Random access DNA memory using Boolean search in an archival file storage system

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DNA is an ultrahigh-density storage medium that could meet exponentially growing worldwide demand for archival data storage if DNA synthesis costs declined sufficiently and if random access of files within exabyte-to-yottabyte-scale DNA data pools were feasible. Here, we demonstrate a path to overcome the second barrier by encapsulating data-encoding DNA file sequences within impervious silica capsules that are surface labelled with single-stranded DNA barcodes. Barcodes are chosen to represent file metadata, enabling selection of sets of files with Boolean logic directly, without use of amplification. We demonstrate random access of image files from a prototypical 2-kilobyte image database using fluorescence sorting with selection sensitivity of one in 10^6 files, which thereby enables one in 10^6 selection capability using N optical channels. Our strategy thereby offers a scalable concept for random access of archival files in large-scale molecular datasets.

While DNA is the polymer selected by evolution for the storage and transmission of genetic information in biology, it can also be used for the storage of arbitrary digital information at densities far exceeding conventional data storage technologies such as flash and tape memory, at scales well beyond the capacity of the largest existing data centres1. Recent progress in nucleic acid synthesis and sequencing technologies continues to reduce the cost of writing and reading DNA, foreshadowing future commercially competitive DNA-based information storage2–4. Demonstrations of its viability as a general information storage medium include the storage and retrieval of books, images, computer programs, audio clips, works of art and Shakespeare’s sonnets using a variety of encoding schemes5–9, with data size limited primarily by the cost of DNA synthesis. In each case, digital information was converted to DNA sequences composed of ~100–200 nucleotide data blocks for ease of chemical synthesis and sequencing. Sequence fragments were then assembled to reconstruct the original, encoded information.

While considerable effort in DNA data storage has focused on increasing the scale of DNA synthesis, as well as improving encoding schemes, an additional crucial aspect of data storage systems is the ability to efficiently retrieve specific files or arbitrary subsets of files. To date, molecular random access has largely relied on conventional polymerase chain reaction (PCR)10,11,12,13, which uses up to ~20–30 heating and cooling cycles with DNA polymerase to selectively amplify specific DNA sequences from a DNA data pool using primers. Nested addressing barcodes14–16 have also been used to uniquely identify a greater number of files, as well as biochemical affinity tags to selectively pull down oligos for targeted amplification17.

While powerful demonstrations of PCR have shown successful file retrieval from a 150 GB file system18, notable limitations include, first, the length of DNA needed to uniquely label DNA data strands for file indexing, which reduces the DNA available for data storage. For example, for an exabyte-scale data pool, each file requires at least three barcodes19, or up to sixty nucleotides in total barcode sequence length, thereby reducing the number of nucleotides that can be used for data encoding. Second, PCR-based retrieval requires an aliquot of the entire data pool to be irreversibly consumed for random access, and therefore additional PCR amplification of the entire data pool may periodically be needed to restore this loss of data. In this case, each PCR amplification may introduce stochastic variation in copy number of the file sequences, leading to up to 2% data loss per amplification20 if using tenfold physical redundancy, as recently suggested21. Finally, avoiding spurious amplification of off-target files due to crosstalk of PCR primers with incorrect barcodes or main file sequences requires careful primer design20. While strategies exist to circumvent these preceding challenges, they generally reduce data density and might not be easily scalable to exabyte and larger file systems. For example, data loss due to periodic PCR amplification of the entire data pool20 may be reduced by increasing the physical redundancy of the files in the main data pool, and PCR crosstalk can be mitigated by spatial segregation of data into distinct pools21 or extraction of selected DNA using biochemical affinity22,23.

As an alternative to PCR-based approaches, here we introduce a direct random access memory approach that retrieves specific files, or arbitrary subsets of files, directly using physical sorting, without a need for amplification, and without any potential for barcode-memory crosstalk, while also preserving non-selected files intact by recycling them into the original memory pool. To realize this file system, we first encapsulate DNA-based files physically within discrete, impervious silica capsules22,24, which we subsequently surface-label with unique single-stranded DNA barcodes that offer Boolean-logic-based selection on the entire data pool via simple hybridization. Downstream file selection may then be optical, physical or biochemical, with sequencing-based read-out following de-encapsulation of the memory DNA from the silica capsule. Each ‘unit of information’ encoded in DNA we term a ‘file’, which includes both the DNA encoding the main data as well as any additional components used for addressing, storage and retrieval. Each file contains a ‘file sequence’, consisting of the DNA encoding the main data, and ‘addressing barcodes’, or simply ‘barcodes’, which are additional short DNA sequences used to identify the file in solution.

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using hybridization. We refer to a collection of files as a ‘data pool’ or ‘database’, and the set of procedures for storing, retrieving and reading out files is termed a ‘file system’ (Supplementary Section 0 for a full list of terms).

As a proof-of-principle of our archival DNA file system, we encapsulated 20 image files, each composed of a ~0.1 kilobyte image file encoded in a 3,000-base-pair plasmid, within monodisperse, 6 μm silica particles that were chemically surface labelled using up to three 25-nucleotide single-stranded DNA (ssDNA) oligonucleotide barcodes chosen from a library of 240,000 orthogonal primers, which allows for individual selection of up to ~10^15 possible distinct files using only three unique barcodes per file (Fig. 1). While we chose plasmids to encode DNA data in order to produce microgram quantities of DNA memory at low cost and to facilitate a renewing ability, closed-cycle write–store–access–read system using bacterial transformation into bacteria to replenish the molecular file database thereby closed the write–access–read cycle.

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**File synthesis**

Digital information in the form of 20 icon-resolution images was stored in a data pool, with each image encoded into DNA and synthesized on a plasmid. We selected images of broad diversity, representative of distinct and shared subject categories, which included several domestic and wild cats and dogs, US presidents and several human-made objects such as an airplane, boats and buildings (Fig. 1 and Supplementary Fig. 1). To implement this image database, the images were substituted with black-and-white, 26 × 26 pixel images to minimize synthesis costs, compressed using run-length encoding and converted to DNA (Supplementary Figs. 1 and 2). Following synthesis, bacterial amplification and sequencing validated the images to minimize synthesis costs, compressed using run-length encoding and converted to DNA (Supplementary Figs. 1 and 2).

To synthesize multiple layers of encapsulated DNA to increase the data density per capsule (Supplementary Fig. 4) showed full encapsulation of plasmids without residual DNA in solution.

To investigate the fraction of capsules that contained plasmid DNA, we compared the fluorescence intensity of the intercalating dye TO-PRO when added pre- versus post-encapsulation (Fig. 2b). All capsules synthesized in the presence of both DNA and TO-PRO showed a distinct fluorescence signal, consistent with the presence of plasmid DNA in the majority of capsules, compared with a silica particle negative control that contained no DNA. In order to test whether plasmid DNA was fully encapsulated versus partially exposed at the surface of capsules, capsules were also stained separately with TO-PRO post-encapsulation (Fig. 2b). Using qPCR to measure released DNA after de-encapsulation, we estimated 10⁶ plasmids per capsule (Supplementary Fig. 5 and Supplementary Table 3). Because encapsulation of the DNA file sequence relies only on electrostatic interactions between positively charged silica and the phosphate backbone of DNA, our approach can equally encapsulate any molecular weight of DNA molecule applicable to megabyte and larger file sizes, as demonstrated previously, and is compatible with alternative DNA file compositions such as 100–200-base oligonucleotides that are commonly used or multiple layers of encapsulated DNA to increase the data density per capsule.
Next, we chemically attached unique content addresses on the surfaces of silica capsules using orthogonal 25-nucleotide ssDNA barcodes (Supplementary Fig. 6) describing selected features of the underlying image for file selection. For example, the image of an orange tabby house cat (Supplementary Fig. 1) was described with ‘cat’, ‘orange’ and ‘domestic’, whereas the image of a tiger was described with ‘cat’, ‘orange’ and ‘wild’ (Supplementary Fig. 1 and Supplementary Table 2). To attach the barcodes, we activated the surface of the silica capsules through a series of chemical steps. Condensation of 3-aminopropyltriethoxysilane with the hydroxy-terminated surface of the encapsulated plasmid DNA provided a primary amine chemical handle that supported further conjugation reactions (Fig. 2c).

We modified the amino-modified surface of the silica capsules with 2-azido acetic acid N-hydroxysuccinimide (NHS) ester followed by an oligo(ethylene glycol) that contained two chemically orthogonal functional groups: the dibenzocyclooctyne functional group reacted with the surface-attached azide through strain-promoted azide–alkyne cycloaddition, while the NHS-ester functional group was available for subsequent conjugation with a primary amine. Each of the associated barcodes contained a 5′-amino modification that could react with the NHS-ester groups on the surface of the silica capsules, thereby producing the complete form of our file. Notably, the sizes of bare, hydroxy-terminated silica particles representing capsules without barcodes were comparable with complete files consisting of capsules with barcodes attached, confirmed using scanning electron microscopy (Fig. 2d,e). These results were anticipated, given that the encapsulation thickness was only on the order of 10 nm (ref. 19) and that additional steps to attach functional groups minimally increase the capsule diameter. We also observed systematic shifts in the surface charge of the silica particles as different functional groups were introduced onto their surfaces (Fig. 2e).

Using hybridization assays with fluorescently labelled probes35–37, we estimated the number of barcodes available for hybridization on each file to be on the order of 108 (Supplementary Fig. 7). To select the ‘Airplane’ file, we hybridized an AFDye-647-labelled ssDNA probe that is complementary to the barcode ‘flying’, which is unique to ‘Airplane’. We were able to detect and select the desired ‘Airplane’ file through FAS even at a relative abundance of 10−6 compared with each other file (Fig. 3c). While comparable in sensitivity to a nested PCR barcoding data indexing approach17, unlike PCR, which requires 20–30 rounds of heating and cooling to selectively amplify the selected sequence, our approach selects files directly without the need for thermal cycling and amplification. This strategy also applies to the gating of N barcodes simultaneously in parallel optical channels, which offers a file selection sensitivity of one in 106 total files, where common commercial FAS systems often reach up to N = 17 channels38,39. For example, comparison of the retrieved sequences between the ‘flying’ gate and the ‘NOT flying’ gate after chemical release of the file sequences from silica encapsulation revealed that 60–95% of the ‘Airplane’ files were sorted into the flying gate (Supplementary Figs. 18–21), where we note that any sort probability above 50% indicates enrichment of ‘Airplane’ within the correct population subset (flying) relative to the incorrect subset (‘NOT flying’). In general, a file with sort probability p has sort error rate 1 – p if it is a target file and error rate p if it is an off-target file. In this case, for example, a sort probability of 100% for ‘Airplane’ into the ‘flying’ subset would indicate ideal performance. Besides single-file selection, our approach allows for repeated rounds of FAS selection, as well as Boolean logic, described below.

### Boolean search

Beyond direct selection of one in 10^6 individual random files, without thermal cycling or loss of fidelity due to primer crosstalk, our system offers the ability to apply Boolean logic to select random file subsets from the data pool. AND, OR and NOT logical operations were applied by first adding to the data pool fluorescently labelled ssDNA probes that were complementary to the barcodes (Fig. 4, left). This hybridization reaction was used to distinguish one or several files in the data pool, which were then sorted using FAS.

We used two to four fluorescence channels simultaneously to create the FAS gates that executed the target Boolean logic queries (Fig. 4, middle).

To demonstrate a NOT query, we added to the data pool an AFDye-647-labelled ssDNA probe that hybridized to files that contained the ‘cat’ barcode. Files that did not show an AFDye 647 signal were sorted into the ‘NOT cat’ subset (Fig. 4a and Supplementary Fig. 14a).

An example of an OR gate was applied to the data pool by simultaneously adding ‘dog’ and ‘building’ probes that both had the tetramethylrhodamine (TAMRA) label (Fig. 4b and Supplementary Fig. 14b). All files that showed a TAMRA signal were sorted into the ‘dog OR building’ subset by the FAS.

Finally, an example of an AND gate was achieved by adding ‘fruit’ and ‘yellow’ probes that were labelled with AFDye 647 and TAMRA, respectively. Files showing a signal for both AFDye 647 and TAMRA were sorted into the ‘fruit AND yellow’ subset in the FAS (Fig. 4c and Supplementary Fig. 14c). For each example query, we validated our sorting experiments by releasing the file sequence from silica encapsulation and sequencing the released DNA with Illumina MiniSeq (Fig. 4, right). Sort probabilities of each file for each search query are shown in Supplementary Figs. 22–24.

The preceding demonstrations of Boolean logic gates enable file sorting with a varying specificity of selection criteria for the retrieval of different subsets of the data pool. FAS can also be used to create multiple gating conditions simultaneously, thereby increasing...
the complexity of target file selection operations, as noted above. To demonstrate increasingly complex Boolean search queries, we selected the file containing the image of Abraham Lincoln from the data pool, which included images of two presidents, George Washington and Abraham Lincoln. The ‘president’ ssDNA probe, fluorescently labelled with TAMRA, selected both ‘Lincoln’ and ‘Washington’ files from the data pool. The simultaneous addition of the ‘18th century’ ssDNA probe, fluorescently labelled with AFDye 647 (Fig. 5a, left), discriminated ‘Washington’, which contained the ‘18th century’ barcode, from the ‘Lincoln’ file (Fig. 5a, middle, and Supplementary Fig. 15a). The combination of these two ssDNA probes permitted the complex search query ‘president AND (NOT 18th century)’. Sequencing analysis of the gated populations after reverse encapsulation validated that the sorted populations matched search queries for ‘president AND (NOT 18th century)’, ‘president AND 18th century’, and ‘NOT president’ (Fig. 5a, right, and Supplementary Fig. 25).

To demonstrate the feasibility of performing a Boolean search using more than three fluorescence channels for sorting, we also selected the ‘Wolf’ file from the data pool using the query ‘dog AND wild’, and used the ‘black & white’ probe to validate the selected file (Fig. 5b, left). Because conventional FAS software is capable of sorting only using one-dimensional (1D) and two-dimensional (2D) gates, we first selected one out of the three possible 2D plots...
‘dog’–TAMRA. We examined the ‘black-and-white’-TYE705 emission channel on members of the ‘dog AND wild’ subset (Fig. 5b, left and bottom, and Supplementary Fig. 15b). Release of the encapsulated file sequence and subsequent sequencing of each gated population from the dog versus wild 2D plot validated sorting (Fig. 5b, right, and Supplementary Fig. 26).

Our use of plasmids as a substrate for encoding information offered the ability to restore selected files back into the data pool after retrieval. In cases where single images were selected (Figs. 4c and 5a), we were able to transform competent bacteria from each search query that resulted in a single file (Supplementary Fig. 27). Amplified material was pure and ready for re-encapsulation into silica particles, which could be reintroduced directly back into the data pool; thus, this represents a complete write–store–access–read cycle that in principle may be applied to larger-scale datasets, with periodic renewal of ssDNA barcodes and bacterial replication of DNA data following reading17–27. Notably, in a practical implementation of our file system, only the selected files must be restored via amplification, encapsulation and barcoding, because files that are not selected can be returned to the data pool immediately following FAS. While sort probabilities were typically below the optimal 100% targeted for a specific file or file subset query, future work may characterize sources of error that could be due to sample contamination or random FAS errors. The latter type of error may be mitigated through repeated cycles of file selection in series.

**Discussion and outlook**

We introduced a direct random access molecular file system for the retrieval of arbitrary files and file subsets from an archival DNA data store. Our file system overcomes several challenges associated with preexisting PCR-based file systems, including obviating the need for numerous heating and cooling cycles and enzymatic synthesis, and eliminating non-specific crosstalk between file sequences and barcodes17,19,21, while enabling arbitrary Boolean...
logical search queries. Notwithstanding, several technical limitations also exist for our system.

First, while our file system supports theoretical storage densities that are orders of magnitude higher than archival file storage systems based on hard disk, magnetic tape or similar archival media, commonly used today, silica encapsulation of DNA file sequences lowers overall data storage density compared with bare DNA memory (Supplementary Section 6). Second, latency associated with DNA-based barcoding of silica capsules may limit its application to archival data storage and retrieval applications for the foreseeable future. Specifically, advanced liquid handling or microfluidics will be needed to manipulate large numbers of distinct files during the encapsulation and barcoding steps, and latency associated with the encapsulation process itself might need to be reduced from days, as currently implemented, to several hours or less, for example, using encapsulant alternatives such as alginites or synthetic polymers, in order to realize a practical DNA data storage device (Supplementary Section 13). This contrasts with existing file systems, such as PCR and nested file addressing, wherein barcodes are instead written directly with the DNA file sequences using solid-phase phosphoramidite synthesis, for example, using DNA microarrays.

Third, retrieval time is currently limiting for our system in an exabyte-scale data pool because FAS scales linearly with the size of the pool. For example, in our proof-of-concept system, data reconstruction was successful with 100 copies of a file (Fig. 3 and Supplementary Fig. 13), so that our file size of 0.1 kB and search rate of 1,000 files per second yield a final sorting rate of only \( \sim 1 \text{ KB per second} \), however, because our proof-of-concept system did not make use of the vast majority of DNA stored in each capsule, the data stored in each capsule can in theory be substantially increased to yield a search rate of \( \sim 1 \text{ GB per second} \), or \( \sim 15 \text{ days to search through 1 PB of data} \) (Supplementary Section 10). Additional technology development such as custom flow nozzles to accommodate high flow rates close to the limit of a typical FAS instrument (\( \sim 10^4 \) particles per second or \( \sim 10 \text{ GB per second} \)); parallel flow cytometry to achieve search rates of \( \sim 10^6 \) particles per second per device (\( \sim 1 \text{ TB per second} \)); or physical sorting strategies such as direct biochemical pulldown using magnetic extraction, may therefore be required to implement a
practical exabyte-scale or larger memory storage and retrieval device with our system.

Finally, while DNA file protection by silica encapsulation offers millennium-scale protection and storage of the encapsulated file, unprotected DNA barcodes and their covalent linkers are susceptible to hydrolysis and will therefore require either periodic renewal or additional protection, as noted at the outset of this article. As also noted there, similar to PCR-based approaches, barcodes can be protected by dehydrating the data pool or immersing files in hydrophobic liquids. Alternatively, secondary encapsulation with chemistries orthogonal to silica de-encapsulation, such as cleavable synthetic polymers or salts, may be explored.

The latency times associated with file selection and recovery of DNA from files renders our file system ideally suited to archival data storage in which data are written once, stored for decades or longer and accessed rarely. File protection by silica encapsulation offers millennium-scale storage of immutable data, such as astronomical image databases, high-energy physics datasets or high-resolution deep ocean floor mapping. Finally, because our system is not limited to synthetic DNA, it also applies to compact and energy-efficient long-term archival storage of bacterial, human and other genomes for archival sample preservation and retrieval.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41563-021-01021-3.

Received: 20 April 2020; Accepted: 26 April 2021; Published online: 10 June 2021

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Methods
Generating file sequences. Twenty 26 × 26 pixel, black-and-white icon bitmaps were generated as representations of 20 high-resolution colour images (Supplementary Fig. 1) encompassing a broad range of subject matter. Each black-and-white icon was converted to a length-64 bitstring in a column-first order, with each black or white pixel encoded as a 0 or 1, respectively. This bitstring was compressed via run-length encoding, replacing each stretch of consecutive 0s or 1s with a two-tuple (value, length) to generate a list of two-tuples describing the entire bitstring. The maximum length is 15; runs longer than 15 bits are encoded as multiples of the same value. This run-length encoding was converted to a sequence of base-4 digits as follows:

1. Begin with an empty string, and set the current run to the first run of the list.
2. Append the value of the current run (0 or 1).
3. Using two base-4 digits, append the length of the current run.
4. If the length of this run was 15, encode the next run starting with step (2). Otherwise, encode starting at step (3). If no runs remain, the process is complete.

This process produces a quaternary string describing the entire run-length encoding of the image. To avoid homopolymer runs and repeated subsequences in the final nucleotide sequence, each digit is offset by a random number generated from a linear congruential random number generator (LCG) beginning with a random seed (that is, the rth value generated by the LCG is added, modulo 4, to the rth base-4 digit of the quaternary string). We used an LCG of multiplier 5, modulus 2^31-1 and increment 0, although any LCG parameters with a period longer than the length of the sequence would be suitable.

The final value of the quaternary string is converted to nucleotides by a direct mapping (0 = G, 1 = A, 2 = T, 3 = C). The number used to seed the LCG is prepended to this sequence by converting it into a ternary string of length 15 whose digits are encoded in nucleotides via a base transition table, as done previously by Goldman et al.7 (0 = AG, TA, CT). The first digit is encoded directly (0 = AG, TA, CT). The second digit is encoded indirectly as the base-4 digit of the quaternary string. This sequence was modified with additional flanking sequences added to the beginning and end. A 64-bit wavelet hash of each bitmap was calculated using the 'whash' function provided by the ImageHash Python package, available through the Python Package Index (https://pypi.org/project/ImageHash/). The 64-bit hash was split into two 32-bit halves, each of which was represented in a length-24 ternary string that was subsequently converted to nucleotides through the same process as that applied to the seed. The two 24-nucleotide regions were appended to the beginning and end of the sequence (Supplementary Table 1). The sequence containing the image hash, seed and image encoding was additionally flanked on the 5' and 3' ends by sequences (5'-CGTGGTGTCGCCCTCAACAT-3' and 5'-GGTGGAAATGTGGCCGATC-3'), respectively, that allow amplification from a 'master primer' pair that would amplify every sequence in the molecular plasmid database (Supplementary Table 1 and Supplementary Fig. 2).

The final sequence was checked for problematic subsequences, specifically, GGGG, CCCG, AAAAA, TTTTT and the restriction enzyme recognition sites GAATTC and CTGCAG. If any of these subsequences were found outside of expected occurrences in the constant flanking regions, the entire sequence was regenerated with a new random seed until no such subsequences appeared.

The generated sequences were cloned on a pUC19-based vector. The software for sequence encoding and decoding is publicly available on Github at https://github.com/lcb/pDNA-Memory-Blocks (ref.7), and the plasmids are publicly available from Addgene (https://www.addgene.org/dnaencode/articles/). Each master primer and the hash barcode pairs were verified by PCR and agarose gel analysis, and the PCR bias was checked by qPCR (Supplementary Fig. 3).

Generated DNA sequences were ordered as custom genes synthesized, cloned, sequenced and validated by Integrated DNA Technologies (IDT) into a pUC19-based vector. Each plasmid was designed with flanking single EcoRI and PstI cut sites, sequences that had been designed against in the inserts, enabling future subcloning to alternative vectors using standard digestion and ligation. All plasmids were amplified in bacterial cultures and purified using Qiagen (Venlo) HiSpeed Midi or Maxi kits following the protocol provided by the manufacturer. Briefly, DH5α cells were made chemically competent by growing 100 ml of cells to reach 0.5–1.0 optical density at 600 nm. The cells were gently pelleted and incubated in ice on 100 mM CaCl2, for 30 minutes. The pelleted cells were then brought up in 100 mM CaCl2, with 10% glycerol. Transformations from each plasmid were accomplished by incubation of 20 µl chemically competent cells with 1 ng of plasmid DNA for 30 minutes on ice, followed by heating to 42°C for 45 seconds, placing on ice for 2 minutes and then addition of 1 ml of prewarmed S.O.C. Medium (Thermo Fisher; catalogue number, 15544034). The mixture was shaken for 1 hour at 250 r.p.m. at 37°C. A volume of 20 µl of transformed cells was streaked to single colonies on agar plates that were supplemented with 100 µg/ml ampicillin. Single colonies were chosen for growth in 100 ml lysogeny broth (LB) with 100 µg/ml ampicillin (Millipore Sigma; catalogue number, A5354) overnight in a 1.5 ml microcentrifuge tube and then followed by the cell culture. Pelleted cells were lysed and treated with Qiagen purification buffers. Plasmid DNA was then purified on the provided silica matrix support and eluted in 1 ml of TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0; Tris-HCl, tris(hydroxymethyl) aminomethane hydrochloride; EDTA, ethylenediaminetetraacetic acid). Plasmid preparations were verified by PCR, Sanger sequencing and short-read sequencing using Illumina MiniSeq for quality control (Supplementary Fig. 3).

DNA encapsulation. A volume of 1.0 ml of 50 mmol/l-1 fluorescent-core 5µm silica particles (Creative Diagnostics; catalogue number, DNG-L034) was added into a 2.0 ml DNA/RNA LoBind Eppendorf tube. The particles were centrifuged at 1,000 r.p.m. for 10 seconds using a benchtop centrifuge. The particles were redispersed in 1.0 ml anhydrous ethanol with vigorous vortexing. The particles were centrifuged and redispersed five times. We then added 50 µl of 50% TMAPS in methanol (Alfa Aesar; catalogue number, H66414) to the dispersed 5µm silica particles (50 mg/ml −1 in ethanol). The mixture was stirred overnight at room temperature using a thermal mixer (Thermo Fisher; catalogue number, 13687711) at 1,200 r.p.m. The mixture was centrifuged at 1,000 r.p.m. and washed with ethanol five times to remove any unreacted TMAPS. The functionalized particles were finally redispersed in 1.0 ml DNAse/RNase-free water (Thermo Fisher; catalogue number, 10977015). The particles were stored at room temperature until further use.

For each data-encoding plasmid, a mass of 1.0 mg of TMAPS-functionalized, fluorescent 5µm particles was added into a 2.0 ml LoBind Eppendorf tube containing 15 µg of plasmid DNA dissolved in 1 ml of water. The mixture was mixed gently using a tube mixer (Thermo Fisher; catalogue number, 88881001) at 30 r.p.m. and at room temperature for 5 minutes. A volume of 10 µl of 50% TMAPS in methanol was then added to the mixture and stirred for 10 minutes at 1,000 r.p.m. and 25°C using a thermal mixer. After 10 minutes, a volume of 2 µl of aminomethylsilane (TEOS; Millipore Sigma; catalogue number, 1333859) was added, and the mixture was stirred for 24 hours at 1,000 r.p.m. and 25°C using a thermal mixer (Thermo Fisher). An additional 5 µl of TEOS was then added, and the mixture was stirred for 4 days at 1,000 r.p.m. and 25°C, which formed the DNA capsules. The mixture was centrifuged at 2,000 r.p.m. for 3 minutes to sediment the DNA capsules, and then the supernatant was removed without disturbing the sediment pellets. The DNA capsules were washed repeatedly five times by redispersing the particles with 1 ml of water, sedimenting the particles with a centrifuge at 2,000g for 3 minutes and removing the supernatant. After the final wash, the DNA capsules were redispersed in 1 ml of ethanol with 30 seconds of vortex mixing. A volume of 20 µl of 2-aminomethylpropiolysilane (Millipore Sigma; catalogue number, 4401409) was then added, and the mixture was stirred for 18 hours at 1,000 r.p.m. and 25°C using a thermal mixer. The mixture was centrifuged at 2,000g for 3 minutes to sediment the amino-modified DNA capsules, and then the supernatant was removed without disturbing the particle sediment pellet. The particles were washed repeatedly five times by redispersing the particles with 1 ml of N-methyl-2-pyrrolidone, sedimenting the particles with a centrifuge at 2,000g for 3 minutes and removing the supernatant. After the final wash, the DNA capsules were redispersed in 1 ml of N-methyl-2-pyrrolidone with 30 seconds of vortex mixing, and the resulting colloidal suspension was then transferred into a clean 2 ml Eppendorf LoBind tube.

Barcoding DNA capsules. Each encoded image was annotated with three semantic metadata descriptors (Supplementary Table 2) associated with the original image (Supplementary Fig. 1). A table was then generated to associate each descriptor with a unique sequence chosen from a list of 240,000 orthogonal barcode sequences7. Hexylamine-modified ssDNA oligonucleotides that represented each descriptor were purchased from IDT and dissolved in nuclease-free water with a final concentration of 100 µM.

Using all the DNA capsules from the previous step, a mass of 5 mg of 2-azido acetic acid N-hydroxysuccinimide ester (Click Chemistry Tools; catalogue number, 1070) and 5 µl N,N-diisopropylaminostyryl (Millipore Sigma; catalogue number, D125806) were added, and the mixture was stirred for 2 hours at 1,000 r.p.m. and 25°C using a thermal mixer. The azido-modified DNA capsules were washed repeatedly five times by redispersing the particles with 1 ml of N-methyl-2-pyrrolidone, and the mixture was stirred for 30 minutes at 1,000 r.p.m. and 25°C using a thermal mixer. The mixture was centrifuged at 2,000g for 3 minutes to sediment the amino-modified DNA capsules, and the supernatant was removed without disturbing the particle sediment pellet. The particles were washed repeatedly five times by redispersing the particles with 1 ml of N-methyl-2-pyrrolidone, sedimenting the particles with a centrifuge at 2,000g for 3 minutes and removing the supernatant. After the final wash, the DNA capsules were redispersed in 1 ml of N-methyl-2-pyrrolidone with 30 seconds of vortex mixing, and the resulting colloidal suspension was then transferred into a clean 2 ml Eppendorf LoBind tube.
particles with a centrifuge at 2,000g for 3 minutes and removing the supernatant. After the final wash, the particles were resuspended in 50 µl of saline TAE buffer with 30 seconds of vortex mixing and 1 minute sonication. All the files were then pooled together, forming the file pool or molecular file database with an estimated final concentration of 2.0 mg ml⁻¹ in 10.0 ml of saline TAE buffer.

Silica particle characterization. Surface zeta potentials were measured using a Malvern Zetasizer Nano ZSP. All samples for surface zeta potentials were prepared and measured in a standard fluorescence quartz cuvette (catalogue number, 3-Q-10) from StaRna Cells at a concentration of 0.1 mg ml⁻¹ with a volume of 700 µl. A universal ‘dp’ probe (Malvern Panalytical; catalogue number, ZEN10002) was used to measure the zeta potential of particles. Scanning electron microscopy of the particles was performed using a Zeiss Gemini 2 field emission scanning electron microscope. Samples were mounted on silicon substrates for non-DNA-modified particles and glass for DNA-modified particles.

Querying molecular file database using fluorescently labelled probes. The molecular file database was vortexed for 10 seconds, sonicated for two minutes and revortexed for another 10 seconds to redisperse the settled particles. For all sorting experiments except Fig. 3, a volume of 100 µl and revortexed for another 10 seconds to redisperse the settled particles. For all sorting experiments except Fig. 3, a volume of 100 µl of the molecular database (2 mg ml⁻¹) was added into a 1.5 ml Eppendorf LoBind tube. Dye-labelled probes (Supplementary Section 7) for querying the molecular file database were added such that the final concentration of the DNA-dye ssDNA in solution was 5 µM. The resulting mixtures were incubated at 70°C for 1.200 r.p.m. using a thermal mixer for 5 minutes. The mixtures were then cooled to 20°C at 1,200 r.p.m. using a thermal mixer over 20 minutes and then centrifuged at 10,000g for 1 minute. The supernatant was discarded, and the pelletified particles were washed with 500 µl of saline TAE buffer. The sedimentation and washing process was repeated five additional times to remove non-specifically bound dye-DNA. The particles were finally resuspended in 500 µl of saline TAE buffer.

For the sorting experiments performed in Fig. 3 and Supplementary Fig. 13, the workflow for querying the molecular file database remained the same, except for the composition of the molecular file database:

| Ratio of ‘Airplane’/19 other files | Volume of ‘Airplane’ stock solution (µl) | Volume of stock solution of 19 other files (µl) |
|----------------------------------|----------------------------------------|-----------------------------------------------|
| 1:1                              | 500µl                                  | 0                                             |
| 1:10²                            | 5µl                                    | 495                                           |
| 1:10⁴                            | 5µl                                    | 495                                           |
| 1:10⁶                            | 5µl                                    | 495                                           |

*C:Concentration of stock solution is 2 mg ml⁻¹.

*C:Concentration of stock solution is 2×10⁻⁶ mg/ml⁻¹ obtained from serial dilution.

*C:Concentration of stock solution is 2×10⁻⁶ mg/ml⁻¹ obtained from serial dilution.

FAS. All FAS experiments were performed on a BD FACSARia III flow cytometer. Samples were filtered through a Corning 70 µm cell strainer (Fisher Scientific; catalogue number, 07-201-431) prior to particle sorts. Samples were flowed into the instrument with phosphate-buffered saline (PBS) as the sheath fluid at a flow rate that maintains an events detection rate of 1,200 events per second and below. We found that performing sorting at a flow rate that exceeds this events rate clogged the FAS instrument intermittently. All sorts were accomplished with a standard 70 µm nozzle. The sample was held at room temperature and agitated periodically every 5 minutes by pausing the sort and vortexing the sample vigorously with a vortex mixer. The internal agitator in the flow cytometer with a 300 r.p.m. agitator speed was not sufficient to prevent the silica particles from sedimenting over time, and we found that periodically agitating the sample tube every 5 minutes with a vortex mixer was more effective. Since all files must contain a fluorescent core, all particles were gated by default using the ‘FITC’ laser and detector settings, which is defined by gating the majority population in the ‘FITC-A’ channel histogram, in addition to standard forward scatter (FSC) and side scatter (SSC) gates, to minimize the sorting of doublets (Supplementary Fig. 11). All FAS experiments were performed at room temperature and at a 500 µl constant volume, given that sorting rates, detection rates and flow rates can vary across different flow cytometers.

Release of DNA from sorted files. Sorted populations were centrifuged at 10,000g for 1 minute. The supernatant was carefully removed with a pipette to avoid disturbing the silica pellets. A volume of 45 µl of electronics-grade 5:1 buffered oxide etch (VWR; catalogue number, JT5192-3) was then added. The mixture was vortexed for 5 seconds to resuspend the pellet, and the mixture was statically incubated at room temperature for 5 minutes. A volume of 5 µl of 1 M phosphate buffer (0.75 M NaH₂PO₄, 0.25 M Na₂HPO₄, pH 7.5 at 0.1 M) was then added, vortexed for 1 second and desalted twice with an Illustra MicroSpin S-200 HR column (Millipore Sigma; GE27-5120-01).

Sequence of retrieved files. For Illumina MiniSeq and MiSeq sequencing, the master primer pair with 5’ extensions matching Illumina Nextera sequencing adapters was used to amplify all plasmids simultaneously (Supplementary Fig. 9).

Template amounts were adjusted based on concentrations determined with Qubit fluorescence assay (Thermo Fisher) or qPCR. If required, the amplification was simultaneously followed by qPCR, and enough cycles were used to rise above the cycle threshold, or alternatively obtain a final concentration of 2 ng ml⁻¹. Dual sequence indices were then added to the adaptor-modified inserts at the 5’ and 3’ ends, associating the sequencing lane with a particular selection experiment, which was followed by solid-phase reversible immobilization bead cleanup (Beckmann Coulter). A 25 µl PCR mixture was added to the material stabilized using KAPA HiFi polymerase (Roche) with 1 ng of template and 1 µM forward and reverse primers. After amplification, the PCR mixture was combined with 20 µl of SPlRSelect beads (Beckman Coulter), mixed and left to stand for 5 min. The mix was then separated by magnetic plates and washed twice with 150 µl 80% ethanol, dried for 2 min and eluted in 20 µl Qiagen TE buffer. Samples were quantified using the Qubit fluorescence assay with the provided high-sensitivity buffer and standards. A sequencing pool was generated to approximately equimolar amount per index pair. Illumina MiniSeq with 150 x 150 read lengths was used to read out the start and end of each sequence. Sequences were demultiplexed, and sequence clustering was used to count the number of occurrences of each image (Supplementary Section 9). The sort probability for a file into a particular fraction was calculated as the count associated with that file divided by the sum of the counts for that file over all fractions generated from an initial sample. This metric is also referred to as ‘enrichment’ throughout the text. In Figs. 3–5 in the main text, the enrichment of each file is indicated by the percent opacity of the images displayed on the grid.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Gene sequences and plasmid maps are available from AddGene (https://www.addgene.org/browse/article/28120796/). Insert sequences and barcoding sequences are given in Supplementary Tables 1 and 2. All the data files used to generate the plots in this manuscript are available from M.B. upon request.

Code availability

Software for sequence encoding and decoding is publicly available on GitHub (https://github.com/lcbib/NA-DNA-Memory-Blocks/).

References

51. Banal, J. L. et al., DNA-Memory-Blocks v.2.0 https://doi.org/10.5281/zenodo.4586900 (Zenegro, 2021).

Acknowledgements

We gratefully acknowledge discussions with C. Leiserson and T. B. Schardl on the scalability and generalizability of our barcoding approach. We thank G. Paradis, M. Jennings and M. Griffin of the Flow Cytometry Core at the Koch Institute at the Massachusetts Institute of Technology (MIT) and P. Rogers of the Flow Cytometry Facility at the Broad Institute of Harvard and MIT for assistance and discussions in developing the flow cytometry workflow. We also thank D. Mankus of the Nanotechnology Materials Core Facility at the Koch Institute at MIT for assistance in the imaging of the particles using the scanning electron microscope and A. Leshinsky of the Biopolymer and Proteomics Core at the Koch Institute at MIT for assistance in mass spectrometry characterization. M.B., J.L.B., T.R.S. and J.B. gratefully acknowledge funding from the Office of Naval Research (N00014-17-1-2609, N00014-16-1-2506, N00014-12-1-0621 and N00014-18-1-2290) and the National Science Foundation (CCF-1564025, CCF-1956054, HDR OAC-1940231 and CBET-1729397). Research was sponsored by the US Army Research Office and accomplished under cooperative agreement W911NF-19-2-0026 for the Institute for Collaborative Biotechnologies. Additional funding to J.B. was provided through a National Science Foundation Graduate Research Fellowship (grant no. 1122374). P.C.B. was supported by a Career Award at the Scientific Interface from the Burroughs Wellcome Fund. C.M.A. was supported by National Institutes of Health grant F32CA236425.

Author contributions

J.L.B., T.R.S. and M.B. designed the file labelling and selection scheme. J.L.B., T.R.S. and C.M.A. implemented the file selection scheme using FAS. J.J.B. and T.R.S. developed the encoding scheme and metadata tagging of the images to DNA. T.R.S. designed the plasmid for encoding imaging. H.H. and T.R.S. performed the cloning, transformation and purification of the plasmids. J.L.B. synthesized and purified all the TAMRA- and AF647-labeled DNA oligonucleotides. J.B. characterized the particles. J.L.B. developed the synthetic route to attach DNA barcodes on the surface of the particles. J.L.B. performed the encapsulation, barcoding, sorting, reverse encapsulation of the particles after sorting and desalting. T.R.S., H.H. and M.R. performed the sequencing. J.B. performed the computational validation of the orthogonality of the barcode sequences, and J.L.B. performed the experimental validation of the orthogonality of barcode and probe sequences. J.B. developed the computational workflow to analyse
the sequencing data, including statistical analyses. M.B. conceived the file system and supervised the entire project. P.C.B. supervised the FAS selection and supervised the sequencing workflow. All authors analysed the data and equally contributed to the writing of the manuscript.

Competing interests
The Massachusetts Institute of Technology has filed patents covering the encapsulation-based file system (US application number 16/097594) and microfluidics-based storage, access and retrieval of biopolymers using the same file system (US application number 16/012583) on behalf of the inventors (J.L.B., T.R.S., J.B. and M.B.) M.B. is the founder of Cache DNA and is a member of its Scientific Advisory Board. P.C.B. is a member of the Scientific Advisory Board of Cache DNA. H.H., M.R. and C.M.A. declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41563-021-01021-3.

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Peer review information Nature Materials thanks Reinhard Heckel, William L. Hughes and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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Software and code

Policy information about availability of computer code

**Data collection**

All fluorescence-activated sorting (FAS) experiments were performed on a BD FACSARia III flow cytometer that are running BDFACS Diva software. Three methods were used for verification of retrieval of DNA, which included quantitative PCR (qPCR), short-read sequencing, and Sanger sequencing. We performed qPCR experiments on a Thermo-Fisher QuantStudio 6 Flex. We used Illumina MiSeq and Miniseq for short-read sequencing. Finally, we sent samples to GeneWiz for Sanger sequencing.

**Data analysis**

FAS data were analyzed using Flowjo (version 10) and MATLAB (R2019b). We analyzed qPCR data using the proprietary software that was installed with the instrument (QuantStudio Software v1.3). Sequencing data were analyzed using custom python scripts which are publicly available on Github [https://github.com/lcb/0DNA-Memory-Blocks/](https://github.com/lcb/0DNA-Memory-Blocks/).

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Life sciences study design

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| Sample size | We arbitrarily chose to encode 20 different images in 20 plasmids to demonstrate our file system. We were able to retrieve, i.e., sort and sequence, each file individually, as well as several collections of file described by various Boolean queries. |
|-------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Data exclusions | Sequencing reads that do not align with any of the file sequences were discarded.                                                                                                                 |
| Replication  | All sorting experiments were done at least in triplicates and were successful in all cases. Each replicate was performed on a molecular database that was synthesized on a different day, i.e., each file sequence was re-encapsulated, re-tagged, and pooled on different days. |
| Randomization | No randomization required for this work as it was not intended to find differences among samples. Rather, the work's main focus was on demonstrating the file system's operational characteristics through retrieval of specific samples using different probe combinations and gating strategies using fluorescence-activated sorting. |
| Blinding     | The decoding step was performed without a priori information on the sorted files.                                                                                                                  |

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| n/a | n/a |
| ☑ Antibodies                     | ☑ Involved in the study |
| ☑ Eukaryotic cell lines          | ☐ CHIP-seq |
| ☑ Palaeontology                  | ☐ Flow cytometry |
| ☑ Animals and other organisms    | ☐ MRI-based neuroimaging |
| ☑ Human research participants    |         |
| ☑ Clinical data                  |         |

Flow Cytometry

Plots

- The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
- The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
- All plots are contour plots with outliers or pseudocolor plots.
- A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation

The molecular file database was vortexed for 10 seconds, sonicated for two minutes, and re-vortexed for another 10 seconds to re-disperse the settled particles. A volume of 100-μL of the molecular database (2 mg mL⁻¹) is added into a 1.5 mL Eppendorf LoBind tube. Dye-labelled probes for querying the molecular file database were added such that the final concentration of the DNA-dye single-stranded DNA in solution is 5 μM. The resulting mixtures were mixed at 70 °C at 1,200 rpm using a thermal mixer for 5 minutes. The mixtures were then cooled to 20 °C at 1,200 rpm using a thermal mixer over 20 minutes and then centrifuged at 10,000 g for 1 minute. The supernatant was discarded, and the pelleted particles were washed with 500 μL of 40 mM Tris, 20 mM acetic acid, 2 mM EDTA, 1.0% Tween-20, 1.0% sodium dodecyl sulfate, 500 mM NaCl. The sedimentation and washing process was repeated for five additional times to remove non-specifically bound dye-DNA. The particles are finally re-suspended in 500 μL of 40 mM Tris, 20 mM acetic acid, 2 mM EDTA, 1.0% Tween-20, 1.0% sodium dodecyl sulfate, 500 mM NaCl. All fluorescence-
activated sorting (FAS) experiments were performed on a BD FACSAria III flow cytometer. Samples were filtered through a Corning® 70-μm cell strainer (Fisher Scientific) prior to particle sorts. Samples are flowed into the instrument with 1x PBS as sheath fluid at a flow rate that maintains an events detection rate of 1,200 events per second and below. We found that performing sorting at a flow rate that exceeds this events rate clogged the FAS instrument intermittently. All sorts were accomplished with a standard 70 μm nozzle. The sample was held at room-temperature and agitated periodically every 5 minutes by stopping the sort and vortexing the sample vigorously with a vortex mixer. We note that the internal agitator in the flow cytometer with a 300 rpm agitation speed was not sufficient to prevent the silica particles from sedimenting over time and we found that periodically agitating the sample tube every 5 minutes with a vortex mixer was more effective.

| Instrument          | BD FACSAria III flow cytometer |
|---------------------|--------------------------------|
| Software            | BDFACS Diva                    |
| Cell population abundance | At least 100,000 particles were sorted every run. |
| Gating strategy     | Since all files must contain a fluorescein core, all particles were gated by default using the ‘FITC’ laser and detector settings, which is defined by gating the majority population in the ‘FITC-A’ channel histogram, in addition to standard FSC and SSC gates to minimize sorting of doublets. We also ran positive controls for every dye before every FAS experiment to validate that there is no significant spectral crosstalk during the sorting process and to validate that we have a distinguishable fluorescence signal in the presence of other fluorescent dyes. For example, because all our files have FITC, we validated that there is no fluorescence spillover of FITC in the TAMRA (PE-Texas Red channel), At647 (APC channel), or YTE705 (Alexa Fluor 700 channel) that would otherwise make it difficult to distinguish the different particle populations. Boundaries of positive and negative in single-color experiments were determined by measuring the intensity populations of positive control and negative controls (silica particles only) for each dye channel and then creating gates on their relative intensity distribution. |

☑ Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.