Detection of BCR/ABL Fusion Gene in Chronic Myeloid Leukaemia Using Reverse Transcriptase Polymerase Chain Reaction

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**Abstract:**

Background: Chronic myeloid leukemia (CML) is a clonal myeloproliferative expansion of transformed, primitive hematopoietic progenitor cells. It involves myeloid, monocytic, erythroid, megakaryocytic, B-lymphoid, and occasionally T-lymphoid lineages.

Aim: This research was aimed at detecting the BCR-ABL fusion gene in patients with chronic myeloid leukaemia using reverse transcriptase polymerase chain reaction technique.

Material and Method: Eight adults (aged 45 years and above) blood samples were collected from the clinics of Lautech Teaching Hospital, Osogbo, Osun State, Nigeria and University College Hospital, Ibadan, Oyo State, Nigeria. RNA was extracted and purified from CML blood samples using Axyprep multisource Total RNA miniprep kit, USA procedures were followed.

Result: At the end of the reaction, amplifications were not observed at the annealing temperature of 55 ºC. The only bands were the primer sets and some DNA contaminations at the RNA controls for both multiplex and nested PCRs.

Conclusion: Total RNA extraction from the blood sample is crucial to the success of RT-PCR, without this, the study cannot run to completion. Meanwhile, in this experiment, it was observed that there were no bands when the eluted total RNA was run on gel electrophoresis. This indicates that there were no extraction of RNA. Rapid degradation of RNA is the most important factor impeding the analysis of gene expression in human cells and tissues.

**Keywords:** CML, BCR-ABL, RNA, electrophoresis, RT-PCR

1. Introduction

Chronic myeloid leukemia (CML) is a clonal myeloproliferative expansion of transformed, primitive hematopoietic progenitor cells. It involves myeloid, monocytic, erythroid, megakaryocytic, B-lymphoid, and occasionally T-lymphoid lineages[1]. Chronic myeloid leukemia was the first human disease in which a specific abnormality of the karyotype – the Philadelphia (Ph) chromosome – could be linked to pathogenetic events of leukemogenesis [2]. It was among the first neoplastic diseases in which therapy which a biologic agent (interferon) was found to suppress the leukemic clone and prolong survival [3].
The median age at presentation is 53 years, but all age groups, including children, are affected. Most patients also have thrombocytosis, which is consistent with the presence of a defect in a pluripotent hematopoietic stem cell. The typical symptoms at presentation are fatigue, anorexia, and weight loss, but about 40 percent of patients are asymptomatic, and in these patients, the diagnosis is based solely on an abnormal blood count. The most common abnormality on physical examination is splenomegaly, which is present in up to half of patients. The natural history of chronic myeloid leukemia is progression from a benign chronic phase to a rapidly fatal blast crisis within three to five years. The blast crisis is often preceded by an accelerated phase in which increasing doses of hydroxyurea or busulfan are required to lower the neutrophil count. In contrast to the maturation of chronic myelogenous leukemia cells during the chronic phase, during blast or lymphoblasts found in patients with acute leukemias [4].

The diagnosis of chronic myeloid leukemia is usually based on detection of the Philadelphia (Ph) chromosome. This abnormality, first described as a shortened chromosome 22 in 1960 [2] and then as at (9;22) translocation in 1973 [5], is present in 95 percent of patients. Another 5 percent have complex or variant translocations involving additional chromosomes that have the same end result, which is fusion of the BCR (breakpoint cluster region) gene on chromosome 22 to the ABL (Abelson leukemia virus) gene on chromosome 9. The Philadelphia chromosome is found in cells from the myeloid, erythroid, megakaryocytic, and B-lymphoid lineages, indicating that chronic myeloid leukemia is a stem cell disease [6].

Chronic Myeloid Leukaemia has been the paradigm for the integration of molecular biology with clinical care. In future gene expression array studies will classify the biologic state of disease, and suggest routes of tailored therapy based on the profile or response genes. Chronic myelogenous leukemia (CML) constitutes a clinical model for molecular detection and therapy surveillance of malignant disease since this entity was the first leukemia shown to be associated with a specific chromosomal rearrangement, the Philadelphia (Ph) translocation t(9;22)(q34;q11),[2][5][4 ]; which generates two chimeric genes: BCR-ABL[7][8] on the derivative chromosome 22, and ABL-BCR on the derivative chromosome 9. BCR-ABL is transcribed and translated in most patients into a 210 kDa fusion protein with deregulated tyrosine kinase activity. ABL-BCR is expressed in about 60% of patients with CML, but probably lacks any biological function[9].

After the first discovery of translocation t(9;22)(q34;q11) which is between breakpoint cluster region (BCR) gene on chromosome 22 and ABL gene on chromosome 9, many chromosomal aberrations causing fusion genes in cancer have been shown by cytogenetic techniques[10][11][12]. Discovery of molecular techniques have led to more precise determination of percentage of cases with certain chromosomal aberrations. Translocation t(9;22) is observed in 95% of CML (chronic myelogenous leukemia) patients, in 2-10% of pediatric AML (acute myeloid leukemia) cases, and in20-50% of adult ALL (acute lymphoblastic leukemia) cases In addition, this translocation might be seen in less than2% of AML cases and rarely seen in lymphoma and myeloma cases. Different transcription products, namely p190, p210, and p230, might arise from different breakpoints on BCR gene [13][14][15].

Inv(16)(p13;q22) is found in almost all AML-M4Eo subgroup patients. This translocation is formed due to fusion of core binding factor beta (CBFB) gene and myosin heavy chain 11 (MYH11) gene (9). Translocation t (15;17) (q22; q21) is detected in acute promyelocytic leukaemia (APL) M3 patients which constitute 10% of all AML cases. In t(15;17) translocation, a chimerical protein is formed by fusion of promyelocytic leukaemia (PML) gene on 15q22 and retinoic acid receptor alpha (RARA) gene on 17q21. This fusion protein is observed in 98% of APL cases[16]. Translocation t(8;21)(q22;q22) is the product of fusion of eight-twenty-one (ETO) gene on chromosome 8 and acute myeloid leukemia 1 (AML1) gene on chromosome 21. This translocation is found in 18% of AML-M2 cases (Miyoishi et al., 1991). Translocation t (12;21) (p13; q22) is formed by fusion of TEL/ETV6 gene on chromosome 12 and AML1/CFBFA2 on chromosome 21 and is observed in 22% of pediatric ALL cases [17]. Fusion of AF4 gene on chromosome 4 with MLL gene on chromosome 11 causes formation of t (4;11) (q21; q23) translocation, which is seen in 50-70% of infant ALL cases and in approximately 5% of adult ALL cases [18][13]. Translocation t(1;19) (q23;p13) is formed as a result of fusion of PEB1 gene on chromosome 1 and E2A gene on chromosome 19 and is observed in 3% of adult ALL cases[19].

The degree of tumor load reduction is an important prognostic factor for patients with CML on therapy[20]. Response is expressed at three levels: (1) hematological response, defined as the normalization of the peripheral blood values and of spleen size; (2) cytogenetic response defined as the proportion of residual Ph-positive metaphases; and (3) molecular response defined according to the method used as the proportion of the residual BCR-ABL gene, transcript, or protein.

Attempts at designing therapeutic tools for CML based on our current knowledge of the molecular and cell biology of the disease have concentrated on 3 main areas—the inhibition of gene expression at the translational level by "antisense" strategies, the stimulation of the immune system’s capacity to recognize and destroy leukemic cells, and the modulation of protein function by specific signal transduction inhibitors.

2. Materials and Methods

2.1. Study Sites

Eight adults (aged 45 years and above) blood samples were collected from the clinics of Lautech Teaching Hospital, Osogbo, Osun State, Nigeria and University College Hospital, Ibadan, Oyo State, Nigeria.

The samples were analyzed at the Molecular Biology Laboratory, Ladoke Akintola University of Technology, Mercyland Wing, Osogbo, Osun State, Nigeria.
2.2 Materials

PCR machine, Axygen RNA kit, water bath, eppendorf tubes, micropipettes, microcentrifuge, primers NB1, AB13, CA3 and B2A, Oven, distilled water, Agerose powder, Boric acid, Taq polymerase, Tris – Borate EDTA buffer, electrophoretic machine, Ultraviolet light and a computer system.

2.3 RNA Extraction and Purification

RNA was extracted and purified from CML blood samples using Axyprep multisource Total RNA miniprep kit, USA and the following procedures were followed.

Exactly 50µl of blood sample was added to 400µl of RNA extraction Buffer R-1 and immediately homogenized by vortexing for 5 minutes and the supernatant was discarded. Then 150µl of Buffer R-11 was added to pellet and vortexed for 30seconds at room temperature. The clarified supernatant was transferred into a 1.5ml microfuge tubes provided and 250µl of isopropanol was added and was mixed by vortexing for 30seconds.

A spin/vac column was placed into a 2ml microfuge tube, the binding solution of isopropanol was transferred into it and was centrifuged at 6000xg for 1minute at room temperature. The filtrate was discarded from the spin/vac column and the spin/vac column was placed back into the same 2ml microfuge tube then 500µl of Buffer W1A was added to the spin/vac column and was centrifuge at 12,000xg for 1minute. Again, the filtrate was discarded and the spin/vac column was placed into another 2ml microfuge tube and 700µl of Buffer W2 was added and centrifuged at 12,000xg for 1minute. This process of washing was repeated for the second time with 700µl of Buffer W2.

The filtrate was discarded from the 2ml microfuge tube and the spin/vac column was place back into the 2ml microfuge tube. This is then centrifuged at 12,000xg for 1minute to remove residual wash solution. Finally, the spin/vac column was transferred into a clean 1.5ml microfuge tube (provided). 20µl of Buffer TE (nuclease free) was added to the center of the membrane and was allowed to stand for 1minute at room temperature and then centrifuged at 12,000xg for 1 minute to elute the total RNA.

2.4 Removal of DNA using DNA se I

The DNA was removed with the following preparations:

| 10µl of the total RNA extracted |
| 2µl of BS |
| 2µl of DNase I buffer |
| 2µl of DNase I |
| 4µl of PCR water |

All these made up 20µl of solution, which was then incubated 37ºC for 30 minutes and then 65ºC for 10 minutes.

2.5 c DNA synthesis (Reverse transcriptase)

RNA/primer/dNTP mix was made by combining the following components in a sterile RNase – free microfuge tube:

| Total RNA | 3µl |
|---|---|
| Primer dT23VN | 2µl |
| dNTP mix | 4µl |
| nuclease-free water | 9µl |
| Total volume | 16µl |

Table 1

The constituted RNA/primer/dNTP mix was incubated at 70ºC for 5 minute, spun briefly and then promptly chilled on ice.

The following components were added to the 16µl RNA/primer/dNTP solution and was mixed well by pipetting up and down.

| 10xRT Buffer | RT test | RT control |
|---|---|---|
| Murine RNase inhibitor | 2µl | 2µl |
| m-MuLV reverse transcriptase | 0.5µl | 0.5µl |
| Nuclease-free water | 1µl | 1.5µl |
| Final volume | 2µl | 20µl |

Table 2

The 20µl cDNA synthesis reaction was incubated at 42ºC for one hour and enzyme inactivation at 80ºC for 5 minutes. The reaction volume was brought to 50µl with water and was then stored at -20ºC. This then ready for PCR amplification.
2.6. Preparation for Multiplex PCR

PCR master mix for four primer sets; c5e, ca3, bcr-c and b2a were prepared in an eppendorf tubes for all cDNA samples to be processed.

C5e  5’ataggaTCTTTTGCAACCGGGTCTGA3’
Ca3  5’TGGTGAACCTGGCTGATGTAATTGCTTGG 3’
Bcr-c  5’ACCGCATGTTCCGGGACAAAAG 3’
Ba2  5’TTTACAGCTTTCCCTGTGACAT 3’

The master mix for these primers contained; 54µl of PCR water, 18µl of c5a, 18µl of ca3, 18µl of bcr-c, 18µl of b2a and 36µl of taq mix. 27µl of the master mix was aliquotted onto the 3µl of the cDNA in PCR tubes.

2.7. Multiplex PCR

Each PCR experiment was repeated at least once and included a negative and three positive controls. DNA derived from K562 cell line was used as one of the positive controls. Two other positive controls used were from known ALL and CML patients with breakpoints at e1a2 and b2a2 respectively. PCR parameters were 35 cycles at 96 ºC for 1min, 55 ºC for 50seconds, and 72 ºC for 1min, followed by 10 min extension at 72 ºC. This was carried out a GeneAmp PCR System 9700 by Applied Biosystem.

2.8. Agarose Gel Electrophoresis

Electrophoresis was carried out in a 2.0% agarose slab gels in Tris-borate buffer (89 mM Tris, 89mM boric and EDTA pH 8.0). About 2.0g of agarose was weighed into a clean flask and dissolved with 95ml of distilled water and 5ml of tris-borate buffer. This was heated on the microwave oven to dissolve the agarose and shaken frequently to prevent charring, and then cooled to 50 ºC. 0.5µg/ml of ethidium bromide was then added in order to stain and enable visualization of DNA bands.

A dry Perspex plate (20 X 20cm) was sealed with paper tape to form a mould, the warm agarose was poured into the mould and the comb was well positioned immediately, positioned and clamped in the mould ensuring that there was about 0.5-1.0mm of agarose between the bottom of the teeth and the base of the so that sample wells are completely sealed. The gel was allowed to set (20-30 minutes at room temperature), the comb and autoclave tape were removed and mount in the electrophoresis tank and electrophoresis buffer was added to cover the gel. The PCR products were mixed with 10µl of loading buffer, loaded into the slots of the submerged gel using a micropipette and run on the electrophoresis machine at 70 V for 45 minutes. The 100bp DNA ladder was mixed with 10µl of loading buffer and also loaded on a slot of the agarose gel. Sigma Gel Loading Solution Type I was used as the loading buffer. It contains 0.25% (w/v) Bromophenol blue, 0.25% (w/v) Xylene cyanole FF and 40% (w/v) sucrose in water.

cDNA bands were then visualised and photographed by filtered ultraviolet illumination on the syngene gel documentation system (Syngene, UK). The sizes of DNA product (310, 385, 481 and 808bp) were compared against the 100bp DNA ladder and the results were obtained.

3. Results

At the end of the reaction, amplifications were not observed at the annealing temperature of 55 ºC. The only bands were the primer sets and some DNA contaminations at the RNA controls for both multiplex and nested PCRs.

The experiment was tried onStaphylococcus aureus as when cultured in the presence of antibiotics, oxacyclinin melaitin broth, produces a Methycilline Resistance Staphylococcus aureus gene which is expressed by RNA. After the RNA extraction from the bacteria, RNA gel electrophoresis could not reveal 18s and 22s bands of RNA to confirm successful extraction of total RNA.

Legend: SM: Size Marker, R1 to R8 are sample 1 to 8 respectively.
4. Discussion and Conclusion

Total RNA extraction from the blood sample is crucial to the success of RT-PCR, without this, the study cannot run to completion. Meanwhile, in this experiment, it was observed that there were no bands when the eluted total RNA was run on gel electrophoresis. This indicates that there were no extraction of RNA.

Rapid degradation of RNA is the most important factor impeding the analysis of gene expression in human cells and tissues. Once a tissue has been removed from its normal environment, the relative rates of RNA synthesis and degradation change. This in turn gave rise to changes in the relative proportions of various RNA species within a tissue, as well as to an overall reduction of RNA concentration. Contamination of samples with exogenous rib nucleases (RNases), particularly those used in the laboratory during RNase treatment of DNA isolates, is universally recognized as an as important cause of RNA loss. The use of meticulous laboratory technique to reduce this contamination absolutely essential to preserve RNA substrates, not sufficient to permit analysis of all RNA target as proven by [21].

It may be due to the potent RNase, eosinophil-derived neurotoxin which is the major factor in reducing the sensitivity of RT-PCR as demonstrated by [22]. [23] also observed in his study that, several endogenous non-secretory RNases complicate the molecular analysis of human tissue. Failure of RNA extraction from method used could be as a
result of conditions like temperature, prolong time of extraction, centrifugation, vortex time and manufacturer’s procedure. Thus, in this study, environmental factors such as irregular power supply in this parts of the world (southwestern, Nigeria) may have made the extraction procedure for RNA isolation and purification unsuccessful. The insensitivity of Manufacturer’s procedure has also contributed to the failure. Therefore, it is highly recommended that this study should serve as a template for developing a quicker, more sensitivity and more reproducible method that will be suitable for this condition than previous RT-PCR based assays for Philadelphia chromosome (bcr/abl translocation) that uses either cell lysate or isolated RNA.

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