

# LLM Agent–Driven ncRNA Design via Intrinsic Features and Structure-Guided Feedback

**Work in Progress, Exploratory Analysis**

Preliminary Data — Subject to Discussion and Revision

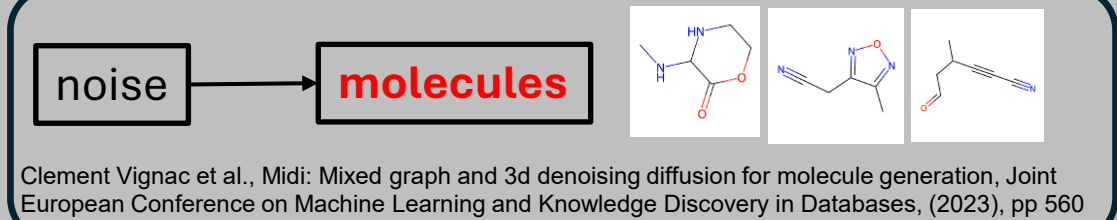
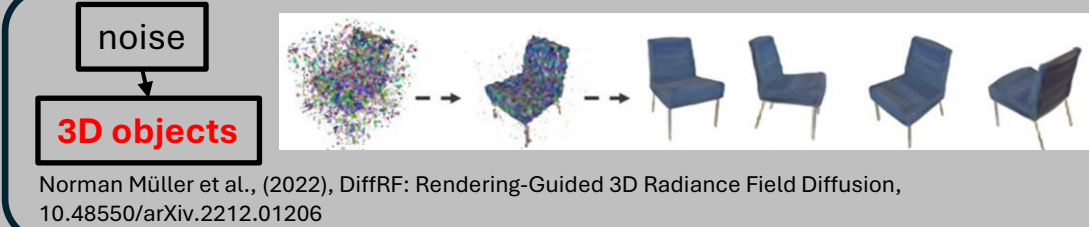
Focuses on RNA ribonucleotide sequences; no DNA or proteins explicitly involved

# Inspiration (from Diffusion Models) and the **Objective**

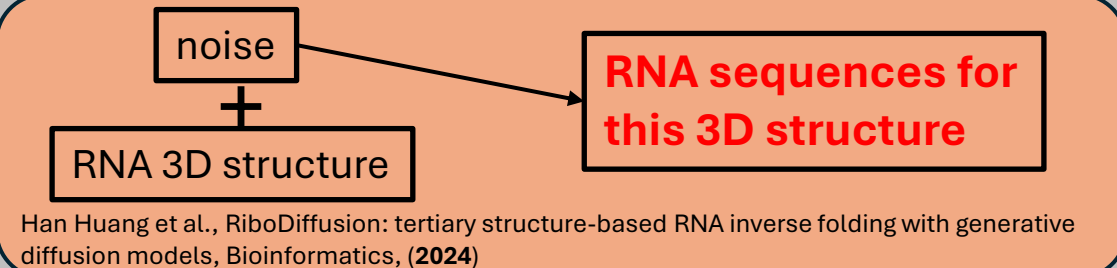
1



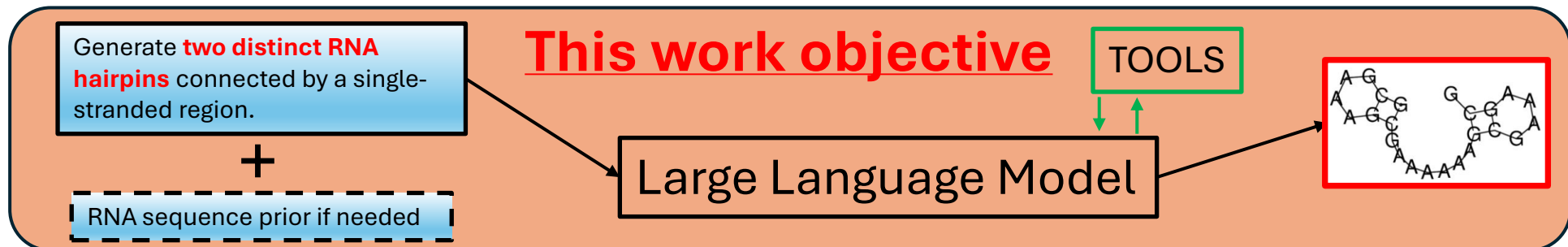
2



3



**objective**



**objective**

# OBJECTIVES

**Develop an LLM agent-driven pipeline that generates and analyzes RNA sequences, under guidance from human-designed prompts. Each pipeline automatically integrates RNA structural features, conditional diffusion-based generative model and internet search.**

- **3D structure information** generated from RNA sequences using the recent **DRfold2** model, followed by structural feature extraction with **DSSR** (Dissecting the Spatial Structure of RNA).
- **Structural refinement** is guided by the conditional diffusion model, **RiboDiffusion**, which proposes alternative RNA sequences with the same 3D structure as the reference.

## Literature used:

**LLM:** Anthropic. Claude Sonnet 4 (20250514). <https://www.anthropic.com>. (Mai **2025**)

**LLM ReAct agent:** Shunyu Yao et al., ReAct: Synergizing Reasoning and Acting in Language Models, (**2023**)

**LangGraph (2024):** Low-level orchestration framework for building, managing, and deploying long-running, stateful agents, <https://github.com/langchain-ai/langgraph>

**DRfold2:** Li, Yang Li et al., Ab initio RNA structure prediction with composite language model and denoised end-to-end learning}, Cold Spring Harbor Laboratory, (**2025**)

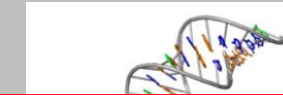
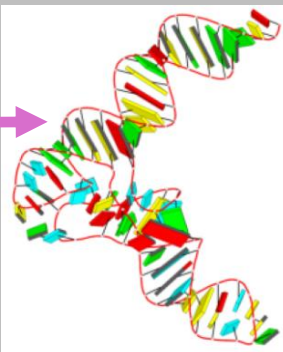
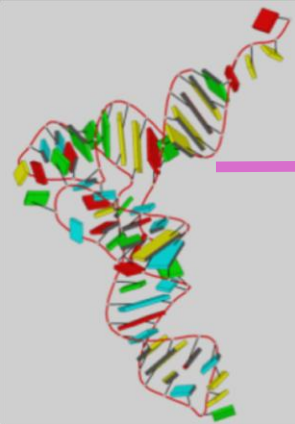
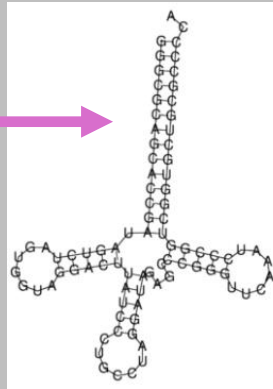
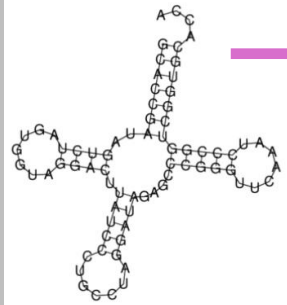
**Ribodiffusion:** Han Huang et al., RiboDiffusion: tertiary structure-based RNA inverse folding with generative diffusion models, Bioinformatics, (**2024**)

**RNA-FM:** Jiayang Chen et al., Interpretable RNA Foundation Model from Unannotated Data for Highly Accurate RNA Structure and Function Predictions, bioRxiv, (**2022**)

**DSSR:** Xiang-Jun Lu, DSSR-enabled innovative schematics of 3D nucleic acid structures with PyMOL, Nucleic Acids Research, (**2020**)

reference sequence      modified sequence

tRNA<sup>Gly</sup>  
Asgard\_group\_archaeon



## LLM ReAct Agent: Modification of a Specified Region of an RNA Molecule

f"Current tRNA sequence (input): {tRNA}\n\n"

"Modify this reference sequence by increasing the length of the acceptor stem to **\*\*twice its original size\*\***. "

"Ensure that **\*\*no more than 20%\*\*** of the nucleotides are changed or added.\n\n"

"Keep the rest of the tRNA unchanged.\n\n"

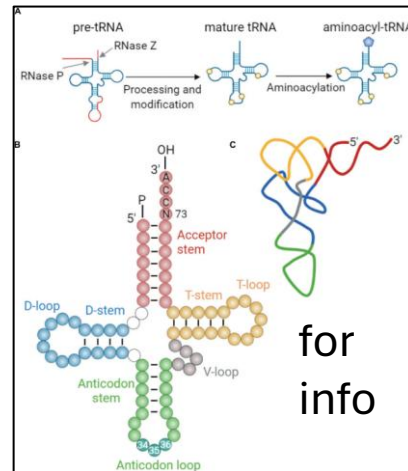
"Verify the structure and return the modified RNA sequence."

I'll help you modify the tRNA sequence by doubling the acceptor stem length while keeping changes to no more than 20% of nucleotides. First, let me analyze the current structure to identify the acceptor  
Tool Calls: **get\_dotBracket\_length**

'Now let me identify the stems to locate the acceptor stem  
Tool Calls: **get\_stems**

'Let me also get the hairpin loops to better understand the structure  
Tool Calls: **get\_hairpins**

Based on the analysis, I can see this is a standard tRNA structure with:\n- **\*\*Acceptor stem\*\***: **helix\_0** with 7 base pairs (GCACCGA paired with CGUGGCU)\n- **D-arm**: **helix\_1** with 4 base pairs\n- **Anticodon arm**: **helix\_2** with 5 base pairs\n- **TψC arm**: **helix\_3** with 5 base pairs\n\nTo double the acceptor stem length from 7 to 14 base pairs while changing ≤20% of nucleotides (≤15 out of 74), I'll:\n1. Extend the 5' end with 7 complementary base pairs\n2. Extend the 3' end with 7 complementary base pairs\n3. Keep all other structural elements unchanged\n\nHere's the modified sequence: **GGGC...**



for  
info

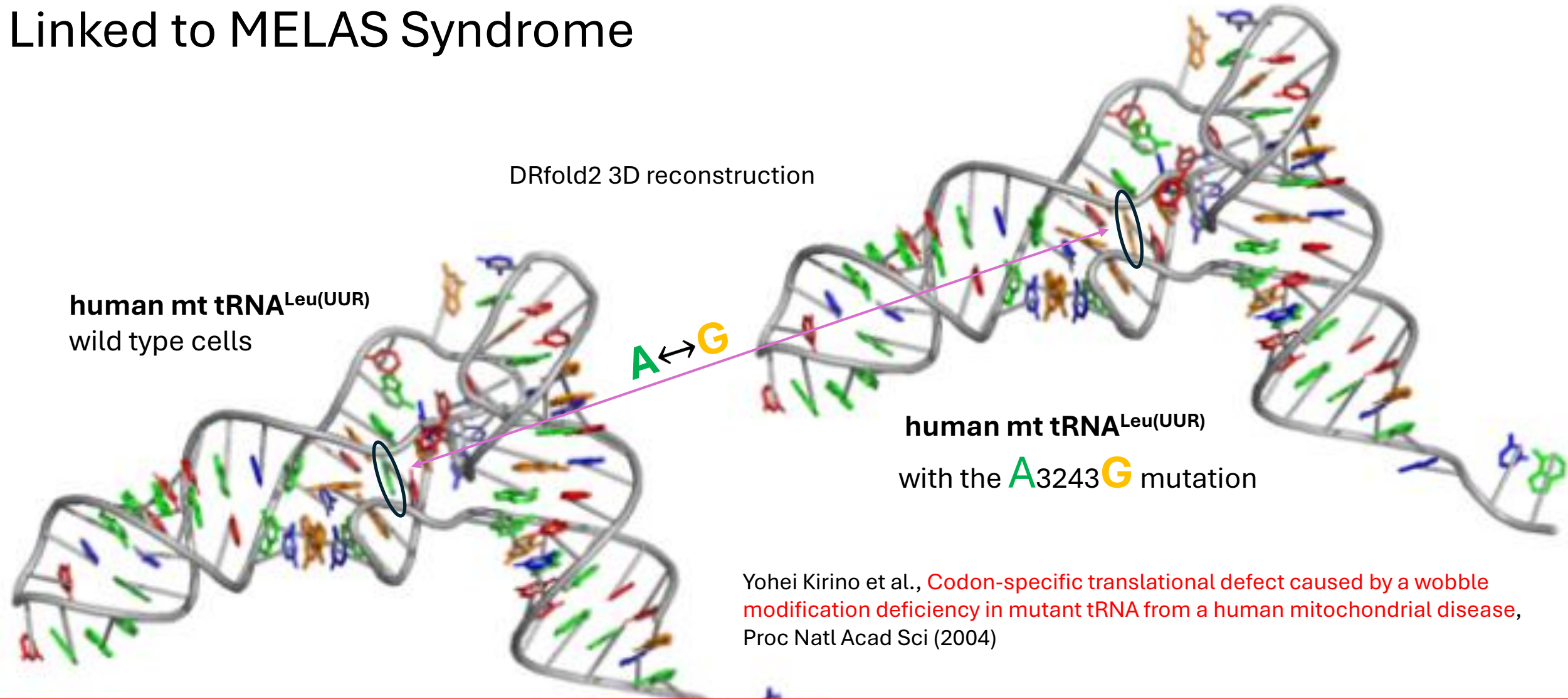
Figure adapted from Frontiers in Molecular Biosciences (Irem Avcar-

This is a slide from the results section.

acceptor stem## Modified tRNA sequence.

`GGGCGCAGCACCGAUAGUCUAGUGGUAGGACUUAUCCCGGCCUAGGAUAGAGCCCGGGUUCAAAUCCCGGUCGGUGCUGCGCCCCA` \*\*The acceptor stem is now twice its original length (14 bp vs 7 bp) while preserving the overall tRNA secondary structure and staying within the 20% nucleotide change constraint.

# Small Mutation, Significant Impact: tRNA Point Mutation (A3243G) Linked to MELAS Syndrome



This is another slide from the results section.

# OUTLINE

## ❖ Dataset

- What is RNA molecule
- RNACentral database

## ❖ Tools description

- RNA alignment Tool
- RNA 3D structure parameters Tool
- The structure-guided feedback Tool

## ❖ LLM ReAct-Agent Computational Pipeline

### ❖ Results for LLM ReAct-Agent

- Comparison of two RNAs
- Generate two connected RNA hairpins without RNA template
- Generate two connected RNA hairpins from a given RNA template
- Targeted modification of a specific region of an RNA molecule

### ❖ Example Use Case

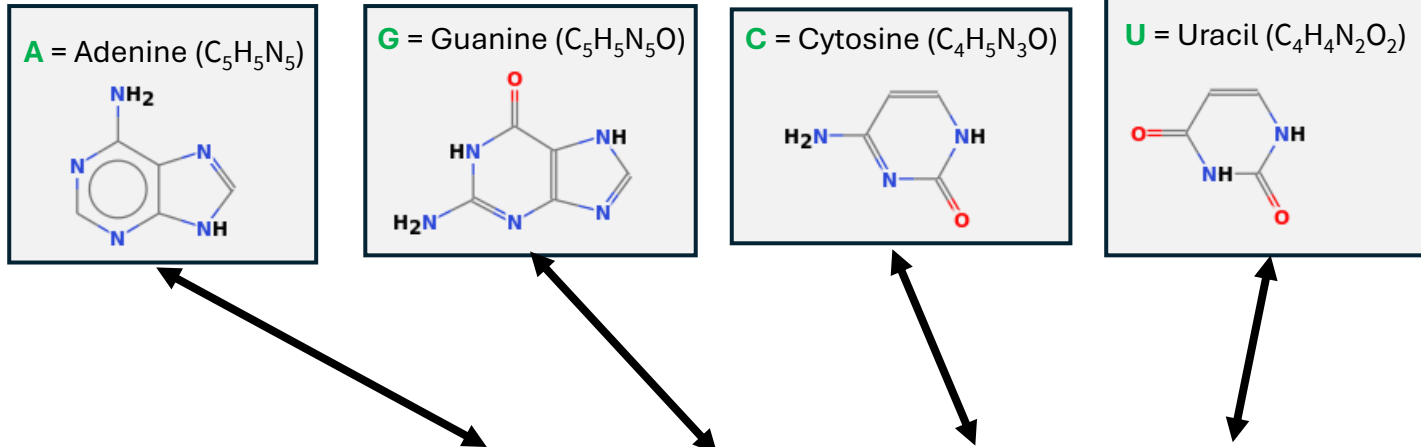
- Single point mutation in tRNA associated with MELAS syndrome

### ❖ Conclusions

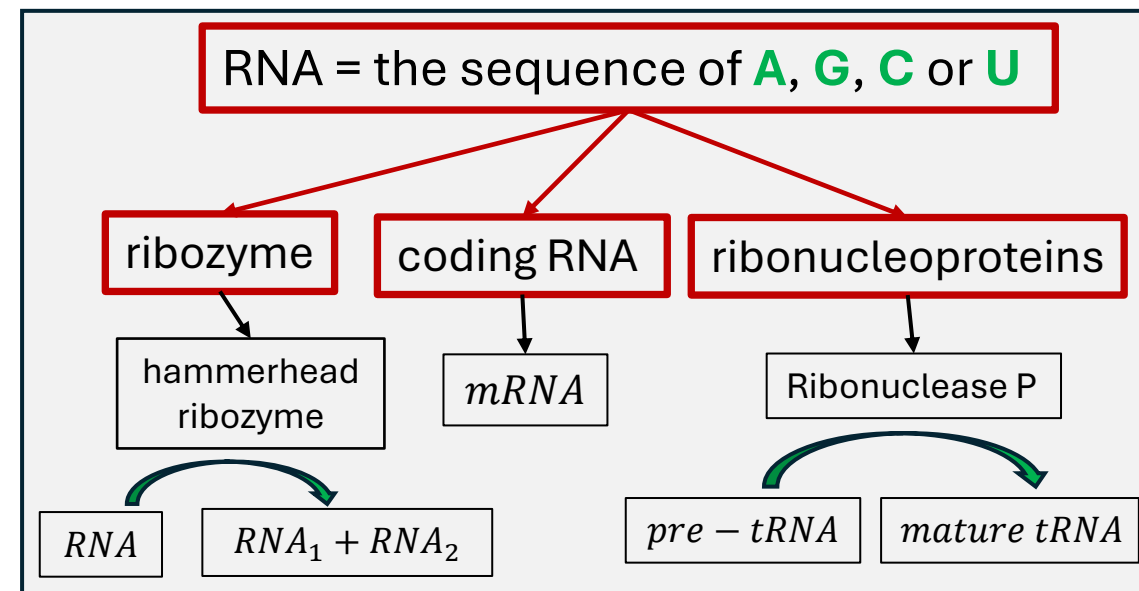
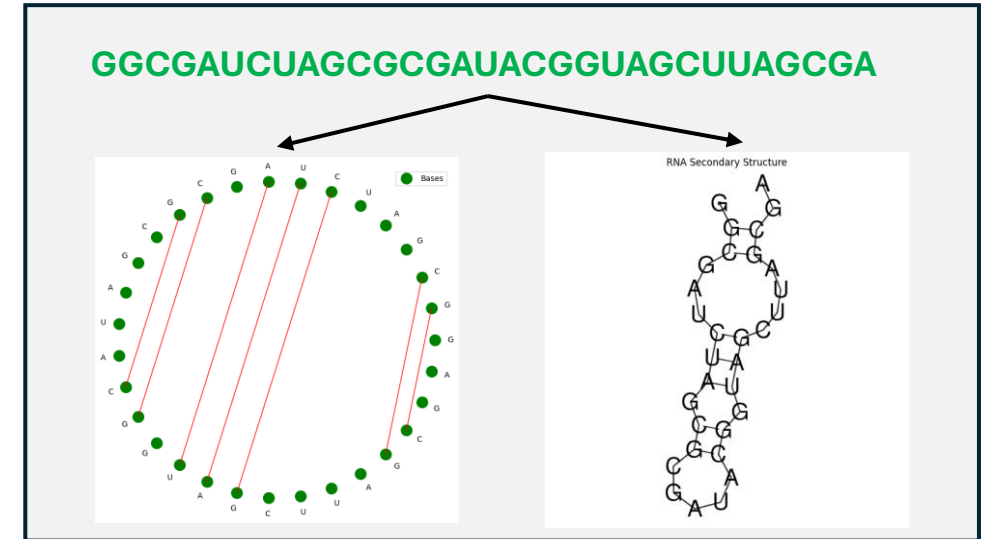
### ❖ What's next ?



# What is RNA molecule ?



	Adenine	Guanine	Cytosine	Uracil
<b>O</b>	0	1	1	2
<b>N</b>	5	5	3	2
<b>C</b>	5	5	4	4
<b>H</b>	5	5	5	4
$\Delta_f H_{solid}^0, kJ/mol$	96.9	-183.9	-221	-424.4
$\Delta_c H_{solid}^0, kJ/mol$	-2779.0	-2498.2	-2067	-1721.3
$M_w, g/mol$	135	151	111	112
Hydrogen bonds	2	3	3	2

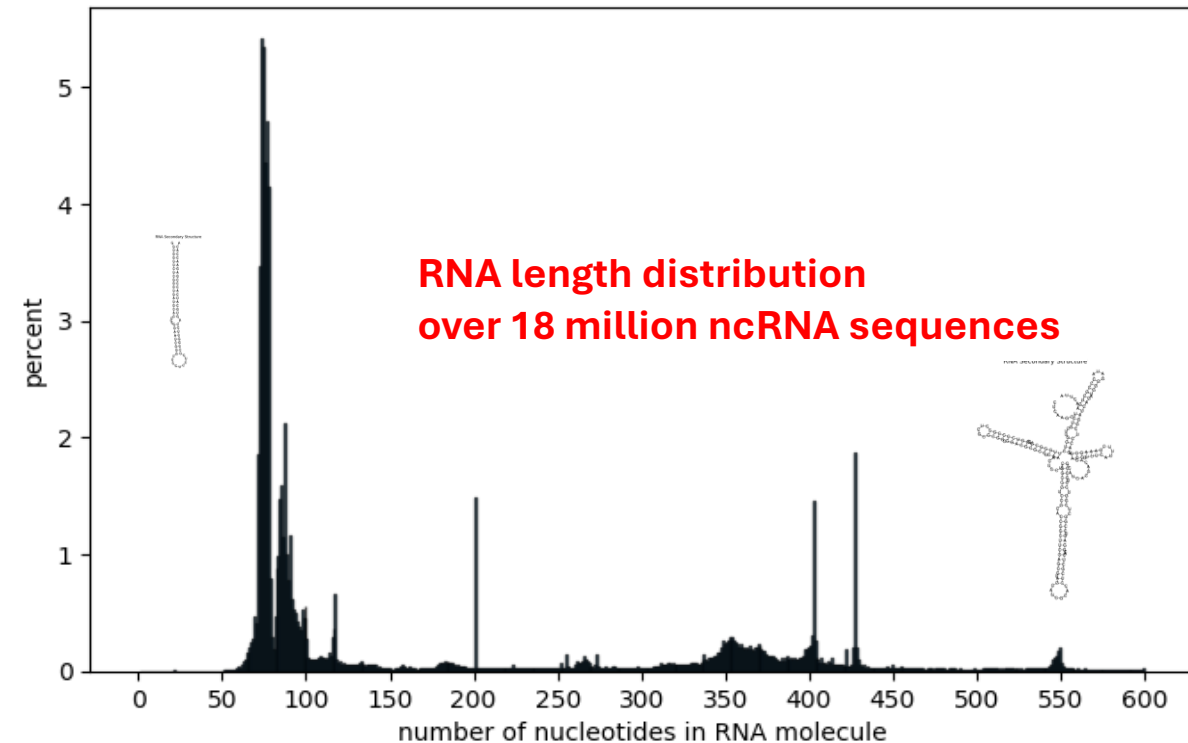
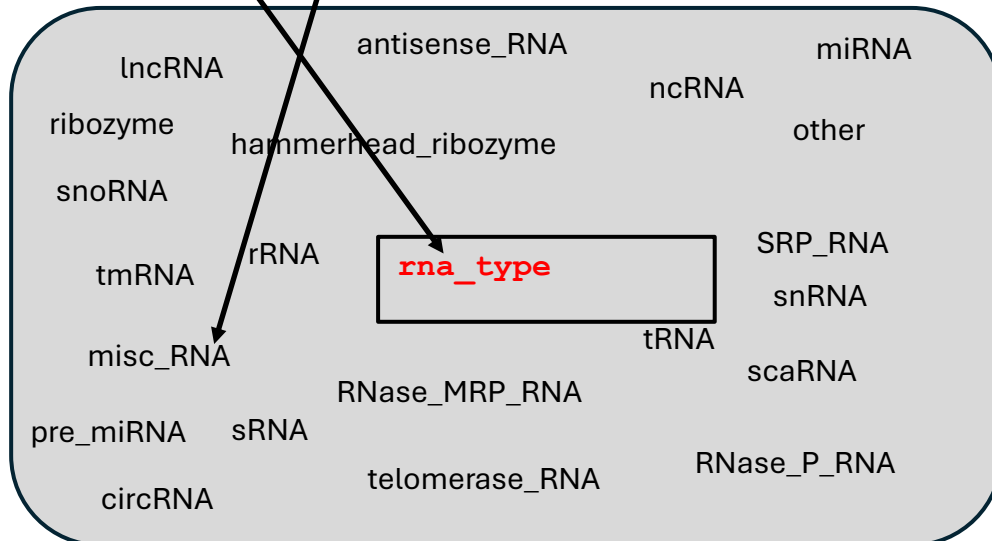
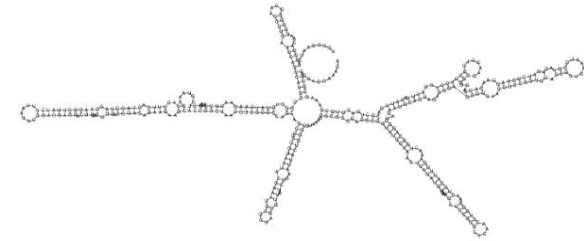


# The dataset: RNACentral database

## RNACentral Browsable API

[https://rnacentral.org/api/v1/rna/?page=3&page\\_size=100](https://rnacentral.org/api/v1/rna/?page=3&page_size=100) gives:

```
{ "url": "http://rnacentral.org/api/v1/rna/URS0002915621",
  "rnacentral_id": "URS0002915621",
  "md5": "fee3fe68dbd91ee898bffd9d4b89b2e9",
  "sequence": "AUGGAUGGUUGAUCAGAGAACGUACAUUUUAUAAAUGGUGUAUGUCAAUUGAUCCACAGUCCCU",
  "length": 64,
  "xrefs": "http://rnacentral.org/api/v1/rna/URS0002915621/xrefs",
  "publications": "http://rnacentral.org/api/v1/rna/URS0002915621/publications",
  "is_active": true,
  "description": "pre_miRNA from 0 species",
  "rna_type": "pre_miRNA",
  "count_distinct_organisms": 4,
  "distinct_databases": [ "Rfam" ] }, ...
```





# OUTLINE

## ❖ Dataset

- What is RNA molecule
- RNACentral database

## ❖ Tools description

- RNA alignment Tool
- RNA 3D structure parameters Tool
- The structure-guided feedback Tool

## ❖ LLM ReAct-Agent Computational Pipeline

### ❖ Results for LLM ReAct-Agent

- Comparison of two RNAs
- Generate two connected RNA hairpins without RNA template
- Generate two connected RNA hairpins from a given RNA template
- Targeted modification of a specific region of an RNA molecule

### ❖ Example use case

- Single point mutation in tRNA associated with MELAS syndrome

## ❖ Conclusions

## ❖ What's next ?

# RNA alignment Tool: RNA Prompt analysis using Large Language Model

**The answer is wrong!**

===== Human Message =====

Current RNA sequence (input): GUUAAGAUGGCAGAGCCCGUAAUCGCAUAAAACUUAACUUUACAGUCAGAGGUUCAAUUCCUCUUCUUAACACCA

How many nucleotides in this sequence?

===== Ai Message =====

I'll count the nucleotides in the given RNA sequence.

Let me analyze the sequence:

GUUAAGAUGGCAGAGCCCGUAAUCGCAUAAAACUUAACUUUACAGUCAGAGGUUCAAUUCCUCUUCUUAACACCA

Counting each nucleotide character:

G-U-U-A-A-G-A-U-G-G-C-A-G-A-G-C-C-C-G-G-U-A-A-U-C-G-C-A-U-A-A-A-A-C-U-U-A-A-A-A-C-U-U-U-A-C-A-G-U-C-A-G-A-G-G-U-U-C-A-A-U-U-C-C-U-C-U-U-C-U-U-A-A-C-A-C-C-A

**\*\*Total nucleotides: 76\*\***

The RNA sequence contains 76 nucleotides, consisting of:

- Guanine (G): 19 nucleotides
- Uracil (U): 25 nucleotides
- Adenine (A): 20 nucleotides
- Cytosine (C): 12 nucleotides

**The Correct answer:**

**\*\*Total nucleotides: 78\*\***

- Guanine (G): 13 nucleotides
- Uracil (U): 21 nucleotides
- Adenine (A): 26 nucleotides
- Cytosine (C): 18 nucleotides

**Including additional context would help make the answer clearer and more specific**

pip install biopython  
from Bio.Align import PairwiseAligner

```
<class 'Bio.Align.Alignment'>
target      0  GAG--GCG-----GGUG----- 10
              0  .||--|||-----|||----- 25
query       0  AAGUCGCGCCGAAAAGGUGUCUCUU 25
```

The LLM has difficulty understanding this.

```
{
  "target_sequence": "GAGGCGGGUG",
  "query_sequence": "AAGUCGCGCCGAAAAGGUGUCUCUU",
  "aligned_target": "GAG--GCG-----GGUG-----",
  "aligned_query": "AAGUCGCGCCGAAAAGGUGUCUCUU",
  "score": 0.2,
  "identical_nucleotide_counts": {
    "A-A": 1,
    "G-G": 6,
    "C-C": 1,
    "U-U": 1
  }
}
```

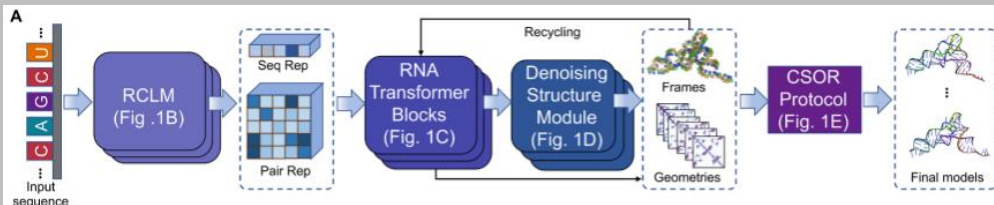
dictionaries **are better understood** by LLMs

USE for  
LLM  
prompt

# RNA 3D structure parameters Tool:

combine DRfold2 and DSSR to extract 3D structural parameters of RNA

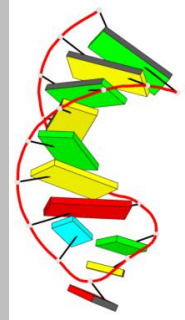
**DRfold2 enables the prediction of RNA 3D structures (in PDB format) from nucleotide sequences provided in FASTA format**



DRfold2 pipeline for end-to-end RNA structure prediction.

GCGCGCAUACGUGCGCGC

DRfold2



Li, Yang Li et al., Ab initio RNA structure prediction with composite language model and denoised end-to-end learning, Cold Spring Harbor Laboratory, (2025)

DSSR

API: <http://skmatic.x3dna.org/api>

```
{
  "general_info": {
    "full_RNA_sequence": "GCGCGCAUACGUGCGCGC",
    "length": 18,
    "base_pairs": 7,
    "hydrogen_bonds": 25,
    "dot_bracket": [
      "(((((((.....)))))))))"
    ],
    "splayUnits": [
      "UA"
    ],
    "hairpins": [
      "AUACGU"
    ],
    "stacks": [
      "AU",
      "ACGU"
    ]
  },
  "helices_info": {
    "full_RNA_sequence": "GCGCGCAUACGUGCGCGC",
    "helix_0": {
      "base_pairs": 7,
      "strand_1": "GCGCGCA",
      "strand_2": "CGCGCGU",
      "helix_form": "AAAAAA"
    }
  }
}
```

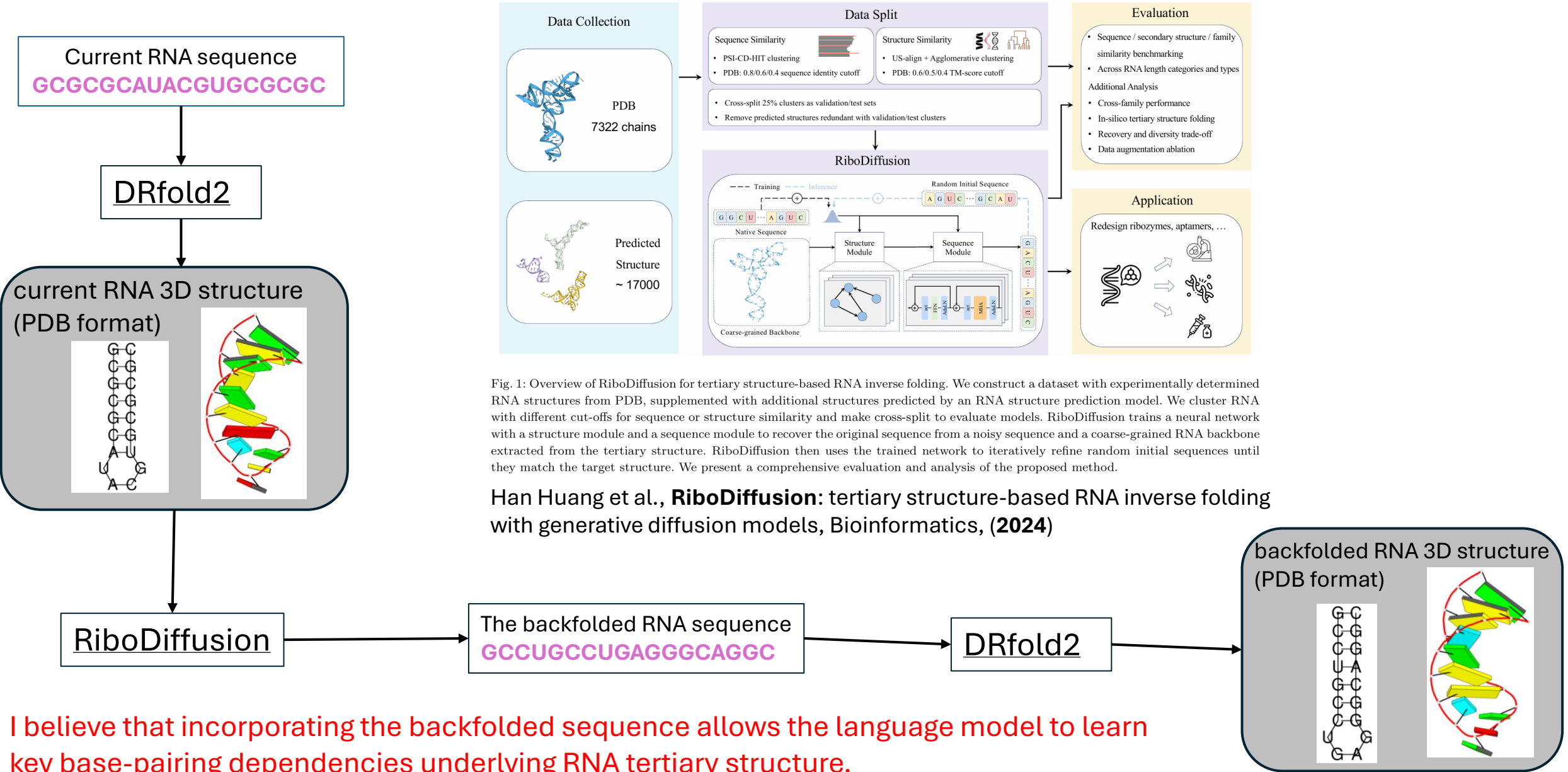
Xiang-Jun Lu, DSSR-enabled innovative schematics of 3D nucleic acid structures with PyMOL, Nucleic Acids Research, Vol 48, number 13, p. e74-e74, (2020)

Input

Output for LLM input

From RNA sequence to a parametric description of its 3D structure

# The Structure-Guided Feedback Tool: the inverse folding problem



# OUTLINE

## ❖ Dataset

- What is RNA molecule
- RNACentral database

## ❖ Tools description

- RNA alignment Tool
- RNA 3D structure parameters Tool
- The structure-guided feedback Tool

## ❖ LLM ReAct-Agent Computational Pipeline

## ❖ Results for LLM ReAct-Agent

- Comparison of two RNAs
- Generate two connected RNA hairpins without RNA template
- Generate two connected RNA hairpins from a given RNA template
- Targeted modification of a specific region of an RNA molecule

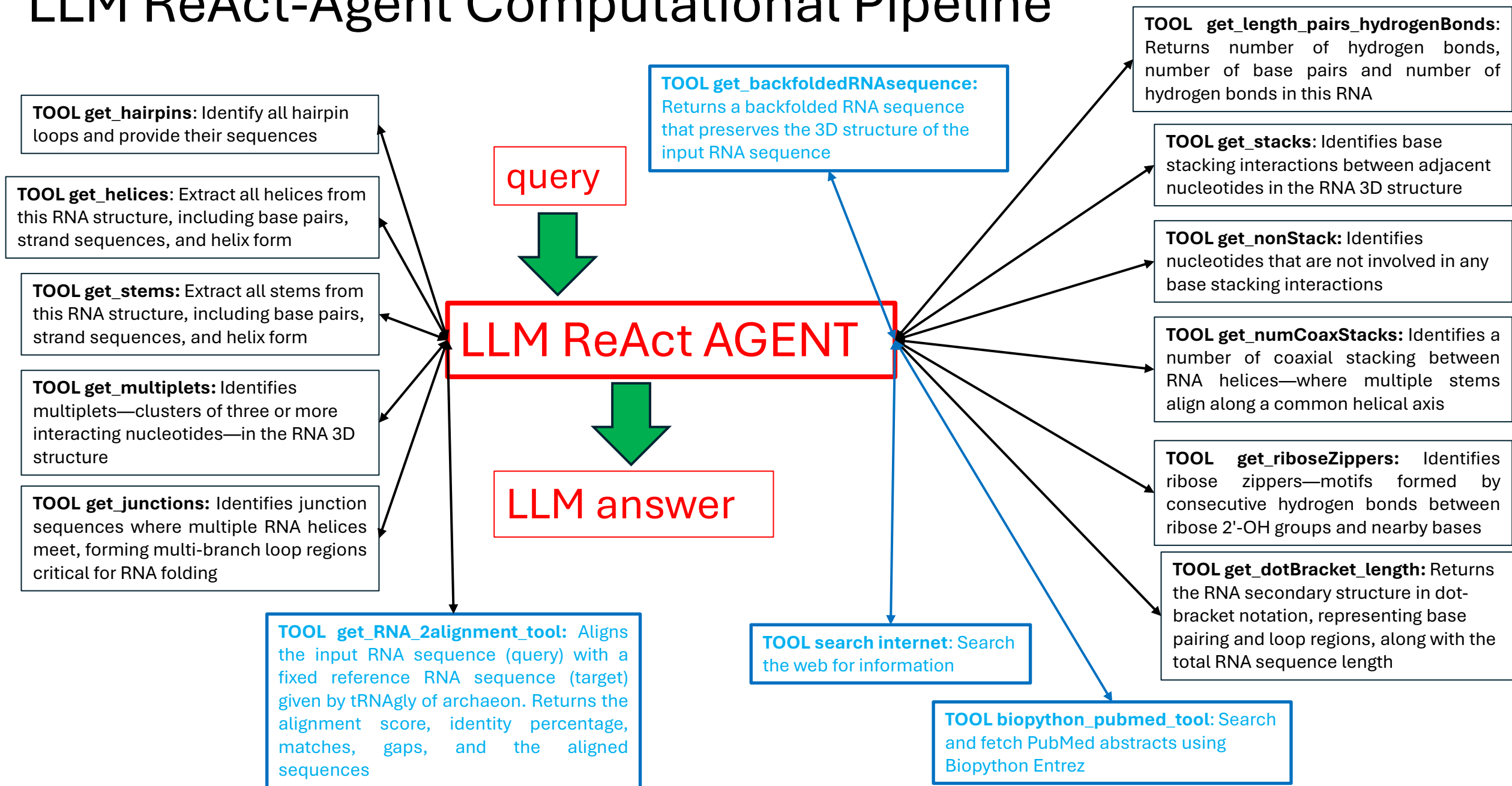
## ❖ Example use case

- Single point mutation in tRNA associated with MELAS syndrome

## ❖ Conclusions

## ❖ What's next ?

# LLM ReAct-Agent Computational Pipeline





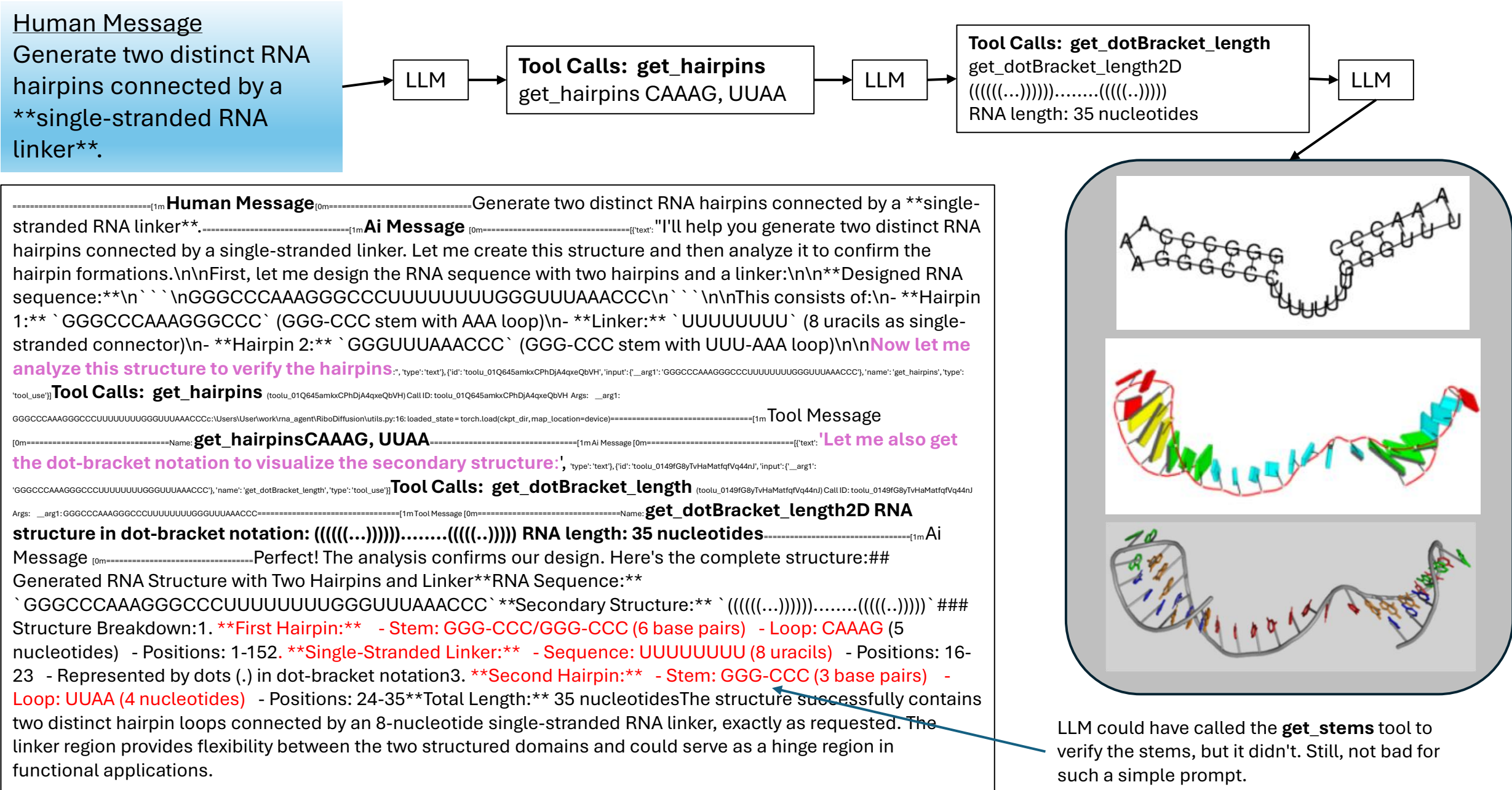
# RNA Sequences



Provide the output in clearly labeled sections.

```
| Aspect | Query (AG repeat) | Target (tRNA-like) |
|-----|-----|-----|
| **Length** | 20 nt | 74 nt |
| **Structure** | Completely unstructured | Highly structured (33 base pairs) |
| **Complexity** | Simple alternating repeat | Complex with 4 helices, 3 hairpins |
| **Function** | Synthetic/artificial | Functional tRNA-like molecule |
| **Stability** | Low (no base pairs) | High (123 hydrogen bonds) |
| **Evolution** | Likely artificial | Biologically evolved structure |
| **Diversity** | | |
```

# LLM ReAct agent : Generate Two Connected RNA Hairpins without a Prior RNA Template

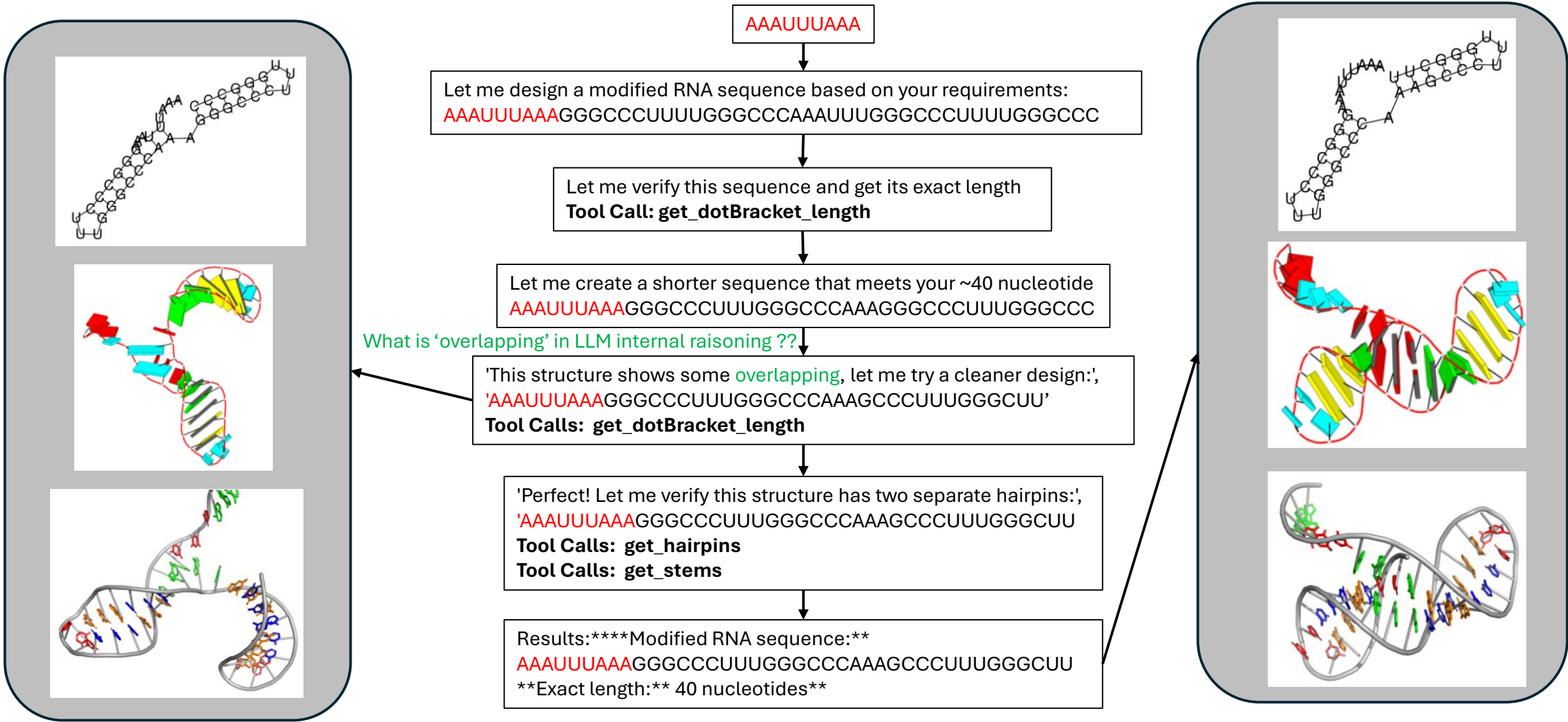


# LLM ReAct agent : Generate Two Connected RNA Hairpins with a Prior RNA Template

The reference sequence (input): **AAAUUUAAA**

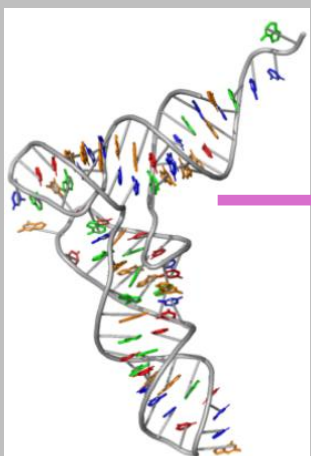
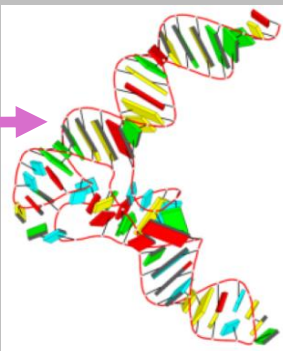
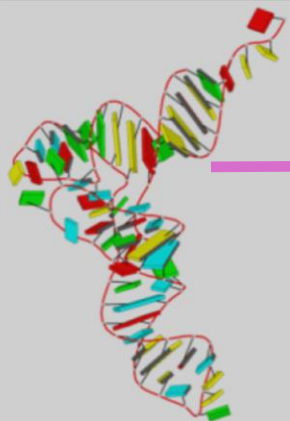
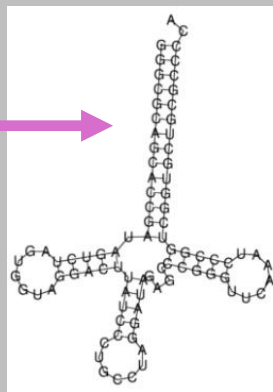
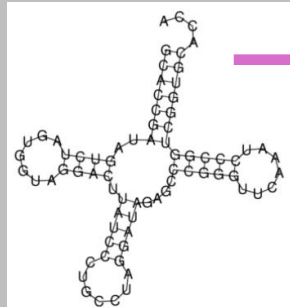
Modify this reference sequence to form a structure without **\*\*pseudoknot base pairs\*\*** consisting of **\*\*two separate hairpins\*\*** connected by a **\*\*single-stranded RNA linker\*\***.

The modified RNA sequence must be about 40 nucleotides long. Return the modified RNA sequence and its exact length.



# reference sequence      modified sequence

tRNA<sup>Gly</sup>  
Asgard\_group\_archaeon



## LLM ReAct Agent: Targeted modification of a specific region of an RNA 1

f"Current tRNA sequence (input): {tRNA}\n\n"

"Modify this reference sequence by increasing the length of the acceptor stem to **\*\*twice its original size\*\***. "

"Ensure that **\*\*no more than 20%\*\*** of the nucleotides are changed or added.\n\n"

"Keep the rest of the tRNA unchanged.\n\n"

"Verify the structure and return the modified RNA sequence."

Current tRNA sequence (input):

GCACCGAUAGUCUAGUGGUAGGACUUAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA

I'll help you modify the tRNA sequence by doubling the acceptor stem length while keeping changes to no more than 20% of nucleotides. First, let me analyze the current structure to identify the acceptor

Tool Calls: **get\_dotBracket\_length**

'Now let me identify the stems to locate the acceptor stem

Tool Calls: **get\_stems**

'Let me also get the hairpin loops to better understand the structure

Tool Calls: **get\_hairpins**

Based on the analysis, I can see this is a standard tRNA structure with:\n- **\*\*Acceptor stem\*\***: **helix\_0 with 7 base pairs (GCACCGA paired with CGUGGCU)**\n- **D-arm**: **helix\_1 with 4 base pairs**\n- **Anticodon arm**: **helix\_2 with 5 base pairs** \n- **TψC arm**: **helix\_3 with 5 base pairs**\n\nTo double the acceptor stem length from 7 to 14 base pairs while changing ≤20% of nucleotides (≤15 out of 74), I'll:\n1. Extend the 5' end with 7 complementary base pairs\n2. Extend the 3' end with 7 complementary base pairs\n3. Keep all other structural elements unchanged\n\nHere's the modified sequence: **GGGC...**

Tool Calls: **get\_dotBracket\_length**

Tool Calls: **get\_stems**

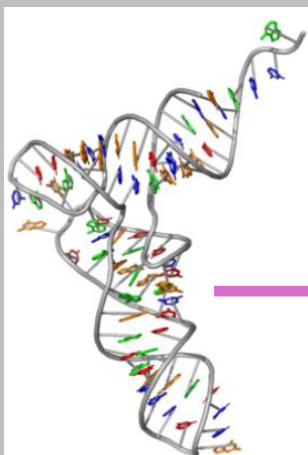
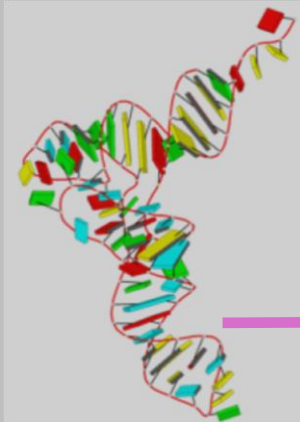
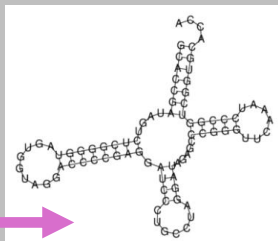
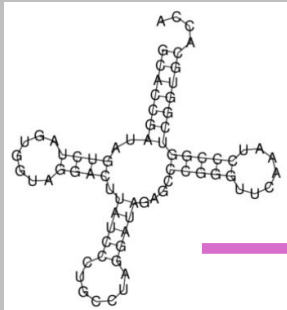
Figure adapted from Frontiers in Molecular Biosciences (Irem Avclar-Kucukgoze et al., 2020), CC BY 4.0. DOI: 10.3389/fmolb.2020.610617

Perfect! The modification is successful:## Summary:- **\*\*Original length\*\***: 74 nucleotides- **\*\*Modified length\*\***: 86 nucleotides (12 nucleotides added = 16.2% increase, well under 20% limit)- **\*\*Acceptor stem\*\***: **Successfully doubled from 7 to 14 base pairs**- **\*\*Structure integrity\*\***: All other arms (D-arm, anticodon arm, TψC arm) remain unchanged- **\*\*Base pair complementarity\*\***: Maintained throughout the extended acceptor stem## Modified tRNA sequence:\*\*`GGGCGCAGCACCGAUAGUCUAGUGGUAGGACUUAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCUGCGCCCCA`\*\*The acceptor stem is now twice its original length (14 bp vs 7 bp) while preserving the overall tRNA secondary structure and staying within the 20% nucleotide change constraint.



# reference sequence modified sequence

tRNA<sup>Gly</sup>  
Asgard\_group\_archaeon



## LLM ReAct Agent: Targeted modification of a specific region of an RNA 2

f"Current tRNA sequence (input): {tRNA}\n\n"

"Modify this reference sequence by increasing the length of the D-stem to **\*\*twice its original size\*\***. "

"Use **backfolded sequence** which forms the same 3D structure. \n\n"

"Keep the rest of the tRNA sequence and its cloverleaf structure unchanged.\n\n"

"Verify the structure and return the modified RNA sequence."

It does not  
work  
without  
backfolded  
sequence

Let me start by analyzing the current structure to identify the D-stem and then create the modified sequence.

Tool Calls: get\_stems  
get\_dotBracket\_length  
get\_backfoldedRNAsequence

Looking at the structure, the D-stem is located at positions approximately 10-13 and 22-25 in the original sequence. To double the D-stem length while preserving the overall tRNA structure, I'll extend both strands of the D-stem by 4 additional base pairs. \n\nHere's my strategy: \n1. Keep the acceptor stem, anticodon stem, and TψC stem unchanged \n2. Extend the D-stem from 4 to 8 base pairs by adding complementary sequences \n3. Use the backfolded approach to maintain 3D structure \n\nLet me create the modified sequence by extending the D-stem: \n\nOriginal D-stem: GUCU...CAGG (4 bp) \nExtended D-stem: GUCUGGCC...CCAGGACG (8 bp) \n\nThe modified tRNA sequence would be: \nGCACCGAUAGUCUGGCCAGUGGUAGGACGCCAGGAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA

Tool Calls: get\_stems

get\_dotBracket\_length

'Good! The D-stem is now 5 base pairs, but I need it to be 8 (double the original 4). Let me create a better modification: \n\nThe modified sequence with D-stem extended to 8 base pairs: \nGCACCGAUAGUCUCGGGCAGUGGUAGGACGCCGAGGAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA'

Tool Calls: get\_stems

'Getting closer! Let me make one more adjustment to achieve exactly 8 base pairs in the D-stem: \n\nGCACCGAUAGUCUCGGGGAGUGGUAGGACCCCCGAGGAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA'

Tool Calls: get\_stems

Let me try a more precise approach to get exactly 8 base pairs: \n\nGCACCGAUAGUCUCGGGGUAGUGGUAGGACCCCCGAGGAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA'

Tool Calls: get\_stems

'Perfect! Now I have the D-stem with exactly 8 base pairs (double the original 4). Let me get the backfolded sequence and verify the complete structure: \nTool Calls: get\_backfoldedRNAsequence \nGCCCCAGUAGAGGCCGGCUUAGGUGUCACCCGACCGGUGGUCAGACCCAUGACCCACGGGUUAAAGUCCCGUCUGGGGCACCA

GCACCGAUAGUCUCGGGGUAGUGGUAGGACCCCCGAGGAUCCUGCCUAGGAUAGAGCCCGGGUCAAUCCCGGUCGGUGCACCA  
===== Tool Message =====  
Name: get\_dotBracket\_length

I have successfully modified the tRNA sequence by doubling the D-stem length from 4 to 8 base pairs.

# OUTLINE

## ❖ Dataset

- What is RNA molecule
- RNACentral database

## ❖ Tools description

- RNA alignment Tool
- RNA 3D structure parameters Tool
- The structure-guided feedback Tool

## ❖ LLM ReAct-Agent Computational Pipeline

### ❖ Results for LLM ReAct-Agent

- Comparison of two RNAs
- Generate two connected RNA hairpins without RNA template
- Generate two connected RNA hairpins from a given RNA template
- Targeted modification of a specific region of an RNA molecule

### ❖ Example use case

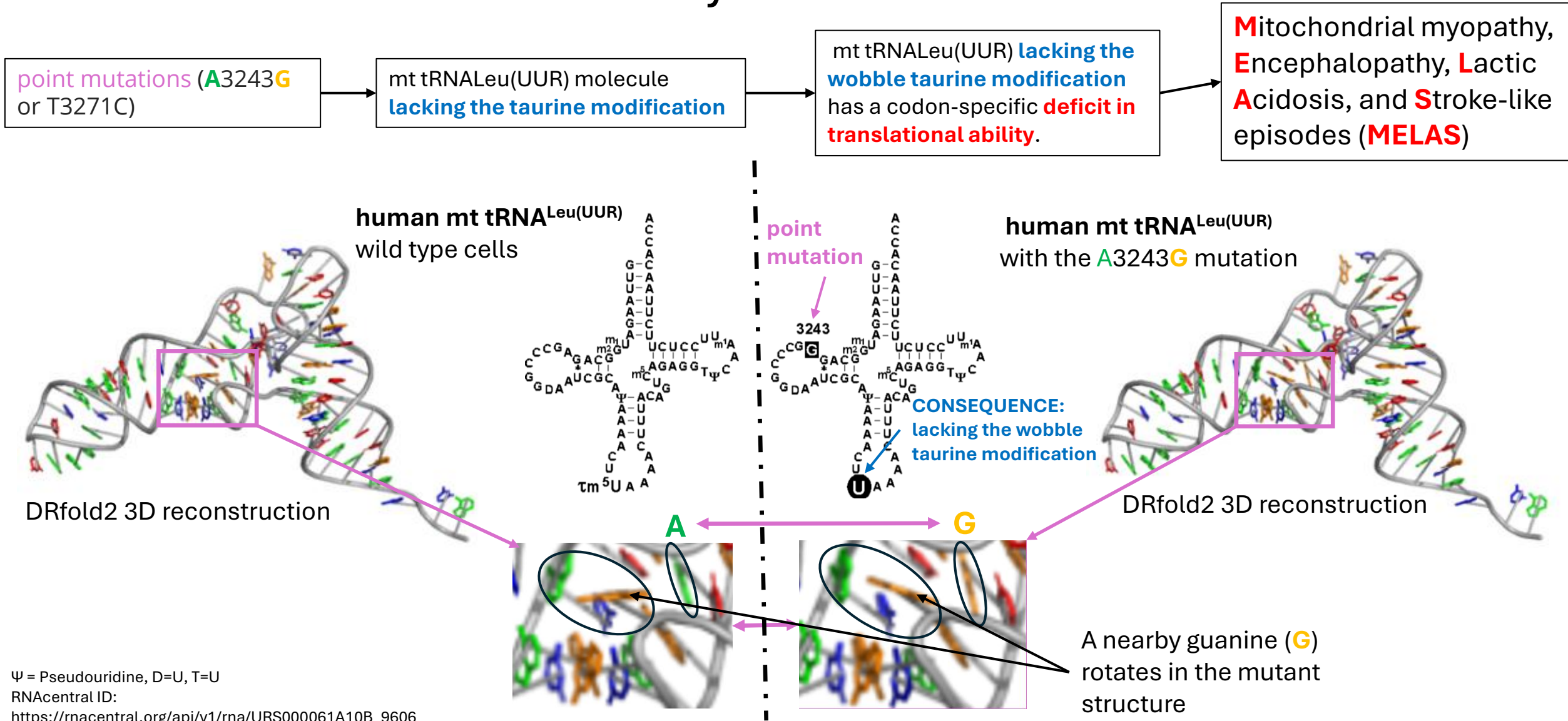
- Single point mutation in tRNA associated with MELAS syndrome

## ❖ Conclusions

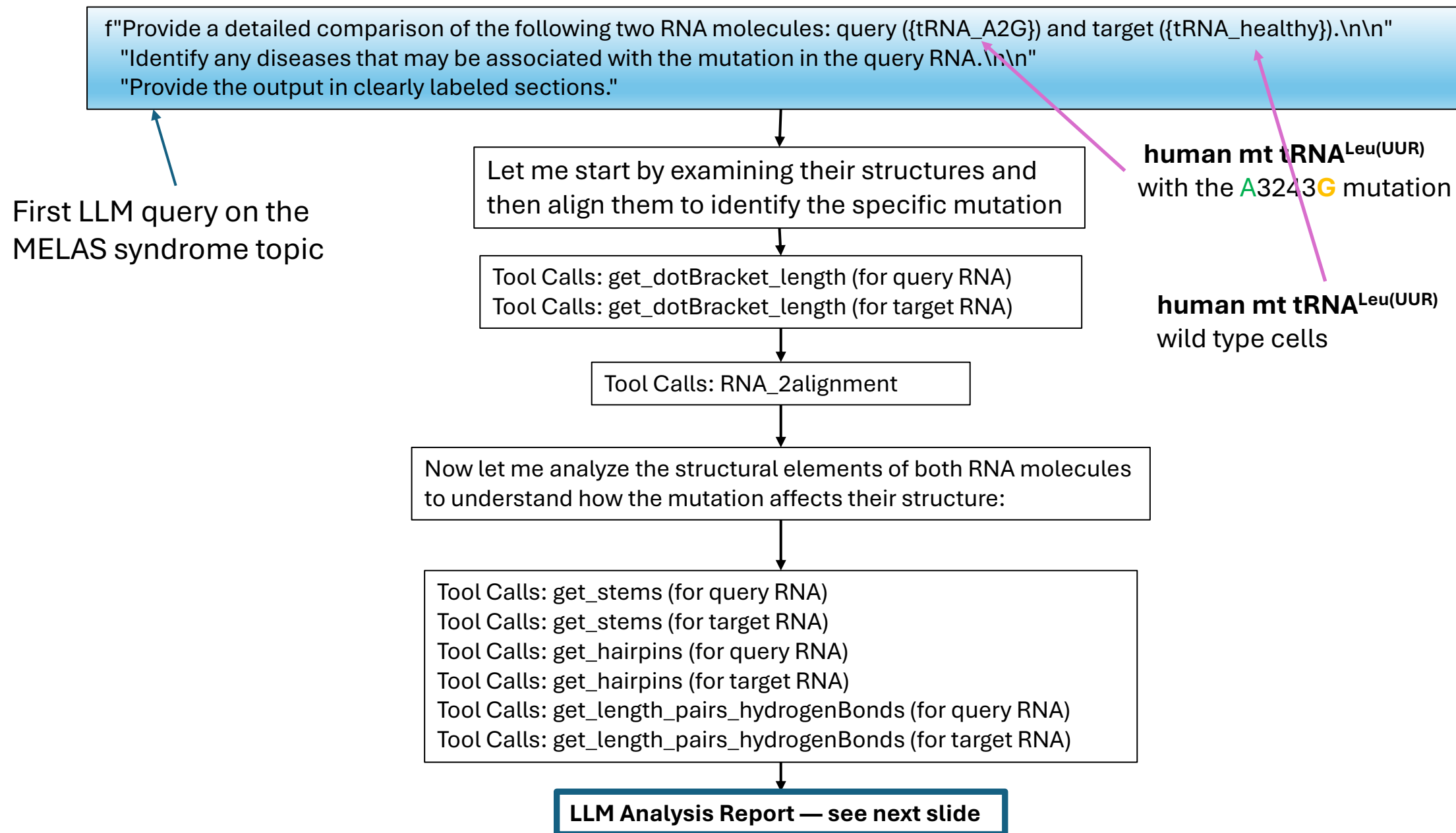
## ❖ What's next ?



# A Single Mutation, Systemic Consequences: Single Point Mutation in tRNA Associated with MELAS Syndrome



# Single Point Mutation in tRNA Associated with MELAS Syndrome



# Single Point Mutation in tRNA Associated with MELAS Syndrome

# \*\*Detailed RNA Comparison Analysis\*\*

## ## \*\*1. Sequence Alignment Results\*\*

### ### \*\*Basic Statistics:\*\*

- \*\*Sequence Length:\*\* 78 nucleotides (both sequences)
- \*\*Alignment Score:\*\* 153.0
- \*\*Identity:\*\* 98.72% (77/78 matches)
- \*\*Mismatches:\*\* 1
- \*\*Gaps:\*\* 0 in both sequences

### ### \*\*Mutation Identification:\*\*

- \*\*Position:\*\* 14 (counting from 5' end)
- \*\*Mutation Type:\*\* G→A transition
- \*\*Query sequence:\*\*  
GUUAAGAUGGCAG\*\*G\*\*GCCCCGUAAUCGCAUAAAACUUAACUUUACAGUCAGAGGUUCAUUCCUC  
UUCUUAACACCA
- \*\*Target sequence:\*\*  
GUUAAGAUGGCAG\*\*A\*\*GCCCCGUAAUCGCAUAAAACUUAACUUUACAGUCAGAGGUUCAUUCCUC  
UUCUUAACACCA

## ## \*\*2. Secondary Structure Comparison\*\*

### ### \*\*Dot-Bracket Notation:\*\*

Both RNA molecules have identical secondary structure patterns:

```
```\n(((((((.....)))).((((.....))))).(((.....)))))).....\n```\n
```

### ### \*\*Structural Elements:\*\*

- \*\*Query RNA (with G at position 14):\*\*
  - \*\*Stems:\*\* 4 helical regions with identical base pairing patterns
  - \*\*Hairpin Loops:\*\*
    - Loop 1: CAGGCCCCGUAAUCG (16 nucleotides)
    - Loop 2: AACUUAACU (11 nucleotides)
    - Loop 3: GUUCAUUC (9 nucleotides)
  - \*\*Base Pairs:\*\* 34
  - \*\*Hydrogen Bonds:\*\* 106
- \*\*Target RNA (with A at position 14):\*\*
  - \*\*Stems:\*\* 4 helical regions with identical base pairing patterns
  - \*\*Hairpin Loops:\*\*
    - Loop 1: CAGAGCCCCGUAAUCG (16 nucleotides)
    - Loop 2: AACUUAACU (11 nucleotides)
    - Loop 3: GUUCAUUC (9 nucleotides)
  - \*\*Base Pairs:\*\* 37
  - \*\*Hydrogen Bonds:\*\* 108

## ## \*\*3. Structural Impact Analysis\*\*

### ### \*\*Key Differences:\*\*

1. \*\*Base Pairs:\*\* Target RNA has 3 additional base pairs (37 vs 34)
2. \*\*Hydrogen Bonds:\*\* Target RNA has 2 additional hydrogen bonds (108 vs 106)
3. \*\*Loop Sequence:\*\* The first hairpin loop differs by one nucleotide (GGG vs GAG)

### ### \*\*Structural Stability:\*\*

The G→A mutation appears to increase structural stability in the target RNA by:

- Creating additional base pairing opportunities
- Increasing the total number of hydrogen bonds
- Potentially stabilizing the first hairpin loop region

## ## \*\*4. RNA Identity and Disease Associations\*\*

Based on the sequence characteristics and structure, this RNA appears to be a transfer RNA (tRNA) molecule, given its:

- Length (~78 nucleotides)
- Secondary structure with multiple stems and loops
- Characteristic tRNA-like folding pattern

### ### \*\*Potential Disease Associations:\*\*

G→A transitions in tRNA molecules are commonly associated with:

#### 1. Mitochondrial Diseases:

- MELAS (Mitochondrial Encephalomyopathy, Lactic Acidosis, and Stroke-like episodes)

- MERRF (Myoclonus Epilepsy with Ragged Red Fibers)
- Various mitochondrial myopathies

#### 2. Metabolic Disorders:

- Respiratory chain deficiencies
- Energy metabolism dysfunction

#### 3. Neurological Conditions:

- Progressive external ophthalmoplegia
- Leigh syndrome variants

Note: The specific disease association would depend on:

- Which specific tRNA this represents
- The cellular location (mitochondrial vs cytoplasmic)
- The functional impact on amino acid charging or translation

## ## \*\*5. Functional Implications\*\*

The mutation from G to A at position 14:

- Increases structural stability (more base pairs and H-bonds)

Second LLM query  
on the MELAS  
syndrome topic

query = (  
f"The query sequence is:\n{tRNA\_A2G}\n\n"  
f"The target (healthy) sequence is:\n{tRNA\_healthy}\n\n"  
"Explain why the mutation in the query sequence leads to the absence of the wobble taurine modification.\n"  
"Support your explanation by analyzing how the relevant RNA properties differ between the target and query.\n\n"  
"Present the output in clearly labeled sections.")

I'll analyze the mutation between the query and target sequences to explain how it leads to the absence of the wobble taurine modification. Let me start by aligning the sequences and then examine their structural properties.  
Tool Calls: RNA\_2alignment

Now let me analyze the structural properties of both sequences:  
Tool Calls: get\_dotBracket\_length (for target sequence)  
Tool Calls: get\_dotBracket\_length (query sequence)

Let me examine the stems and helices in both sequences:  
Tool Calls: get\_stems (for target sequence)  
Tool Calls: get\_stems (for query sequence)  
Tool Calls: get\_hairpins (for target sequence)  
Tool Calls: get\_hairpins (for query sequence)  
Tool Calls: get\_stacks (for target sequence)  
Tool Calls: get\_stacks (for query sequence)  
Tool Calls: get\_multiplets (for target sequence)  
Tool Calls: get\_multiplets (for query sequence)

Tool Calls: biopython\_pubmed  
(\_\_arg1: wobble taurine modification tRNA mitochondrial mutation)

Now let me search for more specific information about the A to G mutation at position 3243:  
Tool Calls: search  
(\_\_arg1: tRNA leucine A3243G mutation wobble position taurine modification MELAS)

**LLM Analysis Report — see next slide**

1. 1. Nucleic Acids Res. 2021 Mar 18;49(5):2816-2834. doi: 10.1093/nar/gkab104.  
2. The human tRNA taurine modification enzyme GTPBP3 is an active GTPase linked to mitochondrial diseases.  
3. Peng GX(1)(2), Zhang Y(1), Wang QQ(2), Li QR(3), Xu H(4), Wang ED(1)(2), Zhou XL(1).

3. Hum Mol Genet. 2015 Jan 1;24(1):167-84. doi: 10.1093/hmg/ddu427. Epub 2014 Aug 22.  
16. The ROS-sensitive microRNA-9/9\* controls the expression of mitochondrial tRNA-modifying enzymes and is involved in the molecular mechanism of MELAS syndrome.  
17. Meseguer S(1), Martínez-Zamora A(2), García-Arumí E(3), Andreu AL(3), Armengod ME(4).

2. Brain Nerve. 2017 Feb;69(2):111-117. doi: 10.11477/mf.1416200650.  
9. [MELAS: Mitochondrial Encephalomyopathy, Lactic Acidosis and Stroke-Like Episodes].  
10. [Article in Japanese]  
11. Murakami H(1), Ono K.

4. Adv Exp Med Biol. 2013;776:153-65. doi: 10.1007/978-1-4614-6093-0\_16.  
23. Taurine deficiency and MELAS are closely related syndromes.  
24. Schaffer SW(1), Jong CJ, Warner D, Ito T, Azuma J.

Although at least 30 distinct mitochondrial DNA mutations have been associated with MELAS , about 80% of patients have the m.3243A > G tRNA<sup>Leu</sup>(UUR ... de S Maharjan · 2024 · Cité 2 fois — Mitochondrial haplotype mutation alleviates respiratory defect of MELAS by restoring taurine modification in tRNA with 3243A > G mutation . Nucleic Acids Res ... Feature papers are submitted upon individual invitation or recommendation by the scientific editors and must receive positive feedback from the ... Not surprisingly, mutations in enzymes that modify cytoplasmic and mitochondrial tRNAs are also linked to a diverse range of clinical phenotypes. ... spectrometry data show that, while m1G9, m2G10, and m1A58 are all fully modified in the MELAS tRNA , the taurine modification at the normally wobble ...

# LLM (ReAct agent) Analysis Report (see previous slide)

# Analysis Report: Mutation Impact on Wobble Taurine Modification

## ## 1. SEQUENCE ALIGNMENT ANALYSIS

The alignment analysis reveals a **single nucleotide mutation** between the query and target sequences:

- **Position 14**: A → G mutation (A3243G in mitochondrial nomenclature)
- **Sequence context**: CAGAGCC (target) → CAGGGCC (query)
- **Overall similarity**: 98.72% identity (77/78 matches)

## ## 2. STRUCTURAL COMPARISON

### ### Secondary Structure Analysis

Both sequences maintain the same overall secondary structure:

- **Dot-bracket notation**: Both show identical folding patterns with the same stem-loop architecture
- **RNA length**: 78 nucleotides for both sequences
- **Base pairing**: The mutation does not disrupt the major stem regions

### ### Hairpin Loop Differences

The mutation affects the first hairpin loop:

- **Target (healthy)**: CAGAGCCCGGUAAUCG
- **Query (mutated)**: CAGGGCCCGGUAAUCG

### ### Stacking Interaction Changes

The mutation alters local stacking interactions:

- **Target**: Contains UC and UA stacking patterns
- **Query**: Shows altered stacking with different GG-rich regions

### ### Multiplet Formation

Significant changes in three-nucleotide interactions:

- **Target**: UAA, GAC, GUAA multiplets
- **Query**: UGA, GAC, GCU multiplets

## ## 3. MOLECULAR MECHANISM EXPLANATION

### ### Why the A→G Mutation Prevents Taurine Modification

1. **Wobble Position Impact**: The A3243G mutation occurs in the mitochondrial tRNA<sup>Leu(UUR)</sup> and affects the structural context required for taurine modification at the wobble position (position 34).
2. **Enzyme Recognition Disruption**:
  - The mutation alters the local RNA structure and base stacking patterns
  - GTPBP3 and MTO1 enzymes, which cooperatively catalyze 5-taurinomethyluridine (tm5U) biosynthesis, require specific structural motifs for recognition
  - The changed multiplet patterns (UAA→UGA) disrupt the enzyme binding interface
3. **Structural Constraint Changes**:
  - The hairpin loop alteration (CAGAGCC→CAGGGCC) changes the local electrostatic environment
  - Modified stacking interactions affect the accessibility of the wobble position
  - The GG-rich region in the mutant may create stronger internal base pairing that prevents enzyme access

## ## 4. FUNCTIONAL CONSEQUENCES

### ### Translation Defects

- **Codon Recognition**: The absence of tm5U modification specifically affects UUA codon decoding
- **Aminoacylation**: Reduced tRNA<sup>Leu(UUR)</sup> aminoacylation capacity
- **Mitochondrial Protein Synthesis**: Decreased translation of mitochondria-encoded proteins

### ### Disease Pathogenesis

- **MELAS Syndrome**: This mutation is found in ~80% of MELAS patients
- **Oxidative Stress**: Increased ROS production due to respiratory chain deficiency
- **Energy Metabolism**: Impaired ATP synthesis leading to lactic acidosis

## ## 5. CONCLUSION

The A→G mutation at position 14 (corresponding to m.3243A>G) prevents wobble taurine modification by:

1. **Disrupting enzyme recognition sites** through altered multiplet formation patterns
2. **Changing local RNA structure** in the critical hairpin loop region
3. **Modifying base stacking interactions** that are essential for GTPBP3/MTO1 enzyme binding
4. **Creating unfavorable structural constraints** that prevent access to the wobble position

This structural analysis explains why this single nucleotide change has such profound consequences for mitochondrial tRNA function and ultimately leads to the MELAS phenotype through defective protein synthesis.

# OUTLINE

## ❖ Dataset

- What is RNA molecule
- RNACentral database

## ❖ Tools description

- RNA alignment Tool
- RNA 3D structure parameters Tool
- The structure-guided feedback Tool

## ❖ LLM ReAct-Agent Computational Pipeline

### ❖ Results for LLM ReAct-Agent

- Comparison of two RNAs
- Generate two connected RNA hairpins without RNA template
- Generate two connected RNA hairpins from a given RNA template
- Targeted modification of a specific region of an RNA molecule

### ❖ Example use case

- Single point mutation in tRNA associated with MELAS syndrome

## ❖ Conclusions

## ❖ What's next ?



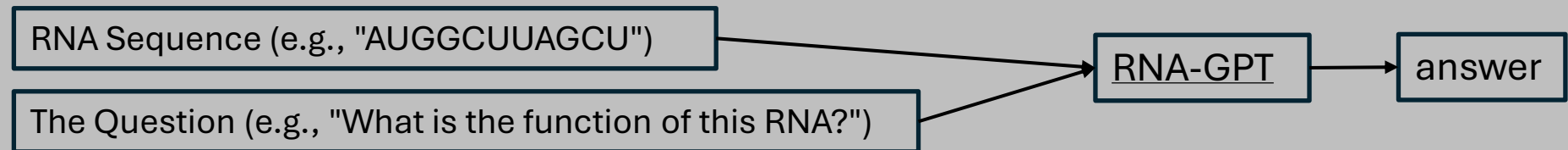
# CONCLUSIONS

- ❖ Although an LLM alone **cannot reliably count** nucleotides without external tools, a simple LLM agent is already capable of effectively answering questions related to non-coding RNA sequences.
- ❖ LLM REAct agent can be used not only for de novo ncRNA sequence **generation** but also for **modifying** existing sequences, providing a potential advantage over conditional diffusion models in guided ncRNA design tasks.
- ❖ A backfolded sequence turns out to be an effective, **LLM-comprehensible** alternative to 3D structural parameters for preserving the original structure of an input (template) sequence during partial modification.

# 1. What's next ?

1. Search for mistakes in the answers and correct the prompt accordingly
2. Comprehensive literature research
3. More tools, leading toward a multi-agent pipeline

## 4. Use RNA-GPT to construct a new tool based on answering questions about given RNA sequence.



The **RNA-FM sequence encoder**, which accounts for 3D structure, **was used** to embed RNA sequences for alignment with natural language, **at this time, RNA-GPT is not available for use**

Yijia Xiao et al. , RNA-GPT: Multimodal Generative System for RNA Sequence Understanding, **2025**

## 5. Fundamental Problems (see next slide) ?

- The oldest part of ribosome: pseudosymmetrical region (~90+90nt)
- The Spiegelman monster: the shortest “self-replicating” RNA molecule (~220nt)

large ribosomal subunit (LSU) rRNA

**the oldest functional part, conserved throughout evolution**



## Last Universal Common Ancestor (LUCA)

**E**

- Phase 6
- Phase 5
- Phase 4
- Phase 3
- Phase 2
- Phase 1

LSU and SSU associated with the end of Phase 3

LSU and SSU evolved independently & uncorrelated through Phases 1-3

PTC Phase

Phase 3

Phase 4

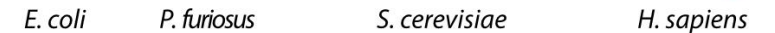
Phase 5

Phase 6

**LSU of tRNA (E. coli) :**

### LSU of tRNA (E. coli):

**Blue part is the oldest one**



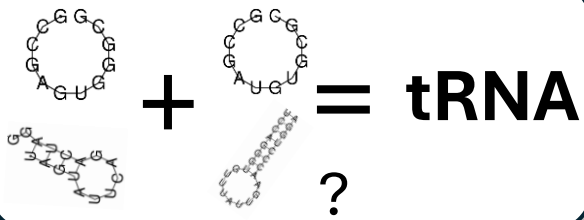
**Peptidyl Transferase Center (PTC) is the oldest part of ribosomes and contains no proteins**

This symmetry (**SymR**) suggests that the ancient ribosome may have been **a dimer of identical or nearly identical RNA molecules**, later evolving into the asymmetrical modern ribosome with **PTC**.

# 3. Peptidyl Transferase Center (PTC) Sequences

## the idea

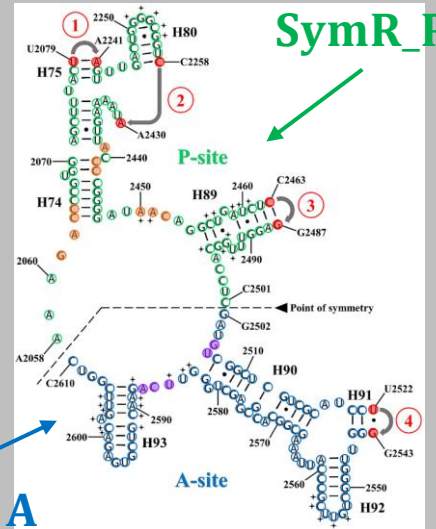
the dimerization of two similar RNA structures



“The peptidyl transferase center (PTC) evolved from a primitive system in the RNA world comprising tRNA-like molecules formed by **duplication of minihelix-like small RNA**”

Tamura, J. Biosci, 2011

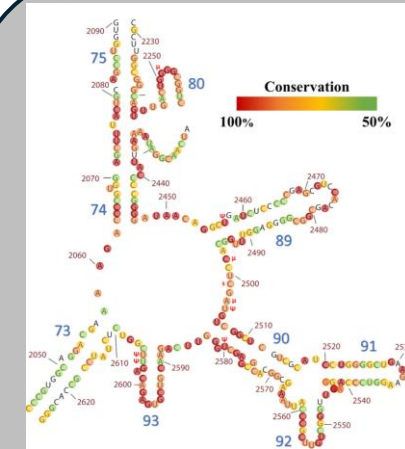
## pseudosymmetrical region



$$\text{SymR}_{\text{PA}} = \text{SymR}_{\text{P}} + \text{SymR}_{\text{A}}$$

Secondary structure of the pseudosymmetrical region (**SymR**; Agmon et al., 2005), derived from the LSU secondary structure of *Thermus thermophilus* (Petrov et al., 2013). (Madhan R. Tirumalai et al., 2021)

## PTC



PTC2 = red  
PTC3 = PTC2 + orange  
PTC4 = PTC3 + yellow  
PTC5 = PTC4 + green

**Nucleotide CONSERVATION level:**  
Red circles: 100% conservation (78 nt).  
Orange circles: 90 to 99.9% conservation (68 nt)  
Yellow circles: 70 to 89.9% (52nt)  
Green circles: 50 to 69.9% conservation (49nt)  
Black letters: less than 50% conservation (35nt)

(Bernier et al.; Faraday Discuss, 2014)  
(Madhan R. Tirumalai et al., 2021)

**SymR\_P** is older than **SymR\_PA**  
**PTC2** is older than **PTC3**, **PTC4**, **PTC5**