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ORIGINAL RESEARCH

MicroRNA Dysregulation in Prostate Cancer

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Abstract: Prostate cancer biology is complex, and needs to be deciphered. The latest evidence reveals the significant role of noncoding RNAs, particularly microRNAs (miRNAs), as key regulatory factors in cancer. Therefore, the identification of altered miRNA patterns involved in prostate cancer will allow them to be used for development of novel diagnostic and prognostic biomarkers. Patients and Methods: We performed a miRNAs transcriptomic analysis, using microarray (10 matched pairs tumor tissue versus normal adjacent tissue, selected based on inclusion criteria), followed by overlapping with TCGA data. A total of 292 miRNAs were differentially expressed, with 125 upregulated and 167 downregulated in TCGA patients' cohort with PRAD (prostate adenocarcinoma), respectively for the microarray experiments; 16 upregulated and 44 downregulated miRNAs were found in our cohort. To confirm our results obtained for tumor tissue, we performed validation with qRT-PCR at the tissue and plasma level of two selected transcripts, and finally, we focused on the identification of altered miRNAs involved in key biological processes. Results: A common signature identified a panel of 12 upregulated and 1 downregulated miRNA, targeting and interconnected in a network with the TP53, AGO2, BIRC5 gene and EGFR as a core element. Among this signature, the overexpressed transcripts (miR-20b-5p, miR-96-5p, miR-183-5p) and the downregulated miR-542-5p were validated by qRT-PCR in an additional patients' cohort of 34 matched tumor and normal adjacent paired samples. Further, we performed the validation of the expression level for miR-20b-5p, miR-96-5p, miR-183-5p plasma, on the same patients' cohort versus a healthy control group, confirming the overexpression of these transcripts in the PRAD group, demonstrating the liquid biopsy as a potential investigational tool in prostate cancer. Conclusion: In this pilot study, we provide evidence on miRNA dysregulation and its association with key functional components of the PRAD landscape, where an important role is acted by miR-20b-5p, miR-542-5p, or the oncogenic cluster miR-183-96-182.

Keywords: prostate adenocarcinoma, microRNA, tissue, plasma, molecular signature, biological network

Introduction

According to Globocan 2020, prostate cancer is the second most frequent malignancy (after lung cancer) worldwide, with 1,414,259 (14.1%) new cases, and causing 375,304 deaths (3.8% of all deaths caused by cancer in men).^{1,2} Around 20% of the new cases will develop invasive disease during their lifetime, most of the cases being diagnosed in advanced stages.³

Current prostate cancer management is based on a series of tests for non-invasive biomarkers.⁴ Till now, the serum markers are not sufficient for both early diagnostic and prognostic of prostate cancer. Patient stratification and identification of high-risk groups of prostate cancer have an important role in therapeutic decision.⁵

A better understanding of the prostate cancer molecular landscape will have an important role in the development of novel biomarkers or identification of therapeutic target.⁶ In this context, in the last decades, the evaluation of non-coding genes opened new avenues in cancer investigation.^{7–10}

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MicroRNAs (miRNAs) are small non-coding RNAs with 19–25 nucleotides in length that regulate the mRNA level of expression by canonical and non-canonical mechanisms^{9,11} and shown an important impact in cellular processes;^{12–14} also miRNAs have and important role in intercellular signaling.¹⁵ Dysregulation of miRNAs has been observed in several human malignancies,^{7,16,17} including prostate cancer. Increasing evidence indicates that the miRNAs can act as either oncogenes or tumor suppressor genes.

miRNAs are mentioned in literature as a potential application of biomarkers for numerous diseases,^{7,9,18} including prostate adenocarcinoma (PRAD).¹⁹ Despite this, there is a current lack of validated studies that evaluated the miRNA expression level and direct connection with clinical variables, particularly from the liquid biopsy.^{20,21} Recent literature data emphasizes the important role of liquid biopsy in patient stratification.^{20,22}

In the present pilot study, using microarray technology, we evaluated the miRNA particularities in PRAD on ten matched paired tumor tissues and normal adjacent tissues samples, followed by overlapping with TCGA data and identification of key pathways involved. The profiling study was followed by validation by qRT-PCR in an additional patient cohort comprising 34 matched PRAD samples, followed by an additional validation step from plasma samples from the same patients' cohort.

Consequently, to these analyses, we performed the exploration of the expression level for the most relevant target genes using UALCAN online tools. miRNA-gene interconnection was analyzed using Ingenuity Pathway Analysis (IPA) software. Our findings contribute to the molecular understanding of the tumorigenesis and for prognostic value in PRAD, and provide a set of miRNAs and their target genes that may be useful for the development of novel diagnostic biomarkers and/or more effective therapeutic strategies, particularly for the identification of high-risk groups of PRAD.

Materials and Methods

Human Subjects, Ethical Approval, and Clinical Data

We designed and performed a microarray experiment for microRNA evaluation on the matched paired samples of PRAD and their normal adjacent tissues (Table 1), age average of 63 ± 4.9 , followed by a validation step by qRT-PCR (Table 2, age average of 66 ± 4.9). Sample collection was done after the approval of the Ethical Committee of The Oncology Institute "Prof. Dr. Ion Chiricuta" no. 5991 from 26.06.2019, following protocols by national and European regulations concerning the use of biological samples and personal data: personal data were anonymized for all cases; all patients and healthy donors were informed about the study, and each provided signed informed consent.

No. Patient	Age	iPSA	тлм	Gleason Score	
I	64	7	20x	7(3+4)	
2	65	5.64	20x	6(3+3)	
3	62	4.94	20x	7(4+3)	
4	66	5.62	2xx	7(3+4)	
5	70	12	40x	7(4.3)	
6	53	31	3b0x	7(4+3)	
7	64	16	3b0x	7(3+4)	
8	56	34	3b0x	8(4+4)	
9	64	19	3b0x	7(4+3)	
10	66	18	3b1x	7(4+3)	

 Table I
 Clinical Data for the Patients Used for the Microarray Study Matched Paired Samples

 from 10
 Patients with PRAD

No. Patient	Age	iPSA	ТММ	Gleason Score
I	64	9.25	2XX	7(4+3)
2	54	19	3bIX	8(4+4)
3	71	4	2c0X	6(3+3)
4	75	-	3aXX	7(3+4)
5	63	7	3a0X	7(3+4)
6	68	13	30X	7(3+4)
7	64	6	2c0C	7(4+3)
8	70	5.5	2c0X	6(3+3)
9	64	47	3b0X	7(4+3)
10	64	7.45	2b0X	7(3+4)
11	60	9	2c0X	7(3+4)
12	65	7	20X	7(3+4)
13	76	5.53	2cXX	7(3+4)
14	68	15	3bIX	7(4+3)
15	68	-	2c0x	6(3+3)
16	65	78	2c0X	7(4+3)
17	61	10	2c0X	6(3+3)
18	68	46	-	9(5+4)
19	64	4.9	3a0X	7(3+4)
20	70	-	3bIX	7(3+4)
21	66	4.26	2cXX	7(3+4)
22	65	8.3	3aXX	7(3+4)
23	63	19.71	3b0X	9(4+5)
24	65	1.1	2cXX	6(3+3)
25	71	25	3a0X	7(4+3)
26	65	5.75	2c0x	7(3+4)
27	63	-	3aXX	7(3+4)
28	76	4.5	2XX	7(3+4)
29	69	8.9	3a00	7(3+4)
30	62	11	2c0X	6(3+3)
31	58	8.09	ЗВхх	7(4+3)

Table 2 Clinical Characteristics of PRAD Patients Used for the qRT-PCR Validation Study inTumor Tissue/Normal Adjacent Tissue and Plasma Samples

(Continued)

Table	2	(Continued).
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No. Patient	Age	iPSA	тлм	Gleason Score
32	64	9.5	20X	9(4+5)
33	70	1.9	3a0X	7(3+4)
34	74	-	20x	7(4+3)

The clinical characteristics for each patient enrolled in the microarray analysis (defined as UMPh patient cohort) are given in Table 1, meanwhile, the clinical data for the validation cohort of PRAD patients used for qRT-PCR from tissue (tumor/normal adjacent) and plasma are given in Table 2. For the qRT-PCR of plasma samples, we used male plasma samples from healthy controls (Table 3), considering the estimated age-specific incidence rates for prostate cancer.²³

Clinical Data of Patients from TCGA Data

The Cancer Genome Atlas (TCGA) is a public database developed to generate complex bioinformatics analysis with important implications in understanding the molecular portraits.²⁴ Therefore, we downloaded the data from the Firebrowse portal (<u>http://firebrowse.org/</u>), comprising data from 494 samples of PRAD and 52 adjacent normal tissue (Table 4).

No. Sample	Age	No. Sample	Age
Ι	44	16	45
2	49	17	41
3	48	18	61
4	41	19	58
5	50	20	53
6	41	21	43
7	58	22	41
8	41	23	44
9	54	24	40
10	43	25	40
11	41	26	41
12	51	27	53
13	41	28	49
14	49	29	57
15	40	30	46

Table 3 Plasma Samples from Healthy Men Selected as the ControlGroup for the Plasma of PRAD Patients Used for the Plasma qRT-PCRValidation Study

Demographics		PRAD (n=494)
AGE	Median, range	61, 41–78
Gleason score	6	45
	7	246
	8	64
	9	136
	10	3
PSA value	0	29
	0.01–323	410
	Unknown	55
т	Т2	187
	ТЗ	291
	Τ4	9
	Discrepancy	2
	Unknown	5
N	N0	344
	NI	78
	Unknown	72
М	M0	454
	МІ	3
	Unknown	37

Table 4	4 Clinical	Data for	· TGCA	Prostate	Adenocarcir	noma (PRA	D)
Patients	5						

MiRNA Analysis Conditions from TCGA

We analyzed the miRNAs using Gene Spring considering FC \pm 1.5 as cut-off value and p-value \leq 0.05.

RNA Extraction and Quality Control

The tissue samples (tumor and normal adjacent tissue) was extracted using Trireagent, and for the plasma was used Plasma/Serum Circulating and Exosomal RNA Purification Kit (Slurry Format) (Cat. 42800), followed by the evaluation of RNA concentration by the use of NanodropND2000 for microarray, respectively qRT-PCR validation step.

miRNA Microarray and Array Data Analysis Evaluation

For miRNA patterns, we used 10 tumor tissue and normal adjacent tissue matched paired PRAD samples. Starting from 100 ng of total RNA, the hybridization was done using Agilent microRNA Spike-In kit and the labeling miRNA Complete Labeling and Hyb Kit, followed by a purification step using Micro Bio-Spin 6 (Biorad, Mississauga, ON, Canada) spin columns according to producer recommended protocol. Also, for the hybridization, Agilent SurePrint Human miRNA v21.0 microarray slides (G4872A) have been used. After the 20 hours hybridization period in the oven at 55°C, the slides have been washed and scanned using an Agilent Microarray Scanner.

The bioinformatics analysis was done for matched paired PRAD samples using the same parameters as those used for TCGA data sets. The data were extracted using Agilent Feature Extraction software (Agilent Technologies); Agilent GeneSpring GX program (Agilent Technologies) was used to obtain differentially expression selected "Filter on Volcano Plot" analysis and moderated *t*-test, with a fold change of 1.5. Benjamini–Hochberg (false discovery rate-FDR) algorithm was applied to reduce the false-positive rates.

Ingenuity Pathway Analyses of Commonly Deregulated miRNAs in PRAD

The identified differentially expressed miRNAs in both patient cohorts (UMPh and TCGA) were uploaded to the IPA software (<u>https://digitalinsights.qiagen.com/products-overview/discovery-insights-portfolio/analysis-and-visualization /qiagen-ipa</u>, QIAGEN). The "core analysis" function of the software was used to analyze the differentially expressed miRNA data, which incorporated the recognition of biological processes, canonical pathways, upstream transcriptional regulators, and interconnected networks.

Online Evaluation of TCGA Gene and miRNA Expression Level in PRAD

Using the UALCAN database (<u>http://ualcan.path.uab.edu/index.html</u>), the expression levels of common miRNA signature in patients with PRAD have been separated on TP53 mutation status and Gleason score. UALCAN is a user-friendly, interactive web resource for TCGA transcriptome data analysis.²⁵ The analytical results were displayed on the webpage (UALCAN); p<0.05 was considered to denote a statistically significant result. The evaluation of the target genes was done using the DIANA-miRPath v3.0 interface.²⁶

miRNA qRT-PCR Evaluation on Tissue and Plasma Samples

Three upregulated transcripts (miR-20b-5p, assay ID:001006; miR-96-5p, assay ID: 000186 and miR-183-5p, assay ID: 002269) and one downregulated transcript (miR-543-5p, assay ID: 002269) were selected from common signature from our microarray data and TCGA in both data analyses (UMPh and TGCA patient cohort). RNA was extracted using the TriReagent based method for qRT-PCR. The validation was performed on 34 matched paired PRAD samples, respectively 34 plasma samples from PRAD and 30 healthy controls using TaqMan MicroRNA Reverse Transcription Kit (Applied Biosystems) protocol as described previously, based on $\Delta\Delta$ Ct method,¹⁶ on VIIA7 system (Life Technologies). U6 (assay ID: 001973) and RNU48 (assay ID: 001006) were used as normalizers. All qRT-PCR reagents were provided by Life Technologies.

Statistically Analysis

Data analysis was done using GraphPad Prism v9 software (GraphPad Software, San Diego, CA, USA). Student's *t*-test was used for the differences among the expression levels of gene and miRNAs, as two-group comparisons (in tumor tissue and normal adjacent PRAD tissue, respective for the expression level in the case of plasma for PRAD patients versus healthy control group); p-value ≤ 0.05 was considered as statistically.

Results

Differential miRNA Expression on PRAD-UMPh Patient Cohort

In the case of our patients' cohort with confirmed histopathological PRAD (10 tumor tissue/ normal adjacent tissue matched paired samples) there has been observed an important number of altered transcripts, only 60 differentially expressed miRNAs (16 upregulated and 44 downregulated), the complete list of the altered miRNAs being displayed in <u>Table S1</u> and Figure 1. Figure 1A presents the UMPh heatmap, in Figure 1C is displayed scatter plot and in the Figure 1E is represented the Volcano plot for the same patient cohort.

Differential miRNA Expression on PRAD-TCGA Patient Cohort

The analysis performed on TCGA samples for PRAD tumor tissue (n=494) versus normal tissue (n=52) reveals a total of 261 miRNAs differentially expressed (208 upregulated and 53 downregulated), displayed as a heatmap in Figure 1B. The



Figure I MiRNA pattern in PRAD in UMPh and TCGA patient cohort. Heatmap of miRNA microarray expression data for (A) UMPh patient cohort, (B)TCGA data; (C) scatter plot used to visualize variations in miRNA expression between tumor values on the X-axes) and normal adjacent tissue (values on the Y axes), using the default fold-change value 1.5, displayed as green lines) of differentially expressed miRNAs for UMPh group (D) scatter plot of differentially expressed miRNAs for TCGA cohort; (E) volcano plot for UMPh group (F) volcano plot for TCGA group, graphical representation generated using Gene Spring (PRAD tumor tissue versus normal adjacent tissues); abscissa represents the log2 transformation value of the differential expression fold change between the tumor and the normal samples.

complete list of the altered miRNAs is presented in <u>Table S2</u>. Additional scatter plot and Volcano plot graphical representation is displayed in Figure 1D, respectively 1F for TCGA dataset.

For common miRNA signature using UMPh and TCGA patient cohort an additional Venn diagram was created for the overexpressed (Figure 2A) and downregulated transcripts (Figure 2B).

Validation of Microarray Data by RT-PCR at Tissue

We performed qRT-PCR for the validation of microarray analysis for 4 differentially expressed miRNAs (miR-20b-5p, miR-96-5p, miR-183-5p, and miR-543-5p) in the expanded collective of 34 PRAD tumor tissues and their adjacent normal tissue. U6 and RNU48 were used as endogenous controls for the normalization of the qRT-PCR data. miR-20b-5p, miR-96-5p, and miR-183-5p-5p levels were overexpressed, while those for miR-543-5p were downregulated in tumor tissues versus normal tissues, data being in accordance with those from the microarray data (Figure 3A). We performed the AUC (area under the curve) analysis showing that the highest AUC value was observed for miR-542-5p (0.8015) and miR-96-5p (0.7355), less sensible as biomarkers being miR-183-5p (AUC=0.6677), respectively miR-20b-5p (AUC=0.6437) (Figure 3B).



Figure 2 Summary of common miRNAs which are differentially expressed in PRAD between UMPh and TCGA. The common miRNAs are presented by overlapping the altered miRNAs as Venny diagram in the two analyzed groups using Gene Spring: (A) overexpressed miRNAs in the two analyzed cohorts and (B) downregulated miRNAs in the two cohorts.

Validation of Microarray Data by RT-PCR at Plasma Level

Additional validation at plasma level using samples from 34 PRAD patients and 30 healthy controls was performed for the selected overexpressed transcripts, confirming the overexpression level retrieved (Figure 4A). AUC values are similar, or even higher than those retrieved in tissue (Figure 4B).

miRNAs as Key Signaling Molecules in PRAD Evaluated Using IPA

The molecular cellular functions identified as significantly enriched ($p \le 0.05$) within the dataset are summarized in Table 5 based on IPA analysis. The most significant altered network identified was "Cellular Development" in both patients' cohorts. We used the common PRAD signature to generate the miRNA-mRNA interaction network using IPA, displayed in Figure 5, revealing the TP53, AGO2, BIRC5, and EGFR as the core element for this network.

Four key representative transcripts (miR-20b-5p, miR-96-5p, miR-183-5p, and miR-543-5p) among the generated IPA network was selected for validation, interconnected with TP53, BIRC5, and EGFR, revealing to have an altered expression level for prostate cancer; meanwhile no statistically significant altered expression level was observed for AGO2 (Figure 6).

miR-20b-5p, miR-96-5p, miR-183-5p and miR-543-5p Involvement in Biological Processes, and Their Target Genes

Identification of the main biological processes and target genes for miR-20b-5p, miR-96-5p, miR-183-5p, and miR-543-5p was done by the use of the DIANA-miRPath v3.0 interface. A heatmap created directly from the DIANA-miRPath v3.0 interface, reveals the main biological pathways targeted by the selected transcripts (Figure 7A). Thus, it has been



Figure 3 Tissue qRT-PCR data validation (A) Graphical representation of the expression levels for miR-20b-5p, miR-96-5p, miR-183-5p and miR-543-5p in tumor versus normal adjacent tissue. The data were normalized using U6 and RNU48 (* $p \le 0.05$, *** $p \le 0.001$) (B) ROC curves for the selected miRNAs, displaying the specificity and sensitivity for the discrimination among the tumoral tissue (TT) versus normal tissue (TN). Abbreviations: ROC, receiver-operating characteristic; AUC, area under ROC curve.

observed the overlapping of 984 target genes for miR-20b-5p, 827 target genes for miR-96-5p, 772 target genes for miR-183-5p, and 76 target genes for miR-543-5p (Figure 7B). The list of the target genes for the selected transcripts (miR-20b-5p, miR-96-5p, miR-183-5p and miR-543-5p) is presented in <u>Table S3</u>.

miRNA Expression Pattern Considering TP53 Mutation Status in PRAD Based on TCGA Data

Additional investigation of miRNA signature based on TP53 status was done using UALCAN. TP53 is a frequently mutated gene in many cancers, the mutation frequency in PRAD being over 20% (Figure 8A). The expression level for miR-20b and miR-183-5p was not affected by the TP53 mutation status (Figure 8B). In the case of miR-96, the expression level was statistically significant much higher in the case of samples with TP53 mutant versus samples with TP53 wild-type. In the case of miR-542, it was statistically downregulated in the TP53 mutant, while the expression level for the TP53 wild-type group was not modified.



Figure 4 Plasma qRT-PCR data validation. (A) Graphical representation of the expression levels for miR-20b-5p, miR-96-5p, miR-183-5p in plasma PRAD patients versus healthy controls plasma. The data were normalized using U6 and RNU48 (** $p \le 0.01$, *** $p \le 0.001$) (B) ROC curves for the selected miRNAs, displaying the specificity and sensitivity for the discrimination among expression levels in plasma for PRAD patients versus healthy controls. Abbreviations: ROC, receiver-operating characteristic; AUC, area under ROC curve.

No.	Name	TCGA		UMPh	
		P-value	#Molecules	p-value	#Molecules
I	Cellular Development	4.88E-02-8.89E-12	65	4.75E-02-1.29E-04	П
2	Cellular Growth and Proliferation	4.88E-02-2.94E-07	52	4.32E-02-1.29E-04	П
3	Cellular Movement	4.61E-02-2.94E-07	38	-	-
4	Cell-Cycle	4.51E-02-1.65E-05	18	4.85E-02-2.44E-04	4
5	Cell Death and Survival	4.88E-02-7.43E-04	29	2.30E-02-8.86E-04	8
6	Cell-To-Cell Signaling and Interaction	-	-	4.04E-02-2.11E-03	3

 Table 5 Molecular and Cellular Functions Altered in PRAD Recognized by IPA (www.ingenuity.com) Considering the Altered miRNA

 Expression Signature for the Two Patient Cohorts

miRNA Expression Pattern Considering Gleason Score in PRAD Based on TCGA Data

Evaluation of the expression levels of microRNAs associated with the Gleason grading system in PRAD based on TCGA data was done using UALCAN (Figure 9). It was observed an increase of the expression level with the Gleason score (particular for Gleason score 10, n=3), these effects being more pronounced for miR-20b-5p and miR-183-5p, but need to be further validated in higher number of cases considering the only 3 cases in Gleason score 10.



Figure 5 miRNA-mRNA interaction network generated using IPA, based on the common miRNA signature among the two-patient cohort; the downregulated miRNAs are presented in green; the overexpressed miRNAs are presented in red; the red circle is highlighted the miRNAs validated by qRT-PCR, blue squares are highlighted the core genes of the network.



Figure 6 Expression level and overall survival of the main genes presented in the miRNA-mRNA network (AGO2, TP53, BIRC, and EGFR), according to TCGA data downloaded from the UALCAN online tool. (A) Expression levels and overall survival graphical representation of AGO2 (or EIF2C2), TP53, BIRC5, and EGFR generated using UALCAN online tool (B) Graphical representation of the overall survival for AGO2, TP53, BIRC5, and EGFR, generated using UALCAN online tool. (B) Overall survival for AGO2, TP53, BIRC5, and EGFR, generated using UALCAN online tool (* $P \le 0.05$, *** $P \le 0.001$).

Discussion

In this study, we evaluated the differentially expressed miRNAs between PRAD tumors and their adjacent normal tissue, using microarray, then we overlapped the date with those from the TCGA. The most relevant transcripts among the common signature were validated by qRT-PCR in both tumor tissue and patients' plasma. The value of liquid biopsy as an alternative to the extremely invasive tissue biopsy is highly important due to the information relieved in plasma from the tumor. This is particularly important when this can be combined with already implemented non-invasive markers, for higher accuracy diagnostic by combining with circulating miRNA,^{20,27} particularly for high risk group identification.²⁸ Circulating miRNAs are considered the next-generation clinical biomarker,²⁸ that promote not only the existence of cancer, but also the dynamics, malignant capacity and drug resistance of tumors.²⁹

Our analysis provides an initial framework to address the long-standing issue of co-operative or Functionally interrelated nature of miRNA targets in a practical setting. Particularly, when correlated with databases related to the



Figure 7 (A) Hierarchical cluster and miRNA/KEGG pathways heat map among the selected miRNAs deregulated in PRAD. The map was generated using the Diana tools miRPath v.3 software (B) overlapping of the target genes for miR-20b-5p, miR-96-5p, miR-183-5p, and miR-543-5p.





Figure 8 The altered transcriptomic pattern was generated using the online tool UALCAN and considering the TP53 mutation status as a clinical variable in PRAD (based on the TCGA dataset). (A) Overall TP53 gene mutations across various PRAD projects data of cBioPortal. (B) miRNA expression level depending on TP53 mutation status, normal adjacent PRAD tissue (n=50), TP53 mutation PRAD cases (n=52), TP53 wild-type PRAD cases (n=438) (*p ≤ 0.05 , ***p ≤ 0.001).

expression levels survival data of miRNAs and their target gene, it can generate functionally relevant biomarkers or therapeutically viable candidates.

Altered miRNA pattern can be considered a valuable molecular characteristic and can impact clinical trajectories and predictions for enhanced responses to specific therapeutics.³⁰ We found that the altered miRNAs are involved in key biological processes (Table 5), with important clinical significance, as was emphasis by IPA analysis.

We identified several miRNAs that were differentially expressed in our patient cohort and TCGA. Among this signature (12 upregulated and 1 downregulated), it was validated by qRT-PCR three transcripts upregulated (miR-20b-5p, miR-96-5p, and miR-183-5p) and one downregulated (miR-542-5p), interconnected with the TP53 and BIRC5, respectively with EGFR downregulated in PRAD, as shows TCGA data downloaded from UALCAN online tool (Figure 4).

Gleason score Gleason score

(n=136)

10 (n=3)

Expression of hsa-mir-96 in PRAD based on Gleason score

Gleason score

(n=243)

TCGA samples

Gleason score

8 (n=63)

Gleason score

(n=45)





Expression of hsa-mir-183 in PRAD based on Gleason score



200

50

Normal

(n=51)

Figure 9 The altered transcriptomic pattern was generated using the online tool UALCAN and considering the Gleason score as a clinical variable in PRAD (based on TCGA dataset)

Our approach selects only tissue-specific mRNA targets in prostate cancer, particularly for specific miRNA signatures. As the reduction in target mRNA expression may not constantly be obvious (as we have assumed), our Research may have minimized the overall effect of miRNAs in prostate carcinogenesis or should be related to downstream/upstream genes, particularly those predicting overall survival rate, highlighted with blue square in Figure 6 (TP53, BIRC5, AGO2 and EGFR), that should be further identified as biomarkers for the identification of high-risks prostate cancer groups.

The established miRNA-gene signatures and the identification of miRNA target gene-related to survival have an important role in the elucidation of molecular mechanisms, underpinning dissimilar molecular profiles, sharing interchangeable prognostic capabilities, and additionally, can serve for prospective applications in drug development. The regulation of miRNAs was interlinked with key regulators of Argonaut 2 (AGO2),¹ sustained also by present study (Figure 5). miRNA association with AGO2 is regulated by TP53, suggesting a new mechanism of miRNA regulation in carcinogenesis through TP53-mediated regulation of AGO2-miRNA interaction.^{31,32} A recent study reveals that AGO2 and miR-96 are incorporated into the cytoplasm.³³ Also, miR-542-5p targeted AGO2, being considered a possible therapeutic target.³⁴ The additional investigation presents important data related to miR-542-5p that targeted AGO2 and EGFR.^{35–37}

Three of the identified transcripts, miR-182-5p, miR-183-5p, and miR-96-5p, are members of the miR-183-96-182 oncogenic cluster.^{38,39} MiR-183-5p is proposed as a specific diagnostic biomarker in prostate cancer.⁴⁰ The well-known prostate tumor suppressor PTEN was demonstrated to be regulated by miR-183-5p and miR-96-5p.³⁸

MiR-96 in prostate cancers is much higher expressed in mutant cases than wild-type p53, as shown UALCAN database. This correlation with TP53 mutation status was observed in the head and neck squamous cell carcinoma.⁴¹

MiR-96 is a key regulator of cell-cell adhesion, having the capacity to overexpress E-Cadherin and EpCAM.⁴² Other studies reveal that this transcript governs androgen signaling and prostate cancer progression,⁴³ and regulation of cell proliferation by targeting FOXO1 and FOXO3a.⁴⁴ miR-20b was identified as overexpressed in prostate cancer tissues, promoting the regulation of key mechanisms like cellular proliferation, EMT, and migration.^{45,46} miR-542 is involved in the regulation of EMT and metastasis,⁴⁷ in our case was downregulated, particularly in TP53 mutated samples. TP53 is positively regulated by these transcripts.⁴⁸ Also, inhibition of the expression level for miR-542 was related to hypoxic condition⁴⁹ or regulated DNA repair/Notch pathway.⁵⁰

Mutations in TP53 are among the most common abnormalities in many cancers,^{30,51–53} observed in Figure 3 as a core element. miRNAs are altered in tumors, underlining the fact that a possible connection between miRNAs and TP53 could be critical in tumorigenesis, considering that TP53 is a frequently overexpressed and mutated gene in the prostate cancer.^{54–56} Additional investigation of miRNA interplay should be investigated, particularly for therapeutic application of miRNAs targeting TP53 pathway,^{57,58} especially for PRAD.

The limitation of the present study is related to the reduced number of cases, this being minimized by the overlapping of TCGA miRNA signature and validation of part of the common signature in an additional patient cohort by qRT-PCR at tissue and plasma level. At the same time, this pilot profiling study provides valuable data related to the complex pathophysiological mechanism that will serve as a basis for further detailed investigations.

Conclusions

Overlapping of two miRNA independent patient cohorts can identify a common signature, that can be further explored for biomarker discovery. Additionally, this common miRNA signature targets key genes related (TP53, BIRC5, EGFR, and AGO2 correlated with overall survival in PRAD) to the key mechanism involved in prostate cancer, where an important element is associated with TP53 signaling and its related miRNAs. Additional investigations in larger sample size is also required to validate the outcomes of this pilot study, and to be able to identify the high risks groups of prostate cancer based on altered miRNA patterns.

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Disclosure

The authors declare no conflicts of interest in this work.

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