Expression of Selected Inflammatory Cytokine Genes in Bladder Biopsies

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ABSTRACT

Besides the deregulation of oncogene and tumour suppressor gene, bladder carcinoma can also be caused by inflammation. To date, the association of inflammatory cytokines with carcinoma of the bladder (especially the transitional cell carcinomas) is not fully understood. In this study, we report an attempt to examine expression patterns of pro- and anti-inflammatory cytokine genes from normal and tumour tissue biopsies of the human bladder. Our molecular assays involved the use of the GeneXP™ Human Cytok3 kit and the Reverse Transcription – Polymerase Chain Reaction test. Due to limitation in our experimental process, mainly attributed by inconsistencies in the results obtained between the two assay systems, we cannot reach a conclusion regarding the association of the six selected inflammatory cytokine genes (IL-8, IL-12A, IL-18, TGF-β1, TGF-β2, and TGF-β3) with bladder carcinoma. However, our data provided early novel evidence of expression of four inflammatory cytokine genes, namely IL-12A, TGF-β1, TGF-β2, and TGF-β3 in tissues derived from the human bladder.

Keywords: Expression analysis, inflammatory cytokines, bladder biopsies

INTRODUCTION

Studies have shown that over-proliferation of cells due to deregulation of oncogenes and tumour suppressor genes, and certain epigenetic mechanism are not the only causative factors of cancer (Coussens & Werb, 2001). In fact, inflammation and infections has been suggested to be amongst the triggers of tumour initiation (Lin & Karin, 2007). Inflammation-mediated carcinogenesis can be explained, in part, by the activation of the inflammatory cells that release mutagenic oxidant-generating enzyme (Michaud, 2007), which may cause irreversible genomic alterations in proliferating epithelial cells (Coussens & Werb, 2007).

According to Michaud (2007), the association between inflammation and carcinogenesis in squamous cell carcinoma of the bladder accounts for the high occurrence of this subtype of bladder cancer in many developing countries where parasitic infection by Schistosoma haematobium is common. However, evidence of such association in cases of transitional cell carcinoma (TCC) of the bladder, a predominant subtype in most developed country and also in Malaysia, is less clear. In this short report, we reveal an attempt to investigate inflammation as a potential contributing event in local cases of transitional cell carcinoma of the bladder. We sought the strategy of simultaneous expression analysis of selected pro- and anti-inflammatory cytokine genes in normal and tumour bladder biopsies. Our hypothesis was that differential expression of inflammatory cytokine genes between normal and tumour biopsies, and also between pro- and anti-inflammatory cytokines would form the basis to suspect a link between inflammation and carcinogenesis of TCC.

MATERIALS & METHODS

Sample collection and total RNA isolation

Normal bladder and tumour tissue biopsies were from patients admitted to the Sarawak
General Hospital (Kuching, Sarawak) who have provided informed consent. Collected tissues were stored in cryovials and snap frozen in liquid nitrogen immediately after excision. They were subsequently stored at -80°C freezer in the laboratory until use. Twenty one tumours and four normal tissues were used. Ten to twenty miligrams (0.5 cm$^2$) of tissue specimen were used in the isolation process. The total RNA isolation was conducted in an RNase-free environment. TRIzol Tri Reagent was used to lyse the tissues and extract the RNA and these were carried out according to the manufacturer’s protocol (Invitrogen, USA). The RNA pellet was dissolved in 30 to 50 µl of nuclease-free water (Promega, USA) depending on the size of the pellet. After DNase treatment, the total RNA extracted was quantified using UV spectrophotometer (Ultrospec 1100) at $A_{260}$ and $A_{280}$ wavelength.

**Gene XP™ Human Cytokine Assay and RT-PCR**

A total of 1 µg of total RNA was used in reverse transcription reaction. An M-MLV reverse transcriptase (Promega, USA) was used in this reaction to a final volume of 20 µl, and this was done according to the manufacturer’s protocol. For the Gene XP™ Human Cytokine Assay kit, six genes were designed to be amplified simultaneously. The target amplicons consist of IL-8 (~590 bp), TGF-β3 (~470 bp), IL-12A (~360 bp), TGF-β1 (~290 bp), TGF-β2 (~240 bp) and IL-18 (~160 bp). Briefly, the PCR was carried out by mixing 10 µl of 2X master mix PCR, 4 µl of 5X human cyto-3 primer, 200 ng of cDNA and sterile dH$_2$O to a final volume of 20 µl. The kit was also supplemented with positive control DNA. Six microlitres of the DNA was used in the PCR reaction as positive control to the PCR reaction. The mixture was the subjected to PCR amplification at 94°C for 15 mins, 40 cycle of 94°C for 30 s, 63°C for 90 s, 72°C for 90 s and final extension at 72°C for 10 mins. Ten microlitres of PCR products was resolved in 2% agarose gel. The intensity of the bands was quantified using AlphaEase® FC Imaging Software (Alpha Innotech Corporation, California, USA). In this study, the quantification was done by using 50 bp DNA marker as a reference marker. The GAPDH gene was used as an internal control to affirm the equimolar concentration of our cDNA. The intensity value of each band was normalized against that of GAPDH, and the final value considered as normalised value.

**RESULTS**

**Cytokine profiling by Gene XP™ Human Cytokine Assay**

As shown in Figure 1, amplification using the cytokine kit yielded six bands as stated in the manufacturer’s protocol. The amplified targets consist of IL-8 (~590 bp), TGF-β3 (~470 bp), IL-12A (~360 bp), TGF-β1 (~290 bp), TGF-β2 (~240 bp) and IL-18 (~160 bp). Positive control DNA provided in the kit was used. However, no bands for IL-8 and IL-12A were visible. Only one of our tumour samples (S6T) shows the presence of all six bands of the targeted transcripts. The TGF-β3 amplicon was not detected in Samples S12T and S26T, while IL-12A is absent in Samples S7T, S21T and S26T.
Table 1. List of primers used in RT-PCR confirmation test.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Primer Sequence (5’-3’)</th>
<th>Primer Length (bases)</th>
<th>Optimized Annealing Temperature (°C)</th>
<th>Expected size of amplicons (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TGF-β1 Fwd</td>
<td>gga cta tcc acc tgc aag act atc</td>
<td>24</td>
<td>60</td>
<td>448</td>
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<td></td>
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<tr>
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<td>23</td>
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<td>515</td>
</tr>
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<td>TGF-β2 Rev</td>
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<td></td>
<td></td>
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<tr>
<td>TGF-β3 Fwd</td>
<td>ctc ttc cca gct cac aca tga a</td>
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<td>60</td>
<td>397</td>
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<td>TGF-β3 Rev</td>
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<td></td>
<td></td>
<td>21</td>
</tr>
<tr>
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<tr>
<td>IL-8 Rev</td>
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<td>466</td>
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<tr>
<td>IL-12A Rev</td>
<td>tca gat agc tca tca ctc tat cca tag tc</td>
<td></td>
<td></td>
<td>29</td>
</tr>
</tbody>
</table>

Figure 1. The amplification of selected cytokines via GeneXP™ Human Cytokine 3 assay kit. M represents 50 bp DNA marker. Band 1 represents IL-8 gene, Band 2: TGF-β3; Band 3: IL-12A; Band 4: TGF-β1; Band 5: TGF-β2 and Band 6: IL-18. Lane M represents 50 bp DNA marker, Lane 1: S10N, Lane 2: S14N, Lane 3: S27N, Lane 4: S29N, Lane 5: S2T, Lane 6: S3T, Lane 7: S4T, Lane 8: S6T, Lane 9: S7T, Lane 10: S9T, Lane 11: S12T, Lane 12: S16T, Lane 13: S19T, Lane 14: S20T, Lane 15: S21T, Lane 16: S24T, Lane 17: S25T, Lane 18: S26T, Lane 19: S27T, Lane 20: S28T, Lane 21: S29T, Lane 22: S30T, Lane 23: S31T, Lane 24: S33T and Lane 25: S34T.
RT-PCR confirmation test

Figure 2 shows the amplification of TGF-β1, TGF-β2, TGF-β3, IL-8, IL-18, and IL-12A respectively. In RT-PCR confirmation test, the mean of the band intensity from the triplicates was used to plot a graph that summarizes the expression level of all six genes throughout the sample. Figure 3 shows the expression level of the six genes generated from Gene XP Human Cyto-3 kit and that of the same genes from RT-PCR test. The amplification of TGF-β1 at ~450 bp was detected in all samples except in two of the normal tissues (S10N and S27N) (Lane 1 and 3, Figure 2) and two of the tumour tissues (S2T and S21T) (Lane 5 and 15, Figure 2). These results did not tally with the cytokine assay kit (Figure 1) which shows the presence of TGF-β1 in all tissues. For TGF-β2 (~500 bp), the amplicon was detected in minute amount in all samples except in four tumour samples S30T, S31T, S33T and S34T (Lane 22-25, Figure 2). These results also did not tally with the cytokine assay kit which shows the presence of this amplicon in all samples (Figure 1). For TGF-β3, the amplicon is observed in two of the normal tissues (S27N and S29N, Lane 3 and 4) and five of the tumour tissues (S3T, S30T, S31T, S33T and S34T) (Lane 6, 22-25, Figure 2). Again, this is not in agreement with the cytokine assay kit in which this amplicon is absent in two tumour sample, S21T and S26T (Figure 1). IL-8 is consistently detected in all samples except in S7T (Lane 9, Figure 2). This result contradicts the expression profile generated from the cytokine assay kit which shows the presence of IL-8 in only S6T (Figure 1). IL-18 is observed in all samples except in S10N, S14N and S27N. Its presence in the second and third replicate was detected in a minute amount. Its presence in cytokine assay kit however was easily detected throughout the samples studied. Finally, in the case of IL-12A, of which is faintly observed samples S27T and S28T, but clearly observed in S16T, S30T, S31T, S33T and S34T (Figure 2). However, it is detected in all samples except for S7T and S21T in the results from the Gene XP Human Cyto-3 kit (Figure 1).

DISCUSSION

Results from the two assays done did not provide conclusive evidence to support the role(s) of pro-inflammatory cytokine genes (IL-8, IL-12A and IL-18) and anti-inflammatory genes (TGF-β1, TGF-β2 and TGF-β3) in the tumourigenesis of BC. Firstly, results from the RT-PCR assays on all six cytokine genes tested are not consistent with observation of their expression profiles procured via the Gene XP Human Cyto-3 kit assay. Secondly, there is no definitive pattern of expression among any of the six genes that showed differential level between normal and tumour samples. Lastly, unless a quantitative assessment of expression level is investigated (via real time PCR strategy), any consideration on observed expression patterns or differential expression (if any) remains an approximation, at best. Indeed, a repetition of similar experiments is required in order that proper conclusion can be inferred. This is especially so when the test on the positive control for the Gene XP Human Cytok 3 kit also requires redoing.

Except for IL-8 and IL-18, the cytokine genes selected for this study have never been brought into the context of gene expression activities in bladder-derived tissues. Despite inconclusive findings, we did show evidence of expression of IL-12A, TGF-β1, TGF-β2, and TGF-β3 in bladder tissues. In the case of IL-8, Inoue and coworkers (2000) have shown it to be involved in angiogenesis of bladder tumour in nude mice. In addition, Thalmann and coworkers (2000) revealed that the level of IL-8 and IL-18 can be used to monitor the response of superficial bladder cancer to Bacillus Calmette-Guerin (BCG)-based intravesical therapy. Inevitably, more work has to be done, and our experiments has to be repeated in order that rational inferences can be obtained regarding the role(s) of IL-12A, TGF-β1, TGF-β2, and TGF-β3 in bladder carcinoma.
Figure 2. RT-PCR validation on genes from GeneXP assay. The item for each lane is as indicated in Figure 1.
Figure 3. Charts of the expression levels of the amplified genes via GeneXP™ Human Cytokine assay kit and selected inflammatory cytokine genes as tested via RT-PCR. This graph was plotted based on the means of the normalised value.
REFERENCES


