Cotranscriptional Folding of a Bio-orthogonal Fluorescent Scaffolded RNA Origami

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ABSTRACT: The scaffolded origami technique is an attractive tool for engineering nucleic acid nanostructures. This paper demonstrates scaffolded RNA origami folding in vitro in which, for the first time, all components are transcribed simultaneously in a single-pot reaction. Double-stranded DNA sequences are transcribed by T7 RNA polymerase into scaffold and staple strands able to correctly fold in a high synthesis yield into the nanoribbon. Synthesis is successfully confirmed by atomic force microscopy, and the unpurified transcription reaction mixture is analyzed by an in-gel-imaging assay where the transcribed RNA nanoribbons are able to capture the specific dye through the reconstituted split Broccoli aptamer showing a clear green fluorescent band. Finally, we simulate the RNA origami in silico using the nucleotide-level coarse-grained model oxRNA to investigate the thermodynamic stability of the assembled nanostructure in isothermal conditions over a period of time. Our work suggests that the scaffolded origami technique is a viable, and potentially more powerful, assembly alternative to the single-stranded origami technique for future in vivo applications.

KEYWORDS: cotranscriptional folding, scaffolded RNA origami, bio-orthogonal, split Broccioli aptamer, oxRNA simulation

Over the past few years, the emerging field of RNA nanotechnology has progressed rapidly with demonstrations of a variety of roles and sophistication of function that RNA devices might display. These include the engineering of essential coding and noncoding functions, mRNAs, tRNAs, ribozymes, aptamers, CRISPR RNAs, etc. As a consequence, a variety of RNA nanostructures with different functionalities, sizes, and shapes have been created to investigate and successfully demonstrate their potential in nanobiomedicine and synthetic biology.1–12

Different self-assembly strategies have been adopted to design RNA nanostructures13 ranging from RNA architectures14–18 to single-stranded self-assembly.19

Furthermore, taking advantage of the sequential transcription reaction by bacteriophage RNA polymerase, single-stranded RNA origami has been synthesized from long ssRNA molecules. In vitro transcribed and purified RNA sequences were self-folded into hearts, rectangles, and rhombus shapes adapting the ssDNA origami design strategy and considering the helical periodicity difference between DNA B-type and RNA A-type helix. Large and complex ssRNA origami were synthesized using partially complemented double-stranded RNA and parallel crossover cohesion without limitation due to RNA kissing loop interactions.20 However, these multi-kilobase ssRNA nanostructures were folded using a thermal annealing ramp gradient (from 85 to 25 °C), thereby limiting potential in vivo applications, such as the scaffolding of enzymes.21

On the other hand, previous work showed the synthesis of smaller ssRNA origami tiles and hexagonal lattices made by annealing and/or in vitro cotranscriptional folding that should be compatible with in vivo folding when genetically encoded and expressed in cells.22 These authors developed a strategy based on the combination of hairpins, kissing loops, and "dovetail seam" to promote and stabilize the folding during the T7 RNA polymerase in vitro transcription reaction.22,23 More recently, the ssRNA 2H-AE-ST tile scaffold presented by Geary et al.23 was used to create an aptamer-based FRET system where RNA tile synthesis in vivo was demonstrated by measuring FRET outputs without a direct atomic force microscopy (AFM) visualization, due to the small construct dimension.24

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Li et al.\textsuperscript{25} developed a different strategy in which the design concept was similar to the approach reported above but avoiding the use of short “dovetail seams”. The RNA nanostructures were designed based on natural motifs: the folding pathway was based on hairpin formation and tertiary interactions of unpaired residues. In detail, an RNA double-square was designed using a 3-way loop observed in pH29 pRNA and a 90°-kink from hepatitis C virus RNA genome. These thermodynamically stable and kinetically favorable RNA nanostructures were folded both in vitro and in vivo: nonetheless, the combination of phage and viral derived structural motifs can limit the structure design, the attachment of functional units, the creation of reconfigurable and dynamic nanostructures or their suitability for future theranostic in vivo applications.

While architectonics, single-stranded self-assembly methods and single-stranded origami strategies, are well explored, scaffolded (and multistranded) RNA origami approaches are still in their infancy. Drawing from DNA origami techniques in which several short staple strands sequences promote the folding of a longer single stranded scaffold into a specific shaped structure,\textsuperscript{26} we designed and synthesized RNA origami following a similar strategy. This approach can provide several advantages such as the high synthesis yield, the possibilities to produce reconfigurable nanostructures and to incorporate multiple and different functionalities in a precise position. Indeed, previous works demonstrated the synthesis of chemically modified and siRNAs functionalized RNA origami following a thermal gradient annealing of RNA staple strands and scaffold.\textsuperscript{27,28}

In our previous work, we went a step further to develop a biologically afunctional (i.e., bio-orthogonal by design) RNA origami able to fold at constant temperature (37 °C) after an initial denaturation step,\textsuperscript{29} which lights-up when folded to its target configuration. Seven RNA staple strands promoted the folding of a RNA bio-orthogonal synthetic De Bruijn scaffold sequence (DBS) that does not contain genetic information, restriction enzyme sites and reduces ambiguity in the addressability.\textsuperscript{30} We demonstrated the possibility to combine scaffold bio-orthogonality, physiologically compatible folding, and assembly monitoring using a new RNA-based reporter system. The folding was monitored using our new split Broccoli aptamer system.\textsuperscript{29}

Motivated by our previous study, and with the ultimate goal of enabling RNA origami expression in living cells, here we demonstrate a full isothermal protocol for scaffolded RNA origami assembly via cotranscriptional folding. We maintain the nanoribbon design simplicity and the scaffold nucleotide composition, while changing each staple in order to guarantee a reasonable yield of the desired transcripts and a low aberrant products synthesis during the in vitro transcription by T7 RNA polymerase. The RNA origami is cotranscriptionally folded into a nanoribbon shape at 37 °C: in detail, during the scaffolded RNA assembly, scaffold, and staple strands are in vitro transcribed and folded in a one-pot reaction. The RNA origami assembly is verified by gel assay and well characterized by AFM. The RNA nanostructures self-assembly is also successfully confirmed and selectively detected using our split Broccoli aptamer system: the transcription mix is analyzed by in-gel imaging and the tagged RNA origami shows a clear fluorescent band. Finally, the assembled RNA origami nanoribbon is visualized and simulated at equilibrium using the oxRNA coarse-grained model.

The in vitro transcribed and folded RNA origami described here can be compatible with expression in bacterial cells, and its self-assembly can be monitored with a protein-free fluorescence detection system using an in-gel imaging assay as a rapid and specific prescreening method.

\section*{METHODS}

\textbf{Materials and Reagents.} (SZ)-5’-[(3,5-Difluoro-4-hydroxyphenyl)methylene]-3,5-dihydro-2-methyl-3-(2,2,2-trifluoroethyl)-4H-imidazol-4-one (DFHBI-1T) was purchased from Tocris Biotechnique. A DFHBI-1T stock solution (20 mM) was prepared in DMSO, stored in the dark at −20 °C and used within 2 weeks. All RNA oligonucleotides were purchased from Eurogentec and resuspended from Ultrapure distilled water to give stock solutions of 100 μM and stored at −80 °C. All DNA oligonucleotides and gBlocks Gene Fragment were purchased from IDT. The DNA oligonucleotides were resuspended in Ultrapure distilled water to give stock solutions of 100 μM and stored at −20 °C. The gBlocks Gene Fragment was resuspended at a final concentration of 10 ng μL\textsuperscript{−1} and stored at −20 °C. EDTA 0.5 M pH 8.0, Ultra Pure 10× TBE buffer, Ultra Pure 1 M Tris-HCl pH 7.5, Ultra Pure distilled water, 5 M NaCl (0.2 μm filtered), SYBR Gold, dithiothreitol molecular biology grade, GlycoBlue Coprecipitant (15 mg/mL), inorganic pyrophosphatase, NTP (100 mM each), 6% Novex TBE gel, 10% Novex TBE gel and 10% TBE-Urea gel were purchased from Thermo Fisher Scientific. Spermidine BioUltra, HEPES 1 M pH 7 Bioreagent, KCl 1 M BioUltra, MgCl\textsubscript{2}, 1 M BioUltra, 3-aminopropyltriethoxysilane (APTES), and agarose were purchased from Sigma-Aldrich.

\textbf{Scaffold and Staples Design.} The synthetic RNA scaffold and RNA staples were generated with the computer code presented by Kozya et al.\textsuperscript{30} The RNA origami ribbon (Figure S1) has been designed as previously described.\textsuperscript{25} In the present study, RNA staple strands were changed (r1, n1, r2, l2, f, and s1) or elongated (s2) at the S′ end of few bases (1 up to 3 bases) in order to start with -GG or -GA (first and second nucleotides of the transcribed region) necessary for an efficient T7 transcription yield. Furthermore, all sequences were checked to reduce aberrant products synthesized during the transcription reaction. The RNA staple strands sequences and the scaffold sequence were placed downstream from the T7 promoter (′TTCTAATACGACTCACTATA-′3′). The dsDNA templates for the transcription were two annealed oligonucleotides (for staple strands transcription) or a gBlocks Gene Fragment sequence (for scaffold transcription): all the dsDNA templates carried the T7 promoter sequence and the template to be transcribed (Table S1).

\textbf{OxRNA Simulation.} We simulated the RNA ribbon/split aptamer system using a nucleotide-level coarse-grained RNA model, oxRNA\textsuperscript{31,32} In particular, we investigated thermodynamic stability of a preassembled nanoribbon in isothermal conditions over a period of time.

In order to simulate the system, the caDNAno design file\textsuperscript{33} were used in combination with caDNAno interface scripts (available with oxRNA software) to generate the initial topology of the simulated system. Preceding the simulation, the system was first relaxed, and the interaction type changed to RNA. To form initial configuration mutual traps were used according to the software documentation. Briefly, a harmonic force was introduced to pull selected particles from one strand of split aptamer toward reference particles on the other strand. This process used low stiffness parameter (0.1) until a
predefined equilibrium distance of the trap was reached, in this case, 1.5 simulation units of length, corresponding roughly to the hydrogen bonding potential. A short simulation was run, and after the equilibrium was reached and the split aptamer structure formed, the mutual traps were removed.

The system was then simulated over a longer time period and the trajectory file recorded. Molecular Dynamics (MD) was the selected simulation algorithm and the simulation was run for $2 \times 10^7$ simulation steps which, in physical units, corresponds to 61.2 $\mu$s. The temperature of the system was set to 310 K ($\sim 37^\circ C$). The John thermostat, which is an Andersen-like thermostat, was used since it is the optimal thermostat that emulates Brownian dynamics.34

**RNA Scaffold Synthesis and Purification.** Double-stranded gBlocks Gene Fragment containing T7 promoter (gBlocks DBS scaffold, Table S1) was amplified using Phusion DNA polymerase (NEB) and DBS forward/DBS reverse primers (Table S1), as previously described. Briefly, an initial denaturation at 98 °C for 30 s was followed by 15 cycles of denaturation at 98 °C for 10 s, annealing at 60 °C for 20 s and extension at 72 °C for 15 s. Finally, an additional extension was achieved for 5 min at 72 °C. The PCR product was purified using Monarch PCR & DNA Cleanup kit (NEB) and the DNA concentration was measured on a NanoDrop spectrophotometer.

The size of purified ampiclon was evaluated on 1.5% agarose gel in TBE for 1 h 40 min at 110 V: the gel was pre-stained with Nacy-S20 and visualized under UV illumination. The low molecular weight DNA ladder (NEB) was used as molecular weight marker.

The purified template was transcribed in vitro at 37 °C for 1 h and 30 min using Amplicscribe T7-Flash Transcription kit (Epicenter). After DNase treatment at 37 °C for 15 min, the RNA transcript was purified using RNA Clean & Concentrator (Zymo Research), quantified using a NanoDrop spectrophotometer and used as scaffold sequence for the RNA origami assembly reaction.

Alternatively, the amplified and purified scaffold template was used for cotranscriptional folding.

**Conventional Folding of RNA Origami Nanoribbon.** The RNA origami nanoribbon was synthesized using the conventional protocol.29 The transcribed and purified RNA scaffold was mixed with an excess of RNA staple strands in 10 mM MgCl$_2$, 20 mM Tris-HCl pH 7.5, 1 mM EDTA pH 8.0.27 After an initial thermal denaturation step at 75 °C for 1 min and a snap cooling ($-1^\circ C/0.42 s$), the mixture was subjected to a folding step at 37 °C for 20 min. Folding was performed with a SensoQuest Labcycler GeneFlow thermocycler. Samples (5 μL of a 50 μL reaction mixture) were run on 6% Novex TBE gel in 1× TBE buffer at 100 V for 40 min at low temperature (below 10 °C). After staining with SYBR Gold in 1× TBE for 5 min, the gels were visualized using Typhoon laser scanner and ImageQuant TL software (normal sensitivity; GE Healthcare Life Sciences). The low range ssRNA ladder (NEB) and ZR small-RNA ladder (Cambridge Bioscience) were used as molecular weight marker.

**Double-Stranded DNA Template Annealing, Purification, and Transcription.** RNA staples to be transcribed are less than 100 bases, ranging from 26 to 51 nucleotides. For this reason, DNA templates for in vitro transcription were obtained from two annealed single-stranded DNA oligonucleotides for each staple strand sequence: complementary sense and antisense strands (RP-HPLC or PAGE purified) containing the T7 promoter sequence and the RNA sequence to be transcribed were annealed by incubating equimolar concentrations (5 μM) of forward and reverse strand in TE supplemented with 12.5 mM MgCl$_2$ (r1, r2, l1, and l2) or with 50 mM NaCl (s1, s2, and f). The samples were heated at 95 °C for 2 min and slowly cooled down at 25 °C (72 cycles, 38 s/cycle, $-1^\circ C$/cycle): all annealing processes were performed with a Biometra TRIO Analistik jena thermocycler. Samples were loaded and run on 10% Novex TBE gel in 1× Tris borate EDTA (TBE) buffer at 200 V for 45 min. After staining with SYBR Gold in 1× TBE for 5 min, the gels were visualized using Typhoon laser scanner and Image Quant TL software (normal sensitivity; GE Healthcare Life Sciences). The low molecular weight DNA ladder (NEB) was used as molecular weight marker. Annealed dsDNA were purified from polyacrylamide gels by the “crush and soak” method, as previously described35 with some modifications. The bands of interest were cut out and the gel slices were transferred to a 0.5 mL tube that was pierced with a 20-G needle and placed inside a 1.5 mL microcentrifuge tube. The tube was centrifuged at 20,000g for 3 min to force the gel through the needle hole. 0.67 mL of DNA soaking buffer (0.3 M NaCl, 10 mM Tris-HCl pH 7.5, 0.97 mM EDTA) were added to the recovered gel and incubated overnight with agitation at room temperature. Gel and soaking buffer were purified from gel debris using a Freeze’n Squeeze DNA gel extraction spin column (Bio-Rad) that was centrifuged at 20,000g for 3 min at 4 °C to recover the soaking mixture. The eluted DNA was transferred to a fresh microcentrifuge tube and precipitated at least 1 h at $-20^\circ C$ using 0.68 mL isopropanol supplemented with 1 μL GlycoBlue Coprecipitant. After centrifugation at 4 °C (30 min at 20,000g), the pellet was washed in 0.75 mL of 80% ice-cold ethanol, air-dried, and resuspended in 12 μL of 10 mM Tris-HCl pH 7.5. DNA samples were run on 10% Novex TBE gel and visualized as described above. Purified and unpurified samples were measured using NanoDrop One/OneC spectrophotometer (Thermo Scientific).

Each purified or unpurified dsDNA template encoding the RNA staples strands were used for RNA synthesis. Transcription reactions were performed in 20 mM Tris-HCl pH 7.6, 10 mM MgCl$_2$, 2.5 mM of each rNTPs, 4 mM DTT, 2 mM spermidine, 2 units/μL T7 RNA polymerase (NEB). The transcription mix was incubated at 37 °C for 3 h and treated with 1 μL DNase RNase free (NEB) at 37 °C for 15 min. After dilution, samples were run on 10% Novex TBE gel in 1× TBE buffer at 200 V for 45 min. After staining with SYBR Gold in 1× TBE for 5 min, the gels were visualized using Typhoon laser scanner and ImageQuant TL software (normal sensitivity; GE Healthcare Life Sciences). The low range ssRNA ladder (NEB) and ZR small-RNA ladder (Cambridge Bioscience) were used as molecular weight marker.

**Cotranscriptional Folding of RNA Origami Nanoribbon.** The concentrations of each dsDNA template encoding the RNA staples strands and scaffold were measured using NanoDrop One/OneC spectrophotometer (average concentration values were calculated from 3 measurements for each sample). Different molar concentrations of DNA templates were used for RNA synthesis and RNA origami folding during the T7 transcription: 5 nM gBlock Gene Fragment (scaffold), 5 nM of each s1, s2, and f dsDNA, 10 nM of each r1 and l2 dsDNA, 20 nM 11 dsDNA, 30 nM r2 dsDNA. Transcription reaction was performed in 20 mM Tris-HCl pH 7.6, 10 mM MgCl$_2$, 2.5 mM of each rNTPs, 4 mM DTT, 2 mM spermidine, 2 units/μL T7 RNA polymerase (NEB). The
transcription mix was incubated at 37 °C for 3 h and treated with 1 μL DNase RNase free (NEB) at 37 °C for 15 min.

**Cotranscribed RNA Origami Native PAGE and In-Gel Imaging.** After dilution, samples were run on 6% Novex TBE gel in 1x TBE buffer at 100 V for 40 min at low temperature (below 10 °C). After staining with SYBR Gold in 1x TBE for 5 min, the gels were visualized using Typhoon laser scanner and ImageQuant TL software (normal sensitivity; GE Healthcare Life Sciences). The low range ssRNA ladder (NEB) was used as molecular weight marker.

To confirm the RNA origami self-assembly by incorporation of Split Broccoli aptamer system, in-gel imaging with fluorophore DFHBI-1T was performed with some modifications. Briefly, RNA origami sample, partially folded samples and Broccoli aptamer (prepared as previously described) were loaded in the polyacrylamide gels. The gels were washed three times for 5 min in RNase free water and then stained for 20–25 min in aptamer buffer containing 1.26 μM DFHBI-1T, 40 mM HEPES pH 7.4, 100 mM KCl, 1 mM MgCl₂. The gels were imaged using Typhoon laser scanner (excitation 488 nm, emission 532 nm): bands were analyzed using ImageQuant TL software. Then, the gels were washed three times with Ultra Pure distilled water, stained with SYBR Gold in 1x TBE for 8 min, and visualized using Typhoon laser scanner. The low range ssRNA ladder (NEB) was used as molecular weight marker.

**Atomic Force Microscope (AFM) Imaging.** All the experiments were conducted on a commercial AFM Cypher ES (Asylum Research, Oxford Instruments, Santa Barbara, CA) and with the tip and cantilever fully immersed into the liquid. The vertical oscillation of the tip was controlled by photothermal excitation (Blue Drive) and the experiments were conducted at 25.0 ± 0.1 °C. All measurements were conducted in Amplitude Modulation (AM-AFM) mode using SCANASYST-Fluid+ (Bruker, Camarillo, CA) cantilever and with a set point ratio between the free amplitude and imaging amplitude of ~0.8.

Freshly cleaved mica was passivated for 5 min with 10 μL of 0.1% APTES in water to ensure the adhesion of the negative charged RNA origami structures on the negative mica surface. After three washing steps with 10 mM MgCl₂, 20 mM Tris-HCl pH 7.5, 1 mM EDTA pH 8.0, the transcription reactions were diluted 1:100 in the same buffer, added (2 μL) to the passivated mica surface, allowed to adsorb for 5 min in a chamber and imaged immediately. When purified RNA origami samples were imaged, 8 μL of sample were added to the passivated mica surface and allowed to adsorb for 5 min in a chamber as above.

All the images were corrected for tilt (line or plane flattening) and lightly low-pass filtered to remove grainy noise using the WSXm software (Nanotec Electronica, Madrid, Spain).37

### RESULTS AND DISCUSSION

**Staple Strands and RNA Origami Design.** Recently, a small functional scaffolded RNA origami made from just few sequences demonstrated gene knockdown functionality. It was a RNA nanostructure synthesized from a short mRNA fragment scaffold and nine synthetic RNA strands.28

Here, our aim is to understand if scaffolded RNA origami, decorated with a split Broccoli aptamer, could be made by in vitro cotranscriptional folding. More precisely, we investigate the cotranscriptional folding of both the scaffold itself and its staples, which has not yet been explored. As a proof of principle, we decided to design a simple RNA origami that involves seven staple strands and a scaffold sequence. Our approach uses all the components as products of the transcription machinery rather than of chemical synthesis. Moreover RNA self-assembly occurs during the transcription reaction. With the ultimate future goal to construct a “bio-circuit board” to spatially organize biomolecules and thus regulate metabolic activity, a simple ribbon nanostructure has been chosen and designed. The design includes the precise organization of a split aptamer system that can allow to monitor the assembly inside a living cell without using fluorescent reporter proteins and thus reducing the energetic burden to the cells.

Recently, we designed a 212 nt biologically inert (i.e., bio-orthogonal) and uniquely addressable De Bruijn scaffold sequence (DBS) characterized by lack of genetic information, restriction enzyme sites, and reduced ambiguity in the addressability: the bio-orthogonality of the RNA scaffold designed in silico was demonstrated in E. coli cells.39

Here, we used our previous optimized RNA scaffold and a new set of seven staple strands, starting with -GG or -GA nucleotides at the 5’ end (Figure S1 and Table S1) in order to promote a reasonable transcriptional yield and allow an efficient control of the 5’ sequence content.38 In detail, transcription reactions of each RNA sequence were carried out with T7 RNA polymerase and synthetic DNA containing the T7 promoter: the above-mentioned 5’ end starting nucleotides can guarantee efficient transcription and efficient control of the 5’ sequence content during the T7 polymerase reaction.38

Furthermore, although T7 RNA polymerase is a highly specific enzyme, the desired short RNA is usually accompanied by undesired products that are longer or shorter than the expected transcript.39–44 Besides its higher DNA affinity, T7 RNA polymerase is able to synthesize RNA from single or double-stranded RNA template.42,43 It has been demonstrated that RNA extension can occur if the 3’ end has a self-complementarity (trans mechanism) or folds back on itself in cis to form extendible duplexes.40,42,43 Next generation RNA-Seq analysis has demonstrated that primer extension occurs predominantly via a cis self-primer mechanism and strongly depends on pairing A at the 3’ end with the U, 9 bases upstream;56 as a result, partially double-stranded RNA byproducts are synthesized. In order to reduce the amount of longer undesired RNA extension products catalyzed by T7 RNA polymerase, all staple strands sequences were examined to take account of the 3’ end self-complementarity and pairing between the 3’ end adenine and uracil, 9 bases upstream; the analysis revealed that all RNA sequences were lacking of these distinct 3’ end characteristc.

In addition, false transcriptions are mainly observed when the correct product is free of stable secondary structures at the 3’ end: the aberrant transcripts are longer than the coded RNA.40 It has been reported that improved transcription of the correct length RNA can be obtained by adding hairpins to the 3’ and 5’ ends; these hairpins decrease the transcription of incorrect-length products, but still a series of bands can remain.55,46 In our work, all the sequences were optimized to weaken secondary structures and avoid hairpin formation.30

**DNA Template Preparation and Transcription.** The scaffold sequence and each of the staple strands sequences were placed downstream from the T7 RNA polymerase promoter and the linear double-stranded DNA templates were
characterized by blunt ends in order to reduce the synthesis of spurious transcripts.

Scaffold DNA template was purified from an amplified gBlocks Gene Fragment (Figure S2). As staple strands to be transcribed were shorter (ranging from 26 to 51 bases), DNA templates were obtained from an annealing step of two RP-HPLC or PAGE purified oligonucleotides.

Annealed DNA templates corresponding to staple strands showed multiple bands (Figure S3) presumably due also to the presence of byproducts in addition to the desired chemically synthesized oligonucleotide (RP-HPLC and PAGE purification usually can provide a purity level up to 85% and around 85—90%, respectively).

To increase the purity, annealed DNA templates were purified from polyacrylamide gel by the “crush and soak” method using GlycoBlue as coprecipitant to facilitate the DNA recovery and reach a concentration compatible with the subsequent transcription reaction avoiding any mixture dilution. All samples were run and checked on native PAGE (Figure S4).

The use of GlycoBlue coprecipitant at final concentrations (22 μg/mL) below the recommended value (50—150 μg/mL) did not allow an accurate DNA quantification due to the blue dye absorbance and the low A260/A230 ratio value (0.8). Furthermore, there were no differences in the transcription products when purified and unpurified dsDNA samples were used as template. For these reasons, we decided to in vitro transcribe unpurified DNA templates.

Each RNA sequence was synthesized during T7 polymerase in vitro transcription from dsDNA templates. T7 polymerase is usually produced by expressing the RNA polymerase gene of bacteriophage T7 in E. coli cells and is used to study transcription mechanism in vitro or as a convenient approach to synthesize RNA strands. Once the enzyme is isolated from E. coli cultures for in vitro reactions, the TRIS working buffer composition is optimized considering the transcriptional activity of the enzyme in vitro and the specific application. The in vitro transcription system is affected by stimulatory natural polyamines, particularly spermidine, and requires freshly prepared dithiothreitol (DTT). The polyamine additive can activate T7 polymerase reaction and increases the fidelity of RNA transcripts: the activation effect is concentration-dependent in a wide range.  The transcription TRIS buffer requires also magnesium ions and usually free ions concentration is adjusted according to the nucleotide concentration. Finally, in the enzymatic reaction, RNA polymerase produces inorganic pyrophosphate (PPI) as byproduct: PPI released by the reaction binds magnesium ions to form Mg2PPI, which can influence the RNA synthesis. Inorganic pyrophosphatase catalyzes the hydrolysis of the pyrophosphate to two orthophosphates and can be used to improve the yield of the in vitro transcription reaction.

Considering all the above-mentioned factors (magnesium ions, spermidine, DTT, and inorganic pyrophosphatase), the RNA origami folding buffer composition, and in order to improve the transcription of each dsDNA template, different DTT and spermidine concentrations were tested with or without inorganic pyrophosphatase. In detail, the initial folding buffer (10 mM MgCl2, 20 mM Tris-HCl pH 7.5, 1 mM EDTA  pH 8.0) was supplemented with 2.5 mM of each rNTPs, while EDTA was removed to avoid inhibition of the DNase treatment performed at the end of the transcription reaction. Then the transcription buffer was supplemented with (i) 10 mM DTT and 2 mM spermidine; (ii) 10 mM DTT, 2 mM spermidine and 0.2 U inorganic pyrophosphatase; (iii) 4 mM DTT and 2 mM spermidine; or (iv) 4 mM DTT and 0.2 U inorganic pyrophosphatase.

In vitro transcribed RNA sequences were analyzed by denaturing PAGE. The transcription of each staple strand dsDNA template in 4 mM DTT and 2 mM spermidine showed a distinct main band corresponding to the full-length transcript (Figure S5). For this reason we decided to use the same concentrations for scaffold dsDNA template transcription: the transcribed scaffold resulted in a main band of the expected size (Figure S6). In addition to the full-length transcript, the synthesis of a pattern of products with a higher molecular weight was also observed as previously described. Triana-Alonso et al. demonstrated the synthesis of longer transcripts.
as a secondary process following an induction time that is different for every template: a minimal concentration of transcript is required for aberrant transcripts that accumulated at the end of long incubations as a result of a late event during the transcription. As RNA staple strands were designed to not support RNA-primed RNA extension, we hypothesized that, after the production of a certain amount of transcript, T7 polymerase accepted template RNA sequences that were not folded in 3′ end stable secondary structures. We then assumed that during the cotranscriptional folding, duplex formation related to the RNA origami folding can sequester the staple strands 3′ end reducing RNA-templated extension and undesired byproducts.

Finally, abortive synthesis was observed in T7 RNA polymerase transcription as a fundamental and major feature of early stages: several short transcripts were produced (maximum length of 12 nt) due to a kinetic competition between dissociation of the enzyme-DNA-RNA complex and incorporation of successive nucleotides. During the scaffold and staple strands transcription, a low amount of abortive products (Figure S5 and Figure S6) were produced without influencing the transcription efficiency. Previous studies showed that both abortive products and intermediate products were observed as an unavoidable result of the in vitro transcription.41,49

OxRNA Simulation. The simulated RNA ribbon/split aptamer system stayed at equilibrium and the split aptamer structure was preserved at 37 °C. The two bulge loops (3 nt and 2 nt-long, respectively) were visible in the aptamer structure (Figure 1, red arrows). The hybridization between split aptamer strands and the origami scaffold allowed for a degree of flexibility between the aptamer and the ribbon. Interestingly, the simulation suggested that the internal loop (located between the red arrows) formed a double-helical structure despite the mismatched nucleotides. However, the Gibbs’s free energy of that interaction was higher than a typical complementary sequence of similar length and thus less stable.

RNA Origami Cotranscriptional Folding and AFM Imaging. Self-assembly at physiologically compatible temperatures can expand and improve the RNA nanotechnology applications under intracellular conditions. To date, the ability to design RNA nanostructures that cotranscriptionally fold into the target shape has been demonstrated using different approaches.19,22,50,51

As a proof of principle, here we demonstrate the cotranscriptional folding of a genetically encoded scaffolded RNA origami using our previously designed bio-orthogonal DBS scaffold. All double-stranded DNA templates encoding scaffold and seven staple strands were mixed in a one-pot reaction and used to transcribe RNA sequences able to fold into the desired nanostructure. The cotranscriptional folding was performed at 37 °C for 3 h using T7 RNA polymerase in the transcriptional buffer containing 4 mM DTT and 2 mM spermidine. Furthermore, two partially folded RNA scaffold samples were transcribed from dsDNA template encoding respectively: (i) scaffold, Staple s1 and s2; (ii) Staple s1, s2, 11, and r1. These samples were considered to demonstrate not only their different electrophoretic migration patterns, but also their different fluorescence band intensities in the following in-gel imaging experiments.

After a DNase treatment to digest template dsDNA, the unpurified transcription mixtures were directly analyzed by nondenaturing polyacrylamide gel electrophoresis: the gel image showed a distinct band around the expected size (black arrow in Figure 2) with a different migration distance compared to that of partially folded products, suggesting the correct folding. Bands corresponding to scaffold and partially folded products were almost not present in the RNA origami sample. The origami yield percentage was estimated from native polyacrylamide gel25 and was approximately 20% of all in vitro transcribed RNA.

In order to compare cotranscriptional and conventional prepared RNA nanostructures, RNA origami was also synthesized from transcribed and purified RNA scaffold and RNA staple strands, and used as control. After an initial thermal denaturation, the sample solution was subjected to a folding step at 37 °C for 20 min and verified by gel electrophoresis (Figure S7) as previously described.29

The cotranscriptional folded nanostructure band showed a slight difference in the migration distance (Figure 2) when compared to conventional assembled nanostructure (Figure S7), as previously observed.19 the electrophoretic migration of unpurified transcribed samples was influenced by the different and more complex reaction mixture composition.52

Finally, we found a higher band closed to the well (Figure 2): as a result of both thermal ramp or cotranscriptional folding, higher bands can appear19,22,28,53 and can correspond to aggregates,52 spurious assemblies53 or undesired kinetically trapped products.52

In detail, abortive and elongated RNA products of incorrect length are an unavoidable result of the in vitro transcription41,49 and they can contribute to the formation of spurious assemblies.53 Stewart et al.53 showed that despite the presence of these transcription byproducts able to generate unknown folding, the one-pot method yields the desired assemblies. The latter authors showed examples of gel electrophoresis images with a significant amount of abortive and elongated products. Furthermore, AFM images of tubular assemblies and flat lattices revealed a “noisy” environment. In the perspective of future in vivo isothermal folding, it should be noted that aberrant products produced in vitro by T7 RNA polymerase are not commonly synthesized inside living cells.40
The unpurified RNA origami sample was characterized by AFM immediately after the cotranscriptional folding. Samples were diluted and deposited on a passivated mica surface using a 0.1% APTES solution in water instead of Mg²⁺. Indeed, it has been suggested that the different phosphate groups’ orientation in dsRNA compared to dsDNA can be a possible reason for the difficult dsRNA adsorption on mica using magnesium ions.⁵⁴ The estimated RNA nanostructures’ dimensions were approximately 27 nm × 5 nm, as the A-form RNA helix revealed a rise per base pair of 0.28 nm.⁵⁴ AFM images confirmed the correct folding of the cotranscribed RNA nanostructures with average lengths of 27.2 ± 3.5 nm and 8.4 ± 1.8 nm (Figure 3 and Figure S8), calculated considering shape closed and consistent with the design, as previously reported.⁵⁵ Cotranscribed RNA origami images were comparable to that of purified conventional RNA origami sample (Figure 4 and Figure S8) previously well characterized,²⁹ indicating that the RNA scaffold and staple strands folded into the desired shape also during the transcription reaction. RNA nanostructures showed a double-stranded RNA mean periodicity of 3.1 ± 0.3 nm (Figure 3, right) or as a thin protrusion (Figure 4 and Figure S9) depending on the orientation during the deposition on the mica surface: the aptamer can either stack with the left or the right duplex, and also be out of plane and not necessarily always in the position pictured in Figure 1. It has been already noted that not all the modifications of the hexagonal mini-lattices with aptamers are well visible by AFM imaging due to the extension orientation with respect to the scanning direction.⁵⁵

Interactions between RNA nanostructures were also observed in AFM images (Figure S9), presumably due to interaction between left (5’-UGUUUAUACAG-3’) and right (5’-CCGGUCCGGC-3’) ssRNA scaffold regions, as previously noted.²⁹

Furthermore, agglomerates were imaged during AFM scanning (Figure S9) and this was consistent with the higher band visible near the gel well (Figure 2). As we discussed above, the formation of undesired assembly can be promoted by runoff elongated RNA molecules produced during the transcription process, as recently described.⁵³ Finally, smaller structures were imaged: it should be noted that RNA nanostructures can be damaged during deposition on mica and by AFM tip, resulting in artificial structure. Previous studies underlined that structural heterogeneity can be observed as the result of an unavoidable fragmentation during sample deposition and imaging.²⁵,⁵⁶

**Light-up RNA Origami and In-Gel Imaging.** Light-up RNA aptamers have been selected to induce a fluorescence emission upon binding specific small molecules allowing a protein-free RNA tagging.⁵⁷ Since their development, fluorescent aptamers have been used in vitro and in vivo as a valid alternative to other imaging strategies (e.g., fluorescent in situ hybridization and molecular beacons⁵⁷,⁵⁸). Malachite green, Mango, Spinach, and Broccoli aptamers have been inserted in the design of different RNA nanostructures to successfully demonstrate the self-assembly and folding through their functional activation and fluorescence emission.¹⁹,²⁴,²⁹,⁵⁵ Considering characteristics such as the short size, the robust folding under physiological conditions, and the specificity for a nontoxic and cell permeable dye, Broccoli aptamer⁵⁹ was recently chosen to functionalize an RNA origami nano-ribbon.²⁹ The Broccoli aptamer was divided into two nonfunctional halves each of which was elongated in the 5’ or 3’ end with two sequences complementary to the bio-orthogonal RNA scaffold. When the RNA origami self-

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**Figure 3.** High-resolution AFM images of cotranscriptional folded RNA origami. The images were taken in solution immediately after cotranscription. The white arrows show the dsRNA helical pitch (3.2 ± 0.3 nm⁵⁴). Scale bar: 10 nm.

**Figure 4.** High-resolution AFM image of conventional folded RNA origami assembled from transcribed and purified RNA scaffold and RNA staple strands. Scale bar: 20 nm.
nanostructures are expressed in bacterial cells. A method to monitor and check if genetically encoded aptamer can act as a prescreening strategy to control metabolic processes in engineered bacterial cells. Toward this perspective, we present a cotranscriptionally folded RNA origami as a further step along this direction.

In the context of in vivo origami expression to precisely control biomolecular processes, our aim is not focused on an efficient way to produce complex RNA nanostructures in large scale. Indeed, in vivo expression of RNA assembly should not impose a cost to the host cell consuming excessive resources for expression and interfering with endogenous molecules. Nonetheless, several factors can be considered and evaluated when a high-yield production of more complex nanostructures is investigated during in vitro transcription. For example the transcription incubation time, [Mg$^{2+}$], [Mn$^{2+}$], [DTT], [NTP], and Tritox-X100 addition represent few parameters that can be tested.

We have demonstrated the cotranscriptional folding of a bio-orthogonal scaffolding RNA origami in a one-pot reaction, revealing that the self-assembly of scaffold and staple strands can occur also in the transcription reaction mixture at 37 °C from double-stranded templates. In alternative to other strategies previously reported (i.e., single-stranded origami and strands self-assembly), the scaffolded RNA origami technique can be successfully used to design and synthesize a desired nanostructure at constant, physiologically compatible temperature, overcoming the use of synthetic purified RNA sequences not possible in living cells.

Split Broccoli aptamer functionalization was introduced into the RNA origami design allowing a simple and specific screening of the well folded nanostructures by in-gel imaging using a cell permeable and compatible dye. This approach can further suggest and confirm the use of our split light-up aptamer as a reporter system to monitor the folding, avoiding the use of fluorescent proteins. Furthermore, the fluorogenic aptamer can be replaced with other functional sequences required for different purposes.

In conclusion, our results represent a further step toward in vivo self-assembly of nucleic acid nanostructures, and expand the design strategies used to synthesize nanostructures tailored to specific applications. Future works can be focused on demonstrating that more complex nanostructures self-assemble by the same cotranscriptional method.
In a wider picture, genetically encoded scaffolded RNA origami could represent a pathway to construct a nanostructure able to direct specific cell metabolic pathways through an orthogonal spatial and post transcriptional control.

■ ASSOCIATED CONTENT

◆ Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssynbio.0c00009.

Details about the sequences, design of the cotranscriptional folded RNA origami nanoribbon, gel images, and AFM images (PDF)

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The manuscript was written by N.K. and E.T. Sequence design and oxRNA simulations were performed by J.W.K. B.S.E. conducted sequence analysis. AFM imaging was conducted by L.P.; all images were discussed with E.T. All other experiments were conceived by N.K. and E.T. All authors have given approval to the final version of the manuscript.

Notes
The authors declare no competing financial interest.

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