RNA origami design tools enable cotranscriptional folding of kilobase-sized nanoscaffolds

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RNA origami is a framework for the modular design of nanoscaffolds that can be folded from a single strand of RNA and used to organize molecular components with nanoscale precision. The design of genetically expressible RNA origami, which must fold cotranscriptionally, requires modelling and design tools that simultaneously consider thermodynamics, the folding pathway, sequence constraints and pseudoknot optimization. Here, we describe RNA Origami Automated Design software (ROAD), which builds origami models from a library of structural modules, identifies potential folding barriers and designs optimized sequences. Using ROAD, we extend the scale and functional diversity of RNA scaffolds, creating 32 designs of up to 2,360 nucleotides, five that scaffold two proteins, and seven that scaffold two small molecules at precise distances. Micrographic and chromatographic comparisons of optimized and non-optimized structures validate that our principles for strand routing and sequence design substantially improve yield. By providing efficient design of RNA origami, ROAD may simplify the construction of custom RNA scaffolds for nanomedicine and synthetic biology.

The field of RNA nanotechnology began by extracting RNA structural modules from natural RNA molecules and connecting them to create engineered constructs. This approach was enabled by the structural determination of biological RNA molecules such as the ribosomal subunits, which provided a large library of RNA modules from which to build. With these modules, architectures ranging from multi-stranded tiles to single-stranded origami have been explored. Of particular recent interest are RNA structures designed to fold cotranscriptionally during their synthesis by RNA polymerase. These have the benefit that they can be genetically expressed and folded within cells. Previously, we introduced the RNA origami method—a highly regular architecture that arranges RNA helices into parallel arrays held together by crossovers and kissing loops (KLs)—which is compatible with cotranscriptional folding, but several bottlenecks in computational design methods have limited the size (450 nt) and folding yield. Later studies constructed somewhat larger (715 nt) wireframe structures that fold cotranscriptionally by composing complex tertiary motifs in vitro and in vivo. The largest currently achieved structures (6,000 nt) require long (~18 h) thermal anneals, making them incompatible with cotranscriptional folding in cells.

RNA nanostructures can serve as functional scaffolds by directly incorporating RNA-protein binding domains, small-molecule aptamers, biosensors, ribozymes, small interfering RNAs or combinations of such modifications to create multifunctional nanoparticles. RNA nanostructures that fold cotranscriptionally have been expressed in cells, where they have the potential to be used as biosensors, scaffolds or regulators for synthetic biology applications—for example, to control product formation from colocalized enzymes and perform gene regulation via recruitment of transcription factors. To verify that two proteins are located on the same scaffold, split fluorescent proteins or Förster resonance energy transfer (FRET) between fluorescent proteins are often used. Similarly, fluorescent RNA aptamers (split-Spinach and apta-FRET) have been used to verify scaffolding effects. RNA origami structures may incorporate tertiary motifs such as the IRES or bKL motifs to produce ~90° bends that allow out-of-plane functionalization. 2′-Fluoro-modified RNA origami scaffolds carrying the thrombin aptamer have been used to produce a potent therapeutic anticoagulant.

Computational methods have played a central role in developing RNA nanotechnology by facilitating core tasks. Dedicated software has been developed to ease the construction of RNA nanostructures from three-dimensional (3D) structural motifs. However, no software exists for the interactive 3D modelling of large and regular RNA scaffolds such as the RNA origami architecture. Algorithms simulating RNA cotranscriptional folding have been developed for predicting folding pathways, which for small structures enables designers to verify that their sequences will avoid kinetic traps, but it has not been possible to do this for RNA origami. RNA sequence design algorithms were originally developed based on secondary structure thermodynamic folding algorithms, but these lack the ability to efficiently predict pseudoknots (such as KLs). RNA origami, which are stabilized by numerous KL interactions along their strand path, necessarily contain numerous pseudoknots and are therefore not easy to design. Another important element for RNA sequence design is the ability to incorporate numerous sequence constraints to allow RNA sequence and structural motifs to be added, but current design pipelines lack the ability to simultaneously incorporate the multiple constraints necessary for the design of RNA origami structures.

In this Article, we introduce the RNA Origami Automated Design (ROAD) software—a computer-aided design software to automate the 3D modeling of structures, analyze folding paths and design sequences and KLs that fold into the designated structures—allowing us to greatly extend the scale and diversity of RNA origami scaffolds. ROAD allows us to rapidly prototype multiple distinct scaffolds and investigate the effects of different design parameters.
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KLs that interact at an angle of 120° are predicted to fold into the desired structure. Constraints and secondary structure from the blueprints and uses random mutation to generate sequences that simultaneously satisfy the sequence and structural constraints. Revolvr takes sequence constraints and secondary structure from the blueprints and produces a visualization of the order in which helical domains and KLs form, flagging regions that might be susceptible to misfolding.

RNApath analyses the blueprints and produces pseudoknots. Dashed lines indicate all other base pairs. The blue arrow denotes the KL interaction, and the pink and orange arrows denote helices in the loop region. RNAbuild parses blueprints into modules and produces molecular models in PDB format. RNAbuild, RNApath and Revolvr, which take a user-specified ‘RNA blueprint’ as input. RNA blueprints are text-based diagrams that encode all Watson–Crick base pairs, sequence constraints, pseudoknots, base stacking at junctions, and 5′ to 3′ strand orientations (Fig. 1g). RNAbuild uses a module library to build atomic models according to specifications in the blueprint (Fig. 1h). The automated atomic modelling helps the user to design curvature and avoid steric clashes within larger RNA structures that are otherwise not apparent in the RNA blueprint. RNApath analyses the folding path for potential topological barriers that may arise during with a short design cycle. We study the effect of curvature and crossover placement within RNA origami structures by atomic force microscopy (AFM), allowing us to greatly increase the scale of the structures. To study the effect on yield, we then constructed a set of non-optimal designs and analysed yields by size-exclusion chromatography (SEC) and negative-stain transmission electron microscopy (TEM). Finally, we tested the ability of ROAD to design RNA origami scaffolds embedded with aptamers for binding fluorescent proteins and small-molecule fluorophores, and used FRET between aptamers and otherwise not apparent in the RNA blueprint. RNApath analyses the blueprints and produces a visualization of the order in which helical domains and KLs form, flagging regions that might be susceptible to misfolding. Revolvr takes sequence constraints and secondary structure from the blueprints and uses random mutation to generate sequences that simultaneously satisfy the sequence constraints and are predicted to fold into the desired structure.

**Results**

**Design tools for creating RNA origami scaffolds.** We developed the ROAD software package (‘Code availability’ section) to automate the main design steps for RNA origami: model building, folding path analysis and sequence design. ROAD is based on a library of compatible structural modules used to construct RNA origami structures. Core modules such as helices, junctions and 180KLs (KLs that interact at an angle of 180°) are used to build the central scaffold (Fig. 1a), and peripheral modules such as tetraloops, 120KL connectors (KLs that interact at an angle of 120°), light-up aptamers and protein-binding aptamers are used to add functionality (Fig. 1b). Schematic representations of the core modules can be used like Lego bricks to compose a large diversity of different designs (Fig. 1c) that directly translate to atomic coordinates (Fig. 1d). Closely spaced crossovers between three helices result in ‘dovetail’ (DT) junctions (Fig. 1e), which is an important design parameter for RNA origami, because the DT length (in base pairs, bp) changes the dihedral angle between connected helices (Fig. 1f). To avoid steric clashes between helices, DTs are restricted to certain lengths and are named sDT, where the spacing s can have values from −5 to +2 bp (ref. 14; Supplementary Fig. 4).

The ROAD software package consists of three main algorithms: RNAbuild, RNApath and Revolvr, which take a user-specified ‘RNA blueprint’ as input. RNA blueprints are text-based diagrams that encode all Watson–Crick base pairs, sequence constraints, pseudoknots, base stacking at junctions, and 5’ to 3’ strand orientations (Fig. 1g). RNAbuild uses a module library to build atomic models according to specifications in the blueprint (Fig. 1h). The automated atomic modelling helps the user to design curvature and avoid steric clashes within larger RNA structures that are otherwise not apparent in the RNA blueprint. RNApath analyses the folding path for potential topological barriers that may arise during...
the cotranscriptional folding process (Fig. 1i). Topological barriers can arise if a KL interaction (Fig. 1g, blue arrow) forms before the formation of helices in the loop region (Fig. 1g, pink and orange arrows), because the formation of a double helix may be sterically hindered by the closed-loop region. RNApath determines topological barriers based on the relative rates of KL and helix formation, as well as the speed of synthesis, and generates plots and 3D folding animations (Supplementary Videos 1–6) to help the user avoid topology-based misfolding. Revolver is a sequence design algorithm that uses a multi-stage sequence optimization procedure involving positive design by minimum free energy (MFE)13 prediction, negative design by sequence symmetry minimization (SSM)14, and KL orthogonalization to develop a sequence that folds into the target structure (Fig. 1j). The ROAD package and the analysis scripts are described in the Methods, a tutorial is provided as Supplementary Note 1, and a web server has been established to make the software easily accessible ('Code availability' section).

Design of multivalent interfaces for RNA origami tiles. The ROAD software was used to design three-helix (3H) RNA origami tiles with edge interactions to form fibres or rings, which make them easier to observe by AFM imaging. To make the interactions stronger, the 3H tiles were connected by two 120KL interactions15. Their relative in-plane positioning defines the tile–tile interaction angle \( \theta \) (Fig. 2a), which can deviate from 120° because the KL motif is flexible enough to accommodate a range of angles16. Using RNAbuild, we designed conformations (cf. the two stacking isomers observed in four-arm 0 and +11-bp DTs) result in minimum curvature of the RNA origami. Three 120KLs added to the edges of these tiles were programmed to join the tiles in a trans configuration, resulting in zigzag-shaped filaments (Fig. 3b,c and Supplementary Fig. 8). Three 120KLs added to the edges of these tiles were programmed to join the tiles in a trans configuration, resulting in zigzag-shaped filaments (Fig. 3b,c and Supplementary Figs. 15 and 16). A few alternative 6H tiles that contained isolated 0-bp DTs were shown to fold well (Supplementary Fig. 7, combining +11, 0, −2, −11 and −13-bp DTs). To create still taller tiles, two copies of ZigZag-A-1X were merged, via −11-bp DTs, to create the core of a nine-helix tile (Supplementary Fig. 8). Addition of 120KLs resulted in trans connections and filaments of alternating up–down orientations for ZigZag-B-9H tiles (Fig. 3d). Addition of 180KLs resulted in cis connections and filaments of consistent orientation for Ribbon-9H tiles (Fig. 3e). The 9H tiles showed more partial structures and had a reduced folding yield, estimated to be 51–62% (Supplementary Figs. 17 and 18), which could be caused by topological folding barriers (marked in red and orange in Supplementary Video 1), as suggested by RNApath analysis.

We used lateral duplication and fusion (Supplementary Fig. 8) of ZigZag-B-1X to create tiles with two repeats (ZigZag-B-2X in Fig. 3f and Supplementary Fig. 10; Supplementary Fig. 11 shows unexpected edge interactions) or four repeats (ZigZag-B-4X in Fig. 3g and Supplementary Fig. 10). The 2X duplication did not seem to affect yield (estimated to be 78%), whereas the 4X duplication had a reduced yield of 58% (Supplementary Figs. 19 and 20). The reduction in yield of the large 12×48-nm ZigZag-B-4X could not be explained by RNApath analysis (Supplementary Video 2), but is most likely to be caused by the misfolding and aggregation of its long transient 5′ single-stranded end. Tiles with alternating −2-bp and +/−11-bp DTs will be flat but have steeply sided tops. To obtain a more rectangular tile, we replaced each −2-bp DT with a +9-bp DT (−2-bp offset by +11 bp), so that every repeat unit had a counterbalanced set of +9-bp and −11-bp DTs. As an example of this architecture we designed the three-repeat Ribbon-5H-3X with 180KLs connectors, resulting in straight linear chains as observed by AFM with a yield of 42% (Fig. 3h and Supplementary Fig. 21). As a second example, the tile was extended to nine helices tall and designed without intermolecular connections, as a standalone scaffold, reaching a length of 2,360 nt and a size of 20×36 nm (Fig. 3a, Rectangle-9H-3X). However, the expansion resulted in only a few examples of rectangular shapes, which all had folding defects (Fig. 3i). Finally, we designed RNA origami with shorter or longer double crossover spacing: ZigZag-B-2X-Mini with two turns between crossovers (Supplementary Fig. 7) and Ribbon-5H-3X-bumps with four turns between crossovers (Supplementary Figs. 7 and 10). The latter was designed with six out-of-plane dumbbells placed in the middle of the four-turn stem regions; however, the complexity of the design resulted in a low observed yield of 30% (Supplementary Fig. 22), and the three-dimensionality of the design resulted in poor imaging by AFM.

The manual evaluation of folding yields from the AFM images is summarized in Supplementary Table 3. The folding yield negatively correlates with increasing length of the RNA origami structures tested, and with RNApath-predicted topologically blocked positions. This observation is supported by an apparent correlation between the number of observed misfolded structures and RNApath-predicted topologically blocked positions. The data
indicate that folding topology is important and that increasing the
height of the tile results in increased occurrence of predicted topo-
logical barriers, which arise because of the longer delay between
the synthesis of KL partners. The large Rectangle-9H-3X was pre-
dicted to have several topological barriers, and this correlated with
the larger folding defects observed (Fig. 3i and Supplementary
Fig. 23). Another example is a merged version of ZigZag-A-1X and
ZigZag-B-1X that is 10 helices tall, where we observe partly formed
tiles that again have large defects that seem to correspond to the
regions with predicted topological barriers (Supplementary Fig. 23).

**Effects of design parameters on folding yield.** To support our
AFM yield analysis, we performed negative-stain TEM imaging
of SEC-purified monomer RNA origami structures. A monomeric

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**Fig. 2 | Design of RNA origami curvature and tile–tile interfaces.**

**a.** Schematic model tile–tile interaction and angle θ.

**b.** Protocol comprising isothermal synthesis at 37 °C by T7 polymerase, cotranscriptional folding, and sample preparation for AFM. NiCl2 binds RNA strongly to mica, improving image quality.

**c–e.** RNAbuild models of three-helix designs with DTs of different length (named 3H–sDT, where s indicates the length of the DT). Each 1-bp increase in
DT length decreases the dihedral angle between the three helices (black cross-sections) by 32.7°. Insets: secondary structures, with DT marked in blue.

The most prevalent closed polygons are shown: hexamer for 3H–2DT (c), octamer for 3H–3DT (d), pentamer for 3H–4DT (e). f–h, AFM images of RNAs
(corresponding to 100-nm and 1-µm fields. Three-helix designs 3H–2DT (f), 3H–3DT (g) and 3H–4DT (h) are shown. The thin filaments in the background
are DNA templates from which the RNA structures are transcribed. The yields of tiles shown in the top AFM images are from Supplementary Table 3.
five-helix scaffold (5HS, Fig. 4a), based on one of our best performing RNA tiles, ZigZag-B-1X (85% yield by AFM), resulted in 86% yield of monomer as determined by SEC analysis (Supplementary Fig. 24), and TEM images of the monomer sample revealed homogeneous and monodisperse particles with class averages displaying highly resolved details of tight helix packing (Fig. 4c and Supplementary Fig. 24). The TEM analysis revealed a clear preference for observing either front- or back-face views of the 5HS...
structure (Fig. 4c; Supplementary Fig. 27 provides a plot of orientation distribution) even though a few edge views were observed as well (Supplementary Fig. 24). Although we were not able to obtain an ab initio model, a 3D reconstruction could be made by using the theoretical model as the input search volume (Fig. 4b and Supplementary Fig. 24).

To investigate the robustness of the RNA origami method in relation to the core design parameters, we generated a challenging monomeric design with five-helix rows and two-KL columns, with an unconventional meandering strand path, and generated two different versions with different 5’ start sites (Path1 shown in Fig. 4d and Path2 shown in Fig. 4e). The two strand paths are equivalent in 3D structure, but the different positioning of the 5’ start sites (Fig. 4d,e, blue circles) has a large effect on folding topology as predicted by RNApath. During transcription, Path1 has a long transient 5’ single strand but no predicted topological barriers (Fig. 4f and Supplementary Video 3), whereas Path2 has no transient 5’ single strand but has substantial topological barriers predicted (orange and red regions, Fig. 4g and Supplementary Video 4). Previously, we have avoided designs with a long 5’ transient single-stranded region, because transient single strands are expected to increase aggregation (stage 5 of Revolvr). As expected, these designs displayed a substantial amount of aggregation that resulted in a relatively low yield (stage 5 of Revolvr). To investigate the effect of sequence design optimization, a third design was created based on Path1 satisfying the MFE structure (stages 1–4 of Revolvr, Methods and Supplementary Figs. 1 and 2) but lacking the final KL optimization (stage 5 of Revolvr). As expected, these designs displayed a substantial amount of aggregation that resulted in a relatively low yield of monomers of 26–44% as determined by SEC analysis (Fig. 4k,l), which can be compared to 5HS, which displays 86% monomer yield by SEC analysis (Fig. 4l and Supplementary Fig. 24). The monomers were observed to be stable post SEC purification (Fig. 4k), indicating that aggregation is happening during the cotranscriptional folding process and is not the result of a subsequent equilibration process.
TEM imaging was performed on the purified monomer and aggregate peaks and the monomers were observed to be monodisperse (Supplementary Fig. 25). To be able to address the folding yield, the TEM grids were prepared from the same concentration of purified RNA samples and quantified from the same number of acquired images. Unbiased blob picking was used to identify particles, and 2D class averages showed that the number of face views of the RNA origami structure was very different in the three samples, and that several alternative particle views and shapes were observed (Fig. 4h-j and Supplementary Fig. 26). Because each design has the same predicted 3D structure, they should have the same angular distribution on the foils. We used the number of easily recognizable face views observed in the 2D class averages as an estimate of the cotranscriptional folding yield of the three samples (Fig. 4m; see Methods for a description of the folding yield calculation). Path1 with an optimized sequence had a 30% folding yield. Path2 with an optimized sequence had 25% folding yield, but, of these, only 3/4 adopted the designed structure, whereas 1/4 displayed a ‘pursehandle’ phenotype (Fig. 4i, blue arrowhead), which we suggest corresponds to distortions in the long topologically blocked helix (Fig. 4e) due to partial inhibition of Watson–Crick base-pairing (Fig. 4g, blue arrowhead). Path1 with non-optimized KL sequences had a reduced folding yield of 6% (Fig. 4m) and a large fraction of alternative shapes (Fig. 4, right and Supplementary Fig. 26).

From the limited, but equivalent, datasets acquired for each design, only the particles picked from the Path1 data produced a reasonable ab initio reconstruction (Fig. 4n). As observed previously in the TEM analysis of the 5HS, the Path1 structure had preferential face adsorption to the carbon foil, but in this case one face was strongly preferred (Supplementary Fig. 27 provides a plot of orientation distribution), which indicates that the larger monomer structure has an asymmetric shape in solution that affects adsorption to the carbon. Although the tested designs can all fold into the correct 3D structure, the choice of strand path and sequence optimization have large effects on both the yield and structural homogeneity of the origami particles.

Scaffolding of proteins and small molecules. To test the ability of RNA origami to scaffold proteins, we used the high-yield 5HS scaffold (Fig. 4a) containing 10 hairpin sites that can be used for functionalization (Fig. 5a). RNAbuild was used to design a series of five scaffolds that positioned two different protein-binding aptamers at increasing distances of 2.5, 5, 7.5, 10 and 22 nm using scaffolds named MxP\(x\)y, where \(x\) refers to the position of the MS2 aptamer and \(y\) to the position of the PP7 aptamer (Fig. 5b,c). All scaffolds were designed by Revolv to have unrelated sequences, except for the fixed sequence of the aptamers. Similar to a previous scaffolding study, we fused mTurquoise2 (a cyan fluorescent protein, CFP) and YPet (a yellow fluorescent protein, YFP) with the viral coat proteins MS2 coat protein (MCP) and PP7 coat protein (PCP), respectively (Fig. 5b,c and sequences in Supplementary Table 8). When the M5P3 scaffold was transcribed in the presence of excess fluorophores it resulted in a FRET signal that reached saturation after 20 min (Supplementary Fig. 28), showing that the scaffold cotranscriptionally folds and brings the two proteins together within FRET distance. To compare several RNA scaffolds, we normalized concentrations of cotranscriptionally folded RNA products and incubated them with excess amounts of fluorescent proteins. The FRET signal was observed to generally decrease with increasing distance between aptamers (Fig. 5d and full spectra in Supplementary Fig. 30); however, some constructs with a spacing differing by 2.5 nm were not significantly different in FRET signal (M5P4 ≈ M5P3; M5P2 ≈ M5P1, \(P > 0.05\), Student’s t-test), and the control constructs (M5P10 with a nominal distance beyond the Förster radius and 5HS with no aptamers) showed measurable levels of FRET. These non-ideal effects may be explained by the large size of the fusion proteins with long linkers used as well as the documented tendency of the fluorescent proteins to form dimers in a colocalized context. In general, the results may also be affected by scaffold flexibility and sequence-specific conformations of particular constructs.

RNAbuild was used to design two series of scaffolds that positioned the fluorescent aptamers Spinach and Mango in various structural contexts (Fig. 5e-h and Supplementary Fig. 29). The first series was based on a two-helix scaffold S2T (short, two turns) with short stems to position Spinach and Mango aptamers and two helical turns between crossovers (Fig. 5e), which was previously shown to produce a strong FRET signal between the fluorophores DFHBI-1T and YO3-biotin. Two variations of the S2T scaffold were produced: S3T (short, three turns) with wider crossover spacing (Fig. 5f) and L3T (long, three turns) with longer stems for positioning fluorescent aptamers and wider crossover spacing (Fig. 5g). The S2T scaffold transcribed in the presence of fluorophores shows slowly increasing fluorescence and FRET signals over at least 90 min (Supplementary Fig. 28), which is probably caused by the slow folding of the fluorescent aptamers. To compare several RNA scaffolds, we normalized the RNA concentrations before incubation with an excess amount of fluorophores. Fluorescence measurements showed ~35% FRET for S2T, ~30% FRET for S3T and ~5% FRET for L3T scaffolds (Fig. 5i; full spectra are provided in Supplementary Fig. 30). Although RNAbuild models predict that all three scaffolds have the same distance and orientation between donor and acceptor fluorophores (Fig. 5e-g), the large decrease in FRET signal with increasing construct size suggests that scaffold flexibility (due to longer stems and to a lesser extent larger crossover spacing) strongly influences the FRET signal. The second series was based on the three-helix scaffold from Fig. 2 with fluorescent aptamers placed on the top and bottom helices and two turns between crossovers (L2T3DT in Fig. 5h and Supplementary Fig. 29). This scaffold is able to tune fluorophore spacing (from 1.3 to 3.2 nm in increments of 0.6 nm) by changing DT length (from \(s = -2\) bp to \(s = -2\) bp in increments of 3.2 bp), respectively. Fluorescence measurements for the L2T3DT scaffolds show a decrease in FRET signal as the predicted distance between the fluorophores is increased (Fig. 5i; statistically significant \(P < 0.05\) in Student’s t-test, except for L2T–3DT). Within this series, care was taken to maintain the relative orientation of the Spinach and Mango aptamers to avoid the possible effects of oriented dipoles on FRET (Supplementary Fig. 29). Comparing between series, we attribute the low FRET signal of the sterically overlapped construct L2T–5DT relative to construct S2T, which shares a similar crossover spacing, primarily to the flexibility contributed from a longer aptamer-bearing arm.

Discussion

The design and synthesis of cotranscriptional RNA structures in high yield is very challenging. In our previous work we were only able to achieve cotranscriptional folds of 440 nt in length with yields so low that only a few correctly formed objects could be identified. In the current work we have improved the RNA origami method to greatly expand both the size and functional complexity of RNA nanostructure designs, as well as dramatically improving the yields of correct products that are able to be produced by cotranscriptional folding. We have rapidly prototyped 32 different RNA origami designs in this work, allowing us to explore the effect of multiple RNA origami design parameters: DT geometry, multivalent interfaces, taller and wider structures, different strand routing strategies, as well as designs incorporating aptamers for scaffolding proteins and small molecules. The achievements were enabled by the development of the ROAD software package, comprising the programs RNAbuild, RNAPath and Revolv, which work together to facilitate the design of large and complex RNA structures and were all found to be crucial for obtaining high-yield RNA scaffolds.
RNAbuild automates the rough 3D modelling of RNA origami structures, which were previously constructed by hand, allowing us to design much larger and more sophisticated designs than before. In this work, we have demonstrated that the DT seam can be used to adjust the curvature of RNA origami structures to tune the tile–tile interaction angle to form rings of defined size (Fig. 2) and to tune the distance between attached fluorescent aptamers (Fig. 5h,i).

RNAbuild further allowed us to expand the RNA origami architecture by domain duplication and fusion (Supplementary Fig. 8), reaching sizes of ~2,000 nt, albeit with decreasing yields as estimated from AFM images (Fig. 3 and Supplementary Table 3). Interestingly, TEM analysis revealed preferred landing of larger RNA origami structures on the carbon film (Fig. 4h and Supplementary Fig. 27), which indicates that larger RNA origami structures may have a curved structure in solution. Even though ab initio reconstruction from TEM images showed that the RNA origami structures are flat, this may be an artefact of the deposition on the carbon film. RNAbuild can in the future be improved by extending the library of functional motifs and by supporting alternative architectures such as parallel crossover RNA origami and wireframe RNA origami as well as allowing physical simulation of the structures to address strain-induced distortions (using, for example, oxRNA).

The recently developed program RNAmake—which specializes in grafting and stabilizing tertiary motifs onto an input model—could complement and extend RNAbuild.

RNAPath makes a simple folding path analysis based on the RNA blueprint (while not taking into account the designed sequence) to predict possible topological barriers for the cotranscriptional folding process. Comparing the number of predicted topological barriers to the folding yield estimation from AFM images revealed a strong correlation, where the most severe cases did not result in any correctly folded objects (Supplementary Table 3). However, the effect of size and number of topological barriers could not easily be separated in this evaluation, because topological barriers arise when designs become larger (and especially taller). The effect of folding path choice was investigated further by designing an RNA origami structure with two alternative folding paths. TEM analysis revealed that there was ~30% decrease in folding yield for the path with topological barriers (Fig. 4m) and that misfolds could be observed with severe distortions of the topologically trapped helix (forming a ‘purse handle’) (Fig. 4i). The observation that only the structure without topological barriers resulted in a reasonable 3D reconstruction further underscores the importance of taking this design parameter into account (Fig. 4n). The kinetic folding analysis of RNAPath may be improved by using thermodynamic kinetic folding algorithms like Kinefold or by using coarse-grained molecular simulations such as oxRNA.

Revolvr designs sequences for RNA blueprints with a high content of pseudoknots—a task that has not been approached by any other RNA design program. Revolvr solves this task by using a multi-stage sequence optimization procedure involving MFE-based positive design, SSMBased negative design and KL orthogonalization, which makes it very efficient in the use of computational time (Supplementary Fig. 31). Sequence design by Revolvr has a high success rate, with most of the structures presented in this study working on the first try. The high success rate prompted us to design new...
sequences for each new RNA origami for scaffolding of proteins and small molecules with the assumption that the geometry of the RNA origami and not the precise sequence was important, which was verified to some extent by the overall ability to control distances on the scaffolds. The effect of sequence design was investigated by designing a non-optimal sequence, where only the last stage of the five-step design procedure (KL orthogonalization) was omitted. The folding yield was observed to decrease from 30% to 6%, showing that this sequence design step has a substantial effect. Future improvements to Revolver sequence design could be to include an RNA secondary structure partition function in the design optimization (for example, NUPACK\textsuperscript{18}). The partition function optimization may become especially important when designing more challenging RNA structures with smaller stem regions. A great future challenge would be to include pseudoknot-prediction and kinetic folding simulation directly in the sequence design algorithm.

Although RNA nanostructures have previously been used to scaffold protein-binding aptamers\textsuperscript{9,10,11} and small-molecule aptamers\textsuperscript{12,13,14}, here we demonstrate distance control by changing the position of aptamers on parallel helix ends and by tuning the DT length to gradually change the distance between helices. Our FRET studies highlight a potential size- flexibility tradeoff in RNA scaffold design: based on our current architecture, larger structures enable complex spatial arrangements of proteins to be constructed, but smaller, more rigid structures are required if more precise distances are desired. The elaboration of RNA origami to multilayer 3D structures may obviate this tradeoff by achieving simultaneously large and rigid structures, as has been achieved for DNA origami design\textsuperscript{15}. Rigidity and the precision of arrangement will also be improved by exchanging large, flexible, dimeric linkers such as MS2 and PP7 aptamer-protein constructs with smaller, monomeric RNA-binding proteins or peptides such as L7Ae\textsuperscript{16} or BIV-Tat\textsuperscript{17}. With improved protein scaffolding methods, the RNA origami scaffolds may be used to control product formation from colocalized enzymes\textsuperscript{18,19,20} and perform gene regulation via recruitment of transcription factors\textsuperscript{21,22}.

In this study, we have improved the RNA origami method to allow the design of cotranscriptionally folding RNA nanostructures approaching the size of ribosomal RNAs. However, the structural complexity and strategies for cotranscriptional folding of RNA origami and the ribosome are very different. The ribosome is constructed from a high percentage of tertiary structural motifs, with almost 50% non-Watson–Crick base pairs. By contrast, the RNA origami architecture is mainly constructed from Watson–Crick base pairs formed by secondary structure elements and pseudoknots. The cotranscriptional folding of the ribosome involves transiently stable helices, protein chaperones and structural switches to guide the strand into a final native state that does not correspond to the MFE. However, RNA origami takes advantage of a very different, very unnatural design construction, in which every helix of the design is able to rapidly find its MFE structure during the kinetic folding process. Thus, we are engineering very smoothed-out folding landscapes, with strand paths designed to minimize the possibility of the strand misfolding during the process. A recent computational study\textsuperscript{23} suggests a general method for choosing strand paths that minimizes the risk of topological barriers, and finds that KIs arranged into columns connected by a single common helix will result in the fewest topological barriers. Although many of our designs have this property (minimized topological barriers), this work deserves to be further explored quantitatively, and determining whether or not RNA origami contain minor misfolded elements may require the adaptation of SHAPE-seq\textsuperscript{24} or other techniques to very large RNA structures, or perhaps high-resolution cryo-electron microscopy. Future challenges for the cotranscriptional RNA origami method will be to increase the structural complexity with 3D architectures and tertiary motifs. Likewise, yet to be explored is the ability to program RNA origami scaffolds with functional, dynamic features and molecular computing elements (like strand displacement logic gates), as achieved with DNA origami structures, to create biosensor devices and nanorobots.

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Methods
The ROAD package. To automate the key processes of RNA origami design, a set of algorithms were made that together constitute a design pipeline. RNABuild builds a PDB structure from an RNA blueprint. RNAPath analyses the folding path of the given RNA sequence and highlights topological barriers, which can be generated in the form of a series of keyframes and a Chimera command file for automatic generation of a video. Revolvr designs RNA sequences that fold into target structures (requires installation of the Vienna RNA package https://www生物.univie.ac.at/RNA/). Trace takes an RNA blueprint without a sequence assigned to it, and a candidate sequence and creates a new blueprint with the candidate sequence threaded onto the blueprint. Trace_pattern converts RNA blueprint diagrams into dot-paren notation and candidate sequence for Revolvr. Trace_analysis analyzes an input blueprint and annotates it with features such as duplicated sequences, unintended complementary sequences, GC content, restriction sites and so on. Flp_trace flips a blueprint horizontally, vertically and in both directions, which aids in the design of more complex patterns using domain duplication and fusion. The analysis package is available for download at GitHub (https://github.com/esa-lab/ROAD) and has been made available as a web server with accompanying tutorials (https://bioan.de/software/rnaa-design/).

RNABuild. The RNABuild algorithm automates structural modelling of RNA origami structures. As input the algorithm takes an RNA blueprint and parses the contents of the套路 into a set of predefined 2D motifs. The 3D atomic model is constructed from the套路 to 3D ‘end by serial addition of 3D structural modules from a library that matches the 2D motifs and 3D structural motifs and modules. When each structure is added to a sequence, it is rotated and translated to the correct position using a single reference base or base pair having A-form parameters (Supplementary Table 1 and Supplementary Fig. 3); these reference bases are added to modules when they are put in the library (Supplementary Fig. 1). For building RNA helices, there are four different trivial nucleotide modules, simply the PDB coordinates for each RNA nucleotide. Modules for helices, terminal tetraloops, fluorogenic aptamers and the RNA protein binding domains are all based on known crystal structures. The helix axes of the crossovers module are modelled as parallel, rather than at the 60°–70° angle found in crystal structures (PDB 1HP0), under the assumption that the coupling of adjacent crossovers forces them flat. The spacing of crossovers between three or more parallel helices defines the geometry of the DT function, and RNABuild helps model these.

RNAPath. The RNAPath algorithm analyses the folding path of RNA origami designs to identify possible topological barriers during the cotranscriptional folding process. The algorithm takes an RNA blueprint as input and analyses the RNA structure as it is being extended from the 5′ end to the 3′ end. For each subsequence of length k, RNAPath takes the fold computed for the k – 1 subsequence, and decides what new base pairs can be added to the fold. RNAPath adds a secondary structure (for example, a particular hairpin) to the fold for the subsequence having the smallest k possible, which models the situation where the secondary structure folds immediately, as soon as the necessary sequence is transcribed. By default, a particular KL is added to the fold of a subsequence k only when k is at least 150 nt longer than the smallest subsequence that contains both halves of the KL. This feature roughly captures the KL formation time, modelling the situation where the KL formation is delayed by ~0.7 s (assuming a transcription speed of 3 nt/s that after the KL sequence has been transcribed. When the folding of KLS might topologically clash with secondary structure formation, RNAPath labels barrier loops by ‘−’ and topologically blocked nucleotides by ‘X’ in an analysis blueprint output. It additionally outputs a list of substructures in dot-paren where transient single strands are shown as ‘(‘, and crossovers as ‘)’. The delay is adjustable from 0 nt (for which almost all KLS cause clashes) to 5 nt (for which no clashes will occur). In addition, RNAPath can output a series of PDB models that can be rendered to create a video in UCSF Chimera v1.10 where pseudoknot loops are coloured in orange and topologically blocked nucleotides in red (Supplementary Videos 1–6 and Supplementary Note 1). Alternatively, the program trace_analysis provides a fast summary of any patterns in the sequence as well as positions of wrinkles within the design, and lastly the strand path analysis.

Revolvr. The Revolvr algorithm designs sequences for target structures by using a five-stage variant of stochastic gradient descent where each stage has an increasingly restrictive cost function (see the algorithm flowchart in Supplementary Figs. 1 and 2). The cost function is a score that combines the MFE folding prediction and different measures of sequence symmetry, and for each successive round of design becomes stricter. The input file defines the target secondary structure, pseudoknots and sequence constraints, and is initially seeded with a random sequence that satisfies the constraints (or with a user-inputted sequence). The first stage optimizes the MFE structure over five rounds of positive selection to stabilize the ends and multi-junctions at a cost of raising the GO content. The current sequence’s MFE structure, as computed by the ViennaRNA packagea, is used to calculate the Hamming distance of the current structure to the target structure, which is used as the cost function score for this round of design. The second stage applies an alternating positive and neutral design for a variable number of rounds, until the target structure is achieved: mutations are either targeted to regions that misfolded in the previous round (two out of every three rounds) or spread randomly throughout the sequence to enable neutral drift (every third round). To increase the speed of design, we scale the rate of mutation per design based on the success/failure of each iteration. A design round is considered successful when the cost function remains the same or decreases. The third stage uses negative design to decrease the probability of misfolding, prevent the inclusion of particular sequences (for example, restriction sites) where undesired, and make the DNA template from which RNA origami are transcribed easier to synthesize and PCR-amplify. SSMa is applied to remove all repeated sequences or regions of undesired complementarity above a threshold length (default setting of 10 nt). Similarly, removal of long homopolymer stretches, and a known transcriptional pause siteb, may help to reduce the frequency of unwanted transcriptional termination. The fraction of GC base pairs is reduced to below 55% to increase folding efficiency. Finally, GU doublets are introduced to simultaneously preserve the helix within the desired RNA structure, and weaken it within the corresponding DNA templates. All of the above constraints are applied through successive rounds of targeted mutation, until they and the MFE fold simultaneously satisfied. The fourth stage eliminates repetition from the set of KLS. Repeated (and thus also palindromic) KL sequences are targeted for mutation in successive rounds until all KL interactions are unique. The fifth stage optimizes the sequences of the KLS to have uniform binding energy and greater specificity. Energies for all possible KL interactions are estimated with the Duplex function of ViennaRNAa, and KLS are targeted for mutation until all desired KLS have energies in the range −10.7 kcal mol−1 and all non-energetically important energies greater than −6.0 kcal mol−1. Revolvr enables potentially conflicting requirements for positive versus negative design, sequence versus secondary structure constraints, and pseudoknotted versus non-pseudoknot structure to be balanced and satisfied. User-specified sequence constraints supersede user-specified secondary structure, which supersedes all other constraints. Sequence explicitly specified in the blueprint (such as aptamers) is left unmutated, even if the secondary structure specified for them cannot be achieved in an MFE structure, or if they violate a sequence symmetry constraint. Upon termination, Revolvr outputs an analysis of the designed sequence, which includes a blueprint populated with the sequence, KL energies, and the positions of potential topological clashes, violations of sequence symmetry constraints and GU wobbles.

Synthesis of RNA origami structures. DNA templates were commercially synthesized (Integrated DNA Technologies) as double-stranded gBlocks. DNA gBlocks were PCR-amplified using 19–20-nt primers (76°C) complementary to the ends of the gBlock, using standard Taq DNA polymerase, and purified using a Qagen PCR purification kit. RNAs were transcribed and cotranscriptionally folded in one-pot reaction containing template DNA (−4 ng/µl final of PCR amiplon), 6 mM Mg(OAc)2, 40 mM Na OAc, 40 mM KCl, 50 mM Tris-OAc (pH7.8), rNTPs (0.5 mM each) and 1 mM dithiothreitol (DTT). Reactions were initiated by adding 17 RNA polymerase (−0.2 U/50 µl). Transcription reactions were carried out in 50–µl volumes at 37 °C for 45 min to 2 h, depending on the sequence length. Larger designs required longer synthesis times (1–24 h), whereas smaller designs (especially for 2A5E) required just a few minutes to reveal multimeric products by AFM.

AFM sample preparation and imaging. A 1–5 µl volume of transcription product was mixed with 40 µl AFM dilution buffer (12.5 mM Mg(OAc)2, 40 mM KCl, 40 mM NaCl, 1 mM spermidine, 0.001% Triton X-100, 100 mM DTT, 12 mM MgCl2, 0.1 mM CaCl2, 0.5X ribolock (Thermo Fisher Scientific) and 40 mM Tris-Cl, pH 8.1) directly on the surface of a freshly cleaved l pipette tip. Most AFM images were collected using a multimode Olympus TR400PSA silicon nitride probes with a spring constant of ~0.08 N m−1. Negative-stain TEM. CF400 Au grids (Electron Microscopy Sciences) were glow-discharged for 45 s at 25 mA before removing and discarding the fluid. The mica was washed with a solution of 60 mM NiCl2. Most AFM images were collected using a multimode AFM (Digital Instruments) with a Nanoscope IIIA controller and a J-scanner. Olympus TR400PSA silicon nitride probes with a spring constant of ~0.08 N m−1 were used for imaging, with a drive frequency of ~6–9 kHz. The AFM results in Supplementary Figs. 10 and 11 were collected with a Bruker Fastscan Bio AFM (Bruker) under buffer using FastScan D-probes (Bruker).

Purification of RNA origami. RNA origami were transcribed from linearized pUC19 plasmid for large-scale synthesis and purification. Briefly, 25 µg of linearized plasmid was used as template in a 0.5 ml reaction containing 40 mM Tris-Cl pH 8.1, 1 mM spermidine, 0.001% Triton X-100, 100 mM DTT, 12 mM MgCl2, 0.1 mM CaCl2, 0.5X ribolock (Thermo Fisher Scientific) and in-house-prepared T7 polymerase. Transcription was carried out at 37 °C for 3 h before the addition of 40 µl of RNase Out (Thermo Fisher Scientific) directly on the surface of a freshly cleaved plate or 1 ml of RNase A (10 mg/ml) and incubation overnight at 37 °C. The RNA product was purified by resin-based chromatography on a Superose 6 column (GE) equilibrated with 25 mM HEPES pH 7.5, 50 mM KCl and 5 mM MgCl2.

Negative-stain TEM. CF400 Au grids (Electron Microscopy Sciences) were glow-discharged for 45 s at 25 mA before application of 3µl of sample and then blotted three times with 3µl of 0.5% uranyl formate. Peak 2 from the 5HS purification was diluted to 25 ng/µl before blotting and peak 2 from the Path2-optimized, Path2-optimized and Path1-non-optimized purifications were...
diluted to 50 nM -1. TEM images were obtained on a 120-kV Tecnai Spirit TEM equipped with a 4-K TVIPS CMOS camera at 67,000 magnification. The images were contrast-inverted and converted from tif to tiff using Emam2 before being imported into cryoSPARC V2.0. CTF correction was applied with CTFFinder. For 3D reconstructions, ~300 particles were manually picked in cryoSPARC and used to generate templates for the first round of templated particle picking. These particles were sorted into 50 2D classes, the best of which were used for 3D reconstructions. Ab initio 3D reconstruction of the Path1-optimized design produced a volume with real space slices similar to what would be expected from our design and was further refined by a non-uniform refinement. Ab initio 3D reconstruction repeatedly failed for the 5HS design and so a homogeneous refinement of the 5HS structure was performed using a 4-Å masked volume of our predicted RNA origami structure as an initial volume. For the comparison of the Path1-optimized, Path2-optimized and Path1 non-optimized designs, blob picking was performed on 88 images of each design with the default settings in cryoSPARC V2.0, followed by a single round of 2D class averaging into 50 classes. Structural deformities observed were measured with Eman2.

Folding yield calculation. The SEC yield was calculated as the monomer-to-aggregate fraction based on the average peak heights of the SEC chromatogram (UV 255 nm) for two or three transcription reactions. The TEM folding yield was estimated by counting the number of face views of the RNA origami particles based on the assumption that correctly folded particles would have a similar preference of adsorbing to the carbon film in this orientation. The number of face views was estimated from the amount (µg) of RNA loaded on the grids, the number of images obtained and the molar mass. The folding yield of the 5HS was assumed to be 95% based on analysis of TEM and AFM images (data not shown) and was used to calculate the relative folding yield of the other samples. The transcription to be 95% based on analysis of TEM and AFM images (data not shown) and was used to calculate the relative folding yield of the other samples. The transcription folding yield was calculated by multiplying the SEC monomer yield by the TEM folding yield. An alternative fold, named the ‘purse handle’, was identified by folding yield was calculated by multiplying the SEC monomer yield by the TEM

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Author contributions
C.G., P.W.K.R. and E.S.A. conceived the project. C.G., G.G. and E.K.S.M. performed the study. P.W.K.R. and E.S.A. supervised the project. C.G., P.W.K.R. and E.S.A. wrote the manuscript. All authors analysed the data and commented on the manuscript.

Competing interests
The authors declare no competing interests.
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