An In-House Method for Molecular Monitoring of BCR-ABL

Hakkı Oğün Sercan, İlgin Öztürk, Ceyda Çalışkan, Melek Pehlivan, Zeynep Sercan
Dokuz Eylül University, School of Medicine, Department of Medical Biology and Genetics, İzmir, Turkey

Abstract

Objective: At present, there are a limited number of facilities in Turkey that can provide reliable real-time quantitative (RQ)-PCR BCR-ABL results. The present study aimed to test a cost-effective, in-house method of BCR-ABL quantification, including verification of the method by RQ-PCR validation tests.

Material and Methods: BCR-ABL and ABL target sequences were cloned into pJET1.2 vectors, from which calibrators were prepared and used as templates in RQ-PCR reactions to generate standard curves. Dilutions of K562 cells (representing an in vitro simulation of BCR-ABL transcript reduction) were analyzed.

Results: Standard curves were generated from calibrators. These curves were then used to calculate the BCR-ABL and ABL copy numbers; in which linear BCR-ABL results were obtained. Repetitive experiments showed that our methodology was able to detect 1 BCR-ABL positive cell from among 1x10^5 cells.

Conclusion: The method described herein is suitable for implementation with any RQ-PCR instrument and/or kit for quantify BCR-ABL transcripts.

Key Words: Chronic myeloid leukemia, BCR-ABL, Real-time PCR, Molecular response

Özet

Amaç: Türkiye’de BCR-ABL gerçek zamanlı kanıtلاف polimeraz zincir tepkimesi ile BCR-ABL düzey tayini yapabilen merkezlerin sayısı, ihtiyaçın altında kalmaktadır. BCR-ABL kanıtlayışı için laboratuarımızda geliştirilmiş olduğumuz yöntem temel alınarak, (PZT) ile kanıtlayışınn basamakları, iç kontrolleri ve geçerlilik testleri anlatılmıştır.

Gereç ve Yöntemler: BCR-ABL ve ABL hedef dizileri pJET1.2 vektörlerine klonlandıktan sonra kalibratörler hazırlanmış ve gerçek zamanlı PZT tepkimelerinde kalıp olarak kullanılarak standart eğriler çizilmiştir. Bu standart eğriler kullanılarak K562 hücre dilüsyonlarının analizi yapılmıştır.

Bulgular: Önceden elde edilen standart eğriler kullanılarak BCR-ABL ve ABL kopya sayıları hesaplanılmış; lineer sonuçlar elde edilmiştir. Tekrarlayan deneylerle yöntem duyarlılığının 1x10^7 hücrede bir BCR-ABL pozitif hücreyi saptayacak düzeyde olduğu gösterilmiştir.

Sonuç: Bu makalede tarif edilen protokollerin geçerliliği/güvenirliliği gösterilmiştir. Kullanılan yaklaşım, BCR-ABL kanıtlayışının hedefleyen diğer gerçek zamanlı PZT kit ve cihazlarına da uyarlantılabilmektedir.

Anahtar Sözcükler: Kronik miyeloid lösemi, BCR-ABL, Gerçek zamanlı PCR, Moleküler yanıt
**Introduction**

Chronic myeloid leukemia (CML) is a clonal myeloproliferative disorder of hematopoietic stem cell origin. The Philadelphia chromosome (Ph)—der(22)t(9;22) (q34;q11)—is the hallmark of CML, and is formed via reciprocal translocation between the long arms of chromosomes 9 and 22. The translocation results in the juxtaposition of the 5’ end of the BCR gene and the 3’ end of the ABL gene, generating a BCR-ABL chimeric oncogene with aberrant kinase activity. The introduction of imatinib mesylate—a small-molecule competitive inhibitor of the BCR-ABL kinase—in 1998 substantially changed the management of CML [1]. Decades of research on CML has led to very effective molecularly targeted therapies.

Since the late 1980’s the percentage of Ph-positive cells in bone marrow based on cytogenetic analysis has been the gold standard for monitoring the response to therapy [1,2,6]. Prior to the use of tyrosine kinase inhibitors for the treatment of CML a molecular response was rarely achieved and it was therefore not feasible to routinely monitor BCR-ABL transcript levels, with the exception of patients that underwent bone marrow transplantation [1-6]. Currently, tyrosine kinase inhibitors are the preferred first-line therapy for CML and based on IRIS trial data a 3-log reduction in BCR-ABL transcript levels after 12 months correlates with progression-free survival [1-6].

Molecular monitoring of CML patients has become clinically very important. Developments in real-time quantitative (RQ)-PCR technology as well as real-time chemistry have enabled routine, high throughput detection of BCR-ABL transcript levels. In turn, there has been an expected increase in the number of centers that provide RQ-PCR results for BCR-ABL mRNA. The rapid increase of these centers revealed that there were significant variations between individual laboratories in reporting RQ-PCR data [5-12]. The International Scale (IS) has been adopted for reporting BCR-ABL values based on the application of laboratory-specific conversion factors that are derived using patient samples [6-12].

Presently, few hospitals in Turkey can provide reliable RQ-PCR BCR-ABL results. Effective implementation of RQ-PCR requires knowledge of reaction kinetics and instrumentation, as well as extensive experience. Lack of experienced personnel could lead to over-dependence on the technical assistance of RQ-PCR kit manufacturers, which in turn may prove to be less than satisfactory. Herein we describe a cost-effective, in-house method developed in our laboratory, and detailed descriptions of validation tests and an in vitro simulation of transcript reduction, all which could be adapted to any RQ-PCR methodology chosen for detecting BCR-ABL transcript levels.

**Materials and Methods**

The study protocol was approved by the Dokuz Eylül University Clinic & Laboratory Research Ethics Committee.

**Primers and probes**

RQ-PCR results for BCR-ABL are usually presented as a ratio of the BCR-ABL transcript level and the reference control gene transcript level. Normalization to a reference gene compensates for variation in the efficiency of the reaction and RNA quality between samples. The most widely used reference control gene in BCR-ABL monitoring is the wild-type ABL gene [9,13].

PCR primers used for BCR-ABL facilitate amplification of b2a2 and b3a2 transcripts, which comprise the BCR exon b2/b3 and ABL exon 2 (primers synthesized by Metabion GmbH, Martinsried, Germany). A dual-labeled (5’ 6-FAM and 3’ TAMRA as quencher) hydrolysis probe (TaqMan Probe, Roche Diagnostics, Mannheim, Germany) was used for real-time detection of product accumulation during each cycle. The primer and probe sequences are shown in Figure 1. We used the wild-type ABL gene as the reference gene. The primers used most widely for quantification of ABL also amplify the BCR-ABL fusion transcript [9,12,13]. It has been reported, and also observed by us, that this may cause a bias in results, especially when a large...
quantity of BCR-ABL is present in the sample [9,12,13].
The primer and hydrolysis probe sequences we used for
ABL (Figure 1) were determined to hybridize only to wild-
type ABL and not BCR-ABL.

**Cell lines**

K562 is a BCR-ABL-positive cell line derived from a
CML patient in erythroid blast crisis. RS4;11 is a BCR-
ABL-negative cell line derived from an acute myelocytic
leukemia (AML) patient and was used as a BCR-ABL-neg-
ative cell line control. Both cell lines were purchased from
DSMZ (Deutsche Sammlung von Mikroorganismen und
Zellkulturen GmbH, Germany) and grown in RPMI-1640
medium supplemented with 15% FBS and 1% L-gluta-
mine in a 5% CO₂-saturated incubator at 37 °C.

**Standard curves and copy numbers**

Calibrators necessary to create the standard curves for
product quantity were generated by cloning the target
DNA into plasmids, followed by copy number determi-
nation and preparation of serial dilutions. RNA or DNA
standards may be used for RQ-PCR, but DNA standards
are reported to have superior stability. The Europe Against
Cancer (EAC) Programme established standardized pro-
tocols for fusion transcript quantification and concluded
that DNA plasmids were appropriate for constructing
standard curves, whereas RNA and cDNA did not provide
the stability of plasmids [10,13].

Total RNA was extracted from K562 cells (Macherey
Nagel-Nucleospin RNAII, Düren, Germany). To avoid
genomic DNA contamination RNA samples were treated
with RNase-free DNase I, according to the manufacturer’s
instructions. cDNA was synthesized using a First Strand
cDNA Synthesis Kit (MBI-Fermentas K1611, St. Leon-
Rot, Germany), 2 μg of total RNA, and random primers,
according to the manufacturer’s instructions. cDNA (5 μL)
was used as the template in a conventional PCR reaction
performed with primers ENF501 and ENR561 (Figure 1)
to amplify the target BCR-ABL region. The PCR product
was cloned into a pJET1.2 vector, in accordance with the
manufacturer’s instructions (CloneJET PCR Cloning Kit,
MBI Fermentas, St. Leon-Rot, Germany).

Following colony selection, plasmid DNA was iso-
lated (NucleoBond® PC20 Macherey Nagel, Düren, Ger-
many) and analyzed spectrophotometrically. Similar pro-
cedures were performed for the ABL gene. Primers AblF
and ENR561 (Figure 1) were used to amplify the ABL
sequence in a conventional PCR reaction, in which cDNA
from K562 cells were used as template. The amplification
product was cloned into a pJET1.2 vector, followed by
colony selection and plasmid analysis.

To prepare the calibrators the plasmid copy number
μL⁻¹ volume was determined using the following equation
(in brief, the molecular weight of the DNA template can
be determined by multiplying the number of base pairs
(bp) by the weight of 1 mol of a bp estimated to be 650 g;
using 6.022 x 10²³ [Avogadro’s number] molecules mol⁻¹,
the number of molecules of the template can be calculated
by first converting to ng [multiplying 1 x 10⁹] and then
multiplying by the quantity of template) [14]:

\[
copy \text{ number } \mu \text{L}^{-1} = \frac{\text{quantity (ng } \mu \text{L}^{-1}) \times 6.022 \times 10^{23}}{[\text{plasmid length (bp)} \times 1.10 \times 650]}
\]

Calibrators containing 10², 10³, 10⁴, 10⁵, and 10⁶ gene
copies were prepared for both the BCR-ABL and ABL genes.
These calibrators were used as templates in RQ-PCR rea-
tions to generate standard curves. RQ-PCR was carried out
using a LightCycler 2.0 (Roche Diagnostics, Mannheim,
Germany) instrument and LightCycler TaqMan Master Kit
(Roche Diagnostics, 0453528001, Mannheim, Germany).
Reactions were performed in a 20-μL volume with 10
pmol of each primer and probe. The same thermal profile
was optimized for BCR-ABL and ABL: pre-incubation for
10 min at 95 °C, followed by 45 amplification cycles of
denaturation at 95 °C for 10 s, primer annealing at 58 °C
for 40 s, and primer extension at 72 °C for 2 s. dH₂O was
included as a no template control. Fluorescence was mea-
sured during the 72°C segment in each cycle. Data were
analyzed using LightCycler v.4.0.0.23 software (Roche
Diagnostics, Mannheim, Germany). Standard curves for
both BCR-ABL and ABL were generated and saved for fur-
ther use (Figure 2). All samples in glass capillaries were
subsequently run in 2% agarose gel electrophoresis to
check for size and non-specific amplifications. All data
were derived from independent experiments performed in
triciplicate.

**Serial dilutions of K562 cells**

Dilutions of K562 cells were prepared in the back-
ground of the BCR-ABL-negative RS4;11 cell line. Mixtures
of BCR-ABL-positive K562 and BCR-ABL-negative RS4;11
cells were prepared, and the total cell number was always
10⁵. The sample mixtures of K562/ RS4;11 cells were pre-
pared so that they contained 10,000, 1000, 100, 10, 1, or
0 K562 cells. These samples represent in vitro simulation
of BCR-ABL transcript reduction in a CML patient under-
going therapy. Total RNA isolation and cDNA prepara-
tion were performed as described in the previous section.
The cDNA was used as a template for determining the
BCR-ABL transcript levels in quantitative PCR reactions. RQ-PCR was performed in independent triplicate sets, as described in the previous section.

Results and Interpretation

RQ-PCR results for the K562/RS4;11 sample mixtures were analyzed using the previously determined standard curves. These standard curves were used to calculate the quantity of BCR-ABL and ABL transcript (copy numbers) in the samples. Change/reduction in the BCR-ABL transcript level was expressed as the ratio of BCR-ABL:ABL (Figure 3). As expected, amplification product was not observed in the dH₂O-negative controls, whereas amplification was evident in the 10⁶ copies µL⁻¹ BCR-ABL-pos-
Laboratory practice be followed when setting up PCR assays and during post-PCR processing of samples. Though not addressed herein, there are excellent reviews of recommended pre-PCR procedures in the literature, including handling of blood samples, RNA extraction, cDNA synthesis, and reverse transcription [8-10,17]. A dH2O-negative control (BCR-ABL negative cells), along with low and high positive controls should be included in each RQ-PCR run to monitor assay performance. Any changes in technique, protocol, or instrument should be accompanied by a thorough evaluation. Standard curves should be generated after opening a new real-time kit or when a new batch of primer/probe is diluted.

Over the past five years significant advances leading to more wide-spread adaptation of RQ-PCR for BCR-ABL have occurred. Implementation of the IS has improved the comparability of results between laboratories, and recently accredited reference reagents for BCR-ABL quantification have been developed [7]. It is essential that more diagnostic centers in Turkey qualify to report BCR-ABL results according to the IS. In essence, the procedure involves assignment of a laboratory-specific conversion factor (CF) to convert BCR-ABL measurements to the IS [15]. Log BCR-ABL values of the same sample set are compared to reference and local laboratories via linear regression [17]. The results are considered linear when the correlation coefficient of any 2 laboratories is >0.98 [17]. The pre-requisite for the procedure is that all in-house validation tests be performed and confirmed prior to application. In this context, the present study aimed to describe a cost-effective, in house method for BCR-ABL quantification and to illustrate an example for RQ-PCR validation testing, as well as to provide a description of DNA plasmids that may be implemented into any RQ-PCR methodology to quantify BCR-ABL transcripts. The primary advantage of the presented methodology over widely used commercial kits—in addition to being cost effective—is that once optimized and validated, both absolute and relative quantification can be performed, whereas most commercial kits are restricted to providing relative quantification results. In conclusion, the methodology described herein is suitable for implementation into any RQ-PCR instrument and/or kit for quantifying BCR-ABL transcripts.

**Discussion**

RQ-PCR is a technically demanding, yet very powerful tool. The exquisite sensitivity of the assay is also its weakness. It is essential that extreme care be taken to avoid false-positive results due to cross-contamination and carry-over of RNA/DNA from a previous amplification. The calibrators themselves are the greatest threat of contamination, as they are cloned DNA containing DNA target regions. Their preparation, storage, and handling should not be performed in the same laboratory in which patient samples are processed. All equipment, including pipette sets, kits, reagents, paper, pens, workbooks, and lab coats should be dedicated for use only in that particular laboratory. It is of great importance that the rules of good laboratory practice be followed when setting up PCR assays and during post-PCR processing of samples. Though not addressed herein, there are excellent reviews of recommended pre-PCR procedures in the literature, including handling of blood samples, RNA extraction, cDNA synthesis, and reverse transcription [8-10,17]. A dH2O-negative control (BCR-ABL negative cells), along with low and high positive controls should be included in each RQ-PCR run to monitor assay performance. Any changes in technique, protocol, or instrument should be accompanied by a thorough evaluation. Standard curves should be generated after opening a new real-time kit or when a new batch of primer/probe is diluted.

**Conflict of Interest Statement**

The authors of this paper have no conflicts of interest, including specific financial interests, relationships, and/or affiliations relevant to the subject matter or materials included.
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