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Research Article

Highlighting the Role of Cdk6 Associated MicroRNAs in Cancer Treatment Using *In Silico* Approaches

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ABSTRACT

MicroRNAs (miRNAs) are small non-coding RNA's that controls the regulation of a gene. Due to the over expression or under expression of miRNAs it leads to cause tumor or any other type of cancers such as, melanoma, lymphoma, cardiovascular issue, breast cancer etc. So, miRNAs can be used as a drug target for cancer therapy. This study aimed to check binding cavities of microRNA's involved in regulation of CDK6 protein. There are 23 different families of miRNAs that are involved in regulation of CDK6. Each family has one or more miRNAs. All these miRNAs are involved in the up regulation or downregulation of a gene, which lead to different type of cancers. All miRNAs of each family docked with mRNA CDK6 protein. After performing in silico analysis of binding interactions of mRNA with miRNAs the results were further refined by their comparison with information regarding their energies, interaction of the mRNA and miRNAs. The results show that all miRNAs lie in Protein Kinase domain, but the residues that lie is different within the families and across the families.

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Introduction

MicroRNAs (miRNAs) are non-coding RNAs (ncRNAs) that control gene expression. miRNAs range from 17 to 24 nucleotide (nt) long RNAs. It has been known that miRNAs had ability to increase the expression of a target mRNA. Every single miRNA may target many different transcripts [1]. The alterations in miRNAs are associated with different human ailments, including cancer, immune disorders or cardiovascular issue. Their abnormalities are connected to start, leading towards metastases of human cancer. In human tumors it was found in no less than two autonomous reports distributed in the vicinity of 2004 and 2009 that 192 miRNAs were strangely communicated in tumor cells, including 168 overexpressed miRNAs, implying that high expression of miRNAs is a sign of harmful phenotype [2, 3]. As a treatment the principle RNA hindrance operators utilized till now in pre-clinical and clinical investigations include: antisense oligonucleotides (ASOs),

ribozymes and the DNazymes, small interfering RNAs (siRNAs) and short hairpin RNAs (shRNAs), and hostile to miRNA agents, for example, ASOs-against miRNAs, and Locked nucleic acids (LNA) against miRNAs or antagomirs [2, 3].

The process to discover any drug is costly and lengthy. In the present era by the use of computers it becomes easier to improve the traditional strategies for RNA target lead identification [4-6]. There are different programs used which is called docking programs i.e. Auto Dock and Dock, it has been used for predicting the binding sites and affinities of ligands. Different RNA could be targeted with small molecules including ribosome, tRNA, mRNA, and so forth. The 3D structure of miRNA is essential for the in-silico studies. Once the miRNA structure is acquired, molecular docking-based virtual high-throughput screening (vHTS) strategies will be utilized to improve the miRNA drug discovery process in light of RNA-perfect scoring functions, in which it is

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The aim of study was to perform in silico analysis of the binding interactions of mRNA and miRNAs is to know the binding sites of miRNA that binds with mRNA and with their ligands.

According to the literature there are many genes that are involved in the cancer. The genes that are involved in cancer causing in humans are

mostly the Cell Dependent Kinases (CDKs) present in Cell Cycle. There are different genes that are present in CDKs at various phases. From the 38 genes checked in Diana Micro T web server, only CDK2, CDK4, CDK6, CDK9 and ATM have miRNAs that are associated with these genes, while rest of the genes does not find any miRNA that are associated with them. Among these genes CDK6 is the most significant gene that are involved in cancer. Cell Dependent Kinase 6 (CDK6) gene is present on the chromosome 7 in human. It has 326 amino acid residues which are encoded by 978 nucleotide bases. Table 1 shows the important regions of CDK6 while (Figure 1) graphically represent the features of CDK6 gene.

Feature keys	Positions	Descriptions
Protein Kinase Domain	13-300	CDK6
Binding site	43	ATP
Active site	145	Proton Acceptor
Nucleotide binding	19-27	ATP
Chain	1-326	CDK6

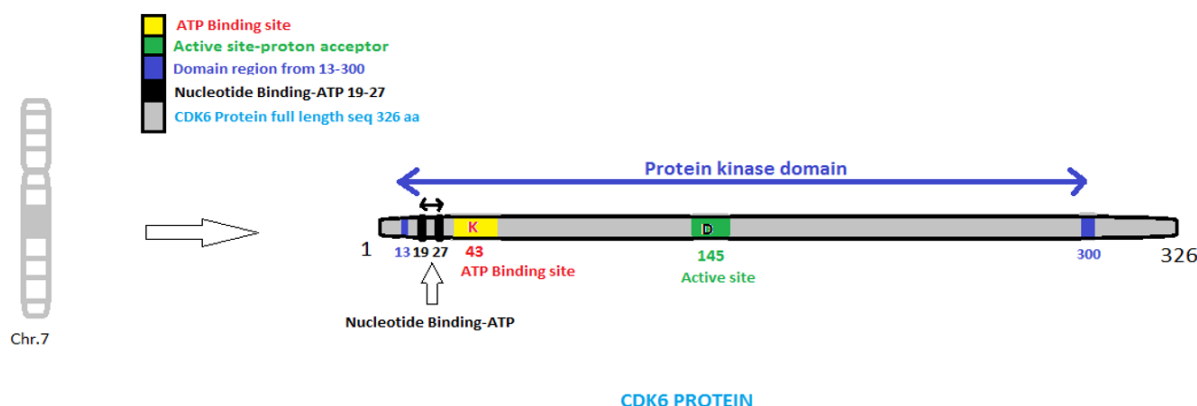


Figure 1: Structure of CDK6 Gene.

All miRNAs that are associated with CDK6 and involved in cancer causing to be searched by web server called Diana micro T version 5.0. There are 23 different miRNAs family that are involved in the regulation

of CDK6 gene. These families have one or more than one miRNAs. All the miRNAs from each family are almost involved in the same pathways. The following (Table 2) shows miRNA's families their members and the pathways in which miRNA's are involved.

Sr #	Family	Members(miRNAs)	Pathways
1	hsa-miR-1305/hsa-miR-1305	hsa-miR-1305	P53 signaling pathway
2	hsa-miR-548c-5p/ miR-548-5p/559	hsa-miR-548a-5p/hsa-miR-548ab/vhsa-miR-548ad-5p/hsa-miR-548ae-5p/hsa-miR-548ak/hsa-miR-548am-5p/hsa-miR-548ap-5p/hsa-miR-548aq-5p/hsa-miR-548ar-5p/hsa-miR-548as-5p/hsa-miR-548au-5p/hsa-miR-548ay-5p/hsa-miR-548b-5p/hsa-miR-548bb-5p/hsa-miR-548c-5p/hsa-miR-548d-5p/hsa-miR-548h-5p/hsa-miR-548i/hsa-miR-548j-5p/hsa-miR-548o-5p/hsa-miR-548w/hsa-miR-548y/hsa-miR-559	Viral carcinogenesis Non-small cell lung Pathways in cancer Chronic myeloid Hepatitis B Melanoma Pancreatic
3	hsa-miR-4429 /miR-320/4429	hsa-miR-320a/hsa-miR-320b/has-miR-320c/hsa-miR-320d/hsa-miR-4429	Leukemia Cell cycle
4	hsa-miR-26b-5p/miR-26-5p/1297/4465	hsa_miR-26a-5p/hsa_miR-26b-5p/hsa_miR-1297/hsa_miR-4465	Measles Glioma
5	hsa-miR-548k/miR-548-5p/8054	hsa miR-548av-5p/hsa miR-548k/hsa miR-8054	Cancer

6	hsa-miR-107/miR103-3P/107	hsa-miR-103a-3p/hsa-miR-107
7	hsa-miR-21-3p/miR-21-3p/3591-p	hsa-miR-21-3p/hsa-miR3-591-3p
8	hsa-miR-590-3p/miR-590-3p	has-miR-590-3p
9	hsa-miR-362-3p/miR-329-3p/362-3p	hsa-miR-329-3p/has-miR-362-3p
10	hsa-miR-335-3p/miR-335-5p	hsa-miR-335-3p
11	hsa-miR-33a-5p/miR-33-5p	hsa-miR33a-5p/hsa-miR33b-5p
12	hsa-miR-3140-3p/miR-3140-3p	hsa-miR-3140-3p
13	hsa-miR-3619-5p/miR-214-3p/761/3619-5p	hsa-miR-214-3p/has-miR-761/hsa-miR-3619-5p
14	hsa-miR-32-3p/miR-32-3p	hsa-miR-32-3p
15	hsa-miR-548e-3p/miR-548ar-3p/548ar-3p/548az-3p/548e-3p/548f-3p	hsa-miR-548a-3p/hsa-miR-548ar-3p/hsa-miR-548az-3p/hsa-miR-548e-3p/hsa-miR-548f-3p
16	hsa-miR-576-5p/miR-576-5p	hsa-miR-576-5p
17	hsa-miR-548o-3p/miR-548-3p/1323	hsa-miR-548o-3p/hsa-miR-1323
18	hsa-miR-320e/miR-320	hsa-miR-320e
19	hsa-miR-424-5p/miR-15-5p/16-5p/195-5p/424-5p/497-5p/6838-5p	hsa-miR-15a-5p/hsa-miR-15b-5p/hsa-miR-16-5p/hsa-miR-195-5p/has-miR-424-5p/hsa-miR-497-5p/hsa-miR-6838-5p
20	hsa-miR-450b-5p/miR-450-5p	hsa-miR-450b-5p
21	hsa-miR-142-5p/miR-142-5p/5590-3p	hsa-miR-142-5p/hsa-miR-5590-3p
22	hsa-miR-29b-3p/miR-29-3p	hsa-miR-29a-3p/hsamiR-29b-3p/hsa-miR-29c-3p
23	hsa-miR-589-3p/miR-589-3p	hsa-miR-589-3p

II miRNA 2D and 3D Structure Prediction

miRNAs 3D structure is predicted by using a web server called RNA Composer (Link) [8]. That server predicts the miRNA tertiary structure by giving input sequence of it. The sequence of miRNAs is present in a database called miRbase (Link) [9]. The tertiary structure of mRNA (CDK6) is also predicted by RNA Composer and using some more additional steps for its prediction, due to a large sequence. Centroid Fold is used for predicting secondary structure of RNA. It is used for predicting the secondary structure of all miRNAs. The following (Figure 2) shows the predicted secondary structure of miRNAs.

III Docking Studies

CDK6 was docked with all the miRNAs of each family for identifying the residues involved in hydrogen bonding and hydrophobic interaction within the families. For this purpose, Auto dock 4 software is used in which all the torsion angles in the small molecules were set free to perform flexible docking [10]. The grid was placed on CDK6 macromolecule that should cover the whole macromolecule and be large

enough to allow the ligand to fully rotate and the empirical free energy function and the Lamarckian genetic algorithm (LGA) were used for docking where number of runs and population size were set to 50 [11]. Out of 23 families, family 4 has a miRNA which has the least binding energy and is considered as best conformation. This PDB file is viewed in LigPlot+ for observing the residues involved in Hydrogen bonding as well as in hydrophobic interactions [12]. Chimera view is used for interactive representation of docking results [13].

Results

I miRNA Secondary Structure Prediction

Secondary structure of all 76 miRNAs of all families are predicted by using a CentroidFold web server. miRNAs have flanking regions, loop, stems and bulges in their structures. The following table shows the important regions of miRNAs which has been observed after predicting their secondary structure. The (Table 3) below shows the important regions of miRNAs that is present in their structure.

Table 3: Different regions present in secondary structure.

Sr #	Family	Members(miRNAs)	Flanking Regions	Stems	Loops	Bulges	Seed Region
1	hsa-miR-1305/ hsa-miR-1305	hsa-miR-1305	1-7/20-22	8-10/17-19	11-16	-	UUUCAAC
2	hsa-miR-548c-5p/ miR-548-5p/559	a) hsa-miR-548a-5p b) hsa-miR-548ab c) hsa-miR-548ad-5p d) hsa-miR-548ae-5p e) hsa-miR-548ak f) hsa-miR-548am-5p	1-2/11-22 1-2/22 1-2/11-20 1-2/11-20 1-6/17-21 1-2	3-4/9-10 3-4/9-13/9-21 3-4/9-10 3-4/9-10 7-9/14-16 3-4/9-10/13-15/20-22	5-8 5-8/14-18 5-8 5-8 10-13 5-8/16-19	- - - - - 11-12	AAAGUAA

		g) hsa-miR-548ap-5p	1-2/11-19	3-4/9-10	5-8	-	
		h) hsa-miR-548aq-5p	1-2/22	3-4/9-12/20-21	5-8/13-19	-	
		i) hsa-miR-548ar-5p	1-2	3-4/9-13/19-21	5-8/14-18	-	
		j) hsa-miR-548as-5p	1-2/22	3-4/9-12/20-21	5-8/13-19	-	
		k) hsa-miR-548au-5p	1-2	3-4/9-12/20-21	5-8/13-19	-	
		l) hsa-miR-548ay-5p	1-2/11-21	3-4/9-10	5-8	-	
		m) hsa-miR-548b-5p	1-2	3-4/9-10/13-15/20-22	5-8/16-19	11-12	
		n) hsa-miR-548bb-5p	1-12	13-15/20-22	16-19	-	
		o) hsa-miR-548c-5p	1-2	3-4/9-10/13-15/20-24	5-8/16-19	11-12	
		p) hsa-miR-548d-5p	1-2	3-4/9-10/13-15/20-22	5-8/16-19	11-12	
		q) hsa-miR-548h-5p	1-3/10-22	4/9	5-8	-	
		r) hsa-miR-548i	1-2/22	3-4/9-12/20-21	5-8/13-19	-	
		s) hsa-miR-548j-5p	1-2/11-22	3-4/9-10	5-8	-	
		t) hsa-miR-548o-5p	1-2	3-4/9-10/13-15/20-22	5-8/16-19	11-12	
		u) hsa-miR-548w	1-4/23	5/12-15/20-22	6-11/16-19	-	
		v) hsa-miR-548y	1-3/22	4/9/14-15/20-21	5-8/16-19	10-13	
		w) hsa-miR-559	1-4/15-21	5-7/12-14	8-11	-	
3	hsa-miR-4429/miR-320/4429	hsa-miR-320a	1-8/21-22	9/11-12/17-18/20	13-16	10/19	AAAGCUG
		a) hsa-miR-320b	1-8/21-22	9/11-12/17-18/20	13-16	10/19	
		b) hsa-miR-320c	1-10/19-20	11-12/17-18	13-16	-	
		c) hsa-miR-320d	1-10/19	11-12/17-18	13-16	-	
		d) hsa-miR-4429	1-9/20	10-12/17-19	13-16	-	
4	hsa-miR-26b-5p/miR-26-5p/1297/4465	a) hsa_miR-26a-5p	1-4/21	5/10-12/19-20	6-9/13-18	-	UCAAGUA
		b) hsa_miR-26b-5p	1-2/15-21	3-5/10-11/14	6-9	12-13	
		c) hsa_miR-1297	1-2/15-17	3-5/10-11/14	6-9	12-13	
		d) hsa_miR-4465	1-2/14-22	3/6/11/13	7-10	4-5/12	
5	hsa-miR-548k/miR-548-5p/8054	a) hsa_miR-548av-5p	1-7/14-18	8/10-11/13	---	-	AAAGUAC
		b) hsa_miR-548k	1-10/22	11-12/20-21	13-19	-	
		c) hsa_miR-8054	1-7/19-21	8-9/17-18	10-16	-	
6	hsa-miR-107/miR103-3P/107	a) hsa-miR-103a-3p	1-3/20-23	4-6/17-19	7-16	-	GCAGCAU
		b) hsa-miR-107	1-3/20-23	4-6/17-19	7-16	-	
7	hsa-miR-21-3p/miR-21-3p/3591-p	a) hsa-miR-21-3p	1-6/21	7-11/16-20	12-15	-	AACACCA
		b) hsa-miR-3591-3p	1-312-22	4-5/10-11	6-9	-	
8	hsa-miR-590-3p/miR-590-3p	hsa-miR-590-3p	1-9/18-21	10/17	11-16	-	AAUUUUA
9	hsa-miR-362-3p/miR-329-3p/362-3p	a) hsa-miR-329-3p	1-3/18-22	4/9-11/16-17	5-8/12-15	-	ACACACC
		b) hsa-miR-362-3p	1-6/18-22	7-9/15-17	10-14	-	
10	hsa-miR-335-3p/miR-335-5p	hsa-miR-335-3p	1-4/21-22	5-7/18-20	8-17	-	UUUUCAU
11	hsa-miR-33a-5p/miR-33-5p	a) hsa-miR33a-5p	1-2/21	3-5/7-9/14-16/18-20	10-13	6/17	UGCAUUG
		b) hsa-miR33b-5p	1-2	3-5/7-9/14-16/18-20	10-13	6/17	
12	hsa-miR-3140-3p/miR-3140-3p	hsa-miR-3140-3p	1/20-22	2-5/16-19	6-15	-	GCUUUUG
13	hsa-miR-3619-5p/miR-214-3p/761/3619-5p	a) hsa-miR-214-3p	1-3/20-22	4-5/18-19	6-17	-	CAGCAGG
		b) hsa-miR-761	1-4/18-22	5-7/15-17	8-14	-	
		c) hsa-miR-3619-5p	1/16-22	2-5/12-15	6-11	-	
14	hsa-miR-32-3p/miR-32-3p	hsa-miR-32-3p	-	-	1-22	-	AAUUUAG
15	hsa-miR-548e-3p/miR-548ar-3p/548ar-3p/548az-3p/548e-3p/548f-3p	a) hsa-miR-548a-3p	1-8	9-12/19-22	13-18	-	AAAACUG
		b) hsa-miR-548ar-3p	1-7	8-11/18-21	12-17	-	
		c) hsa-miR-548az-3p	1-7	8-11/18-21	12-17	-	
		d) hsa-miR-548e-3p	-	-	1-22	-	
		e) hsa-miR-548f-3p	-	-	1-19	-	
16	hsa-miR-576-5p/miR-576-5p	hsa-miR-576-5p	-	-	1-22	-	UUCUAAU
17	hsa-miR-548o-3p/miR-548-3p/1323	a) hsa-miR-548o-3p	1/22	2-6/17-21	7-16	-	CAAACU
		b) hsa-miR-1323	-	-	1-22	-	
18	hsa-miR-320e/miR-320	hsa-miR-320e	1-4/18	5-6/16-17	7-15	-	AAGCUGG
19		a) hsa-miR-15a-5p	1-6	7-10/19-22	11-18	-	AGCAGCA

	hsa-miR-424-5p/miR-15-5p/16-5p/195-5p/424-5p/497-5p/6838-5p	b) hsa-miR-15b-5p	1-3/16-22	4-5/14-15	6-13	-	
		c) hsa-miR-16-5p	1-5/22	6-7/20-21	8-19	-	
		d) hsa-miR-195-5p	1-2	3-4/10/15/20-21	11-14	5-9/16-19	
		e) has-miR-424-5p	1-4/21-22	5/10/12-13/19-20	69/14-18	11	
		f) hsa-miR-497-5p	1-2/15-21	3-6/11-14	7-10	-	
		g) hsa-miR-6838-5p	1-2/15-22	3-4/12-13	5-11	-	
20	hsa-miR-450b-5p/miR-450-5p	hsa-miR-450b-5p	-	-	-	1-22	UUUGCAA
21	hsa-miR-142-5p/ miR-142-5p/5590-3p	a) hsa-miR-142-5p	1-5	6-10/17-21	11-16	-	AUAAAGU
		b) hsa-miR-5590-3p	1-6/20-21	7-9/17-19	10-16	-	
22	hsa-miR-29b-3p/miR-29-3p	a) hsa-miR-29a-3p	1-4/21-22	5-7/18-20	8-17	-	AGCACCA
		b) hsa-miR-29b-3p	1-2/22-23	3-6/19-21	7-18	-	
		c) hsa-miR-29c-3p	1-4/21-22	5-7/18-20	18-17	-	
23	hsa-miR-589-3p/miR-589-3p	hsa-miR-589-3p	1-3/20-24	4-7/16-19	8-15	-	CAGAACA

It is noticed that all the families detailed in (Table 3) have bulges, loops, Flanking regions and stems in their structure. The flanking regions are found at the start and end of the structures while the bulges, loops and

stems are found in between. These secondary structures lie at different positions of the miRNA within and across the families.

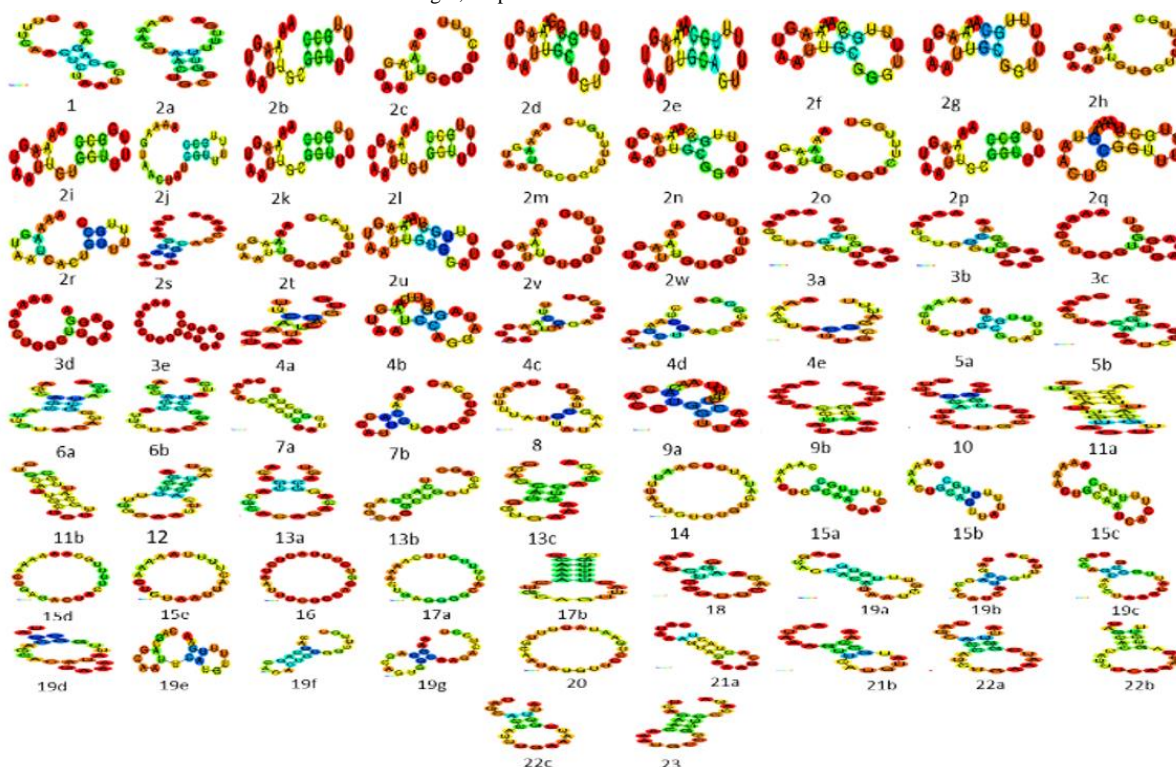


Figure 2: Secondary structure of miRNAs.

II Docking Analysis of All Families

All members of each family have a sequence of not less than 17 nucleotides and not more than 24. The energy values differ within the family and across the families as well, the energy values are not continuous. The protein residues present in all the families are not that much different. Most of the residues repeats in most of the miRNAs. The seed region, which is the important region of miRNAs, that interacts with mRNA is same within all the members of same family while it differs across the families. All the members of each family lie in the Protein

Kinase (PK) domain, which ranges from 13-300. The flanking regions, stems and loops are present in almost all of the miRNAs. The flanking regions must lie in the start and end of the structure, stem, loop and bulges are almost present between the flanking regions.

III mRNA and miRNA Residues Mapping

The (Table 4) has the only miRNAs from each family having least binding energy that interact with CDK6 residues and their region.

Table 4: Mapping of CDK6 residues and miRNAs.

Sr #	Family	miRNA	Energy	Protein Residues Mapping + mRNA Mapping	miRNA Residues Mapping
1	hsa-miR-1305/ hsa-miR-1305	hsa-miR-1305	-0.16	ARG90 = AGA + PK-Domain ARG87 = CGA + PK-Domain ARG78 = AGG + PK-Domain ARG186 = UAC + PK-Domain GLU72 = GAG + PK-Domain PHE135 = GAC + PK-Domain TYR292 = UCU + PK-Domain VAL181 = GUC + PK-Domain ARG144 = CAU + PK-Domain ALA167 = CUU + PK-Domain LEU166 = GGC + PK-Domain GLY165 = UUC + PK-Domain GLU21 = GAG + PK-Domain SER195 = UCC + PK-Domain SER194 = CAG + PK-Domain GLN193 = CUC + PK-Domain	UUUUCAACU A=6,7 U=1,2,3,4 C=5,8
2	hsa-miR-548c-5p/ miR-548-5p/559	hsa-miR-548ap-5P	-7.46	ARG46 = CGG + PK-Domain TYR24 = UAU + PK-Domain + NB-ATP LEU96 = UUA + PK-Domain GLN193 = CAG + PK-Domain SER195 = CUA + PK-Domain TYR196 = CGC + PK-Domain ARG144 = CGC + PK-Domain PRO244 = UAG + PK-Domain	AAAAGUAAU A=2,3,4,7,8 G=5 U=6
3	hsa-miR-4429/miR- 320/4429	hsa-miR-320b	-1.2	VAL179 = GUG + PK-Domain ILE169 = AUC + PK-Domain SER171 = AGU + PK-Domain TYR170 = UAU + PK-Domain TRP184 = UGG + PK-Domain GLN103 = CAA + PK-Domain TYR185 = UAC + PK-Domain ARG168 = CGC + PK-Domain GLN149 = CAG + PK-Domain PRO17 = GCG + PK-Domain LYS216 = GCC + PK-Domain LEU152 = CUG + PK-Domain LEU166 = CUU + PK-Domain GLU18 = GAG + PK-Domain GLY20 = GGG + PK-Domain + NB-ATP ILE19 = AUC + PK-Domain + NB-ATP ALA17 = GCG + PK-Domain LYS111 = AAA + PK-Domain PHE164 = UUC + PK-Domain ASN150 = AAC + PK-Domain PRO148 = CCA + PK-Domain THR106 = ACC + PK-Domain ILE151 = AUU + PK-Domain ASP110 = GAU + PK-Domain HIS100 = CAU + PK-Domain ASP104 = GAC + PK-Domain LEU105 = UUG + PK-Domain VAL101 = GUC + PK-Domain ASP102 = GAU + PK-Domain	AAAAGCUGGG A=2,3,4 G=5,8 C=6 U=7

4	hsa-miR-26b-5p/miR-26-5p/1297/4465	hsa-miR-1297	-10.63	GLU21 = GAG + PK-Domain + NB-ATP LEU166 = CUU + PK-Domain ALA167 = GCC + PK-Domain THR107 = ACU + PK-Domain ASP110 = GAU + PK-Domain LYS111 = AAA + PK-Domain TYR196 = CGC + PK-Domain ALA197 = CAC + PK-Domain ARG140 = CGA + PK-Domain THR198 = CCC + PK-Domain HIS137 = CAU + PK-Domain SER138 = UCA + PK-Domain	UUCAAGUAA A=4,5,8 G=6 U=2,7 C=3
5	hsa-miR-548k/miR-548-5p/8054	hsa-miR-548av-5p	-6.29	ALA167 = GCC + PK-Domain GLN193 = CAG + PK-Domain ILE169 = AUC + PK-Domain VAL181 = GUC + PK-Domain ARG186 = AGA + PK-Domain ARG78 = AGG + PK-Domain HIS139 = CAC + PK-Domain GLU72 = GAG + PK-Domain SER139 = CAC + PK-Domain PHE135 = UUU + PK-Domain TYR292 = CAG + PK-Domain	AAAAGUACU A=2,3,4,7 G=5 U=6 C=8
6	hsa-miR-107/miR103-3P/107	hsa-miR-107	-1.2	VAL179 = GUG + PK-Domain GLN193 = CAG + PK-Domain SER155 = AGC + PK-Domain SER156 = AGC + PK-Domain TYR108 = UAC + PK-Domain LYS111 = AAA + PK-Domain VAL112 = GUU + PK-Domain THR121 = ACC + PK-Domain PRO113 = CCA + PK-Domain ARG140 = CGA + PK-Domain LEU94 = CUA + PK-Domain THR84 = ACA + PK-Domain THR92 = ACC + PK-Domain VAL47 = GUG + PK-Domain PRO118 = CCC + PK-Domain	AGCAGCAUU A=4,7 G=2,5 U=8 C=3,6
7	hsa-miR-21-3p/miR-21-3p/3591-p	hsa-miR-3591-3p	-5.8	GLN193 = CAG + PK-Domain SER194 = UAG + PK-Domain SER195 = CUA + PK-Domain ARG144 = CGC + PK-Domain ALA197 = CAC + PK-Domain TYR292 = CAG + PK-Domain	AAACACCAU A=2,3,5,8 G= -- U= -- C=4,6,7
8	hsa-miR-590-3p/miR-590-3p	hsa-miR-590-3p	-0.93	ARG140 = CGA + PK-Domain SER195 = CUA + PK-Domain ALA197 = CAC + PK-Domain ASN284 = CCC + PK-Domain ALA280 = UUU + PK-Domain ARG78 = AGG + PK-Domain GLY157 = GGA + PK-Domain SER156 = AGC + PK-Domain THR121 = ACC + PK-Domain PRO118 = CCC + PK-Domain GLU120 = GAA + PK-Domain	UAAUUUUUAU A=2,3,8 G= U=4,5,6,7 C=

9	hsa-miR-362-3p/miR-329-3p/362-3p	hsa-miR-329-3P	-2.64	GLU21 = GAG + PK-Domain +NB-ATP LEU166 = CUU + PK-Domain ALA167 = GCC + PK-Domain LYS111 = AAA + PK-Domain PRO113 = CCA + PK-Domain GLN193 = CAG + PK-Domain ARG144 = CGC + PK-Domain SER195 = CUA + PK-Domain VAL142 = GUG + PK-Domain TYR196 = CGC + PK-Domain ALA197 = CAC + PK-Domain THR198 = CCC + PK-Domain HIS137 = CAU + PK-Domain SER138 = UCA + PK-Domain ARG140 = CGA + PK-Domain	AACACACCU A=2,4,6, G= U= C=3,5,7,8
10	hsa-miR-335-3p/miR-335-5p	hsa-miR-335-3p	0.12	SER194 = UAG + PK-Domain ASP246 = UGU + PK-Domain GLN193 = CAG + PK-Domain VAL179 = GUG + PK-Domain ILE169 = AUC + PK-Domain TYR292 = CAG + PK-Domain	UUUUUCAUU A=7 G= U=2,3,4,5,8 C=6
11	hsa-miR-33a-5p/miR-33-5p	hsa-miR33a-5p	-0.81	ASP246 = UGU + PK-Domain GLN193 = CAG + PK-Domain SER194 = UAG + PK-Domain SER195 = CUA + PK-Domain VAL190 = GUC + PK-Domain ARG144 = CGC + PK-Domain VAL181 = GUC + PK-Domain GLU189 = GAA + PK-Domain ALA197 = CAC + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP GLY22 = GGC + PK-Domain + NB-ATP ARG140 = CGA + PK-Domain VAL142 = GUG + PK-Domain VAL141 = GUA + PK-Domain HIS143 = CAU + PK-Domain	GUGCAUUGUAGUUGCAUU GCA A=5,16 G=3,8,14,19 U=2,6,7,13,17,18 C=4,15
12	hsa-miR-3140-3p/miR-3140-3p	hsa-miR-3140-3p	-6.79	VAL16 = GUG + PK-Domain VAL17 = GCG + PK-Domain PRO74 = CCC + PK-Domain TYR292 = CAG + PK-Domain SER155 = AGC + PK-Domain SER156 = AGC + PK-Domain SER293 = UGC + PK-Domain LYS279 = GUG + PK-Domain SER290 = UGC + PK-Domain LYS111 = AAA + PK-Domain ALA286 = CAA + PK-Domain LYS287 = AAG + PK-Domain	AGCUUUUGG A= G=2,8 U=4,5,6,7 C=3
13	hsa-miR-3619-5p/miR-214-3p/761/3619-5p	hsa-miR-214-3p	-2.26	THR84 = ACA + PK-Domain LEU94 = CUA + PK-Domain VAL82 = GUG + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP TYR24 = UAU + PK-Domain + NB-ATP GLY22 = GGC + PK-Domain + NB-ATP ARG140 = CGA + PK-Domain VAL142 = GUG + PK-Domain VAL141 = GUA + PK-Domain	ACAGCAGGC A=3,6 G=4,7,8 U= C=2,5

				ARG144 = CGC + PK-Domain TYR196 = CGC + PK-Domain ARG186 = AGA + PK-Domain VAL181 = GUC + PK-Domain ILE169 = AUC + PK-Domain LEU166 = CUU + PK-Domain ALA167 = GCC + PK-Domain LYS111 = AAA + PK-Domain TYR108 = UAC + PK-Domain VAL112 = GUU + PK-Domain GLY116 = GGA + PK-Domain PRO113 = CCA + PK-Domain PRO115 = CCU + PK-Domain PRO118 = CCC + PK-Domain VAL117 = GUG + PK-Domain THR119 = ACU + PK-Domain	
14	hsa-miR-32-3p/miR-32-3p	hsa-miR-32-3p	-5.18	SER194 = UAG + PK-Domain ASP246 = CUA + PK-Domain GLN193 = CAG + PK-Domain LEU166 = CUU + PK-Domain ALA167 = GCC + PK-Domain ILE169 = AUC + PK-Domain ARG168 = CGC + PK-Domain TYR292 = CAG + PK-Domain	CAAUUUAGU A=2,3,7 G=8 U=4,5,6 C=
15	hsa-miR-548e-3p/miR-548ar-3p/548ar-3p/548az-3p/548e-3p/548f-3p	hsa-miR-548e-3p	-6.04	GLN193 = CAG + PK-Domain SER194 = UAG + PK-Domain SER195 = CUA + PK-Domain TYR196 = CGC + PK-Domain ALA197 = CAC + PK-Domain THR198 = CCC + PK-Domain ASP110 = GAU + PK-Domain LYS111 = AAA + PK-Domain	AAAAACUGA A=2,3,4,5 G=8 U=7 C=6
16	hsa-miR-576-5p/miR-576-5p	hsa-miR-576-5p	-0.47	SER19 = AUC + PK-Domain TYR196 = CGC + PK-Domain ALA197 = CAC + PK-Domain ILE169 = AUC + PK-Domain	AUUCUAAUU A=6,7 G= U=2,3,5,8 C=4
17	hsa-miR-548o-3p/miR-548-3p/1323	hsa-miR-1323	0.16	ARG140 = CGA + PK-Domain TYR196 = CGC + PK-Domain VAL142 = GUG + PK-Domain HIS137 = CAU + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP GLU21 = GAG + PK-Domain + NB-ATP LEU166 = CUU + PK-Domain GLY165 = GGC + PK-Domain ARG168 = CGC + PK-Domain GLN149 = CAG + PK-Domain ASP110 = GAU + PK-Domain LYS216 = GCC + PK-Domain ARG215 = AAA + PK-Domain GLU114 = GAG + PK-Domain THR267 = AGA + PK-Domain	CCAAAACUG A=3,4,5,6 G= U=8 C=2,7
18	hsa-miR-320e/miR-320	hsa-miR-320e	-9.15	LEU166 = CUU + PK-Domain ALA167 = GCC + PK-Domain ILE169 = AUC + PK-Domain LYS111 = AAA + PK-Domain PRO113 = CCA + PK-Domain	AAAGCUGGG A=2,3 G=4,7,8 U=6 C=5

				PRO118 = CCC + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP ARG144 = CGC + PK-Domain SER195 = CUA + PK-Domain VAL142 = GUG + PK-Domain TYR196 = CGC + PK-Domain THR198 = CCC + PK-Domain ARG140 = CGA + PK-Domain SER138 = UCA + PK-Domain	
19	hsa-miR-424-5p/miR-15-5p/16-5p/195-5p/424-5p/497-5p/6838-5p	hsa-miR-15b-5p	-5.65	VAL16 = GUG + PK-Domain TYR24 = UAU + PK-Domain + NB-TP ARG44 = CGC + PK-Domain ARG186 = AGA + PK-Domain ASP145 = GAU + PK-Domain HIS143 = CAU + PK-Domain ARG78 = AGG + PK-Domain GLU72 = GAG + PK-Domain LYS111 = AAA + PK-Domain SER155 = AGC + PK-Domain	UAGCAGCAC A=2,5,8 G=3,6 U= C=4,7
20	hsa-miR-450b-5p/miR-450-5p	hsa-miR-450b-5p	-1.22	ARG46 = CGG + PK-Domain VAL47 = GUG + PK-Domain GLN48 = CAG + PK-Domain ALA23 = GCC + PK-Domain LEU94 = CUA + PK-Domain TYR24 = UAU + PK-Domain + NB-ATP VAL82 = GUG + PK-Domain LEU96 = UUA + PK-Domain ARG144 = CGC + PK-Domain ARG186 = AGA + PK-Domain ASP145 = GAU + PK-Domain-Active Site LEU79 = UUG + PK-Domain ARG78 = AGG + PK-Domain GLU72 = GAG + PK-Domain VAL16 = GUG + PK-Domain LYS111 = AAA + PK-Domain THR154 = ACC + PK-Domain GLN18 = GAG + PK-Domain SER155 = AGC + PK-Domain GLY157 = GGA + PK-Domain SER156 = AGC + PK-Domain	UUUUGCAAU A=7,8 G=5 U=2,3,4 C=6
21	hsa-miR-142-5p/miR-142-5p/5590-3p	hsa-miR-142-5p	-3.36	VAL147 = AAA + PK-Domain GLN48 = CAG + PK-Domain VAL45 = GUG + PK-Domain LEU94 = CUA + PK-Domain TYR24 = UAU + PK-Domain + NB-ATP ASP246 = UGU + PK-Domain GLN193 = CAG + PK-Domain ALA248 = CCU + PK-Domain VAL82 = GUG + PK-Domain LEU96 = UUA + PK-Domain LYS43 = AAG + PK-Domain + ATP-Binding Site ASP163 = GAC + PK-Domain VAL247 = UGC + PK-Domain SER194 = CCU + PK-Domain LEU79 = UUG + PK-Domain VAL142 = GUG + PK-Domain ALA162 = GCU + PK-Domain	CAUAAAGUA A=2,4,5,6 G=7 U=3,8 C=

				ARG140 = CGA + PK-Domain HIS143 = CAU + PK-Domain LEU161 = CUC + PK-Domain VAL141 = GUA + PK-Domain LEU136 = CUU + PK-Domain VAL76 = GUG + PK-Domain	
22	hsa-miR-29b-3p/miR-29-3p	hsa-miR-29c-3p	-6.13	VAL47 = GUG + PK-Domain LEU94 = CUA + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP GLU21 = GAG + PK-Domain + NB-ATP GLY22 = GGC + PK-Domain + NB-ATP GLY165 = GGC + PK-Domain VAL181 = GUC + PK-Domain SER223 = AGA + PK-Domain ASP224 = UGU + PK-Domain ARG144 = CGC + PK-Domain ARG186 = AGA + PK-Domain ASP145 = GAU + PK-Domain LYS147 = AAA + PK-Domain VAL82 = GUG + PK-Domain ARG140 = CGA + PK-Domain ASP242 = CUG + PK-Domain THR198 = CCC + PK-Domain PRO199 = CGU + PK-Domain PHE283 = UAA + PK-Domain PRO238 = AGG + PK-Domain ARG288 = AAU + PK-Domain THR282 = AUU + PK-Domain ASN284 = CCC + PK-Domain PRO285 = AGC + PK-Domain ALA286 = CAA + PK-Domain LYS287 = AAG + PK-Domain ILE289 = AUC + PK-Domain LEU237 = CCC + PK-Domain GLY236 = ACU + PK-Domain LYS279 = GUG + PK-Domain	UAGCACCAU A=2,5,8 G=3 U= C=4,6,7
23	hsa-miR-589-3p/miR-589-3p	hsa-miR-589-3p	8.2	LEU94 = CUA + PK-Domain CYS83 = UGC + PK-Domain VAL82 = GUG + PK-Domain THR95 = ACU + PK-Domain ASP81 = GAU + PK-Domain LEU96 = UUA + PK-Domain VAL97 = GUG + PK-Domain LEU79 = UUG + PK-Domain GLN103 = CAA + PK-Domain VAL77 = GUC + PK-Domain ALA41 = GCG + PK-Domain PHE98 = UUU + PK-Domain PHE28 = UUC + PK-Domain LYS43 = AAG + PK-Domain + ATP-Binding Site LEU42 = UUG + PK-Domain ALA23 = GCC + PK-Domain + NB-ATP LYS26 = AAG + PK-Domain + NB-ATP THR106 = ACC + PK-Domain + NB-ATP GLU21 = GAG + PK-Domain + NB-ATP GLU18 = GAG + PK-Domain VAL27 = GUG + PK-Domain + NB-ATP	UCAGAACAA A=3,5,6,8 G=4 U= C=2,7

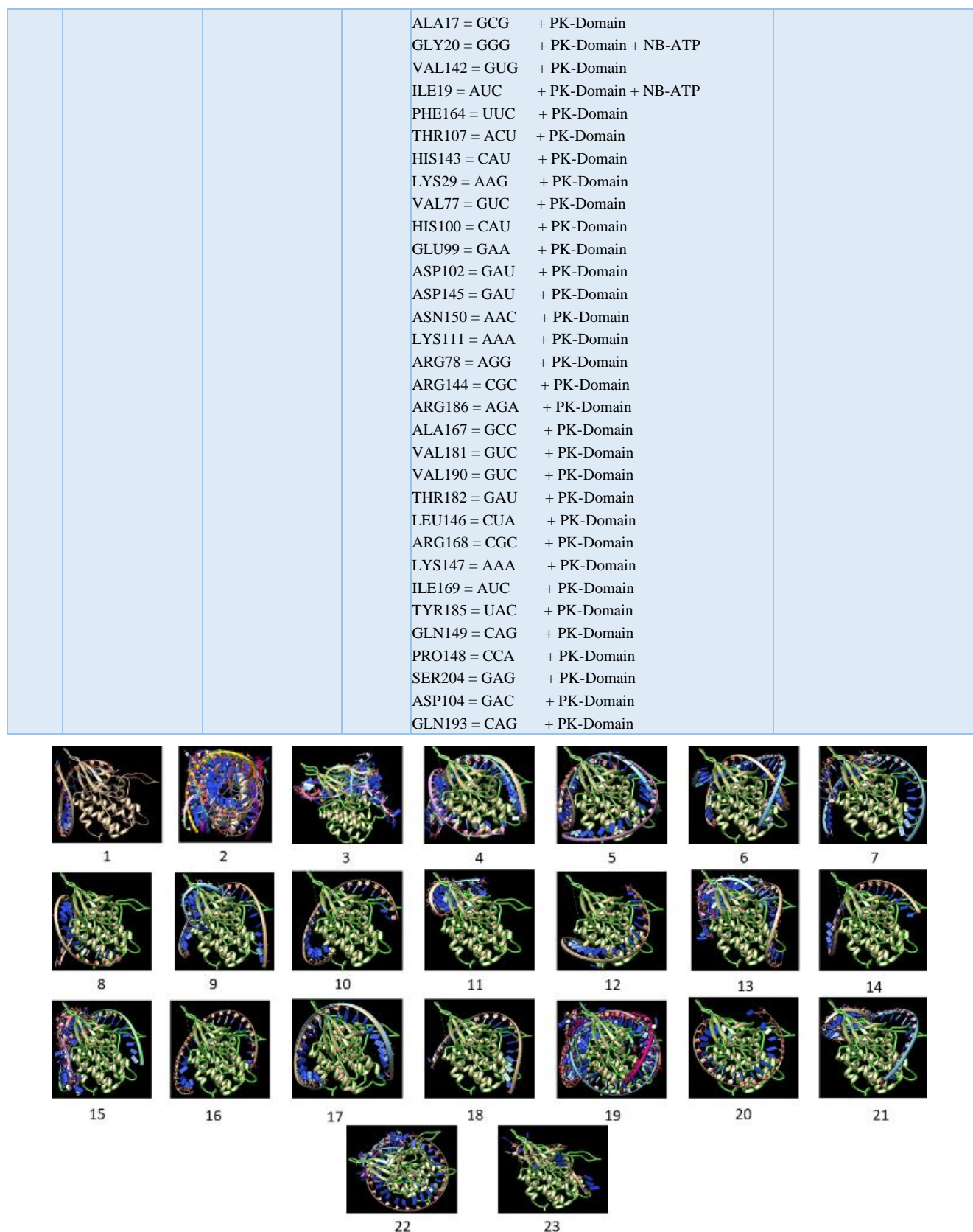


Figure 3: Shows interaction between mRNA (CDK6) with miRNAs of each family.

IV Observed Significant Features of mRNA and miRNAs after Docking

After the analysis of all the above results of mRNA and miRNAs all the complexes have different energy values, structures and their interacting residues. So, it is difficult to get only the most significant miRNA. So, a threshold value according to energy was taken as reference. miRNA which has the least energy value it will be the most significant miRNA because, miRNA with least energy value will have more interaction with ligands. The miRNA hsa-miR-1297 has the least energy value which is 27. So, this could be the significant miRNA and would be better to interact with ligands.

V Significant miRNA(hsa-miR-1297)

As discussed in the previous chapter the miRNA hsa-miR-1297 lies in the fourth family hsa-miR-26b-5p/miR-26-5p/1297/4465, this family

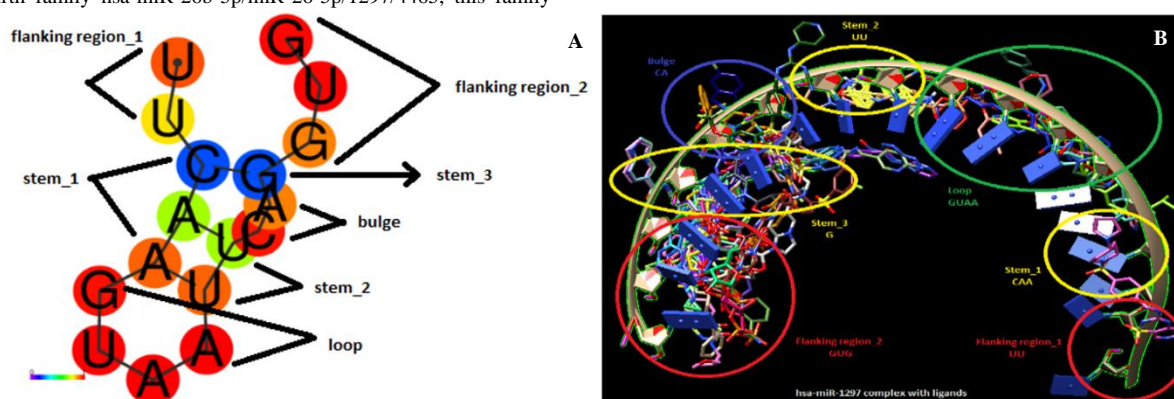


Figure 4: A) Secondary structure of hsa-miR-1297. B) 3D visualization of has-miR-1297 and its interaction with ligands.

Conclusion

This study focuses on microRNA's taking part in regulation of CDK6 gene and involved in Cancer. miRNA, hsa-miR-1297 belongs to the family 4 (hsa-miR-26b-5p/miR-26-5p/1297/4465), this family has four members (miRNAs), miRNA, hsa-miR-1297 has the least binding energy within the family and across the families. Hsa-miR-1297 is the most stable according to energy values to bind with ligands. The ligands are obtained from the PDB. The purpose of studying miRNAs in silico is to know the binding sites, behaviour, structure and the most important is the interaction with their targets. From the above study we came to know that miRNAs can be used as a drug targets, because it has high binding affinity towards their ligands.

Conflicts of Interest

None.

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