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THE INTERACTION OF miR-4258, miR-3960, miR-211-3p AND miR-3155b WITH mRNAs GENES OF NON-POLYGLUTAMINE TRINUCLEOTIDE DISORDERS

Abstract. Trinucleotide repeat expansion disorders constitute a group of dominantly inherited neurological diseases that are incurable and ultimately fatal. In the present work, miRNA binding sites were predicted by the MirTarget program. It was given characteristics of miRNAs binding sites in 5' and 3' UTR mRNAs genes of non-polyglutamine trinucleotide disorders with CGG, GCC, CUG repeats. Binding sites of 2567 miRNAs with mRNAs of 17494 human genes were determined. 206 genes with nucleotide repeats, mRNAs of which are bind with miRNA in the 5'UTR and 3'UTR, were observed. From thus, 2668 miRNAs binding sites are located in the 5'UTR, 3853 – in the 3'UTR with $\Delta G/\Delta G_m$ values equal to 85 % and more. It was found that 34 gene's mRNA having trinucleotide (CGG\GCC\CUG) repeats were targets for miR-4258, miR-3960 miR-211-3p and miR-3155b. miR-4258 binds to mRNA of *ADARB1*, *C11orf87* and *CBFB* genes with free binding energy - 93 kJ/mole and $\Delta G/\Delta G_m$ 91%, to mRNA of *ARHGEF7*, *BCR*, *BRSK2* and *C9orf91* genes with free binding energy - 91 kJ/mole and $\Delta G/\Delta G_m$ 89%. miR-3960 binds in GCC repeats to mRNA of *ABCC1* and *BLMH* genes with free binding energy - 116 kJ/mole. miR-211-3p and miR-3155b interact with mRNA of *ACACA* and *ANKRD13D* genes in 5'-3'untranslated regions. Studying binding characteristics of miRNA and genes will help identify association of miRNAs with genes with trinucleotide repeats for recommending for the diagnosis of nucleotide repeat expansion disorders.

Key words: miRNA, mRNA, binding site, trinucleotide repeat expansion.

Introduction. Trinucleotide repeats are sets of three nucleotides present in succession in various copy numbers throughout the human genome [1]. Repetitive sequences of genetic code are quite common. However, when these sequences grow beyond the scope of what would be considered normal, they cause disease. While the human genome has mechanisms to protect against these expansions, patients present with what can be severe neuromuscular and neurodegenerative disorders. There have been many diseases discovered by TNR (trinucleotide repeat) expansions, but the most prominent are spinocerebellar ataxia, Huntington disease, Fragile X syndrome, myotonic dystrophy, and Friedrich ataxia [2].

Small regulatory RNAs, particularly miRNAs, are known to be dynamically regulated in neurogenesis and brain development. Some recent studies have suggested that the alterations in small regulatory RNAs could contribute to the pathogenesis of several neurodevelopmental disorders [3,4]. miRNA refers to a small non-coding, single stranded RNA molecule comprising of around 22 nucleotides. By base pairing to messenger RNA (mRNA) and triggering translation repression, the miRNAs control gene

expression [5]. The use of miRNA as biomarkers to help diagnose neurodegenerative disorders offers several advantages. As the expression of miRNAs are commonly altered during disease, they have gained much attention for their potential use as biomarkers [6]. With better understanding of the role of miRNAs in neurodegenerative diseases, scientists and researchers may create effective new drugs to treat these devastating human illnesses. However, the biological function of most miRNAs remains to be uncovered [7, 8]. It is therefore important to provide characteristics of miRNA interaction with mRNA genes associated with non-polyglutamine trinucleotide disorders.

Materials and methods. The nucleotide sequences of mRNAs of human genes were downloaded from NCBI (<http://www.ncbi.nlm.nih.gov>). The nucleotide sequences of human miRNAs were downloaded from the miRBase database (<http://mirbase.org>). The miRNAs binding sites in mRNAs of several genes were predicted using the MirTarget program [9]. This program defines the following features of miRNA binding to mRNA: a) the start of the initiation of miRNA binding to mRNAs; b) the localization of miRNA BS in 5'UTRs, CDSs and 3'UTRs of the mRNAs; c) the free energy of interaction miRNA and the mRNA (ΔG , kJ/mole); d) the schemes of nucleotide interactions between miRNAs and mRNAs. For analyzing and formatting sequences of genes, we used the sequence manipulation suite program (<https://bioinformatics.org/sms>). To prediction the secondary structure of RNA, the software RNA fold was used (<http://rna.tbi.univie.ac.at>) [10].

Results and discussion. Using the MirTarget program, the binding sites of 2567 miRNA with the mRNA of 17494 human genes were determined. 206 genes with nucleotide repeats, the mRNAs of which are bind with miRNA in the 5'UTR and 3'UTR, were observed. 2668 miRNAs binding sites are located in the 5'UTR, 3853 – in the 3'UTR with $\Delta G/\Delta G_m$ values equal to 85 % and more. Only miR-4258, miR-3960 miR-211-3p and miR-3155b bind with 34 gene's mRNA having trinucleotide CGG, GCC, CUG repeats causing non - polyglutamine disorders. In table 1-3 are shown characteristics of miRNA binding with mRNA genes having trinucleotide repeats in 5'-UTR and 3'-UTR. The mRNA of *ABL2*, *ACVRIB*, *ADARB1*, *ADRBK1*, *APBA1*, *ARHGEF7*, *FMRI*, *B4GALT2*, *BCL11B*, *BCR*, *BRSK2*, *BRWD1*, *BTBD7*, *C11orf87*, *C9orf91*, *CACNA1A*, *CADM4*, *CAMK4*, *CARM1*, *CBFB*, *CBL* and *CCDC93* genes having trinucleotide repeats interact with miR-4258 in 5'-UTR in regions with CGG repeat. The binding sites of miR-3960 in mRNA of *ABCC1*, *ABCD3*, *AFF2*, *ANKH*, *ANKRD13D*, *BCL11A*, *BCL2L11*, *BLMH*, *C4orf19* and *CA10* genes are located in 5'-UTR in regions with GCC repeat. mRNA of *ACACA* and *ANKRD13D* genes interact with miR-211-3p and miR-3155b in 5'-UTR / 3'-UTR in regions with CUG repeat.

Table 1 – Characteristics of miR-4258 binding sites
in the 5'-UTR mRNA genes having CGG trinucleotide repeat

Gene	Beginning of binding site	ΔG , kJ/mole	$\Delta G/\Delta G_m$, %	Scheme of miRNA binding with mRNA genes
<i>ABL2</i>	21	-89	87	5' – CGCGGGCGGGUGGCAGGCC – 3' 3' – GGUUCCGCCACCGCC-CC – 5'
<i>ACVRIB</i>	46	-89	87	5' – CGCGGGCGGGUGGCAGGCC – 3' 3' – GGUUCCGCCACCGCC-CC – 5'
<i>ADARB1</i>	18	-93	91	5' – CCGUGGGGGCGGGCGGGCGG – 3' 3' – GGUUCCGCCACCGCC-CC – 5'
<i>ADRBK1</i>	7	-87	85	5' – CGCGGGCGGGCGGGCGG – 3' 3' – GGUUCCGCCACCGCC-CC – 5'
<i>APBA1</i>	50	-87	85	5' – UCCCGGGCGGGCGGGCGG – 3' 3' – GGUUCCGCCACCGCC-CC – 5'
<i>ARHGEF7</i>	155	-91	89	5' – GCGAGGGGGCGGGCGGGCGG – 3' 3' – GGUUCCGCCACCGCC-CC – 5'

<i>FMR1</i>	98	-87	85	5' - GCGC GGGCGGGCGGC GGCGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>B4GALT2</i>	133	-87	85	5' - CCCG GGGCGGGCGGC GGCG - 3' 3' - GGUU-CCGCCACCGCCCC - 5'
<i>BCL11B</i>	150	-87	85	5' - CGGCG GGGCGGGCGGC GGCG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>BCR</i>	200	-91	89	5' - CCGAGGAGGCGGCGGC GG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>BRSK2</i>	100	-91	89	5' - CCUC GGGCGGGCGGC GGCG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>BRWD1</i>	175	-87	85	5' - CGGCG GGGCGGGCGG GGCG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>BTBD7</i>	87	-87	85	5' - CGGCG GGGCGGGCGG UGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>C11orf87</i>	11	-93	91	5' - CGAAGGCGGCGGC GG CG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>C9orf91</i>	10)	-91	89	5' - CCGGGGU GGGCGGGCGG GG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>CACNA1A</i>	158	-87	85	5' - UCAG CGGCGGGCGGCGG CG - 3' 3' - GGUU-CCGCCACCGCCCC - 5'
<i>CADM4</i>	27	-87	85	5' - CGGCG GGGCGGGCGG CG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>CAMK4</i>	90	-87	85	5' - CGCGGGCGGCGGGCGGUGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>CARM1</i>	12	-87	85	5' - CAGCGGCGGCGGGCGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>CBFB</i>	39	-93	91	5' - CUGAGGCGGCGGGCGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'
<i>CBL</i>	14	-89	87	5' - CCGGCGGCGGGCGGCG - 3' 3' - GGUU-CCGCCACCGCCCC - 5'
<i>CCDC93</i>	33	-87	85	5' - CGGCGGGCGGCGGGCGG - 3' 3' - GGUUCCGCCACCGCC-CC - 5'

According to the table 1 only mRNA of *ADAR1*, *C11orf87* and *CBFB* genes have miR-4258 binding sites with free binding energy – 93 kJ/mole with the ΔG/ΔGm values equal to 91%. miR-4258 binds to CGG repeats of *ADAR1* gene in repeat nucleotide sequences CGG₁₀. The binding sites are located between 15 and 62 repeat nucleotide sequences with start in 18 nt. The binding sites of miR-4258 in mRNA of *C11orf87* and *CBFB* genes are located in repeat nucleotide sequences CGG₁₀/CGG₈ between 5-40 and 36-65 with start in 11 and 39 nt, respectively free binding energy equal to – 93 kJ/mole (figure 1). The secondary structures given in the figure clearly show the preferential formation of bonds with miR-4258.

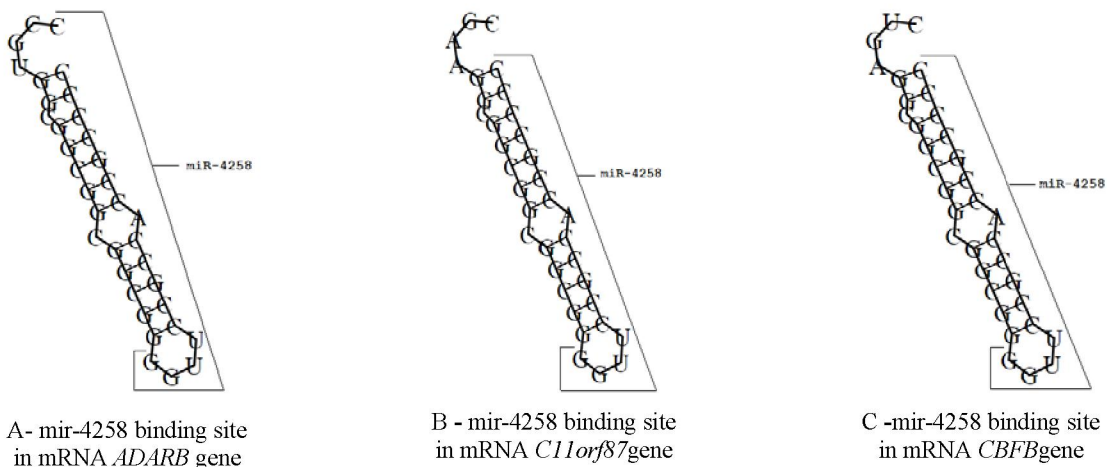


Figure 1 – Secondary structures of location of mir-4258 binding sites in 5' UTR mRNA of *ADARB*, *C11orf87* and *CBFB* genes

It was known that *ADARB1* gene encodes an RNA editing enzyme, expressed mainly in the central nervous system (CNS), which is involved in the downstream regulation of neuro transmitters. *ADARB1* (also known as *ADAR2*) spans a genomic region of 25 kb on chromosome 21q22.3 and comprises 16 exons and regulates its own expression through self-editing [11]. It was demonstrated that mRNA of *ARHGEF7*, *BCR*, *BRSK2* and *C9orf91* genes interact with miR-4258 with free binding energy – 91 kJ/mole and $\Delta G/\Delta G_m$ values equal to 89%.

Table 2 – Characteristics of miR-3960 binding sites in the 5'-UTR mRNA genes having GCC trinucleotide repeat

It can be seen from the table 2 that the free energy of the interaction of the miR-3960 with mRNA of *ABCC1*, *ABCD3*, *AFF2*, *ANKH*, *ANKRD13D*, *BCL11A*, *BCL2L11*, *BLMH*, *C4orf19* and *CA10* genes are constituted more than - 108 kJ/mole with $\Delta G/\Delta G_m$ values equal to 86 -91%. Among mRNA genes having nucleotide GCC repeats only *ABCC1* and *BLMH* genes bind with high free energy - 116 kJ/mole with miR-3960. miR-3960 binding sites are located in region with GCC_7 and GCC_8 repeats between 31 (beginning of binding sites) – 57 and 182 (beginning of binding sites) – 211 nt (figure 2). The secondary structures given in the figure clearly show the preferential formation of bonds with miR-3960. The mRNA gene of *ABCD3* interact with miR-3960 with full GCC_7 repeat. However, the binding sites of miR-3960 and mRNA genes of *ANKH* (GCC_5), *ANKRD13D* (GCC_4) and *BCL11A* (GCC_4) have only four and five GCC repeat.

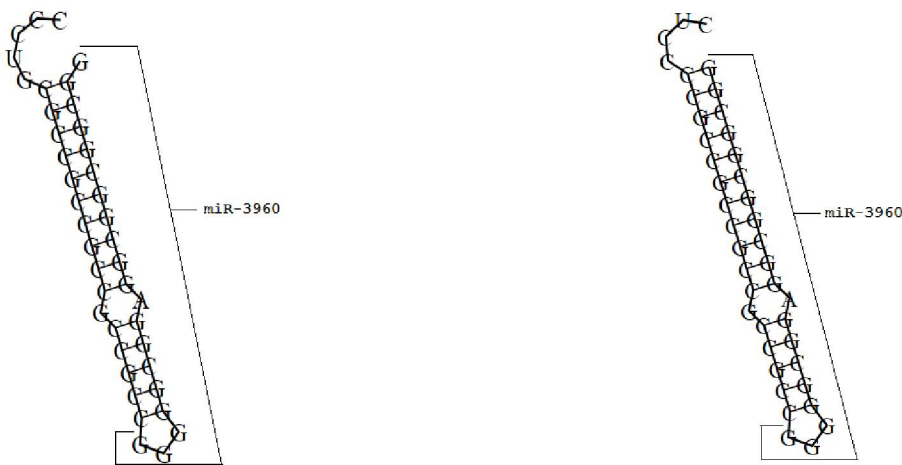


Figure 2 – Secondary structures of location of miR-3960 binding sites in 5'UTR mRNA of *ABCC1* and *BLMH* genes

ABCC1 (MRP1) is well known for its role in rendering cancer cells resistant to chemotherapy. ABCC1 is expressed in brain capillaries on the abluminal surface between the luminal membrane (CD31) and astrocytic end-feet (GFAP). Moreover, the free energy of miR-3960 interaction with mRNA gene of *ABCD3* indicate -114kJ/mole with $\Delta G/\Delta G_m$ values equal to 91%. *ABCD3* is one of the most abundant peroxisomal membrane proteins, at least in hepatocytes, and has been reported to be involved in the transport of various fatty acids. Mutation in *ABCD3* have been found in two individuals affected by Zellweger syndrome.

Table 3 – Characteristics of miR-211-3p, miR-3155b binding sites in 5'-UTR and 3'-UTR mRNA genes having CUG trinucleotide repeat

Gene	miRNA	Region	Beginning of binding site	ΔG , kJ/mole	$\Delta G/\Delta G_m$, %	Scheme of miRNA binding with mRNA genes
ACACA	miR-211-3p	5'UTR	61	-101	85	5' - GCGCGCCUGCUGCUGUCCCCGU - 3' 3' - CGUGGGGA-AACGACAGGGACG - 5'
ANKRD13D	miR-3155b	3'UTR	2056	-87	85	5' - UCUCUGCUGCUGAGCUUUGG - 3' 3' - AGGG-UGACGUCUCGGACC - 5'

From the table 3 obtained data indicate that miR-211-3p and miR-3155b interact with mRNA of *ACACA* and *ANKRD13D* genes in 5'-3' untranslated regions. miR-211-3p binding site in the 5'UTR mRNA of *ACACA* gene is located from 61 to 82 nt with (CUG)₃ repeat. miR-211-3p interact with high free energy -101 kJ/mole mRNA of *ACACA* gene.

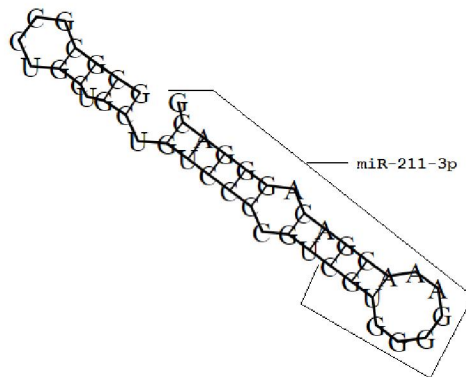


Figure 3 –
Secondary structures
of miR-211-3p binding sites
location in 5' UTR mRNA
of *ACACA* gene

Thus, the obtained results show that the greater number of genes are targets for miR-4258. miR-4258 binds to mRNA of 22 genes (*ABL2*, *ACVR1B*, *ADARB1*, *ADRBK1*, *APBA1*, *ARHGEF7*, *FMR1*, *B4GALT2*, *BCL11B*, *BCR*, *BRSK2*, *BRWD1*, *BTBD7*, *C11orf87*, *C9orf91*, *CACNA1A*, *CADM4*, *CAMK4*, *CARM1*, *CBFB*, *CBL* and *CCDC93*) with free binding energy -89 kJ/mole -(93 kJ/mole) and $\Delta G/\Delta G_m$ value from 85% to 91%. Moreover, the binding sites of miR-3960 in mRNAs of *ABCC1*, *ABCD3*, *AFF2*, *ANKH*, *ANKRD13D*, *BCL11A*, *BCL2L11*, *BLMH*, *C4orf19* and *CA10* genes have highest free binding energy from -108 kJ/mole to -116 kJ/mole and $\Delta G/\Delta G_m$ value from 86% to 93%. The maximum free energy of miR-3960 binding to mRNA is -116 kJ/mole. miR-3960 plays an important role in osteogenic transdifferentiation of vascular smooth muscle cells (VSMCs) and contributes to vascular calcification [12]. miR-3960 has 1100 binding sites on 375 target mRNAs with $\Delta G/\Delta G_m$ values of 90% or more and belong to a group of unique miRNAs [13].

Conclusion. In this paper, we have presented characteristics of predicted binding sites of miRNAs with mRNA genes of non - polyglutamine trinucleotide disorders. The most interesting data concern the analysis of target genes of miR-4258, miR-3960 miR-211-3p and miR-3155b. The identified associations of these miRNAs and target genes can be used to develop molecular methods for the neurological disease diagnosis. Also to date, there is a limited researches on nucleotide repeats. Therefore, further analysis using interaction of miRNAs with mRNA genes of all nucleotide repeats (di-, tri-, tetra-, penta-) including CDS region may be very useful to obtain advances knowledge.

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**ПОЛИГЛУТАМИНДІ ЕМЕС ТРИНУКЛЕОТИДТІК БҰЗЫЛЫСТЫ
ГЕНДЕРДІҢ Mrna-МЕН miR-4258, miR-3960, miR-211-3Р
ЖӘНЕ miR-3155b-ЛАРДЫң ӨЗАРА ӘРЕКЕТТЕСУІ**

Аннотация. Тринуклеотидтік қайталанымды экспансия бұзылыстары емделмейтін, ақыр сонында өлімге әкеліп соғатын неврологиялық түқым қуалайтын аурулар тобын құрайды. Жұмыс барысында miRNA-дың байланысатын сайттары MirTarget бағдарламасы арқылы болжанды. Бұл бағдарлама келесілерді аныктайды: miRNA-ның mRNA-мен байланыстыратын сайттарының басталуын; 5'UTR, CDS және 3'UTR-де mRNA сайттарының орналасуы; бос байланысу энергиясы (ΔG , кДж/моль) және miRNA-ның нуклеотидтерінің mRNA-мен әрекеттесу схемалары. Әр аймак үшін $\Delta G/\Delta G_m$ катынасы (%) есептелді, мұндағы ΔG_m нуклеотидтердің толық тізбегі бар miRNA-ның бос байланыс энергиясына тең. Гендердің барлық нуклеотидтік тізбегі GenBank-тан алынды (<http://www.ncbi.nlm.nih.gov>). miRNA нуклеотидтер тізбегі miRBase мәліметтер базасынан алынды (<http://www.mirbase.org>). miRNA-гендік экспрессияны транскрипциядан кейінгі реттеуде маңызды рөл атқаратын, кодташмаған РНҚ-ның үлкен тобы. CGG, GCC, CUG қайталанатын полиглутамидтік емес тринуклеотидті бұзылыстардың 5' және 3' UTR-де гендердің mRNA-мен miRNA-дың байланысатын сайттарына сипаттама берілді. 2567 miRNA-дың 17494 адам гендерінің mRNA-мен байланысатын

сайттары анықталды. 5'UTR және 3'UTR-де mRNA-лары miRNA-мен байланысқан нуклеотидтердің қайталауы бар 206 ген байқалды. Осылайша, 5'UTR-де 2668, 3'UTR-де 3853 miRNA-мен байланысатын сайттар 85 % және одан да көп ΔG / ΔGm мәндерде орналасқан. Тринуклеотидтік CGG, GCC, CUG қайталауы бар 34 геннің mRNA-лары miR-4258, miR-3960 miR-211-3р және miR-3155b-ларына нысан екендігі анықталды. *ADARB1*, *C11orf87* және *CBFB* гендердің mRNA-ларының бос энергиясы ΔG/ΔGm 91 % мәнінде – 93 кДж/моль-ға тең, сонымен қатар *ARHGEF7*, *BCR*, *BRSK2* және *C9orf91* гендердің mRNA-ларының бос байланысатын энергиясы ΔG/ΔGm 89 % мәнінде – 91 кДж / моль-ға тең, miR-4258-бен байланысады. *ABCC1* және *BLM1* гендерінің mRNA-ларының GCC қайталанымдары бос байланысатын энергиясы – 116 kJ/mole болатын miR-3960-бен байланысады. miR-3960 байланыстыру сайттары *GCC₇* және *GCC₈* қайталанатын 31 аймақта орналасқан (байланыстыру сайттарының басталуы) – 57 және 182 (байланыстыру сайттарының басталуы) – 211 nt. *GCC₇*-қайтальмының толық қайталауымен, *ABCD3* генінің mRNA-сы miR-3960-пен өзара әрекеттеседі. Дегенмен miR-3960 және *ANKH* (*GCC₅*), *ANKRD13D* (*GCC₄*) және *BCL11A* (*GCC₄*) гендерінің байланысу сайттарында тек төрт және бес GCC қайталауы бар. miR-3960-375 гендердің mRNA нысандарында ΔG/ΔGm 90 % улесінде 1100 сайтпен байланысады немесе одан жоғары нысандарға арналған байланысу сайттары бар және ерекше miRNA тобына жатады. 5' және 3'UTR-де *ACACA* және *ANKRD13D* гендерінің mRNA-лары miR-211-3р және miR-3155b-мен өзара әрекеттеседі. *ACACA* гендерінің mRNA-лары 5'UTR-де miR-211-3р-пен байланысуы 61-ден 82 нт аралығында (CUG)₃ қайталауымен орналасқан. miR-211-3р жоғары бос энергиямен mRNA-да – 101 кДж/моль есебімен *ACACA* генімен әрекеттеседі. miRNA-лармен гендердің байланысу сипаттарын зерттеу нуклеотидтік қайталанатын экспансиялық бұзылыстарды диагностикалауда miRNA-мен ассоциациясын анықтауға көмектеседі.

Түйін сөздер: miRNA, mRNA, байланысатын сайт, тринуклеотидті қайталанатын экспансия.

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ВЗАЙМОДЕЙСТВИЕ miR-4258, miR-3960, miR-211-3Р И miR-3155b С mRNA ГЕНОВ НЕПОЛИГЛУТАМИНОВЫХ ТРИНУКЛЕОТИДНЫХ РАССТРОЙСТВ

Аннотация. Расстройства экспансии тринуклеотидных повторов представляют собой группу доминантно-наследуемых неврологических заболеваний, которые неизлечимы и в конечном итоге приводят к летальному исходу. Изменение экспрессии miRNA считается отличительным признаком многих заболеваний, включая нарушения экспансии тринуклеотидных повторов. В настоящей работе сайты связывания miRNA были предсказаны программой MirTarget. Программа определяет: начало сайтов связывания miRNA с mRNA; расположение сайтов в 5'UTR, в CDS и в 3'UTR mRNA; свободную энергию гибридизации (ΔG , kJ/mole) и схемы взаимодействия нуклеотидов miRNA с mRNA. Для каждого сайта рассчитывали отношение $\Delta G/\Delta G_m$ (%), где ΔG_m равна свободной энергии связывания miRNA с полностью комплементарной нуклеотидной последовательностью. Все нуклеотидные последовательности mRNA генов заимствовали из GenBank (<http://www.ncbi.nlm.nih.gov>). Нуклеотидные последовательности miRNA получены из базы miRBase (<http://www.mirbase.org>). miRNA представляет собой большое семейство консервативных некодирующих РНК, играющих ключевую роль в посттранскрипционной регуляции экспрессии генов. Приведены характеристики сайтов связывания miRNA в 5'UTR и 3'UTR mRNA генов неполиглутаминовых тринуклеотидных расстройств с повторами CGG, GCC, CUG. Были определены сайты связывания 2567 miRNA с mRNA 17494 генов человека. Было обнаружено 206 генов с нуклеотидными повторами, mRNA которых связывались с miRNA в 5'UTR и 3'UTR. Таким образом, 2668 сайта связывания miRNAs расположены в 5'UTR, 3853 - в 3'UTR со значениями $\Delta G/\Delta G_m$, равными 85% и более. Было обнаружено, что mRNA 34 генов, имеющих тринуклеотидные CGG, GCC, CUG повторы, были мишенью для miR-4258, miR-3960 miR-211-3р и miR-3155b. miR-4258 связывается с mRNA генов *ADARB1*, *C11orf87* и *CBFB* со свободной энергией взаимодействия - 93 кДж/моль и $\Delta G/\Delta G_m$ 91%, с mRNA генов *ARHGEF7*, *BCR*, *BRSK2* и *C9orf91* со свободной энергией взаимодействия - 91 кДж/моль и $\Delta G/\Delta G_m$ 89%. miR-3960 связывается в повторах GCC с mRNA генов *ABCC1* и *BLM1* со свободной энергией взаимодействия - 116 кДж/моль. Сайты связывания miR-3960 расположены в области с повторами *GCC₇* и *GCC₈* между 31 (начало сайтов связывания) - 57 и 182 (начало сайтов связывания) - 211 нт. mRNA гена *ABCD3* взаимодействует с miR-3960 с полным повтором *GCC₇*. Однако сайты связывания генов miR-3960 и mRNA генов *ANKH* (*GCC₅*), *ANKRD13D* (*GCC₄*) и *BCL11A* (*GCC₄*) имеют только четыре и пять повторов GCC. miR-3960 имеет 1100 сайтов связывания на 375 mRNA - мишениях со значениями $\Delta G / \Delta G_m$ 90% и более и относится к группе уникальных miRNA. miR-211-3р и

miR-3155b взаимодействуют с mRNA генов *ACACA* и *ANKRD13D* в 5'-3'-нетранслируемых областях. Сайт связывания miR-211-3р в 5'UTR mRNA генов *ACACA* расположен от 61 до 82 нт с повторением (CUG)₃. miR-211-3р взаимодействует с mRNA высокой свободной энергии -101 кДж / моль гена *ACACA*. Изучение характеристик связывания miRNA и генов поможет выявить связь miRNA с генами с тринуклеотидными повторами для рекомендации для диагностики нарушений экспансии нуклеотидных повторов.

Ключевые слова: miRNA, mRNA, сайт связывания, экспансия тринуклеотидных повторов.

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