Micro-RNA Biogenesis Genes (AGO1 and GEMIN4) Single Nucleotide Variants of Bad Prognosis and Poor Therapeutic Response in Egyptian Chronic Myeloid Leukemia Patients: Case–control Study

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Abstract

BACKGROUND: Chronic myeloid leukemia (CML) is one of the most common hematological tumors. Gene candidate studies cleared the association of single genetic variants (SNVs) to the risk and progression in CML. MicroRNA biogenesis genes disruption contributes a fundamental role in carcinogenesis.

AIM: We aimed to determine the association between rs638832 and rs2740348 SNVs of AGO1 gene and GEMIN4 gene, respectively, and the risk and prognosis in CML Egyptian patients with 5 years survival estimation.

METHODS: The study was conducted on 110 newly diagnosed CML patients and 110 age and sex healthy matched controls. Real-time polymerase chain reaction utilizing TaqMan probes was operated to demonstrate genetic modalities of rs636832 and rs2740348.

RESULTS: No significance difference was observed between the cases and controls regarding the genotypic and allelic frequencies for both variants. On the other hand, the rs636832 GG genotype was more evident at a younger age of diagnosis and associated with the poor grades of the Sokal and Eutos scores. As well, rs2740348 CC genotype was encountered in high Eutos score levels. Regarding the response therapy, rs636832 GG genotype was associated with the major grades of the Eutos scores.

CONCLUSION: Our study revealed that the major homozygous genotypes of both variants were associated with bad prognostic clinical scores and poor response to therapy but with no role in CML risk.

Introduction

Chronic myeloid leukemia (CML) is BCR-ABL positive myeloproliferative disease affecting hematopoietic stem cell by clonal growth of leukemic progenitor cells with myeloid predominance [1]. It is considered one of the most hematological disorders, accounts 15% of all leukemia affecting the elderly with male predominance. The annual incidence of CML is 0.87/100,000 subjects that increase 1.52 in age more than 70 [2]. The estimated deaths are 1.130 per year to 0.87/100,000 subjects that increase 1.52 in age more. The estimated deaths are 1.130 per year to 0.87/100,000 subjects that increase 1.52 in age more.

Multiple genetic variants showed increased evidence in pathogenesis as well clinical course and prognosis of CML [5]. MicroRNA (miRs) and its biogenesis genetic abnormalities found to be linked to hematological malignancies with little researches on CML [6].

MicroRNAs (miRs) are epigenetic modifiers of genetic expression by translational regulation and mRNA processing. They are a single strand of small non-coding nucleotides (nt) (~22 nt) that started its biogenesis as long primary (pri-miRNA) by RNA polymerase II (Pol II). Pri-miRNA is progressed to precursor miRNA (pre-miRNA) a ~60–70-nt hairpin-shaped as action of RNase III enzyme Drosha and its cofactor in nucleus that extracted to cytoplasm by exportin 5 (Xpo5) to undergo second modification to~22 double-strand microRNA duplex by RNase III enzyme Dicer [7].

One strand of the miRNA duplexes combines into miRNA-induced silencing complex (miRISC) and...
converts to mature miRNA. The miRISC involves proteins as GEMIN3 and 4, and AGO1-4 are encoded by miRNA biogenesis machinery genes. Their genetic abnormalities; deletion, insertion, mutations affect expression of several oncogenes and tumor suppressor genes with cancer risk, and progression susceptibility [8].

AGO1 gene is present on 1p34.3 chromosome, encodes one of the argonaute family proteins that have important actions in RNA interference (RNAi) and RNA silencing [9]. The rs636832 SNP is existed in the intron 8 of AGO1 gene, related to the risk of lung cancer and the lymphatic metastasis of gastric carcinoma [10], [11].

GEMIN 4 gene is subsisted on chromosome 17p13.3, encoding the Gemin4 protein which is essential component of the miRISC complex [12]. The rs2740348 single genetic variants (SNV) of GEMIN4 are found to be related to rise breast cancer risk and with reduced risk for prostate cancer development [9], [13].

AGO1 rs636832 interacts through Wnt pathway that has an important role in renewing of normal cells and its disruption is involved in carcinogenesis risk and behavior [11]. As well AGO1 was found to regulate the transcription activity through stimulation of phosphoinositide 3-kinase (PI3K/AKT) [14]. Wnt pathway activation with subsequent PI3K/AKT signaling; is involved in CML progression and TKIs resistance [15]. Hence, AGO1 variants may participate in CML risk and prognosis.

PI3K/AKT and STAT5 pathways are essentially involved in the progression and TKI resistance in CML [16]. The N-terminal of STAT5a/b pathway proteins is the domain responsible for cytokine receptor binding, suppresses its autoactivation, and interacts with nuclear receptors as glucocorticoid receptors and mineralocorticoid receptor (MR) [16]. GEMIN4 has a regulator repressor function to MR receptor so its disruption can affect CML progression [17].

Galectin-1 (GAL1) is a beta galactoside binding protein that involved in tumor progression and metastasis. It acts as a bad prognostic marker in CML that enhance its transformation and treatment resistance. It has several pathways of action, including PI3K/AKT and binding to GEMIN4 that affects subsequently miRs biogenesis, this suspects GEMIN4 variants role in CML [18], [19].

Our research aimed to investigate the association between rs636832 and rs2740348 SNVs of AGO1 gene and Gemin4 gene, respectively, and risk, progression, and response to treatment in CML Egyptian patients to clarify if they could be considered as one of biomarkers for diagnosis and targeted for therapy in CML disease.

Materials and Methods

Our case–control study enrolled 110 adult Egyptian newly diagnosed Philadelphia (Ph) (p210) positive CML cases. Patients with other myeloproliferative neoplasms or double malignancies were excluded from the study. They were selected from the cases consulted to Medical Oncology department, National Cancer Institute (NCI) as well to Hematology Outpatient Clinic, Faculty of Medicine, Cairo University in the interval from February 2015 to July 2015 with 5 years follow-up.

All CML cases were picked out by following the diagnostic features of CML settled by the World Health Organization (WHO) guidelines 2015 [20]. They were exposed to detailed history taking, complete physical examination, routine baseline laboratory checks, complete blood count (CBC), coagulation profile, and chemistry profile. Bone marrow study (BM), FISH for t(9; 22), and reverse transcriptase quantitative polymerase chain reaction (PCR) for BCR-ABL on peripheral blood (PB) or BM samples as well abdominal ultrasound (US) were performed to our patients.

The CML patient can be present in one of 3 stages; chronic phase (CP), accelerated phase (AP), or blast phase (BP) that also known by blast transformation and blastic crisis. CP is known by blast count less than 10% with the absence of defining features of AP and BP. AP is diagnosed by either one of the following: basophils >20% in PB, blasts =10–19% in PB, and/or BM, persistent platelet count less than 100 independent to treatment, permanent thrombocytosis >1000 resistant to therapy, increase white blood cells >10, spleen size not-responding to therapy or cytogenetic clonal progression. BP is transformation of CML to acute leukemia and diagnosed by the presence of blast cells >20% in PB and/or BM.

The response to treatment was evaluated first for complete hematological recovery by CBC each 2 weeks, then for complete cytogenetic resolution by FISH monitoring each other 3 months followed by quantitative RT-PCR every 3 months until complete molecular response.

Sokal and EUTOS scores were assessed by the clinical data at diagnosis previous to the beginning of therapy. Age, size of spleen, PB blasts, and platelet count were included in estimation of Sokal score. The score is divided into low-risk (<0.8), intermediate-risk (0.8–1.2), while score >1.2 is considered as a high risk. EUTOS score was depended on the size of spleen and basophils % in the PB. Patients were divided between low-risk and high-risk according to the score of either <87 or ≥87, respectively [21].

One hundred and ten age, sex, and race-matched healthy persons were enrolled as a control
group. They were selected among normal subjects coming for a routine check-up with normal investigations and no history of CML or other medical diseases.

All subjects were included after they gave willingly informed consent to participate in the study. Protocol was handled in accordance to the Declaration of Helsinki that involves the ethics of human medical researches. The approval of the study was obtained by the Ethical Committee of Faculty of Medicine, Cairo University.

Genotyping of rs636832 A/G and rs2740348 G/C

Extraction of genomic DNA

Three-milliliter venous PB was collected on a sterile vacutainer tube containing 5% ethylene diamine tetra-acetic acid. Samples were kept at −20°C to the time of DNA extraction. GeneJET™ DNA Purification Mini Kit (ThermoFisher, # K0781) was used for DNA extraction according to instruction’s procedure.

SNP genotyping assay

TagMan SNP assay (Thermo Fisher; #4351379) was utilized with context sequence for rs636832 A/G of AGO1; [VIC/FAM] TCTGATTCCAGAACATATCACTCCT[A/G] AAAGAAAGCTCAGTTAGGACAG. For rs2740348 G/C of GEMIN4; it was [VIC/FAM] AGCAGCCTCAACACCAAGTCTGGCT[C/G] TCGGAAGAGGGCCCTGTTACTCCCC.

DNA amplification

Thermo Fisher TaqMan® Genotyping Master Mix was in use (#4371353). The final volume of PCR reaction was 20 μL; 3 ml extracted DNA, 10 μL Master Mix, 0.5 μL SNP, and 6.5 μL distilled water. The mixture was gently mixed and put on a real-time PCR Thermal Cycler (Step One apparatus; Applied Biosystems). Amplification program started by hold for 10 min at 95°C followed by 45 cycles; each processed as denaturation for 15 s at 95°C then annealing/extension for 1 min at 60°C. Allelic and genotypic identification was done according to the plotted fluorescence signals.

Statistical analysis

Hardy–Weinberg equilibrium (HWE) was measured by a goodness-of-fit χ2 test. Data were evaluated using IBM SPSS advanced statistics version 20 (SPSS Inc., Chicago, IL). Mean and standard deviation (SD) used to demonstrate quantitative data. Qualitative data were expressed as frequency and percentage (%). Chi-square test or Fisher’s exact test was employed to examine the relations between qualitative variables. For quantitative data, two groups comparison was analyzed using Student’s t-test while ANOVA test was operated for three groups’ comparison. Risk estimation was expressed by odds ratio (OR) and 95% confidence interval (CI). Linkage disequilibrium (LD) was mathematically analyzed [22, [23]. CML patients’ survival rates were detected using Kaplan–Meier method and Log Rank Test. A p ≤ 0.05 was regarded as significant.

Results

Demographic and clinical features

Our CML cases were matched with the controls regarding age and sex with p > 0.05. Most of our cases were in the chronic phase at presentation. Eutos and Sokal scores were calculated to CML cases at diagnosis before treatment beginning (Table 1).

| Characteristics            | Cases (n=110) | Controls (n=110) | p-value* |
|----------------------------|--------------|-----------------|----------|
| Age                        |              |                 |          |
| Mean±SD                    | 41.68 ± 12.87| 43.3 ± 12.5     | 0.345    |
| Range                      | 20–75        | 21–75           |          |
| ≤40 n (%)                  | 52 (47.27%)  | 42 (38.18%)     | 0.173    |
| >40 n (%)                  | 58 (52.73%)  | 68 (61.82%)     |          |
| Gender                     |              |                 |          |
| Male n (%)                 | 56 (50.91%)  | 53 (48.18%)     | 0.686    |
| Female n (%)               | 54 (49.09%)  | 57 (51.82%)     |          |
| Clinical manifestations    |              |                 |          |
| CML phase                  |              |                 |          |
| Chronic n (%)              | 106 (96.30%) | -               |          |
| Accelerated n (%)          | 2 (1.82%)    | -               |          |
| Blastic n (%)              | 2 (1.82%)    | -               |          |
| Sokal score                |              |                 |          |
| Mean±SD                    | 0.84 ± 0.38  | -               |          |
| Range                      | 0.4–2.9      | -               |          |
| Low (≤0.8) n (%)           | 48 (43.64%)  | -               |          |
| Intermediate (0.8–1.2) n (%)| 58 (52.73%)  | -               |          |
| High (>1.2) n (%)          | 4 (3.64%)    | -               |          |
| Eutos score                |              |                 |          |
| Mean±SD                    | 61.591 ± 32.293| -              |          |
| Range                      | 18–195       | -               |          |
| Low (<87) n (%)            | 86 (78.18%)  | -               |          |
| High (>87) n (%)           | 24 (21.82%)  | -               |          |
| Treatment lines            |              |                 |          |
| 1st line (Imatinib)        | 110 (100)    | -               |          |
| 2nd line (Nilotinib)       | 41 (37.27)   | -               |          |
| 3rd line (Dasatinib)       | 16 (14.55)   | -               |          |
| Response to therapy        |              |                 |          |
| 1st line response          |              |                 |          |
| Responsive                 | 69 (62.73)   | -               |          |
| Resistant                  | 41 (37.27)   | -               |          |
| 2nd line response          |              |                 |          |
| Responsive                 | 94 (85.45)   | -               |          |
| Resistant                  | 16 (14.55)   | -               |          |
| 3rd line response          |              |                 |          |
| Responsive                 | 8 (50)       | -               |          |
| Resistant                  | 8 (50)       | -               |          |
| MMR at 12 months n (%)     | 68 (61.82)   | -               |          |
| Blastic transformation n (%)| 8 (7.27)    | -               |          |
| Death n (%)                | 8 (7.27)     | -               |          |

* p ≤ 0.05 is statistically significant, SD: Standard deviation, MMR: Major molecular response, PD: Progressive disease.

Our patients received Hydrea (3000 mg/day) at their initial presentation to control disease until Ministry of Health approval for TKIs. The first line of treatment was Imatinib at dose of 400 mg/day. In resistance to 1st line of treatment, shift to the 2nd line in the form of Nilotinib 800 mg/day was occurred that if failed to produce the optimal response; 3rd line (Dasatinib) 100 mg/day was started instead. On the regular follow-up to response to
therapy, major molecular response (MMR) was detected in more than 50% of our cases. Blastic transformation was observed in 8 cases that ended by death in 3 patients from the total 8 deaths on follow-up (Table 1).

**Genotyping of AGO1_rs636832 A/G and Gemin4_rs2740348 G/C**

The observed frequencies of genotypes of both SNVs were found to be concomitant to the expected under HWE in both case and controls with p > 0.05 (Table 2).

### Table 2: Genotypes and alleles frequencies of rs636832 and rs2740348 among cases and controls

| Characteristics | Cases (n=110) | Controls (n=110) | OR (95% CI) | p-value* |
|-----------------|--------------|-----------------|-------------|----------|
| **AGO1 rs636832 A/G** |              |                 |             |          |
| Genotypes       | n | %   | n | % |          |          |
| AA             | 5 | 4.55 | 2 | 1.82 |        | 0.389 |
| AG             | 29 | 26.36 | 25 | 22.73 |        |        |
| GG             | 76 | 69.09 | 83 | 75.45 | Reference |        |
| AA+AG          | 34 | 30.91 | 27 | 24.55 | 1.375 | (0.760–2.489) |
| **Alleles**    |              |                 |             |          |
| A              | 39 | 17.73 | 29 | 13.18 | 1.419 | 0.189 |
| G              | 181 | 82.27 | 191 | 86.82 | Reference |        |
| P HWE          | 0.601 | 0.997 | | | | |
| **GEMIN4 rs2740348 G/C** |              |                 |             |          |
| Genotypes      | n | %   | n | % |          |          |
| CC             | 92 | 83.64 | 97 | 88.18 | 0.435 |        |
| GG             | 1 | 0.91 | 0 | 0 |        | |
| GC             | 17 | 15.45 | 13 | 11.82 | 1.460 | (0.677–3.148) |
| GG+GC          | 18 | 16.36 | 13 | 11.82 | 0.355 |        |
| **Alleles**    |              |                 |             |          |
| G              | 19 | 8.64 | 13 | 5.91 | 1.505 | 0.273 |
| C              | 191 | 91.36 | 207 | 94.09 | 0.977 | (0.724–3.129) |
| **P HWE**      |              |                 |             |          |

* p ≤ 0.05 is statistically significant, HWE: Hardy–Weinberg equilibrium.

No statistically significant difference was observed regarding the genotypic and allelic frequencies between cases and controls for both variants. Using Logistic regression analysis revealed that the rs636832 A/G and rs2740348 G/C genotypes were not significantly associated with CML risk (Table 2).

LD analysis of the two SNPs revealed that there was a modest LD between rs636832 A/G and rs2740348 G/C among CML cases with p < 0.001, D'=0.51, and r²=0.1.

**Genotypes and clinical criteria of CML patients**

For AGO1 rs636832 A/G

GG genotype was the major homozygote genotype in our study. GG genotype in CML cases associated with male predominance even after sex stratification. Moreover, age stratification revealed that GG genotype was more overrepresented at younger age at diagnosis (≤40 years) in CML cases (Table 3 and 5).

GG genotype was detected to be more prevalent in CML cases with higher bad prognostic grades of both Eutos and Sokal scores. GG genotype was found to be more evident in patients not achieving MMR at 12 months than AA and AG genotypes or (AA+AG) genotypes. GG genotype compared to (AA+AG) genotypes showed more existence in CML patients who had no response to the 1st line of treatment with p < 0.05 (Table 3).

No statistical significance was found in CML patients with genetic modalities of rs636832 A/G regarding CML phases, response to 2nd and 3rd lines of treatment as well to the progression to blastic phase (Table 3).

For Gemin4_rs2740348 G/C

CC genotype was the major homozygote genotype in our work. It was found after sex stratification to be less predominant in female CML cases in comparison to GC and GG as well as to (GG+GC) genotypes with p = 0.03 and 0.009, respectively. Higher levels of Eutos score were encountered in CC genotype than those observed in (GG+GC) genotypes with p value=0.037 (Tables 4 and 5).

CC genotype in comparison to (GG+GC) genotypes was more presented in CML patients with suboptimal response at 12 months of therapy as well in unresponsive to 2nd and 3rd lines of TKIs with p < 0.05 (Table 4).

No statistical significance was found in CML patients with different genetic types of rs2740348 G/C regarding age, phases of the disease and grades of prognostic scores (Eutos and Sokal). In addition, no significant difference was detected regarding the progression to blastic transformation (Table 4).

**Overall survival (OS) of CML patients and genotyping**

Over 5 years follow-up, the number of deaths in genotypic analysis revealed no statistically significant difference for genotypic modalities in both variants (Tables 3 and 4). For AGO1_rs636832 A/G, the mean of OS in (AA+AG) and GG were 59.82 and 57.19 months, respectively. While in Gemin4_rs2740348 G/C, the mean of OS in (GG+GC), CC was 58.3 and 57.95 months independently with (p = 0.238 and 0.770, respectively) (Figure 1).

**Discussion**

Although CML was the initial malignancy with a definitive chromosomal specification (Ph chromosome), many transformations were recognized in its pathogenesis [24]. There is increase evidence of the importance of miRs and their biogenesis as well binding site genes in the pathogenesis along with prognosis of hematological malignancies including CML [6].
miRs are essential epigenetic regulators of genetic expression directly by transcription inhibition, mRNA degradation, or indirectly activate expression of mRNA by cleavage their inhibitors. They affect 80% of cellular transcriptome to be essential in the regulation of cellular properties of apoptosis, differentiation, and proliferation [7].

AGO1 and GEMIN4 are essential proteins of miRISC. miRISC has a function as oncogenes or tumor suppressor genes in carcinogenesis [25]. AGO1 and GEMIN4 genetic variants play an important role in the development and prognosis of multiple tumors. AGO1 rs636832 SNV was detected to be related to lung cancer risk, gastric carcinoma metastasis [10], [11]. GEMIN4 rs2740348 variants were found to affect the severity of prostatic cancer as well the risk of bladder cancer and renal cell carcinoma [12], [26], [27].

Table 3: rs636832 genotypic variants compared to clinical criteria of CML patients

| Characteristics | AGO1 rs636832 | n (% ) | GG | AG | AA+AG |
|-----------------|---------------|--------|----|----|-------|
| Sex             |               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Male            | 0 (0)         | 6 (35.3)| 50 (54.3)| 0.209 | 6 (33.3)| 50 (54.3)| 0.103 |
| Female          | 0 (0)         | 11 (64.7)| 42 (45.6)| 12 (66.7)| 42 (45.6)| 12 (66.7)| 42 (45.6)| 12 (66.7)| 42 (45.6)| 12 (66.7)| 42 (45.6) |
| Age group       |               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| ≤40 years       | 0 (0)         | 0 (0)| 48 (100)| 0.0000* | 48 (100)| 0.0000* |
| >40 years       | 1 (100)       | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Phase of disease|               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Chronic         | 0 (0)         | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Accelerated+ Blastic | 0 (0) | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Sokal score level|               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Mean ± SD       | 0.740 ± 0.182 | 0.741 ± 0.031| 0.874 ± 0.400| 0.0000* | 0.741 ± 0.312| 0.741 ± 0.312| 0.884 ± 0.400| 0.0000* |
| Line response (Imatinib) | 0 (0) | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Line response (Nilotinib) | 0 (0) | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Sokal score level|               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Mean ± SD       | 0.735 ± 0.401 | 0.785 ± 0.037| 0.869 ± 0.377| 0.0000* | 0.739 ± 0.390| 0.739 ± 0.390| 0.860 ± 0.377| 0.0000* |
| Low             | 0 (0)         | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Intermediate    | 0 (0)         | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| High            | 0 (0)         | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| EUTOS score level|               |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Mean ± SD       | 28 ± 12       | 69.855 ± 30.822| 43.118 ± 27.688| 0.001* | 69.855 ± 30.822| 43.118 ± 27.688| 0.001* |
| Death           | 0 (0)         | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |

Table 4: rs2740348 genotypic variants compared to clinical criteria of CML patients

| Characteristics | Gemini4 rs2740348 | n (% ) | GG | GC | CC | p-value* |
|-----------------|-------------------|--------|----|----|----|----------|
| Sex             |                    |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Male            | 0 (0)              | 6 (35.3)| 50 (54.3)| 0.209 | 6 (33.3)| 50 (54.3)| 0.103 |
| Female          | 0 (0)              | 0 (0)| 48 (100)| 0.0000* | 48 (100)| 0.0000* |
| Age group       |                    |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| ≤40 years       | 0 (0)              | 0 (0)| 48 (100)| 0.0000* | 48 (100)| 0.0000* |
| >40 years       | 1 (100)            | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Phase of disease|                    |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Chronic         | 0 (0)              | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Accelerated+ Blastic | 0 (0) | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |
| Sokal score level|                    |        | n (%) |      | n (%) |      | n (%) |      | n (%) |
| Mean ± SD       | 28 ± 12            | 17 (94.44)| 89 (96.74)| 0.0000* | 89 (96.74)| 0.0000* |
| Death           | 0 (0)              | 0 (0)| 1 (100)| 0.0000* | 1 (100)| 0.0000* |

* p ≤ 0.05 is statistically significant. SD: Standard deviation, MMR: Major molecular response, PD: Progressive disease.

gemini4 miRISC has a function as oncogenes or tumor suppressor genes in carcinogenesis [25]. AGO1 and GEMIN4, respectively, in prognosis and
response to therapy of Egyptian CML patients as GG genotype of rs636832 and CC genotypes of rs2740348 were appeared to be correlated with the bad clinical prognostic scores as well linked to the resistance to TKIs therapy. Hence, they can be implanted for prognosis prediction as well for the targeted therapy in CML disease.

Regarding AGO1_rs636832, our results revealed no a significance difference between the cases and controls as regard the frequencies for AA and (AG+GG) genotypes (p = 0.293, OR=1.375, and 95% CI = 0.760–2.489) as well for A and G alleles (p = 0.189, OR = 1.419, and 95% CI = 0.842–2.391). In respect to rs2740348, the frequencies for CC and (CG+GG) genotypes, as well C and G alleles frequencies in our cases showed no a statistically significance difference from their frequencies in the controls (p = 0.355, OR = 1.460, and 95% CI = 0.677–3.148) and (p = 0.273, OR = 1.505, and 95% CI = 0.724–3.129), respectively.

Modest LD between both variants was detected in CML patients with p < 0.001. LD is referred to non-random association of alleles at two or more chromosomes not necessary to be on the same chromosome that leads to the SNV alleles to be inherited together [22].

In consistent to our results, Gutiérrez-Malacatt et al. [9] investigated 312 CML patients and 487 healthy controls for rs636832 and rs2740348 by real-time PCR confirmation of their result was done by sequencing of 10% of their analyzed samples. Regarding rs636832, they found that the frequencies of AA, AG, and GG in cases were 0.23, 0.5, and 0.27 near to that in controls 0.26, 0.49, and 0.25, respectively. There was no a statistically significance difference in comparing AG to AA (p = 0.44 and OR [95% CI] = 1.15 [0.81–1.64]) as well as in comparing GG to AA (p = 0.35 and OR [95% CI] =1.21 [0.81–1.81]). Their allelic A and G frequencies were found to be in controls 0.5 and 0.49 similar to that in cases 0.48 and 0.52 independently with (p = 0.35 and OR [95% CI] = 1.10 [0.90–1.35]).

In addition, regarding rs2740348 Gutiérrez-Malacatt et al. [9] reported the frequencies of GG, GC, and CC genotypes in controls were 0.62, 0.35, and 0.04 and in cases were 0.65, 0.3, and 0.04, respectively. No
a statistically significance was found in comparison of CC to GG as well GC to GG genotypes \( (p = 0.66 \text{ and OR} [95\% CI] = 1.18 [0.57–2.44]) \) and \( (p = 0.26 \text{ and OR} [95\% CI] = 0.84 [0.61–1.14]) \) individually. Allelic G and C were 0.79 and 0.21 in controls and 0.8 and 0.2 in cases, respectively, with \( (p = 0.57 \text{ and OR} [95\% CI] = 0.93 [0.72–1.20]) \).

Our results revealed that GG genotype in CML cases showed male predominance compared to AA and AG or to \((AA+AG)\) with \( p < 0.001 \). After sex stratification, GG genotype showed higher frequencies in male cases than in male controls with \( p = 0.027 \). Furthermore, our female cases showed less frequencies of CC genotype of rs2740348 than found in female controls with \( p = 0.03 \).

In contrast to our finding, Gutiérrez-Malacatt et al. [9] showed that frequencies of AA, AG, and GG genotypes in male controls 0.23, 0.53, and 0.24 were near to it in male cases 0.26, 0.5, and 0.24, respectively. There was no statistical significance in the comparison of GG to AA or AG to AA genotypes with \( p = 0.78 \) and 0.49 independently. Furthermore, for rs2740348, they found the frequencies of GG, GC, and CC in female CML cases to be 0.68, 0.27, and 0.05, while in female controls were 0.64, 0.33, and 0.03, respectively, with \( p \) value on comparison of CC to GG and GC to GG genotypes were 0.37 and 0.29, respectively.

Furthermore, our cases after age stratification showed that GG genotype of rs636832 was more presented at younger age at diagnosis (<40 years) in cases than controls with \( p = 0.037 \) that was not obvious in rs2740348 \( (p = 0.183) \). This augments bad prognostic value of rs636832 in CML patients. Gutiérrez-Malacatt et al. [9] did not correlate the genotype frequencies with the age of their patients.

No statistical significance was found in our CML patients with genetic modalities of both variants regarding the phase of disease (CP vs. AP/BP). In homogenous to our results, Gutiérrez-Malacatt et al. [9] showed no difference in genotypic frequencies of both variants between early CML disease (CP) and late stage (AP/BP). In rs636832, frequencies of AA, AG, and GG genotypes in CP were 0.26, 0.48, and 0.26 and in AP/BP were 0.15, 0.63, and 0.22, respectively, with \( p = 0.56 \) and 0.17 in comparing GG to AA and AG to AA genotypes respectively. As well for rs2740348, the frequencies for CC, GC, and GG genotypes showed no statistically significant difference between CML phases with \( p > 0.05 \).

Regarding the prognostic clinical scores of our CML cases, GG genotype of rs636832 was found to be related to high grades of Sokal and Eutos compared to \((AA+AG)\) genotypes with \( p = 0.011 \) and 0.001 individually. As well, CC genotype of rs2740348 compared to \((GG+GC)\) showed high levels of Eutos score with \( p = 0.037 \). Gutiérrez-Malacatt et al. [9] did not relate the genotypic modalities of both variants to CML prognostic scores. We are the first to demonstrate these relations.

In respect to the therapeutic response in our CML cases, GG genotype of rs636832 compared to \((AA+AG)\) genotypes was found to be overexpressed in CML cases that resistant to the 1st line therapy as well in those who failed to obtain MMR at 12 months of treatment with \( p = 0.046 \) and 0.001, respectively. Similarly, CC genotype of rs2740348 compared to \((GG+GC)\) was overrepresented in CML cases that exhibited resistance to the 1st and 2nd line therapy as well in those did not reach the MMR at 1 year of therapy with \( p = 0.002 \), 0.056, and 0.040 independently. No statistical significance between the genetic modalities of both variants regarding response to 3rd line therapy, blastic transformation as well to OS of CML patients. In addition, no significant difference between the genetic modalities of rs636832 for the response to 2nd line therapy with \( p > 0.05 \).

In opposite to our results, Gutiérrez-Malacatt et al. [9] found no difference in response to Imatinib in both variants. As in rs636832, the frequencies of AA, AG, and GG in responsive were 0.24, 0.45, and 0.3 and in resistant were 0.23, 0.56, and 0.21 with \( p = 0.56 \) and 0.51 in comparison GG to AA and AG to AA genotypes, respectively. Furthermore, in rs2740348, the frequencies of GG, GC and CC in responsive were 0.64, 0.32, and 0.05 and in resistant were 0.66, 0.29, and 0.05 with \( p \)-value 0.98 and 0.74 in comparing CC to GG and GC to GG genotypes, respectively. However, they did not follow-up their resistant patients on the other lines of treatment as well at different intervals of routine follow-up therapy. Our study is the initial research to document these finding.

Their opposite results to our study may be related to ethnic difference as well to that the major homozygous genotypes of both variants in our population were different than that found in Mexican-Mestizos ethnic of the case–control study of Gutiérrez-Malacatt et al. [9]. The major homozygous genotypes of rs636832 and rs2740348 in our study were GG and CC, while in Gutiérrez-Malacatt et al. were AG and GG, respectively.

AGO1_rs636832 is an intronic un-translated variant that modifies the DNA three-dimensional structure and function with change in AGO1 expression. This affects the optimal production and action of various miRs with subsequent linkage to risk and prognosis of multiple diseases [28]. AGO1_rs636832 interacts through Wnt pathway that is essential in cancer development and prognosis. In addition, AGO1 regulates the transcription activity by stimulation of PI3K signaling [14]. Wnt pathway through PI3K signaling is involved in CML progression and therapeutic resistance [15]. This can give the basis of considering rs636832 as a bad prognostic as well poor therapeutic marker of CML in the present study.
GEMIN4_ rs2740348 is an exonic non-synonymous variation (Gln439Glu). It is an important component of miRISC that plays a pivotal role in carcinogenesis. GEMIN4 expression is related to the biogenesis of its linked miRs that affect cancer risk and prognosis [13]. GEMIN4 interacts through MR receptor [25]. MR receptor is a target of GAL1 protein which is a bad prognostic marker for CML [18], [19]. This supports the prevalence of rs2740348 in the poor clinical scores and therapeutic resistance in our study.

In homogenous to our finding, Song et al. investigated rs636832 to gastric cancer. They revealed that the AA genotype and A allele had a reduced risk of lymphatic metastasis of gastric cancer with no association to its risk [10]. Furthermore, Horikawa et al. 2008 documented no risk of rs636832 to renal carcinoma [12]. Discordant to the present study, Kim et al. found that the rs636832 AG or GG genotype has a lower risk to lung cancer than the AA genotype [11].

In contrast to our finding, Zhu et al., meta-analysis documented that GC+CC genotypes associated with increased risk to cancer [29]. The absence of risk association to CML to rs2740348 in our study can be explained by the predominance of CC genotype as a major homogenous genotype in our study was with the presence of GG genotype in only one CML case.

It has been found that GC genotype of rs2740348 was related to the increased risk of cancer prostate and cancer breast [12], [26]. However, renal carcinoma risk has been decreased with the combined GC+CC genotypes as well with GC genotype in the haplotype representation of GEMIN4 gene [12].

The variability in the findings of researches regarding these variants may be related to the ethnic difference, the discrete malignancy criteria in each variant, as well presence of other associated natural factors and linked genotypes with cancer risk contribution. Further researches are required to allow their efficient evaluation in different malignancies.

We faced some limitations due to the lack of the financing to do another confirming method of our results, but we relayed on its analysis in other nations studies by sequencing. Our study contributes an important role in clarifying the relationships between microRNA biogenesis genes and CML in Egyptian population.

Conclusion

Our study demonstrates the first record of an association between rs636832 and rs2740348 variants and prognosis of CML disease. The rs636832 GG and rs2740348 CC genotypes are associated with poor clinical prognostic scores and linked to the resistance to TKIs therapy. No association between CML risk and both variants.

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