Determining in vivo RNA structures with conformation capture using clickable psoralen

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RNA structural flexibility underlies fundamental biological processes, including splicing, translation, and the lifecycle of RNA viruses. However, there are no experimental or computational methods to explore the multiple conformations adopted by RNAs in vivo. We developed Crosslinking Of Matched RNAs And Deep Sequencing (COMRADES) for in-depth conformation capture of RNA and retrieval of RNA structural ensembles. Using COMRADES, we determined the architecture of the Zika virus RNA genome inside cells, and revealed multiple interactions with human non-coding RNAs.

Structure probing techniques have been proven valuable for studying cellular and viral RNA structure. Inside cells however, RNA is engaged in multiple processes including splicing, translation, and regulation by RNAi, suggesting high structural plasticity. Recent methods have utilized proximity ligation to reveal RNA base-pairing within cells. Nevertheless, the ability to assess the structural plasticity of RNA and report the ensemble of coexisting conformations has not been reached, due to insufficient probing depth and lack of appropriate computational algorithms. We developed a method—Crosslinking Of Matched RNAs And Deep Sequencing (COMRADES)—that couples an improved in vivo probing of RNA base-pairing with selective RNA capturing. We additionally established an algorithm for assessing the structural complexity of RNA inside cells (Fig 1a).

COMRADES utilizes a cell-permeable, azide modified, psoralen derivative (Psoralen-triethylene glycol azide) to facilitate coupling of two effective affinity capturing steps, while overcoming the limited cell permeability of biotin labelled psoralen (Supplementary Fig. 1a). The azide group does not affect the psoralen
crosslinking properties (Supplementary Fig. 1a). Following in vivo crosslinking, an RNA of interest is selectively captured, reaching enrichment level of nearly 1000-fold (Fig. 1b, Supplementary Fig. 1b-c). The RNA is then fragmented, and a copper free click-chemistry reaction links a biotin moiety to in vivo crosslinked regions, enabling a second streptavidin-based affinity selection of crosslinked regions (Fig. 1c, Supplementary Fig. 1d). The resulting RNA is split: One half is proximity ligated to create RNA chimeras, following reversal of the crosslink to enable high-throughput sequencing and assessment of the base-pairing (Fig. 1d, Supplementary Fig. 1e). The other half is used as a control, in which reversal of the crosslink precedes the proximity ligation. COMRADES and control samples contain essentially identical RNA composition, ensuring accurate assessment of artificial chimeric reads originated from random ligation or reverse transcription errors. COMRADES’s dual enrichment substantially increases structure probing depth of selected RNA, thus enabling an unbiased and global view of coexisting conformations. COMRADES yields high levels of ligated chimeric reads (8%), whereas these are kept 4 fold lower in the controls, or in a non-crosslinked samples (Fig. 1e). We successfully reported on the known ribosomal RNA structure with high sensitivity (Supplementary Fig. 2), while spurious interactions between cytoplasmic and mitochondrial ribosomal RNA subunits occurred at a very low level (Fig. 1f). The robustness of COMRADES is further demonstrated by its high reproducibility (Fig. 1g-h).

RNA viruses utilize RNA base-pairing to regulate various aspects of their life cycle. Inside the host cell however, the full-length architecture of RNA genomes and their interactions with the host transcriptome are largely unknown. We used COMRADES to determine RNA base-pairing along the 10.8 kilobases-long single-
stranded RNA genome of Zika virus (ZIKV) from the Flavivirus genus inside human cells. We identified 1.7 million non-redundant chimeric reads corresponding to the structure of the ZIKV genome (Fig. 1g-h, Supplementary Fig. 3a). This high probing coverage is valuable for analysing multiple coexisting conformations. Previous work mainly identified RNA structures in the untranslated regions (UTRs) of flaviviruses, while leaving 95% of the genome unexplored\textsuperscript{17–21}. COMRADES identified base-pairing along the entire genome and between the open reading frame (ORF) and the UTRs (Supplementary Fig. 3b, 4a). Nearly 80% of the identified interactions span a distance of less than 1,000 nucleotides (nt), implying local structure with a certain degree of three-dimensional compaction (Supplementary Fig. 3c). Both short- and long-range interactions were supported by reproducible, well-defined clusters of chimeric reads, ligated in 5'-3' and 3'-5' orientations (Supplementary Fig. 3a), and showed strong evidence of base-pairing when analyzed with the hybrid-min RNA-folding algorithm as compared to a shuffled-chimeras control (Wilcoxon test p-value < 0.0001). COMRADES therefore enables a deep and comprehensive analysis of RNA base-pairing.

During replication, the genome of flaviviruses undergoes a global conformational change mediated by a long-distance base-pairing between the 5' and 3' cyclization sequences\textsuperscript{17,18} (5' CS and 3' CS respectively), as well as the upstream and downstream of AUG regions (UAR and DAR respectively)\textsuperscript{19,20}. COMRADES detected extensive and highly specific base-pairing between the known cyclization elements, therefore demonstrating genome cyclization inside cells (Fig. 2a-c, Supplementary Fig. 4b). COMRADES further refined the nature of the base-pairing associated with genome cyclization, by identifying contact regions upstream of the 5'
UAR and downstream of the 3' UAR (Fig. 2a,c). We additionally detected an alternative 5' UTR conformation where stem-loops A and B (SLA and SLB respectively) are not formed but rather engaged in long-distance base-pairing with the downstream envelope coding sequence (Fig. 2d-f). COMRADES confirmed the existence of previously defined functional RNA pseudoknots including the dumbbell (DB) pseudoknot\textsuperscript{22}, the downstream of 5' cyclization sequence-pseudoknot (DCS-PK)\textsuperscript{23}, and the SL1 pseudoknot\textsuperscript{24} (Supplementary Fig 4c). Overall, COMRADES identified nearly all previously known flavivirus RNA structures and has further defined critical base-pairing involving the UTRs.

Our intra-viral RNA-RNA interaction map revealed the presence of multiple mutually exclusive RNA structures, where one region alternately base-pairs with several other regions. The averaged Shannon entropy per nucleotide was 5.9 bits, implying high folding plasticity (Supplementary Fig. 5a). We found a strong inverse correlation between the degree of experimental support for base paired regions and their entropy (Supplementary Fig. 4a, 5b-d). To explore the ensemble of alternative structures, we developed an algorithm to computationally fold ~1,000 nucleotide-long regions using randomly selected subsets of high-confidence mutually compatible folding constraints derived from the in vivo data. For each region, a set of 1,000 structures was generated. The validity of this approach is demonstrated by the clear correlation between the thermodynamic stability and the number of reads supporting each structure (Fig 2g, Supplementary Fig. 6). Nevertheless, the most thermodynamically favoured structures gained only moderate experimental support, implying the additional impact of the cellular environment on RNA folding\textsuperscript{3,4,16,25,26}.
We further computed the degree of similarity between all pairs of structures and applied multidimensional scaling to cluster structures based on their similarity. The presence of separated well defined clusters reflects the occurrence of alternative conformations (Fig. 2h, Supplementary Fig. 7a). As a control, we randomly shuffled the interacting RNA partners between the chimeric reads; the resulting shuffled structures clustered separately from the structures recovered by COMRADES (Supplementary fig. 8). A single ZIKV structure typically accounted for ~30% of the in vivo observed interactions, whereas a reduced set of 5 structures was sufficient to capture 80-90% of the in vivo data (Supplementary Fig. 7b). Our analysis suggests that the intracellular folding complexity of the ZIKV genome might be explained by postulating the coexistence of a small subset of alternative conformations.

Viral RNAs have an inherent capacity to form specific interactions through base-pairing with host RNAs\textsuperscript{27}, however little is known about the prevalence of such interactions. COMRADES revealed multiple interactions between the ZIKV genome and human small regulatory RNAs (Fig. 3a). We found site-specific interactions between the ZIKV ORF and the U1 small nuclear RNA (snRNA, Fig. 3b), plausibly affecting host splicing. We similarly detected highly site-specific interactions between the ZIKV ORF and tRNAs Glu\textsuperscript{116} and Gly\textsuperscript{34} (Fig. 3c). We identified several interactions between the ZIKV genome and human microRNAs, including miR-21, miR-19, miR-512, miR-515, and miR-1323 (Fig. 3d). While previous psoralen-based methods failed to effectively capture interactions involving miRNAs\textsuperscript{10–12}, COMRADES successfully discovered miRNA base-pairing (Fig. 3a,d). COMRADES indicated non-canonical base-pairing between the 5' CS of ZIKV and the seed region.
of miR-21 (Fig. 3e, adjusted p-value 1.0E⁻¹³). The miR-21 interaction with the ZIKV genome was further proved significant using an independent analysis pipeline (FDR 3.0E⁻²⁵, supplementary methods). In vitro synthesized miR-21 failed to bind the ZIKV 5' CS on its own, while pre-loading miR-21 onto purified Argonaute2 (Ago2) facilitated a strong and sequence specific interaction (Supplementary Fig. 9), supporting the involvement of Ago2 in this base-pairing. CRISPR/Cas9 deletion or antisense inhibition of miR-21 in human cells decreased the intracellular level of the ZIKV genome (Supplementary Fig. 10a-d) or the ZIKV envelope protein (Supplementary Fig. 10e-j). Abrogating the miR-21 binding ability of a ZIKV replicon through point mutations renders it insensitive to miR-21 antisense inhibition (Supplementary Fig. 10k), indicating that miR-21 acts through direct interaction with the 5' CS.

In summary, COMRADES revealed the highly dynamic nature of an RNA genome inside cells, and its ability to engage in base-pairing with multiple host regulatory RNAs. The involvement of the conserved 5' CS of ZIKV in genome cyclization, capsid translation and miR-21 binding further demonstrates the structural complexity of RNA genomes inside cells (Fig. 3f). The general applicability of COMRADES provides an opportunity to undertake a complete and unbiased analysis of the dynamic nature of RNA inside cells and can be utilized to investigate the structure and interacting partners of any cellular or foreign RNA in any species.
**a** In-vivo RNA crosslinking

- Virus-RNA pulldown
- Biotin-click
- Crosslinked-RNA pulldown
- Ligation
- Crosslink reversal

- Enriched
- Flowthrough
- Control

**b**

- Fluorescent units
- Molecular size, nt

- ZIKV marker

**c**

- Input
- Crosslink
- Enriched
- Flowthrough

**d**

- Randomised parallel RNA folding
- Structure scoring and clustering

**e**

- Chimeric reads, %

- COMRADES
- Control
- Non-crosslinked
- rRNA-RNA
- mtrRNA:mtrRNA
- mtrRNA:rRNA

**f**

- % of total chimeric reads

**g**

- Zika genome coordinate (1-10,807 nt)
- Zika genome coordinate (1-10,807 nt)

**h**

- Replicate 1
- Replicate 2
- Replicate 3

- Control 1
- Control 2
- Control 3

**Ziv et al., Fig. 1**
Ziv et al., Fig. 2
Ziv et al., Fig. 3
Figure legends

**Fig. 1:** COMRADES methodology. a, Experimental and computational workflows (upper and lower panels respectively). B: biotin; b, Virus-RNA pulldown, demonstrated by bioanalyzer profiles of ZIKV enriched RNA (upper panel) and input RNA (lower panel). c, Crosslinked-RNA pulldown, demonstrated by dot blot of biotin labelled RNA from COMRADES experiment. d, Crosslink reversal, demonstrated as described in (c). UVC: short-wavelength UV. e, % of chimeric reads in COMRADES and control libraries. Error bars represent standard deviation of 3 biological replicates. f, Probed interactions among cytoplasmic and mitochondrial ribosomal RNA subunits. rRNA: cytoplasmic ribosomal RNA; mtrRNA: mitochondrial ribosomal RNA. Error bars represent standard deviation of 3 biological replicates. g, Heat map of ZIKV RNA-RNA interactions. Each dot represents an interaction between the genomic coordinates on the x and y axes. Chimeras ligated in 5'-3' and 3'-5' orientations are plotted above and below the diagonal respectively. h, Zoom-in of a selected 1,500 nt region from (a).

**Fig. 2:** The genomic structure of ZIKV inside human cells. a, Heatmap of RNA-RNA interactions between cyclization elements. Rep: biological replicate; cont: control. b, Viewpoint histograms showing binding positions of the cyclization sequences along the ZIKV genome. Viewpoint regions are marked by dashed red lines. c, Probed interactions along the circular genome conformation. Colours representing numbers of non-redundant chimeric reads supporting each interaction. New: newly identified base-pairing. d, Heatmap of RNA-RNA interactions between the 5' UTR and the envelope coding region. e, Viewpoint histogram showing binding of nucleotides at position 2-56 along the ZIKV genome. f, Newly identified 5' UTR
structure. Colour code as described in (c). cHP: capsid-coding region hairpin element; DCS: downstream of 5’ cyclization sequence. g, Computationally predicted structures for nt 2288-3323. dG: folding energy. h, Clustering and prediction of alternative structures of nt 2288-3323. Colour code as described in (c).

Fig. 3: Host-virus RNA-RNA interactions. a, Human RNA species interacting with the ZIKV genome. Error bars represent standard deviation of 3 biological replicates. b-d, Site specific base-pairing between the ZIKV genome and snRNAs (b), tRNAs (c), and specific miRNAs (d) in COMRADES libraries and controls. e, COMRADES determined base-pairing between ZIKV and miR-21. f, A model of the ZIKV 5’ CS engaged in three separate functions. Ribosome and nascent polypeptide are marked in green.
Ziv et al., Supplementary Fig. 1
Ziv et al., Supplementary Fig. 3
Supplementary Fig. 4

**a**

- 5'UTR
- CDS
- 3'UTR

**b**

- 5' CS viewpoint: control
- 3' CS viewpoint: control

Chimeric reads count

- Nucleotide position

**c**

- SLB
- SLA
- cHP
- DCS-PK
- SL1-PK
- SL2
- DB-PK
- sHP
- 3' SL

- 5' CS
- 3' CS

Ziv et al., Supplementary Fig. 4
Shannon entropy

Ziv et al., Supplementary Fig. 5
Fragment 1: nt 108-1275
Fragment 2: nt 1276-2287
Fragment 3: nt 2288-3323
Fragment 4: nt 3324-4521
Fragment 5: nt 4522-5551
Fragment 6: nt 5552-6810
Fragment 7: nt 6811-7757
Fragment 8: nt 7758-8755
Fragment 9: nt 8756-9543
Fragment 10: nt 9544-10379

Ziv et al., Supplementary Fig. 6
**a** labeled miR-21 labeled ZIKV 5' CS

**b** wildtype
Kd = 127±41 nM

**c** mutant 1
Kd = 405±369 nM

**d** mutant 2
Kd = 1969±10312 nM

**e** perfect seed match
Kd = 24±4 nM

**f** No cHP loop
Kd = 78±16 nM

**g** wildtype long
Kd = 74±8 nM

**h** 8mer (seed match)
Kd = 21±1 nM

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**Ziv et al., Supplementary Fig. 9**
**Supplementary Fig. 10**

**a**

miR-21 expression

<table>
<thead>
<tr>
<th>miR-21 WT</th>
<th>miR-21 KO1</th>
<th>miR-21 KO2</th>
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**b**

Viral RNA copies, relative

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**c**

Reporter expression

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<th>miR-21 inhibitor</th>
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**d**

Viral RNA copies, relative

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<th>miR-21 inhibitor</th>
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</table>

**e**

miR-21 wildtype

Cell count

ZIKV positive 29.1%

**f**

miR-21 KO1

Cell count

ZIKV positive 11.6%

**g**

miR-21 KO2

Cell count

ZIKV positive 17.8%

**h**

SSCA

All cells

FSC-A

**i**

Non-infected

488 nM excitation, 610 nM emission

ZIKV positive (0.1%)

**j**

ZIKV-infected

488 nM excitation, 610 nM emission

**k**

Luminescence, normalised

<table>
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<tr>
<th>wt replicon</th>
<th>5' CS-3' CS mutant replicon</th>
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<td>0.2</td>
<td>0.4</td>
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**Ziv et al., Supplementary Fig. 10**
Supplementary figure legends

Supplementary Fig. 1: COMRADES assay development. a, Reverse transcription stalling assay indicating \textit{in vivo} crosslinking positions in the 5.8S rRNA from cells treated with Psoralen (AMT), Psoralen-TEG azide, Psoralen-TEG biotin, or DMSO control. Arrows indicate stalling events. b, Bioanalyzer RNA profiles of ZIKV enriched (upper panel) or input (lower panel) RNA from non ZIKV-infected cells, as a control to Fig. 1b. c, Enrichment of ZIKV RNA measured by TaqMan PCR. Beta-actin serves as a control RNA. d-e, Non-cropped dot-bLOTS corresponding to Fig. 1c-d. A shorter exposure is shown.

Supplementary Fig. 2: COMRADES validation. a, \textit{In vivo} detected interactions overlaid on the Ribovision 18S phylogenetic ribosomal RNA secondary structure. Colour-code is indicative of the number of supporting chimeras for each base-pair. b, Precision and sensitivity of ribosomal RNA base-pairing detection by COMRADES. Analysis is based on ZIKV enriched libraries, therefore the sensitivity of COMRADES is expected to be underestimated by this analysis.

Supplementary Fig. 3: Intra-viral RNA interactions. a, Heat maps of ZIKV RNA-RNA interactions in crosslinked libraries and controls. Chimeras ligated in 5’-3’ and 3’-5’ orientations are plotted above and below the diagonal respectively. b, Arch plot representation of short and long-range RNA-RNA interactions along the ZIKV genome. Colours representing the number of non-redundant chimeric reads supporting each interaction. c, Distribution of RNA-RNA interactions by nucleotide distance between interacting chimeric partners. Left pie chart shows distribution of all
interactions; right pie chart shows distribution of interactions that span less than 1,000 nucleotides.

Supplementary Fig. 4: The genomic structure of ZIKV inside human cells. a, The ZIKV genome conformation with the highest chimeric-reads support. Colour code is indicative of the number of supporting chimeras for each base-pair. *:

Supplementary Fig. 5: Folding entropy. a, Shannon entropy values calculated for each nucleotide along the ZIKV genome. Entropy may range from 0 to 13.4 bits; ZIKV coordinates are indicated by the position of genomic elements below. b, Inverse correlation between the degree of experimental support of base paired regions and their entropy. c, Shannon entropy values for a selected region of the ZIKV genome. d, Number of supporting chimeric reads for each base-pair shown in (c).

Supplementary Fig. 6: Structures prediction. Computationally predicted structures for ~1,000 nucleotides regions along the ZIKV genome using randomly selected subsets of experimentally probed folding-constraints. Each structure is plotted as a dot according to its folding energy (dG) and experimentally supporting
evidence (chimera reads). Red dots indicating the structure with the lowest possible
folding energy for each region.

**Supplementary Fig. 7: Clustering of structures.** a, Clustering of structures based
on degree of similarity, related to Fig. 2h. Example of structures are shown for
selected fragments with a colour code representing the number of non-redundant
chimeric reads supporting each interaction. b, Representation of the % of *in vivo*
probed interactions in use in an ensemble of 5 structures, an ensemble of 1,000
structures, or individual structures. Error bar represent standard deviation of 1,000
structures.

**Supplementary Fig. 8: Co-clustering of non-shuffled and shuffled structures,
related to Fig. 2h.** a, Red scale indicates the number of chimeric reads supporting
each non-shuffled structure. b, non-shuffled and shuffled structures, shown as white
and black dots respectively.

**Supplementary Fig. 9: *In vitro* affinity of Ago2-miR-21 to the ZIKV 5’ CS.** a,
Electrophoretic mobility shift assay (EMSA) of 5’-labeled miR-21 (lanes 1, 2) upon
addition of ZIKV 5’ CS (lane 2); and 5’-labeled ZIKV 5’ CS (lanes 3-5) upon addition
of miR-21 (lane 4) or of Ago2-miR-21 complexes (lane 5). b-h, *In vitro* measured
affinities of Ago2-miR-21 to a wildtype ZIKV 5’ CS (b), mutated 5’ CS (c-d), a fully
seed matched mutated 5’ CS (e), mutated 5’ CS with no cHP stem-loop (f), extended
5’ CS sequence containing the DCS stem-loop (g), and a short target perfectly
complementary to the miR-21 seed (h).
Supplementary Fig. 10: miR-21 interaction with the ZIKV 5' CS. a, TaqMan PCR measurement of mature miR-21 expression level in wildtype cells and CRISPR/Cas9 miR-21 deletion-clones. Values are normalized to spike-in control. b, Intracellular ZIKV RNA in miR-21 knockout and wildtype cells. c, Expression level of control and miR-21 psiCHECK-2 reporters upon treatment with miR-21 or control inhibitors. miR-21 expression values denote for Renilla / Firefly luminescence signals. d, Intracellular ZIKV RNA in miR-21 inhibited and control cells. e-j, Intracellular levels of ZIKV envelope protein, measured by FACS. Representative experiment out of 3 is shown in (e-g); Gating strategy is shown in (h-j). NS1-AF488: Alexa Fluor antibody against ZIKV envelope protein. k, Replication of a ZIKV replicon carrying a wildtype 5' CS or a 5' CS - 3' CS double mutant, pre-treated with miR-21 or non-targeting inhibitors. wt: wildtype; KO1-2: CRISPR/Cas9 miR-21 deletion-clones 1 and 2; cont: control; Error bars represent standard deviation of 3 (a, b, d), 4 (c), and 6 (k) biological replicates; Two-sided Student's t-test p-value: ** <0.005; *** <0.0005; **** <0.00005; n.s.: non-significant.

Online methods

COMRADES. Each biological replicate was carried out independently on a different day, and included crosslinked sample, a control sample, and a non-crosslinked sample.

Psoralen crosslink. JEG-3 cells (~50 million cells per experiment) were inoculated with ZIKV isolate PE243, at MOI: 2 TCID$_{50}$/cell. 20 hours post inoculation cells were washed 3 times in HANKS solution and were incubated for 20 minutes with 0.4 mg/ml Psoralen-triethylene glycol azide (psoralen-TEG azide, Berry & Associates)
dissolved in PBS and diluted in OptiMEM I with no phenol-red (Gibco). Cells were irradiated on ice with 365 nM UV for 10 minutes using a CL-1000 crosslinker (UVP). Prolonged UVA irradiation should be avoided as it might decompose the azide moiety. Cells were lysed using RNeasy lysis buffer. Proteins were degraded by proteinase K (NEB) and RNA was purified using RNeasy midi kit (Qiagen).

**Viral RNA enrichment.** Total RNA was mixed with an array of 50 biotinylated DNA oligos, 20 nucleotides-long each (IDT) designed to capture the ZIKV genomic RNA and was maintained at 37 °C for 6 hours rotating in the following hybridization buffer: 500 mM NaCl, 0.7% SDS, 33 mM Tris-Cl pH 7, 0.7 mM EDTA, 10% Formamide. At the end of incubation Dynabeads MyOne Streptavidin C1(Invitrogen) were added and the RNA was incubated for additional 1 hour at 37 C. Beads were captured on a magnet and were washed 5 times with 2x SSC buffer containing 0.5% SDS. RNA was released from beads by degrading the DNA-probes with 0.1 units/µl Turbo DNase (Invitrogen) at 37 °C for 30 minutes. RNA was cleaned by RNA Clean & Concentrator (Zymo Research).

**Crosslink pulldown.** RNA was fragmented to an average size of 100 nucleotides using RNase III (Ambion) and was cleaned by RNA clean & concentrator (Zymo Research). Copper-free Click reaction was carried at 37 °C for 90 minutes in the presence of 150 µM Click-IT Biotin DIBO Alkyne (Life technologies) and 0.5 units/µl Superase-In (Invitrogen). Reaction was terminated by RNA Clean & Concentrator (Zymo Research). Biotinylated RNA was pulldown using Dynabeads MyOne Streptavidin C1(Invitrogen) at the following reaction conditions: 100 mM Tris-Cl pH 7.5, 10 mM EDTA, 1 M NaCl, 0.1% Tween-20, 0.5 unit/µl Superase-In. Beads were
captured on a magnet and were washed 5 times with 100 mM Tris-HCl pH 7.5, 10 mM EDTA, 3.5M NaCl, 0.1% Tween-20. RNA was eluted by adding 95% Formamide, 10 mM EDTA solution and incubating at 65°C for 5 minutes. To avoid enrichment of small RNA chimeric reads that cannot be double-aligned to the reference ZIKV genome / Human transcriptome, RNA was size fractionated on 10% TBE-Urea gel and fragments corresponding to a size of 100-200 nucleotides were eluted overnight at 4 °C in 10 mM Tris-HCl pH 7.5, 1 mM EDTA, 250 mM NaCl, 0.2% SDS. RNA was concentrated using RNA Clean & Concentrator (Zymo Research).

**Proximity ligation and crosslink reversal.** At this stage, the RNA sample was divided in two. One half was used for proximity ligation and then crosslink reversal (i.e. Crosslink sample), while in the other half, crosslink reversal was done before proximity ligation (i.e. Reverse-control). We included an additional control containing an equimolar concentration (albeit a non-similar composition) of non-psoralen treated, non-crosslinked enriched RNA (i.e., Non-crosslinked sample). Before proximity ligation, the RNA was heated to 85 °C for 2 minutes and was cooldown rapidly on ice. Proximity ligation was done under the following conditions: 1 unit/µl RNA ligase 1 (New England Biolabs), 1x RNA ligase buffer, 50 mM ATP, 1 unit/µl Superase-in (Invitrogen), final volume: 200 µl. Reaction was incubated for 16 hours at 16 °C and was terminated by cleaning with RNA Clean & Concentrator (Zymo Research). Crosslink reversal was done by irradiating the RNA on ice with 2.5KJ/m² UVC.

**Sequencing library preparation.** Library preparation was done as described in with the following modifications: 6N unique molecular identifiers were added to the 5'
end of the 3’ sequencing adapter; primers and adapters concentrations were lowered to match the low RNA input; Agencourt RNaClean XP beads (Beckman Coulter) were used for clean-up and size separation; pre-adenylated 5’ and 3’ adapters were used and all ligation reactions were carried without ATP to reduce ligation artefacts. All libraries and controls went through 13 PCR cycles using KAPA HiFi HotStart Ready Mix (KAPA Biosystems). PCR products were size-selected on a 1.8% agarose gel before loading on a HiSeq 1500 sequencer (Illumina).

**Cell culture.** JEG-3 placental trophoblasts (ATCC) and Hela cells (ATCC) were cultured in Minimum Essential Medium supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate, GlutaMAX, non-essential amino acids and penicillin-streptomycin. Vero cells (Sigma-Aldrich) were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, GlutaMAX and penicillin-streptomycin. All cell lines were cultured in a humidified CO₂ incubator at 37 °C and were regularly examined to exclude mycoplasma contamination.

**Virus inoculation.** ZIKV isolate PE243 was originated from Recife, Brazil in 2015. The virus was propagated in Vero cells and titer was determined by measuring the 50% Tissue Culture Infective Dose (TCID₅₀) in JEG-3 cells. For measurements of virus replication, JEG-3 or Hela cells were inoculated with ZIKV at MOI: 0.1 TCID₅₀/cell for three hours, after which cells were washed 3 times with PBS and supplemented with fresh growth medium. 24 hours post inoculation medium was removed, cells were washed 3 times with PBS and RNA was extracted using RNeasy kit (Qiagen). Virus copy number was determined using a TaqMan real-time PCR Assay (Primerdesign) and was normalized to GAPDH and ribosomal RNA. All
virus work was handled in a containment level 2 facility registered with the HSE under COSHH.

**Replicon assay.** ZIKV wildtype and 5' CS - 3' CS double mutated replicons were described previously. Replicon RNA was synthesized *in vitro* using MEGAscript T7 Transcription Kit (Ambion). Replicon RNA was Capped using the ScriptCap m7G Capping System (Cellscript) and transfected to Hela cells using the TransIT-mRNA Transfection Kit (Mirus). Replicon levels were analyzed after 24-48 hours using a microplate luminometer (Promega) and normalized to baseline luminescence values measured at 6 hours post transfection.

**miR-21 knockout.** JEG-3 cells were transfected with CAS9-gRNA riboprotein complexes using Lipofectamine RNAiMAX (Life technologies) according to the Alt-R CRISPR-Cas9 user guide (IDT). miR-21 knockout clone1 was generated using the following guide RNA: 5'-TCATGGCAACACCAGTCGATGGG-3' and contains a homozygous deletion at the positions 59841310-59841326 on chromosome 17 (GRCh38/hg38 Assembly). miR-21 knockout clone2 was generated using a mixture of two guide RNAs: 5'-ATGTCAGACAGCCCATCGACTGG-3', 5'-CTACCATCGACATCTCCATGG-3', and contain a homozygous deletion at positions 59841249-59841321 on chromosome 17. miR-21 knockout and control clones were validated by sanger sequencing and by TaqMan Advanced miRNA Assay targeting the mature miR-21 (Life Technologies).

**miR-21 inhibition.** Hela cells were transfected with inhibitors targeting human miR-21 or non-targeting control A (Power inhibitors, Exiqon) at a final concentration of 25
nM using Lipofectamine RNAiMAX. 6 hours post transfection medium was replaced and cells were inoculated with ZIKV or re-transfected with ZIKV replicons as described above. miR-21 inhibition was validated using a psiCHECK-2 reporter (Promega) carrying a fully complementary miR-21 site at the 3' UTR of a Renilla luciferase reporter along with a Firefly reporter to normalise transfection efficiency. Luminescence was assessed using the Dual-reporter assay (Promega) and normalized to control psiCHECK-2 without the miR-21 binding site.

**Gel-based Reverse Transcription Stalling (RTS) assay.** RTS assay was performed as previously described using a Cy5-labeled primer targeting the human 5.8S ribosomal RNA: 5'-Cy5-AAGCGACGCTCAGACAGG-3'.

**Dot blot analysis.** 50 ng crosslinked RNA, or the indicated amount of 50 nt-long biotinylated standards were spotted on to a Biodyne B Nylon Membrane (Life technologies) and dried by baking at 80 °C for 10 minutes. Biotinylated RNA was detected using the chemiluminescent nucleic acid detection module Kit (Life technologies) and visualized using ChemiDoc MP Imaging System (Biorad).

**Purification of human Ago2 loaded with miR-21.** Human Ago2 homogeneously loaded with miR-21 was prepared according to a published protocol. Human Ago2 was expressed in Sf9 cells using the Bac-to-Bac baculovirus expression system (Thermo Fisher Scientific). Sf9 cells were lysed and human Ago2 was purified by Ni-NTA affinity chromatography using a His tag. Human Ago2 was loaded with synthetic 5'-phosphorylated miR-21 (IDT), and the His tag was removed using Tobacco etch virus protease. Human Ago2 loaded with miR-21 was captured using
an antisense oligonucleotide (IDT), eluted, and purified by size exclusion chromatography on an ÄKTA FPLC (GE Healthcare Life Science). Protein concentration was measured using absorption at 280 nM with extinction coefficients obtained from the protparam tool (www.expasy.org) and from the ribotask oligocalculator (www.ribotask.com).

**Target RNA labelling.** Synthetic RNA oligonucleotides (IDT) were radiolabelled at the 5'-end using gamma 32P ATP (Perkin Elmer) and T4 polynucleotide kinase (NEB), and purified by denaturing polyacrylamide gel and ethanol precipitation. RNA concentration was determined from absorption at 260 nM using extinction coefficients calculated with the ribotask oligocalculator (www.ribotask.com).

**Electrophoretic mobility shift assay (EMSA).** Binding reactions were prepared in reaction buffer (28 mM Tris pH 8.0, 20 mM KCl, 80 mM KOAc, 1.6 mM Mg(OAc)2, 0.5 mM TCEP, 0.004% NP-40, 0.01 g/l baker's yeast tRNA) with a final volume of 20 µl, and a final concentration of the labeled RNAs of 10 nM and of the non-labeled RNA or Ago2-miR-21 of 100 nM. Reactions were incubated for 10 minutes at room temperature and analyzed on a 15% acrylamide native gel in 0.5x TBE.

**Kd measurements.** Binding experiments were conducted according to the protocol published in20. Ago2-miR-21 (0-200 nM) was incubated with 0.1 nM radiolabeled target in reaction buffer (28 mM Tris pH 8.0, 20 mM KCl, 80 mM KOAc, 1.6 mM Mg(OAc)2, 0.5 mM TCEP, 0.004% NP-40) with a total volume of 25 µl for 45 minutes at room temperature. Filter-binding was performed using a dot-blot apparatus (GE Healthcare Life Sciences) with Protran nitrocellulose membrane (Amersham, GE)
Samples were applied with vacuum and washed with 50 µl wash buffer (30 mM Tris pH 8.0, 100 mM KOAc, 2 mM Mg(OAc)2, 0.5 mM TCEP). After air drying, the membrane strips were used to expose phosphor screens (GE Healthcare Life Sciences) for visualization. Screens were imaged on a Typhoon phosphorimager (GE Healthcare Life Science) and signals were quantified with ImageQuant (GE Healthcare Life Sciences). Dissociation constants were calculated by fitting the data to a single site binding equation:

\[ F = \frac{B_{\text{max}}[\text{Ago2}]}{[\text{Ago2}]+KD} \]

F = fraction target RNA bound, B_{\text{max}} = maximal number of binding sites, [Ago2] = total concentration of the Ago2-miR21 complex, and KD = calculated dissociation constant, using Prism (GraphPad Software). For weakly binding RNAs B_{\text{max}} was constrained to ≤1.

**Processing and visualization of sequencing data.** Sequencing data were preprocessed to combine FASTQ files of two sequencing lanes (cat) and to remove adapters (cutadapt). Paired end reads were merged by paired-end read merger (pear). UMIs were collapsed by collapse.py (T.D. Domenico, https://github.com/tdido). Chimeric reads were called and annotated with the hyb package\(^{31}\), using the command:

```
hyb analyse in=data.fasta db=hOH7_and_Zika format=comp eval=0.001
```

Hyb uses bowtie2\(^{32}\) in local mapping mode to map reads to a transcriptome database and to identify chimeras, and it annotates the chimeras with RNA base-
pairing information generated by hybrid-min\textsuperscript{33}. The transcriptome database used by hyb, “hOH7\_and\_Zika”, consists of human spliced mRNAs and noncoding RNAs described in\textsuperscript{14}, and the genome sequence of the Zika virus (Zika virus isolate ZIKV/H.sapiens/Brazil/PE243/2015, complete genome). To evaluate the folding energy of chimeric reads, we used hybrid-min\textsuperscript{33} with default settings. We then randomly reassigned (shuffled) pairs of fragments found in chimeric reads, and repeated the folding energy analysis. The folding energies of experimentally identified and shuffled chimeras were compared by Wilcoxon test.

Virus interaction heatmaps were plotted using Java Treeview\textsuperscript{34}, such that color intensity represents the coverage of chimeric reads at every pair of positions. The first read of each pair is plotted along the X axis, and the second read along the Y axis. As a result, chimeras found in the 5’-3’ orientation are shown above the diagonal, and chimeras in the 3’-5’ orientation are below the diagonal. Viewpoint histograms were plotted with gnuplot, and arc plots were plotted with R-chie\textsuperscript{35}.

For every pair of positions ($i, j$) along the virus genome we calculated the COMRADES score, $C_{ij}$: the number of chimeric reads that, when analyzed with the program hybrid-min with default settings, indicated base-pairing between positions $i$ and $j$. We used COMRADES scores to calculate per-base Shannon entropy for each nucleotide position along the virus. Shannon entropy of position $i$ is defined as:

$$\text{Entropy}_i = -\sum_{j=1}^{n} P(C_{ij})\log_2 P(C_{ij}),$$

where $n$ is the length of the genome (10,807 nt); and $P(C_{ij})$ is
High entropy indicates flexible positions that may form multiple alternative base-pairs, whereas low entropy indicates positions that always pair with the same nucleotide partner. We visualized RNA structures using VARNA\textsuperscript{36}, where the colour scale represents the COMRADES score for each base pair.

**RNA structure prediction.** For RNA structure predictions, we collected all potential base pairs with a non-zero $C_{ij}$ value, assembled sets of adjacent base pairs into uninterrupted stem structures, and calculated the base-pairing score of each stem as the sum of $C_{ij}$ values of individual base pairs. We then ranked these stem elements by their scores. In a preliminary analysis, we folded the 10,807 nt virus genome in a set of 50 overlapping 1,000 nt fragments, using the hybrid-ss-min program\textsuperscript{33}. Each fragment was folded using a set of 250 top-ranked *in vivo* probed stem elements as folding constraints. Based on this preliminary analysis, we identified high-scoring stem-loop structures that were reproducibly predicted across multiple fragments, and we defined new fragment boundaries to prevent the disruption of these reproducible structural elements. As a result, we obtained fragment sizes that vary in size, but are approximately 1,000 nt long each.

We then performed full folding analysis using the following fragment boundaries:

For each fragment, we assembled a set of folding constraints that represented the 75 top-scoring stem elements within that fragment. We randomly shuffled this set of constraints 1,000 times, and we used the shuffled constraints for folding prediction by hybrid-ss-min. The resulting individual structures typically incorporate 25%-40% of these constraints. We recorded the folding energy of each structure, as predicted by hybrid-ss-min, and we used the sum of $C_{ij}$ values to calculate an overall score for each structure. To assemble the top-scoring full-genome structure shown in Supplementary Fig. 4a, we assembled the top-scoring structures for each coding sequence fragment (F1-F10), and the previously proposed structures of the 5' and 3' UTRs. An additional analysis of folding within and between the 5' and 3' UTRs is shown in Fig. 2a-c and Supplementary Fig. 4c.

We also repeated the folding analysis with shuffled sets of 50-250 top-scoring constraints per fragment. This yielded similar results, but we found that either reducing or increasing the numbers of constraints tended to reduce the number of high-scoring structures.

To explore the sets of alternative structures, we computed pairwise distances between structures as the number of positions with discordant base-pairing. This resulted in a 1,000 x 1,000 matrix of distances, which we then represented on a two-dimensional surface using multidimensional scaling (using the R function cmdscale). Multidimensional scaling, also known as Principal Coordinate Analysis, maps multidimensional objects (in this case, RNA structures) to a set of points on a plane, such that the distances between RNA structures are well-approximated by Euclidean distances between points, by minimization of a stress function:
The RNA structure prediction pipeline can be downloaded from:

https://github.com/gkudla/comrades

**Validation of ZIKV miR-21 interaction with an independent analysis pipeline.**

**Alignment.** The first read of each pair was processed using UMI-tools\(^3^8\) to extract the 6 nucleotide unique molecular identifier (UMI) at the start of the read. Processed reads were aligned using the STAR aligner\(^3^9\), reporting all reads in their original order (--outSAMtype BAM Unsorted --outSAMunmapped Within); only reporting unique alignments (--outFilterMultimapNmax 1); and reporting alignments to individual segments of chimeric reads (--chimOutType WithinBAM --chimSegmentMin 20 --chimScoreJunctionNonGTAG 0 --chimMainSegmentMultNmax 1). The reference consisted of the hg38 build of the human genome, combined with the genome sequence of the PE243 strain of the Zika virus. Each read of the pair was aligned separately to avoid preferencing alignment to the same genomic locus.

For each library, the pair of BAM files were collated and pair information was fixed using samtools\(^4^0\). PCR duplicates were removed on the basis of their UMIs, using UMI-tools in paired mode.

**Detecting significant interactions.** We considered the "interaction space" between the human and Zika genomes, consisting of pairs of 1 kbp bins (one on each genome). For each replicate library in each condition (crosslinked and reversed control), we counted the number of read pairs with one read in each bin using

\[
\text{Stress}_D(x_1, \ldots, x_n) = \left( \sum_{i \neq j=1 \ldots n} (D_{i,j} - \|x_i - x_j\|^2)^{1/2} \right)
\]
diffHic\textsuperscript{41}. This yielded a count matrix that was normalized using the trimmed mean-of-M-values method\textsuperscript{42} to correct for composition biases, under the assumption that most read pairs mapping across the Zika and human genomes was caused by non-specific ligation. We then applied the quasi-likelihood framework in edgeR\textsuperscript{43} to test for significant differences between the read pair counts for crosslinked and the reverse control. This was performed using an additive design matrix that blocked on the batch to reflect the paired-sample design of the experiment. Robust empirical Bayes shrinkage\textsuperscript{44} was also used to stabilise the dispersion estimates in the presence of limited replication. Bin pairs were aggregated into clusters based on whether they overlapped the same human gene. Test statistics were combined for each gene-Zika interaction using Simes' method\textsuperscript{45} prior to applying the Benjamini-Hochberg method. Interactions that were significantly enriched in COMRADES over the reversed control were defined at a false discovery rate threshold of 5%.

References


**Supplementary references**


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Author information All sequencing data sets have been deposited in ArrayExpress under accession number: E-MTAB-6427. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to E.A.M (eric.miska@gurdon.cam.ac.uk) or G.K (gkudla@gmail.com) or O.Z. (omer.ziv@gurdon.cam.ac.uk).