Demonstrated pulse accumulation effect is atoms. Another interesting application of the comb directly by an optical transition in cold sequence of these results is a method to control coherent pulse accumulation may prove the added spectral resolution due to multipulse interference. The precise and phase-coherent pulse accumulation may prove particularly useful in efficiently populating atomic Rydberg states for quantum information processing. Although the current experiment involves two-photon transitions, the advantages of DFCS should apply equally to single-photon and multiphoton excitations. Multiple ultrafast lasers with optical spectra independently tailored for different spectroscopic features could be phase coherently stitched together (27, 28) to further extend the utility of this approach.

References and Notes
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Building Programmable Jigsaw Puzzles with RNA

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One challenge in supramolecular chemistry is the design of versatile, self-assembling building blocks to attain total control of arrangement of matter at a molecular level. We have achieved reliable prediction and design of the three-dimensional structure of artificial RNA building blocks to generate molecular jigsaw puzzle units called tectosquares. They can be programmed with control over their geometry, topology, directionality, and addressability to algorithmically self-assemble into a variety of complex nanoscopic fabrics with predefined periodic and aperiodic patterns and finite dimensions. This work emphasizes the modular and hierarchical characteristics of RNA by showing that small RNA structural motifs can code the precise topology of large molecular architectures. It demonstrates that fully addressable materials based on RNA can be synthesized and provides insights into self-assembly processes involving large populations of RNA molecules.

DNA has been extensively used to generate artificial geometrical objects like polyhedra (1–3), various self-assembling two-dimensional (2D) nanostructures (1, 4–6), and DNA nanomechanical devices (7–9). Seeman, Winfree, and collaborators (1, 4, 5) have shown that DNA tiles based on various “crossover” DNA motifs could assemble in a predictable manner into periodic and aperiodic patterned 2D arrays. These DNA arrays are still made of a limited number of distinct molecular tiles and display rather simple patterning with no finite dimensions. However, their work suggests that versatile programmable molecular systems capable of algorithmic assembly into an infinite variety of 2D or three-dimensional (3D) supra-architectures with increasing pattern complexity, shape, molecular diversity, and size could potentially be generated with nucleic acids (10).

Although more chemically labile than DNA, natural RNAs offer a richer treasure trove of rigid structural motifs (11–14) that can be potential modules for supramolecular engineering (15–20). RNA tectonics (15) refers to the modular character of RNA, which can be decomposed and reassembled to create new RNA nanoscopic architectures. With the idea in mind to generate addressable materials with increasing patterns of complexity and molecular diversity, we have used a sequential stepwise assembly strategy to construct programmable building blocks with RNA tectonics. These molecules behave as “smart” RNA pieces, which could ultimately self-assemble in a predictable manner into any possible 2D architecture with full control over size, shape, and pattern geometry. Thus, the final position of each molecule can eventually be known and, therefore, be addressable, within a molecular jigsaw puzzle of finite size.

At a molecular level, “square-shaped” RNA supramolecules with sticky, interacting tails can potentially be programmed to assemble into many different planar networks of predefined geometries. We chose
two small RNA structural motifs present in the ribosome crystallographic structures (12–14) to guide our design of a self-assembling square made of four similar but nonidentical subunits, called tectoRNAs (Fig. 1, A and B) (21). Each tectoRNA contains two interacting hairpin loops (19, 22) covalently joined by a small structural motif of 11 nucleotides, called the right angle (RA) motif, that specifies 90° angle corners between adjacent helices within the context of the ribosome (12, 13). To avoid homomultimers, the formation of a closed, circular tetramer is directed by four distinct, specific noncovalent loop-loop interactions, called kissing loop (KL) complexes (22), which are expected to adopt collinear extended helical structures according to the crystallographic structures of the ribosome (12) and the dimerization initiation site of human immunodeficiency virus (HIV) RNA (22). A tectosquare 3D model resembles a square when viewed from the top. Nevertheless, it is not flat, because its extended helical sides adopt a log cabin–like conformation at the level of RA corners (Fig. 1C; fig. S1). Rather than being a perfect four-fold pseudosymmetry (C 4), the tetramer has two-than being a perfect four-fold pseudosymmetry (C 4), the tetramer has two- than being a perfect four-fold pseudosymmetry (C 2). Small (ST) and large (LT) tectosquares, with 10-nm and 13-nm side lengths, can be constructed from tectoRNAs with hairpin stems of 9 and 15 base pairs (bp), respectively.

Tectosquares can further self-assemble through specific sticky tail connectors (Fig. 1). The tails of the tectoRNAs can be designed to have a wide variety of sequences. Their precise positioning and orientation are inferred from the RA motif geometry. The 3′ tail, stacked in continuity to the 3′ stem-loop, is expected to be structurally more constrained and directional than the 5′ tail. By swapping the RA motif, the orientation of the 3′ tail can be modified by 90° without changing the overall positioning of the stem-loop arms (Fig. 1D). Moreover, small variations in the tail length can change the overall length of tail connectors by one-half of a helical turn, which positions two adjacent tectosquares in either a cis or trans configuration (Fig. 1E). About 88.5 million distinct tectosquares can be built with a limited set of 12 tail connectors with two different tail orientations and sizes (23).

We constructed two sets of tectoRNAs for building ST and LT tectosquares (21).

Each tectoRNA sequence was optimized to favor folding into a unique, stable secondary structure (17, 24). After being synthesized by run-off transcription, tectosquare modular assembly was monitored by native polyacrylamide gel electrophoresis (PAGE). Magnesium is absolutely required for assembly. At 0.2 mM Mg(OAc) 2, an equimolar mixture of each tectoRNA set forms 60 to 90% of a circular supramolecular species that migrates slower than monomers and linear tetramers lacking one of the four KL motifs (Fig. 2A). Remarkably, both tectosquares can be purified out of native PAGE...
gels without dissociating (Fig. 2). We investigated further tectosquare stability by temperature-gradient gel electrophoresis (Fig. S2), a method for separating different assemblies on the basis of temperature-dependent conformational change (25). In 15 mM Mg$^{2+}$, tectosquares are stable up to 56°C because of the presence of the two structural motifs encoded within their sequence. With equilibrium constants of dissociation ($K_d$) ranging from 1 to 20 nM at 0.2 mM Mg$^{2+}$, KL motifs are more stable than RNA duplexes of identical sequences (19, 26). Moreover, tectosquares that contain the RA motif in each of their units are 4.2°C (ST) to 5.8°C (LT) more stable than those without any RA motif (Fig. S2). Accordingly, the thermal stability of a tectosquare increases with an increasing number of RA motifs present within its assembly.

The overall topology of an LT was investigated by atomic force microscopy (AFM) (21, 27) in air or in 15 mM Mg(OAc)$_2$ solution after deposition of the RNA sample solution on a mica surface. The predicted and observed tectosquare architectures (Fig. 2, B and C) are in remarkable agreement with each other and unambiguously establish the RA and KL motifs as autonomous folding modules outside their natural context. The LT folds into a square shape with stiff, straight sides of $13 \pm 3$ nm and corners between 70° and 110°. The central cavity has an overall size of $8 \pm 3$ nm, and the width of the RNA helical region is estimated to be $3 \pm 1$ nm at half height, the expected width for a double-stranded RNA helix. The seldom-observed rhombus shape for LT is indicative of a tilted rather than flat tetramer that could be deformed when forced to lie on the mica surface.

The stable, rigid characteristics of tectosquares are particularly attractive for building programmable planar supra-architectures. To demonstrate the predicted geometrical properties of tectosquares described above (Fig. 1), we designed 12 specific 3’ tail connectors of 6 bp and used them to program the assembly of nine different sets of tectosquares into distinctive periodic fabrics (Fig. 3A). The connectors have similar free energies of formation, chosen to be less stable by at least two orders of magnitude than those of KL complexes. At 15 mM Mg$^{2+}$, two tectosquares joined by two parallel tail connectors disassemble around 30°C; that is, 25°C below tectosquare melting temperature. A simple monitoring of the RNA annealing temperature can thus be used to hierarchically control the assembly process by uncoupling tectosquare association from fabric formation. Forty-nine tectoRNAs with different sizes, tail sequences, tail lengths, and orientations were synthesized and combined to separately construct a total of 22 tectosquares that were then appropriately mixed to generate the nine tectosquare patterns. Pattern formation was performed at 15 mM Mg$^{2+}$, on the mica surface, by slow cooling from 50° to 4°C and was monitored by AFM under

![Diagram and AFM images of tectosquare nanopatterns generated from 22 tectosquares.](Fig. 3. Diagram and AFM images of tectosquare nanopatterns generated from 22 tectosquares. (A) One micrometer square scale AFM images obtained in solution for: LT1-2, ladder pattern; LT3-4, fish net pattern; LT5-6, diamond pattern; ST3-4, striped velvet pattern; ST3-LT4, basket weave pattern; LT7-8-9, lace pattern; LT10-11-12-13-14, polka dot pattern; LT7-8-15-16, tartan pattern; LT17-18-19-20, cross pattern. Scale bars, 500 nm. (B) Magnification of patterns in (A). Scale bar, 20 nm. (See also fig. S3.)}
aqueous conditions, similar to those used for native PAGE.

We first designed two distinct pairs of large tectosquares, LT1-2 and LT3-4, characterized by the same four tail-connector sequences oriented parallel to “a-d” and “b-c” (LT type I, Fig. 1F), but with tails of 12 and 10 nucleotides, respectively. LT1-2, with connectors of length equivalent to 14 bp, adopts a cis configuration (Fig. 1E) that leads to ladders of 10 to 20 LT (Fig. 3 and fig. S3). The next set, LT5-6, presents a combination of two of the previous pairs of long and short connectors. This molecular arrangement leads to the formation of stable tectosquare dimers as- 

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This molecular arrangement leads to the formation of stable tectosquare dimers assem- 

bly in diamond-like arrays that are as large as LT3-4 arrays but with meshes twice as big (14 ± 2 nm by 27 ± 3 nm) (Fig. 3). STs efficiently assemble into predetermined architectures as well. The ST3-4 pair, with the same connectors as LT3-4, forms large arrays that can span almost 1 μm and can involve more than 500 tectosquares. The plain central cavity of ST3 and ST4, too small to be well resolved by AFM, gives a striped velvet texture to these assemblies (Fig. 3; fig. S3). STs and LTs can also be combined when in the trans configuration. For example, the mix of ST3 and LT4 forms basket weave patterns characteristic of the LT and ST alternating arrangement (Fig. 3).

More complex patterns can be obtained from a greater number of tectosquares by using additional tail connectors or by preventing association at specific positions within the 2D lattice. Both LT7-8-9 and LT10-11-12-13-14 sets take advantage of six connectors to form the lace and polka dot patterns, respectively (Fig. 3 and fig. S3). For LT10-11-12-13-14, the regularly spaced dotted motif is programmed with five LTs assembling symmetrically in an all trans configuration network, two of them lacking one of their tails.

Tectosquare assembly is highly dependent on the directionality of the 3′ tail (Fig. 1F). We observed significantly larger architectures with type I LT networks than with type II (fig. S4, A and B) (28). Nevertheless, patterns taking advantage of eight connectors with various 3′ tail orientations can be obtained. The tartan pattern LT7-8-15-16 is derived from the lace pattern LT7-8-9 (Fig. 3A), by replacement of LT9 with LT15-16, a tectosquare dimer formed by association of type III and type IV LTs (Fig. 1F). The different directionality of LT15 and LT16 tails leads to the formation of a lattice with meshes of 28 ± 3 nm, twice as big as those obtained with LT5-6 and LT7-8-9 (Fig. 3B). LT17-18-19-20, a set of four type V tec- 

tosquares with 3′ tails pointing in the four cardinal directions, assembles into the cross pattern corresponding to a C4 pseudosymmetrical arrangement of LTs (Fig. 3A). This regular lattice has two distinct square-shaped mesh sizes of 12 ± 2 nm and 17 ± 2 nm that correspond to the LT structure and the central hole formed by association of four LTs, respectively (Fig. 3B).

As an initial step toward fully addressable self-assembling materials, we designed three sets of tectosquares to assemble specifically into finite aperiodic nanogrids (Fig. 4; fig. S4). To control the size and shape of the RNA assembly, some LTs lacking a 3′ tail at specific corners are programmed to act as edges. The 2 by 2 grid is a cross of 45 nm formed of four type V LTs linked by four different connectors. The 3 by 3 and 4 by 4 grids are symmetrical, modular arrangements of five and eight different type I LTs, respectively. In the 3 by 3 grid, four edge LTs are linked to a central LT by six different connectors. In the 4 by 4 grid, six edge LTs assemble around a central 2 by 2 cross of two LTs through 12 different connectors. According to our structural models, the 2 by 2 grid is not perfectly flat. This partially explains its infrequent observation on the mica surface and the rhombus, rather than square, shape adopted by the 2 by 2 LT during the AFM imaging process (Fig. 4). By contrast, more than a hundred diamond-shaped 3 by 3 grids are identified on a 16-μm² surface, in perfect agreement with the flat planar arrangement expected for type I LTs (Fig. 4; fig. S4, C and D). The 4 by 4 grid demonstrates that the assembly of up to 27 different tectorRNAs can be hierarchically and reproducibly controlled to form RNA nano- 

scale jigsaw puzzles, which suggests that aperiodic assemblies of even greater molecular diversity can be obtained with additional connectors (29).

We have demonstrated that two rRNA structural motifs participate in a predictable manner to stabilize, position, and pack RNA helices without the need of proteins. The length and geometry, rather than the sequence of loops, predispose the formation of linear coaxial stacks of helices in KL complexes. Similarly, it is the bent geometry of the RA motif that favors stacking of the 3′ end tail connector in continuity with its 3′ stem. The importance of base stacking is emphasized by the inability of unstacked dangling 5′ tail connectors to form any organized networks. Thus, in both the ribosome and tectosquares, RA and KL motifs are likely to assist the assembly process not only by local contributions to a specific RNA fold but also by reducing the entropy cost.

The subtle interplay of enthalpy and entropy that successfully promotes the forma- 

tion of tectosquare assemblies is highly dependent on the strength, length, and orientation of the tail connectors and the environmental cues (RNA and divalent ion concentrations, temperature, and assembly protocols). For instance, thermodynamically stronger tail connectors of the same order of magnitude as KL complexes, as well as a reduced initial temperature of assembly, can negatively contribute to ordered assembly by kinetically trapping tectosquares in wrong configurations. Moreover, variation of the magnesium concentration can be used to switch on and off tectosquare assembly. Understanding phase diagrams for assembly is thus important for finding annealing conditions to self-heal irregular lattice points.

Tectosquares assemble into their respective lattices at nanomolar concentrations, where-
as $K_v$ values measured for single 3’ tail connectors are in the micromolar range. This fact and the small number of overlapping RNA lattices observed by AFM suggest that epitaxial phenomena occurring between RNAs and magnesium ions adsorbed on the negative mica surface might promote assembly. The observed RNA networks grow in a radial fashion. All LT tectosquare arrays involve a similar number of molecules, indicating that LT assembly is independent of the nature of the pattern formed (Fig. 3). However, STs generate significantly larger $d$ values measured for single 3’ tail connectors, or a 3’ tail of $n$ different sequences with different size $l$ and orientation $o$, a total of $4(l+1)$ different a, b, c, and d tecto-RNAs can be combined to construct $(l+1)^2$ tectosquares.

23. Considering that each of the four tectosquare units can either have no tail, or a 3’ tail of $n$ different sequences with different size $l$ and orientation $o$, a total of $4(l+1)$ different a, b, c, and d tecto-RNAs can be combined to construct $(l+1)^2$ tectosquares.


28. Lts associate better when tails are oriented parallel to their a-d and b-c sides. This fact supports our twofold symmetrical models (Fig. 1C; fig. S1), with a-d and b-c being parallel to each other and a-b and c-d being tilted. Assembly through tails oriented parallel to a-b and c-d is nevertheless possible.

29. With 12 different tail-tail connectors, we are presently able to generate a 3 by 3 grid made of nine different tectosquares with the position of each of the 36 constitutive tectoRNAs being fully addressable within the RNA lattice.


35. S. Wolfram, A New Kind of Science (Wolfram Media, Champaign, IL, 2002).

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Materials and Methods
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Translation of DNA Signals into Polymer Assembly Instructions
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We developed a DNA nanomechanical device that enables the positional synthesis of products whose sequences are determined by the state of the device. This machine emulates the translational capabilities of the ribosome. The device has been prototyped to make specific DNA sequences. The state of the device is established by the addition of DNA set strands. There is no transcriptional relationship between the set strands and the product strands. The device has potential applications that include designer polymer synthesis, encryption of information, and use as a variable-input device for DNA-based computation.

We built a DNA nanomechanical device that mimics the translational capabilities of the ribosome. In response to a DNA signal, it aligns a series of molecules in specific positions; these molecules are then fused together in a specific order. For convenience, we have prototyped this system with DNA, so the products are DNA oligonucleotides of a defined sequence. Thus, in this case, the chemistry of the product is similar to that of the signal molecules, but there is no complementary relationship to the signal sequences. By using DNA molecules to set the states of two DNA PX-JX2 devices (1) independently, we programmed the synthesis of four different product molecules.

The PX-JX2 device is a sequence-dependent DNA machine, the state of which is controlled by hybridization topology (1). It can assume two structural states (termed PX and JX2), which differ from each other by a half-turn rotation of one end of the molecule relative to the other end (Fig. 1A). Two different pairs of set strands can bind to the framework of the device, thereby establishing which structural state it adopts. The set strands contain short unpaired segments (“toeholds”) at one end to facilitate their removal by unset strands that bind to the toeholds and then remove the set strands by branch migration (2). In addition to the PX-JX2 device, numerous variants of sequence-dependent control, pioneered in DNA tweezers by Yurke et al. (2), have been reported; these include a DNA