**Study of some MiRNAs in Breast Cancer Patients**

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**Abstract:** Breast cancer (BC) is one of the most common cancers in women around the world and the second leading cause of death worldwide. MicroRNAs (miRNAs) expression participates in breast cancer. The purpose of this study is to investigate the expression of miR-133a and miR-155 in breast cancer serum and study their correlation with tumor suppressor protein (p53), carcinoembryonic antigen (CEA) and cancer antigen-15.3 (CA-15.3) concentrations in serum of breast cancer patients and also study their correlations with clinicopathological features. In this study the expression of miR-133a and miR-155 in serum were measured using quantitative real-time polymerase chain reaction (qRT-PCR), P53 concentration was measured by enzyme-linked immunosorbent assays (ELISA**)**, CEA and CA-15.3 concentration were measured using ARCHITECT immunoassay in women with breast cancer (n=60) and controls (n = 20). MiRNA-155 was significant overexpressed (*P*<0.001) while miR-133a had significant down expression (*P*<0.001) in the serum of breast cancer patients compared to control serum. P53 had no significant correlations with any of the studied miRNAs. Carcinoembryonic antigen and CA-15.3 have significant higher concentration in the serum of breast cancer patients compared to control serum. A significant association was observed between miR-133a with tumor grade (*P*<0.05) and miR-155 with lymph node involvement (*P* <0.05). A significant correlation between miR-155 and CEA (*P* <0.05). No correlations between miR-133a and P53, CEA, CA-15.3. **Our Conclusion** these miRNAs have a significant signature in the pathogenesis of breast cancer and can be used as noninvasive biomarkers for breast cancer detection.

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**Key words:** MicroRNAs; Breast cancer; qRT-PCR; P53; ELISA; CEA; CA-15.3

**1. Introduction**

Breast cancer is one of the most common cancers in women worldwide, and the second cause of death in female cancer patients **(**[**Wang *et al.,* 2016**](#_ENREF_41)**).** A total of 1,665,540 new cases and 585, 720 deaths occurred in the USA during the year 2014, according to the American cancer society **(**[**Hecht *et* *al.,* 2016**](#_ENREF_13)**).** In Egypt, According to the Egyptian National Cancer Institute (NCI), breast cancer is the most common familiar type of cancer between Egyptian women representing 18.9% of total cancer cases **(**[**Elatar, 2002**](#_ENREF_9)**)**. But this ratio was increased so, it accounts for approximately 38% of reported malignancies between Egyptian women **(**[**Rashad *et al.,* 2014**](#_ENREF_35)**).** Worldwide Public health data revealed that more than one million women are diagnosed with breast cancer each year and more than 410,000 women will die from the disease **(**[**Yu *et al.,* 2016**](#_ENREF_44)**)**. Breast cancer is an heterogeneous disease with numerous morphological appearances, behaviors molecular features, and response to therapy **(**[**Polyak, 2011**](#_ENREF_31)**)**. Treatment of breast cancer is based on the accessibility of strong diagnostic, prognostic, and predictive factors to guide the choice of different treatment options **(**[**Carlson *et al.,* 2011**](#_ENREF_6)**)**.

MicroRNAs are highly conserved noncoding RNA molecules that are approximately 17–25 nucleotides in length. They control gene expression at the posttranscriptional level by interacting with a specific target messenger RNA (mRNA) **(**[**Lagos-Quintana *et al.,* 2002**](#_ENREF_18)**)**. They also regulate a variety of cellular processes, such as proliferation, differentiation, metabolism, aging, and cell death. As such, the importance of miRNAs is increasingly recognized in almost all fields of biological and biomedical fields **(**[**Li *et al.,* 2010**](#_ENREF_21)**)**. In humans, it has been estimated that there are more than 1000 miRNAs in the genome, which regulate approximately 30% of all protein-coding genes **(**[**Lewis *et al.,* 2005**](#_ENREF_20)**)**.

The importance of miRNAs in oncogenesis has also been recognized. Dysregulation of miRNA expression plays an important role in cancer development through various mechanisms, such as deletions, amplifications, epigenetic silencing, or mutations in miRNA loci **(**[**Kosaka *et al.,* 2010**](#_ENREF_17)**)**. To date, an association between differentially expressed miRNAs and many clinicopathological features has been shown, including mRNA expression-based classification **(**[**Blenkiron *et al.,* 2007**](#_ENREF_2)**)**, tumor grade, and breast cancer staging **(**[**Iorio *et al.,* 2005**](#_ENREF_14)**)**. The current in vivo diagnostic tools used for the detection of breast cancer at its early stages, e.g., mammography and ultrasound had several limitations, such as breast density or calcification recognition. Other imaging modalities, e.g., magnetic resonance imaging (MRI), have been proposed as complementary diagnostic modalities, with limited sensitivity **(**[**Bertoli *et al.,* 2015**](#_ENREF_1)**).**

Nevertheless, the cost incurred and skill required for a mammogram has hindered a wide acceptance of this method. Thus, there is still need to clarify new mechanisms to develop accurate screening method that can diagnose patients with early cancer or precursor lesions by minimally invasive techniques **(**[**Swellam *et al.,* 2015**](#_ENREF_38)**)**. The emergence of small non-protein-coding RNAs called miRNAs which are stable in serum and playing important roles in oncogenesis has opened new opportunities for early cancer diagnosis **(**[**Calin and Croce, 2006**](#_ENREF_4)**;** [**He *et al.,* 2007**](#_ENREF_12)**).** MiRNA has a great potential to be a novel biomarker for breast cancer and holds a potential for individualizing patients' treatment regimens **(**[**van 't Veer *et al.,* 2002**](#_ENREF_40)**).** However, there is still restricted awareness on the exact mRNA target of the deregulated miRNAs in breast cancer.

In this study, we aimed to explore the efficacy of combining a two of circulating miRNAs (miR133a and miR155) as prospective biomarkers in breast cancer and their relationship with clinicopathological features and concentration of tumor suppressor protein (p53), CEA and CA-15.3. To our knowledge, this is the first study to implicate the role of these miRNAs and p53 in breast cancer patients in Egypt.

**2. Patients and methods**

**Patients:**

The study was performed on 80 females, including 60 with newly diagnosed breast cancer at different disease stages and 20 control age corresponding females. Patients with breast lesions were recruited from the surgery department, National Cancer Institute (NCI), Cairo University during the time period from March 2015 to August 2015. The study was permitted by the Institutional Review Board (IRB) of the NCI, Cairo University. It was permitted according to the Helsinki guidelines of studies performed on human beings and a written conversant permission was obtained from all study subjects. All patients were subjected to ordinary biochemical and hematological investigations and imaging diagnoses and chest x-ray for stage IV. Samples were obtained from all patients previous to any therapeutic or surgical intervention. Participant’s age showed a mean ± SD of (49.4 ± 7.64) years in breast cancer patients and (43.9 ± 9.45) years in control females. Patients’characteristics is presented in **table (1).**

**Methods**

**Blood sampling:**

Ten ml of blood was withdrawn into 2 serum collection tubes, allowed to clot for 30 minutes and centrifuged at 4000 R.P.M for 10 minutes. Yielded serum was divided into 2 micro tubes and stored at -80 ºC till the time of analysis. First tube was used for determination of serum concentration of P53 by ELISA according to the manufacturer’s instructions, for determination of serum concentration of CA-15.3 and CEA by chemiluminescence assays ([ARCHITECT i1000SR Immunoassay Analyzer](https://www.abbottdiagnostics.com/en-us/products/ARCHITECT-i1000SR.html), Abbott, U.S.A) according to the manufacturer’s instructions. The second tube was used for quantification of mature miRNAs (miR-133a and miR-155) in serum by qRT-PCR.

**Determination of CA-15.3 and CEA concentrations:**

Determination of serum concentration of CA-15.3 by ARCHITECT immunoassay **using ARCHITECT CA-15.3 Kit (catalog no.2K44) supplied by Abbott laboratories diagnostics division (USA)** according to the manufacturer’s instructions. Determination of serum concentration of CEA by ARCHITECT immunoassay **using ARCHITECT CEA Kit (catalog no.7K68) supplied by Abbott laboratories diagnostics division (USA)** according to the manufacturer’s instructions.

**Determination of P53 concentration:**

Determination of serum concentration of tumor suppressor protein (p53) by enzyme-linked immunosorbent assay (ELISA) **using the kit (catalog#: ELH-P53) supplied by Ray Biotech, Inc** (**Norcross, GA**) according to the manufacturer’s instructions.

**Serum miRNAs assays:**

**RNA extraction:**

RNA was isolated from 200 µL of serum using **the miRNeasy Mini Kit (cat. no. 217004) Qiagen, Germany**, according to the manufacturer’s instructions. The RNA was eluted in 40 µL of RNAse-free water and was stored at −80◦C until qRT-PCR reaction. Concentrations of all RNA samples were measured using Nanodrop 1000 (Thermo Scientific, Wilmington, DE, USA).

**Reverse transcription (RT):**

The total RNA (100 ng) was reverse transcribed after thawing using a miScriptHiSpec buffer supplied in **miScript II RT Kit (catalog no. 218161) Qiagen, Germany**, according to the manufacturer’s instructions, and cDNA synthesis was performed in a thermal cycler (IGEM: MIT/2005/Thermo cycler), the thermal cycler reaction setting were as follows: incubation at 37°C for 60 min followed by 5 min at 95°C. The cDNA was stored at −20°C until use**.**

**Quantitative Real-time Polymerase Chain Reaction:**

Quantitative real-time PCR was performed by **miScript SYBR Green PCR Kit (200) Qiagen, Germany (Catalog no. 218073**) in compliance with the manufacturer’s instructions. MicroRNA specific primers were provided by Qiagen/Germany, miR133a/ Hs\_miR-133a\_2 miScript Primer Assay (MS00031423) and miR-155/ Hs\_miR-155\_2 miScript Primer Assay (MS000031486). MiRNA expression levels were quantified using Step One (Applied Biosystems, USA). Cycling program: primary activation step for 15 min at 95 °C to stimulate HotStarTaq DNA polymerase, cycling:( first denaturation for 15 s at 94 °C, annealing for 30 s at 55°C, finally extension for 30 s at 70 °C at which fluorescence data collection were performed) ×40 cycles and endogenous control miScript Primer Assay SNORD68 (Hs\_SNORD68\_11 miScript Primer Assay (MS00033712) was used to normalize the data **(**[**Motawi *et al.,* 2016**](#_ENREF_27)**)**. The data obtained from the miRNA expression levels were calculated and evaluated by the cycle threshold (Ct) method, which is the number of cycles required for the fluorescent signal to cross the threshold in RT-PCR. The level of miRNA expression was reported as ΔCt value. The ΔCt was calculated by subtracting the Ct value of miRNA SNORD68 from the Ct values of the target miRNAs [mean value Ct (miR-133a, miR-155) - mean value Ct (housekeeping gene)], the relative expression level of the miRNA of interest corresponded to the 2-∆Ct value. ΔΔCt was then determined by subtracting the average ΔCt of the control from the ΔCt of cases. The fold change in the miRNA expression level was calculated (fold change = 2-∆∆Ct) to determine the relative quantitative levels of individual miRNA **(**[**Livak and Schmittgen, 2001**](#_ENREF_23)**)**. An RT-PCR was done in duplicate, including non-template controls. The qualified expression of the mature miRNA was calculated by the relative cycle threshold (2-∆∆Ct) method. Then the fold change was sharp peak for miRNA and endogenous gene indicating that they were efficiently extracted transformed to the log form and used in statistical analysis. Melting curves analysis was done and showed a single and specifically amplified from serum.

**Statistical analysis:**

In the existing study SPSS software package (version 17 for Windows; SPSS INC., Chicago, IL, USA) was used to perform statistical analysis. To compare the miRNA expression in cancer versus the normal serum; the Wilcoxon’s rank sum test for one sample was used. For comparing two different groups such as metastatic and non-metastatic, the Mann-Whitney U nonparametric test was used, while Kruskal-Wallis nonparametric test was used for more than two independent variables. To find a correlation between two variables Spearman’s rho (r) was calculated. The Receiver Operating Characteristic (ROC) curve was used to determine the cut-off values of miRNAs and to analyze the diagnostic utility of different markers. A p-value of less than 0.05 was considered statistically significant. All P-values are two sided.

**3. Results**

**Serum expression levels of miR-133a and miR-155**

The expression levels of miR-133a and miR-155 were evaluated by qRT-PCR. The serum level of miR-155 was significantly higher in BC serum than in healthy controls (*p* < 0.001), while serum level of miR-133a was significantly lower in breast cancer serum than in healthy controls (*p* < 0.001) as shown in**Figure 1 (A).**

**Serum concentration levels of CEA and CA-15.3**

The serum level of CEA and CA-15.3 were significantly higher concentration in BC than in healthy controls as shown in **Figure 1 (B).**

**Evaluation of the diagnostic accuracy of miR-133a and miR-155**

The diagnostic accuracy of miR-133a and miR-155 were evaluated using ROC curve analysis. ROC curve analysis showed that the two miRNAs can significantly differentiate between breast cancer and healthy controls, showing an area under thecurve (AUC) of 0.950 for miR-133a (95% CI 0.89-1.00, *p* < 0.001) and AUC 0.767(95% CI0.66-0.87, *p* < 0.001) for miR-155. The optimal sensitivity and specificity were (95%and 100%) and (76.7% and 100%), respectively. When the diagnostic significance of serum miRNAs was compared in breast cancer patients, the results of the ROC curve suggested that the diagnostic accuracy of serum miR-133a was superior to miR-155 with AUCs of 0.950 and 0.767, respectively, and total accuracy were 96.3% and 82.5%, respectively as shown in **Figure 2 (A)(B) and Table (2).**

The diagnostic accuracy P53 was evaluated using ROC curve analysis. All diagnostic accuracy for these markers at the selected cutoff values are shown in **(Table 2).** But in comparison to studied miRNAs we found that miR-133a is more sensitive than miR-155 and two miRNAs are more sensitive than P53, as shown in **Table (2)** and **Figure 2 (C).**

**Correlation between the expression levels of miRNAs (miR-133a and miR-155) and concentration of CEA and CA-15.3**

Using Spearman’s correlation coefficient showed that significant correlation between miR-155 and CEA, also there was a significant correlation between CA-15.3 and CEA, but there was not a significant correlation between two miRNAs and no significant correlation between miR-133a and both of CEA, CA-15.3 as shown in **Table (3).**

**Relationship between miRNAs (miR-133a and miR-155) and tumor markers (CEA, CA-15.3) in serum of breast cancer patients with theirclinicopathological features**

The relative expression of serum miRNAs of the breast cancer patients were studied in relation to their clinicopathological data. The level of miR133a (*p* <0.05) was significantly higher in sera of patients with grade (III) than in those with lower grade tumors (II). Also miR-155 had a statistically significant association with lymph node involvement (*p* =0.039), no significant differences between miRNAs (miR-133a and miR-155) expression and (age, tumor size, menopausal states, estrogen receptor and progesterone receptor status). The relative concentration of serum CEA and CA-15.3 of the breast cancer patients were studied in relation to clinicopathological data, which showed that CA-15.3 had a statistically significant association with stages (*p* <0.001) and HER2 (p=0.036), also CEA had a statistically significant association with stages (*p* <0.001), age (*p* =0.018) and menopausal state (*p* =0.013) as shown in **Table (1).**

**P53 concentration:**

P53 had statistically significant low concentration in the breast cancer serum range from (5.04-9.27) than control serum ranges from (6.27-19.70) with (*p* <0.05) **Figure (1B).** At the optimal cutoff value of 6.90, the sensitivity and specificity were 55% and 85% respectively **Figure (2D).** Its serum level showed that no significant associations with any of the clinicopathological parameters **Table (1).** No significant correlation between p53 and expression of miR-133a and miR-155 was found, no significant correlation between p53 and CEA, CA-15.3as shown in **Table (3).**

**Table (1): Association between study markers with different clinicopathological characteristics**

|  |  |  |  |
| --- | --- | --- | --- |
|  |  | Relative expression\* of microRNAs with p value in different groups\*\*. | Relative\*\*\* concentration with *p* value in different groups\*\*. |
| **variable** | **N (%)** | **miR-133a** | *p***value** | **miR-155** | *p***value** | **P53** | *p* **value** | **CA-15.3** | *p* **value** | **CEA** | *p* **value** |
| **Age**≤49>49 | 36(60)24(40) | -1.47-1.81 | 0.561 | 2.242.64 | 0.952 | 6.586.61 | 0.774 | 28.2532.0 | 0.428 | 1.823.26 | **0.018** |
| **Menopausal state**PREPOST | 35(58.3)25(41.7) | -1.53-1.87 | 0.658 | 2.322.16 | 0.793 | 6.586.59 | 0.893 | 28.033.9 | 0.219 | 1.83.22 | **0.013** |
| **Stages**IIIIIIIV | 6(10)28(46.7)5(8.33)21(35) | -1.22-1.76-0.91-2.12 | 0.228 | 1.912.911.141.63 | 0.571 | 6.576.596.456.65 | 0.511 | 24.7523.8521.8039.40 | **0.000** | 2.291.761.714.41 | **0.000** |
| **Grade**IIIII | 45(75)15(25) | -1.41-2.19 | **0.05** | 2.911.21 | 0.107 | 6.586.66 | 0.657 | 29.3028.40 | 0.813 | 2.63.28 | 0.105 |
| **HER2(IHC)**+VE-VE | 23(38.3)37(61.7) | -1.37-1.88 | 0.191 | 2.911.79 | 0.134 | 6.586.44 | 0.373 | 35.2128.10 | **0.036** | 3.282.17 | 0.094 |
| **ER(IHC)**+VE-VE | 52(86.7)8(13.3) | -1.78-1.14 | 0.373 | 2.242.03 | 0.811 | 6.626.47 | 0.184 | 29.9025.00 | 0.794 | 2.672.10 | 0.853 |
| **PR(IHC)**+VE-VE | 53(88.3)7(11.7) | -1.76-1.41 | 0.527 | 2.162.63 | 0.991 | 6.596.52 | 0.375 | 29.328.1 | 0.863 | 2.612.04 | 0.765 |
| **Lymphnode involvement**YESNO | 37(61.7)23(38.3) | -1.87-1.53 | 0.425 | 3.451.44 | **0.039** | 6.646.58 | 0.561 | 31.729.0 | 0.820 | 1.852.6 | 0.242 |

\* Median of relative expression.

\*\*Mann-Whitney U and Kruskal-Wallis nonparametric test were used for comparing different group. \*\*\* Median of concentration. IHC: Immuno-histochemistry

**Table (2): Diagnostic accuracy of the studied miRNA and P53**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
|  | **AUC** | **95%CI (SE)** | **Sensitivity** | **Specificity** | **Cut-off** | *p* **value** | **PPV** | **NPV** | **Total accuracy** |
| **miR-133a** | 0.950 | 0.89-1.00 (0.028) | 95% | 100 % | 0.940 | 0.000 | 100% | 87% | 96.3% |
| **miR-155** | 0.767 | 0.66-0.87 (0.055) | 76.7% | 100 % | 1.006 | 00.00 | 100 % | 58.8 % | 82.5 % |
| **P53** | 0.731 | 0.591-0.872 (0.072) | 55 % | 85 % | 6.900 | 0.002 | 55% | 85% | 77.5% |

 PPV: positive predictive value

NPV: negative predictive value

**Table (3): Correlations among studied miRNAs expression and (CEA, CA-15.3 and p53) in serum of breast cancer**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **miR-133a** | **miR-155** | **CEA** | **CA-15.3** | **P53** |
| **miR-133a (log fold expression)**CCP value | 1.0 | .245.059 | -.131.320 | -.064.627 | -.169.196 |
| **miR-155 (log fold expression)**CCP value | .245.059 | 1.000 | -.295**\*.022** | .064.629 | .014.916 |
| **CEA (ng/ml)**CCP value | -.131.320 | -.295\***.022** | 1.00…. | .613\*\***.000** | -.159.226 |
| **CA-15.3 (IU/ml)**CCP value | -.064.627 | -.064.629 | .613\*\***.000** | 1.000…. | -.057.665 |
| **P53(ng/ml)**CCP value | -.169.196 | .014.916 | -.159.226 | -.057.665 | 1.000.... |

**CC:** Spearman’s correlation coefficient. **\*** Correlation is significant at the 0.05 level (2-tailed). **\*\***Correlation is significant at the 0.01 level (2-tailed).

|  |
| --- |
|  |
| **Figure 1 (A)**: miR-133a and miR-155 expression is dysregulated in breast cancer. This box plot displays the log fold change of relative expression of miR-133a and miR-155 in 60 breast cancer patient’s serum versus 20 control serum. This box plot represent minimum and maximum, the top, the bottom and the band in the box represent the first and third quartile and the median respectively. |
|  |
| **Figure 1(B):** CEA and CA-15.3 had higher concentration in breast cancer than healthy control. P53 had low concentration in breast cancer than healthy control. This box plot displays the concentration of CEA, CA-15.3 and P53 in 60 breast cancer patient’s serum versus 20 control serum. |

|  |  |
| --- | --- |
|  |  |
| **Figure 2(A):** ROC curve of miR-133a. | **Figure 2(B):** ROC curve of miR-155. |



**Figure 2(C):** ROC curve of P53

**4. Discussion:**

MicroRNAs were identified to play vital roles in the pathogenesis of cancer. The up regulation of oncogenic miRNAs or down regulation of tumor suppressor miRNAs can be implicated in tumorigenesis by varying many pathways, including cell cycle, angiogenesis, invasion and metastasis ***(***[***Nana-Sinkam and Croce, 2014***](#_ENREF_28)***)*.** Thus, there is still a pressing need to elucidate novel mechanism of breast cancer development so as to develop a cost effective and accurate screening method for this cancer. Recently, the emergence of small non protein coding RNAs called microRNAs, playing important roles in oncogenesis, has opened new opportunities for early cancer diagnosis **(**[***Calin and Croce, 2006***](#_ENREF_3)***;*** [***He et al., 2007***](#_ENREF_12)***)***. In this study, we analyzed the levels of miR-133a and miR-155 in serum from 60 breast cancer patients and 20 controls. MiR-133a is one of non-coding RNA which plays an important role in preventing progression of breast cancer, as loss of miR-133a expression resulted in aberrant cell invasion and proliferation associated with poor prognosis in breast cancer **(**[***Cui et al., 2013***](#_ENREF_8)***)*.**

Studies have reported that altered expression of miR-133a in several human cancers including esophageal squamous cell carcinoma**,** bladder cancer, and breast cancer ***(***[***Kodahl et al., 2014***](#_ENREF_16)***)****.* MiR-133a affects breast carcinogenesis either through targeting fascin actin-bundling protein 1 (FSCN1) **(**[***Wu et al., 2012***](#_ENREF_43)***)*** or through regulating the cell cycle and proliferation of breast cancer cells by focusing on the epidermal growth factor receptor (EGFR) through the EGFR/Akt signaling pathway **(**[***Cui et al., 2013***](#_ENREF_8)***)***. These previous studies are consistent with our current study. We found that the level of miR-133a was statistically significantly lower in breast cancer serum than in the controls. Hence, supporting the suggestion that miR-133a plays a role as a tumor suppressor gene affecting breast cancer development and progression ***(***[***Qin et al., 2013***](#_ENREF_32)***;*** [***Rao et al., 2010***](#_ENREF_34)***;*** [***Ruebel et al., 2010***](#_ENREF_36)***)***.

Our study data revealed that miR-133a was significantly down regulated in serum of breast cancer than normal control serum which in agreement with ***(Kodahl et al., 2014)*** which state that miR-133a down regulated in breast cancer serum in relation to normal control serum proving that miR-133a consider tumor suppressor gene in breast cancer. Our result in contrast to **(**[***Chan et al., 2013***](#_ENREF_7)***)*** which state that miR-133a up regulated in serum of female breast cancer versus to control.

Another non coding RNA (miR-155), we found that miR-155 had statistically significant higher expression in BC serum than healthy control serum. Our data in harmony with **(**[***Hagrass et al., 2015***](#_ENREF_10)***)*** who found that miR-155 was over expressed in serum of BC patients compared to normal control serum.**(**[***Mattiske et al., 2012***](#_ENREF_25)***)*** state that miR-155 over expressed in serum breast cancer than normal control serum, also miR-155 up regulated in breast cancer tissue than normal tissue and higher expression of miR-155 in breast cancer cell line when compared to normal cell line all these previous results in harmony with our results in which miR-155 over expressed**.** Our result in agreement with **(**[***Zhang et al., 2013***](#_ENREF_45)***)***which state that miR-155 over expressed in breast cancer cell line when compared to normal cell line**.**

In regarding the relation of miRNAs (miR-133a and miR-155) with clinicopathological features of BC, our results clearly showed statistically significant difference down expression of miR-133a in grade III more than grade II in serum of breast cancer, which is in contrast to other study which state that no association between miRNA-133a expression and tumor grade **(**[***Kodahl et al., 2014***](#_ENREF_16)***)***. Also our result in contrast with **(**[***Wu et al., 2012***](#_ENREF_43)***)*** who stated that there had not association between miR-133a expression and tumor grade of breast cancer tissue and cell line. **(**[***Wu et al., 2012***](#_ENREF_43)***)*** state that there was no association between miR133a expression and patient age, tumor size, or estrogen receptor and progesterone receptor status these results symmetric with our result.

Our results clearly showed that statistically significant high mean expression of miR-155 with lymph nodes involvement. In the same line with our results, **(**[***Hagrass et al., 2015***](#_ENREF_10)***)*** found that the levels of miR155 were significantly higher in serum of BC patients with lymphnodes involvement than breast cancer patients without lymph nodes involvement. Our result showed that higher miR-155 expression in breast cancer has been shown to be signiﬁcantly associated with lymph nodes involvement, our result consistent with **(**[***Liu et al., 2015***](#_ENREF_22)***)***, suggesting its potential as a clinical prognostic value,**.** On the contrary to our result, the up regulation of miR-155 was not statistically significant to lymph nodes involvement **(**[***Nassar et al., 2014***](#_ENREF_29)***)*.**

Our result showed that significantly higher serum CEA and CA-15.3 in breast cancer patients in compared to normal control, our result in harmony with **(**[***Wu et al., 2016***](#_ENREF_42)***)***. [***Shao et al., 2015***](#_ENREF_37) stated that serum levels of CEA and CA-15.3 were elevated in preoperative breast cancer patients this results in agreement with our results. There was controversy regarding the use of CEA and CA-15.3 in the diagnosis of breast cancer. The European Society for Medical Oncology (ESMO) and the European Group on Tumor Markers (EGTM) suggested that routine measurement of tumor markers such as CEA and CA-15.3 should be performed in patients with breast cancer **(**[***Molina et al., 2005***](#_ENREF_26)***;*** [***Cardoso et al., 2012***](#_ENREF_5)***)***

However, the American Society of Clinical Oncology (ASCO) does not recommend routine measurement of CEA, CA-15.3 or other tumor markers for patients with breast cancer **(**[***Khatcheressian et al., 2013***](#_ENREF_15)***)***. A report suggested that tumor markers including CEA and CA-15.3 should not be routinely measured in patients with early stage breast cancer **(**[***Ramsey et al., 2015***](#_ENREF_33)***)***. Although the limitation of low sensitivity and speciﬁcity preclude the use of serum tumor marker CEA and CA-15.3 for the detection of early breast cancer, elevated preoperative tumor marker levels at initial presentation may predict poor outcome **(**[***Park et al., 2008***](#_ENREF_30)***)***.

The American Society of Clinical Oncology and the National Comprehensive Cancer Network (NCCN) guidelines do not currently recommend the use of serum CA 15.3 and CEA for breast cancer screening and directing treatment or a routine surveillance tool or for therapeutic response monitoring due to inconsistent ﬁndings of their sensitivity and speciﬁcity **(**[***Harris et al., 2007***](#_ENREF_11)***;*** [***Shao et al., 2015***](#_ENREF_37)***;*** [***Maric et al., 2011***](#_ENREF_24)***)***. Carcinoembryonic antigen and CA-15.3 serum levels may be increased in other benign conditions such as gastritis, gastric ulcer, bronchitis, cholangitis, and liver abscess in cases of CEA and chronic hepatitis, liver cirrhosis, tuberculosisin cases of CA-15.3 **(**[***Lee et al., 2013***](#_ENREF_19)***)***.

In our study we have suggested that there could be a relationship between the expression level of studied miRNAs and tumor suppressor protein 53 (P53) level, based on that miRNAs act as oncogenes or tumor suppressor, but our results revealed that no significant correlation between the studied miRNAs expression level and the p53 serum level. This may be due to the miRNAs not directly targeting P53 as miR-133a by targeting epidermal growth factor receptor (EGFR) **(**[***Cui et al., 2013***](#_ENREF_8)***)*** and finally miR-155 act by negatively regulating a tumor suppressor gene known as suppressor of cytokine signaling 1 (socs1)**(**[***Nassar et al., 2014***](#_ENREF_29)***)***.

Regarding their diagnostic efficacy, miR-133a reported the best sensitivity, specificity, PPV, NPP and total accuracy followed by miR-155 then P53 and finally routine tumor markers (CA-15.3 and CEA) indicating usefulness of miRNAs as molecular markers for diagnosing of breast cancer. In this study, we were also able to identify miRNA signatures that were associated miR-133a with the grade. Circulating miRNAs association with some clinicopathological parameters have been reported in other studies **(**[***Tashkandi et al., 2015***](#_ENREF_39)***)*** suggesting their clinical prognostic value and carrying the possibility of a serologic test that can augment the histologic information of a tumor without the need for biopsy.

**In conclusion**

Due to the fact that miRNAs exist stable in circulating blood, with an easy extraction and quantification methods, this serum miRNA-133a and miR-155 can be used as valuable noninvasive biomarkers for breast cancer detection.

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