**Determination of ctxAB Expression in Vibrio cholerae Classical and El Tor Strains using Real-Time PCR**

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Cholera is an infection of the small intestines caused by the bacterium *V. cholerae*. It is a major cause of health threat and also a major cause of death worldwide and especially in developing countries. The major virulence factor produced by *V. cholerae* during infection is the cholera toxin. Total mRNA extraction and reverse transcription was performed for making ctxAB cDNA. Relative Real-Time PCR analysis showed unequal enterotoxin production in *V. cholerae* strains. The results showed that, classical strain produces cholera toxin more than El Tor strain.

**Key words:** *Vibrio cholerae*, RT-qPCR, ctxAB expression
finally would decrease the ctxAB toxin production (11). The RS$_1$ region contains rstA, rstB and rstC fragments which have positive effect on ToxR gene and therefore increase ctxAB toxin production (12). Bakhshi et al. reported several V. cholerae attacks in Southwest and Southeast of Iran between 2005-2009 (13, 14). As the level of production of a protein is somehow related to its mRNA quantity, we therefore aimed to determine V. cholerae strains that can produce more ctxAB toxin.

**Material and Method**

**Bacterial strains and growth conditions**

We used two standard strains named V.cholerae O$_1$ Classic ATCC 14035 & V. cholerae O$_1$ El Tor N16961. The isolates were confirmed by biochemical and immunological tests. Serotyping was performed using monoclonal O$_1$ antiserum and mono-specific Inaba and Ogawa antisera (Pasteur institute, Paris, France). All selected strains were cultured according to the AKI-SW method and standard growth curve were drawn (15).

**Isolation of RNA and RT-PCR**

Approximately, $2 \times 10^8$ cfu/ml from each strain, was used for total RNA extraction. Total RNA was isolated from the strains isolated randomly from each V. cholerae grown in AKI medium using the RNeasy® Protect Bacteria Mini Kit (Qiagen Inc, GMBH, Germany) and the integrity and purity was checked. Equivalent concentrations of total RNA from each strain were selected as template for RT-PCR. cDNA synthesis and PCR amplification were performed using QuantiTec Reverse Transcription Qiagen kit (Qiagen Inc, GMBH, Germany). RT-PCR was performed in the presence of random primer at 42°C for 10 min. After cDNA synthesis, the ctxAB and recA genes were PCR amplified for checking. PCR amplification was performed for 35 cycles as follows: initial denaturation at 94°C for 5 min, then denaturation at 94°C for 30 sec, annealing at 60°C for 30 sec, extension at 72°C for 30 sec. At the end of the 35th cycle, reaction mixtures were left at 72°C for another 3 min. Five microliters of each reaction mixture was loaded on a 1% agarose gel and subjected to electrophoresis to confirm that the unique amplified fragment correspond to the expected ctxAB gene fragment and recA as housekeeping gene (16).

**Real-Time PCR**

Prepared cDNA was quantified using SYBR green I dye. Four primers were designed by AlleleID 6 software, 5’-CAGTCAGGTGGTTATGC-3’ (ctxAB-F) and 5’-ATCGTGCTAACL-
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AAATCCC-3' (ctxAB-R) for gene of interest and 5' -ATTGAAGCCGAATGGGGCATAG- 3' (recA-F) and 5' -TACACATACAGTTGGATTGCTTG AGG- 3' (recA-R) for housekeeping gene. Those primers were specific to ctxAB and recA and amplified a 115 & 106 bp respectively. SYBR green Real-Time PCR assay was performed with a 20 µl PCR mixture volume containing 2x QuantiTec SYBR Green PCR Master Mix (Qiagen Inc, GMBH, Germany), 0.25 µM specific primer sets, and 2 µl of cDNA sample. Amplification of the primers, data acquisition, and relative analysis were carried out in Chromo4 BioRad Real-Time PCR. PCR reactions were performed as followings: one cycle of 95 °C for 5 min, then 40 cycles of 95 °C for 15 sec, 60 °C for 30 sec. Following the amplification, melting temperature analysis of the PCR products was performed to determine the specificity of the PCR. The standard curve was established by using genomic DNA for each studied gene to confirm that the primers amplified at the same rate and to validate the experiment (55-95°C with warming of 0.2°C per sec). Reverse transcription and PCR positive controls (RNA and DNA, respectively) and negative controls (distilled water) were included in each run. The Real-Time PCR reaction was performed twice assayed in triplicate. Classical V. cholerae O1 ATCC 14035 was used as a standard control.

Results

The specificity of each primer set for V. cholerae was tested by PCR with genomic DNA extracted via boiling. Only one size of amplicon was obtained by PCR reaction for ctxAB and recA genes when DNA from V. cholerae strains was used. The amplicons obtained for each gene were verified by sequencing. The presence of a single PCR product was confirmed by Real-Time PCR for each set of the primers using melting curve analysis that resulted in a single product-specific melting curve (Fig 2). The PCR efficiencies varied between 1.90 and 1.94. The relative expression ratio was calculated for each gene of interest by a mathematical model described by ∆C_T method. The Cycle Threshold (C_T) results are showed in table 1. Histogram and samples C_T values are indicated in Fig 3.

Discussion

In our study, the results are derived by using “relative” method and ∆C_T formula. By considering that ctxAB primers have been carefully designed, the amount of standard deviation results are close to

Fig 2. Melting curves of ctxAB and recA genes.

Fig 3. Level of cholera toxin production in classical and EL Tor strains.
Table 1: Cycle threshold (Ct) results for the V. cholerae O1 Classic ATCC 14035 & V. cholerae O1 El Tor 62013

<table>
<thead>
<tr>
<th>Strain</th>
<th>Mean Ct recA±SD</th>
<th>Mean Ct ctxAB±SD</th>
<th>Mean ∆∆Ct</th>
<th>Mean Ratio*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classic</td>
<td>25.58±0.14</td>
<td>24.35±0.39</td>
<td>1.23</td>
<td>1.0</td>
</tr>
<tr>
<td>El Tor</td>
<td>22.85±0.13</td>
<td>26.02±0.15</td>
<td>4.35</td>
<td>0.06</td>
</tr>
</tbody>
</table>

*∆∆Ct was calculated as: ∆Ct (test) - ∆Ct (calibrator). Ratio=efficiency^-∆∆Ct.

zero, the primer-dimer bands were not seen, because the concentration of participating primers in the reaction had been set up. Also, for more accuracy and sensitivity, the PCR efficiency in Real-Time PCR reactions were calculated and replaced with ratio 2 in computational equations.

Dirita et al. showed that the level of cholera toxin was higher in the classical strain compared to El Tor strain because of the influence of toxR on cholera toxin production in the classical strain (17). In our study, in addition to Dirita et al. results, we determined quantitative cholera toxin production between classic and El Tor strains. Both classical and El Tor strains have been shown to express equivalent levels of ToxR. In contrast, the classical strain expresses more ToxT, which has a higher binding affinity to toxbox region, resulting in higher expression of cholera toxin (18).

The comparison of the Ct of the El Tor strain with classical strain shows that toxin production in El Tor strain is approximately 16–17 times lower than in the classical strain (P Value<0.05 for each). This result is consistent with other reports because the amount of pathogenicity in classic strain is more than El Tor strains (18). Furthermore, histogram and samples Ct results indicated that toxin production in classical strain is higher than El Tor strain (Fig 3). In conclusion, the results of our study suggest that other factors modulate the production of cholera toxin by regulating the CTX cassette, supporting the idea that cholera toxin production in V. cholerae classical and El Tor strains is a multi-factorial phenomenon.

Acknowledgment

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References

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