. & Shrouf, T. R. Ultrafast strain and piezoelectric behavior in relaxor based ferroelectric single crystals. J. Appl. Phys. 82, 1804 (1997).
25. Hu, J.-M. Spin-transfer torque MRAM (STT-MRAM): challenges and prospects. AAPP Bulletin 18, 33–40 (2008).
26. Chung, A., Deen, J., Lee, J.-S. & Meyyappan, M. Nanoscale memory devices. Nanotechnology 21, 412001 (2010).
27. Zhang, J.-X. & Chen, L.-Q. Phase-field microlactycity and micromagnetic simulations of domain structures in giant magnetostriective materials. Acta Mater. 53, 2845–2855 (2005).
28. Khachatryan, A. G. Theory of Structural Transformation in Solids (Wiley, 1983).
29. Pertsev, N. A., Kohlstedt, H. & Dkhil, B. Strong enhancement of the direct magnetoelectric effect in strained ferroelectric–ferromagnetic thin-film heterostructures. Phys. Rev. B 80, 054102 (2009).
30. Schalom, D. G. et al. Strain tuning of ferroelectric thin films. Annu. Rev. Mater. Res. 37, 589–626 (2007).
31. Hu, J.-M., Sheng, G., Zhang, J.-X., Nan, C.-W. & Chen, L.-Q. Phase-field simulation of strain-induced domain switching in magnetic thin films. Appl. Phys. Lett. 98, 112505 (2011).
32. Chen, L.-Q. Phase-field method of phase transitions/domain structures in ferroelectric thin films: a review. J. Am. Ceram. Soc. 91, 1835–1844 (2008).
33. Stroh, A. N. Steady state problems in anisotropic elasticity. J. Math. Phys. 41, 77–103 (1962).
34. Chen, L.-Q. & Shen, J. Applications of semi-implicit Fourier-spectral method to phase field equations. Comput. Phys. Commun. 108, 147–158 (1998).
35. Vaz, C. A. F. et al. Magnetism in ultrathin film structures. Rep. Prog. Phys. 71, 056501 (2008).
36. Walowski, J. et al. Intrinsic and non-local Gilbert damping in polycrystalline nickel studied by Ti: sapphire laser Is spectroscopy. J. Phys. D: Appl. Phys. 41, 164016 (2008).
37. Kohlstedt, H. et al. Current status and challenges of ferroelectric memory devices. Microelectron. Eng. 80, 296–304 (2005).
38. Li, Y. L., Hu, S. Y., Liu, Z. K. & Chen, L.-Q. Phase-field model of domain structures in ferroelectric thin films. Appl. Phys. Lett. 78, 3878 (2001).
39. Schroder, K. Stress operated random access, high-speed magnetic memory. J. Appl. Phys. 53, 2759 (1982).
40. Mohammad Salehi, F. et al. Magnetization dynamics, Bennett clocking and associated energy dissipation in multiferroic logic. Nanotechnology 22, 155201 (2011).
41. International Technology Roadmap for Semiconductors, 2010 Future Memory Devices Workshop Summary, http://www.itrs.net/ (accessed October 2011).
42. Atulasimha, J. & Bandyopadhyay, S. Bennett clocking of nanomagnetic logic using multiferroic single-domain nanomagnets. Appl. Phys. Lett. 97, 173105 (2010).
43. Dean, J., Bryan, M. T., Schrefl, T. & Allwood, D. A. Stress-based control of magnetic nanowire domain walls in artificial multiferroic systems. J. Appl. Phys. 109, 023915 (2011).
44. Callegaro, L. & Puppin, E. Stress dependence of coercivity in Ni films: thin film to bulk transition. Appl. Phys. Lett. 68, 1279 (1996).
45. Natarajan, S., Chung, S., Paris, L. & Keshavarzi, A. Searching for the dream embedded memory. IEEE Solid-State Circuits Magazine 1, 34–44 (2009).
46. Driskill-Smith, A. et al. Latest advances and roadmap for in-plane and perpendicular STT-RAM. in Memory Workshop (IMW), 3rd IEEE International. 1–3 (2011).

Acknowledgements
This work was supported by the NSF (grant nos. DMR-1006541 and DMR-0820404), the National Basic Research Program of China (grant no. 2009CB623303) and the NSF of China (grant nos. 50832003 and 50921061). The computer simulations were carried out on the LION and Cyberstar clusters at the Pennsylvania State University, supported in part by NSF Major Research Instrumentation Program through grant OCI-0821527 and in part by the Materials Simulation Center and the Graduated Education and Research Services at the Pennsylvania State University.

Author contributions
J.M.H. conceived and performed the simulations. J.M.H., Z.L. and C.W.N. designed the device structure. C.W.N. and L.Q.C. directed the work. J.M.H., L.Q.C. and C.W.N. wrote the paper. All contributed to the analysis.

Additional information
Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Hu, J.-M. et al. High-density magnetoresistive random access memory operating at ultralow voltage at room temperature. Nat. Commun. 2:553 doi: 10.1038/ncomms1564 (2011).

License: This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivative Works 3.0 Unported License. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/3.0/