Cost-effectiveness analysis of lamivudine, telbivudine, and entecavir in treatment of chronic hepatitis B with adefovir dipivoxil resistance

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Abstract: The purpose of this study was to analyze the cost-effectiveness of lamivudine (LMV), telbivudine (LdT), and entecavir (ETV) in treatment of chronic hepatitis B with adefovir dipivoxil (ADV) resistance. Two hundred and fifty-two patients were recruited and screened for resistance to ADV and randomly assigned into three groups: LMV + ADV, LdT + ADV, and ETV + ADV. The ratio of biochemical response, virological response, seroconversion of hepatitis Be antigen (HBeAg)/hepatitis Be antibody (HBeAb), viral breakthrough, and the cost and effectiveness of treatments were analyzed. A comparison of the results of the ratio of biochemical response, virological response and seroconversion of HBeAg/HBeAb, showed no statistical difference between the three groups, with the economic cost of LMV + ADV the lowest, LdT + ADV the middle, and ETV + ADV the highest. The side effects of the three plans are all rare and tolerable. LMV + ADV is the optimal rescue strategy, and LdT + ADV the alternative selection in the economically less developed regions, while ETV + ADV was used in the economically developed regions.

Keywords: chronic hepatitis B, adefovir dipivoxil, lamivudine, telbivudine, entecavir, resistance

Introduction
Chronic infection with the chronic hepatitis B virus (HBV) is a leading cause of liver-related morbidity and mortality.1,2 It is estimated that approximately two billion people are infected with HBV in the world, which results in approximately 500,000 deaths every year, mainly due to its complications including cirrhosis and hepatocellular carcinoma (HCC).3,4 Evidence-based medicine has demonstrated that effective antiviral treatment of chronic hepatitis B (CHB) can reduce the risk of long-term complications and improve patient survival.5,6 Current nucleos(t)ide analogs (NUCs) approved by the US Food and Drug Administration (FDA) for CHB patients include lamivudine (LMV), adefovir dipivoxil (ADV), entecavir (ETV), tenofovir disoproxil fumarate (TDF), and telbivudine (LdT).7,8 Though ETV and TDF have been recommended as the first-line options for treatment of naïve CHB patients, they are not in widespread used in countries with limited health resources due to the high daily cost or difficulty of availability,9,10 and therefore LMV and ADV are still widely used in the world, especially in the economically less developed regions due to their low cost and easy availability.11,12 ADV was recommended by certain scholars because it has less drug resistance ratio than LMV, while LMV was the earliest available pharmacon and was still used frequently in many economically less developed countries. The ratio of therapy with ADV for 1-3 years associated with mutations in the polymerase (particularly in N236T or N181T) is about 0%, 1.6%, and 3.1%, respectively.13 If the HBV develops a mutant, ADV is
recommended to be used continuously and combinative with a second drug without cross-resistance such as LMV, LdT, and ETV. This is because the HBV reproduce quickly and it is making it very difficult for sequential monotherapy to prevent the virus reproduction, which results in significantly lower probability of virologic response and significantly higher risk of virologic breakthrough in the “switch” group than in the “add-on” group.14 Because the cost of the LMV, LdT, or ETV is greatly different from each other, we aim to compare the cost-effectiveness of the three pharmacons for treatment of chronic hepatitis B with adefovir dipivoxil resistance.

Materials and methods

Patients

This is a prospective double-blind study, and a total of 252 patients diagnosed with CHB who were admitted to our hospital between January 2006 and August 2014 were recruited and screened for resistance to ADV. CHB patients who met the following criteria were included in the study: serum chronic hepatitis B e antigen positive and serum HBV DNA level ≥500 copies/mL. ADV resistance was defined on the basis of genetic testing (RTN236T or RTN181T) or presence of virological breakthrough,15 which we defined as a documented rise in serum HBV DNA by ≥1 log10 IU/mL above the nadir, or to a detectable level (≥500 copies/mL) after achieving virological response while continuing ADV therapy. Exclusion criteria were: previous treatment for chronic hepatitis B with LMV, LdT, or ETV; coinfection with hepatitis C or human immunodeficiency virus; other forms of liver disease; breast-feeding, pregnancy or inadequate contraceptive measures; coexisting serious medical disease; and evidence of hepatic decompensation or HCC. Demographic data, factors of sex, age, alanine aminotransferase (ALT), and HBV DNA were not statistically different between the three subgroups (Table 1).

Table 1 Baseline characteristics of the study population

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Age (mean:SD)</th>
<th>Males/ females</th>
<th>ALT (IU/L)</th>
<th>HBV DNA (log10 copies/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LMV + ADV</td>
<td>88</td>
<td>41±11</td>
<td>45/43</td>
<td>137.9±37.1</td>
<td>4.98±1.34</td>
</tr>
<tr>
<td>LdT + ADV</td>
<td>84</td>
<td>38±13</td>
<td>43/41</td>
<td>146.7±30.9</td>
<td>5.14±2.13</td>
</tr>
<tr>
<td>ETV + ADV</td>
<td>80</td>
<td>40±12</td>
<td>42/38</td>
<td>138.9±32.4</td>
<td>5.02±1.36</td>
</tr>
<tr>
<td>P-value</td>
<td></td>
<td>0.644</td>
<td>0.718</td>
<td>0.857</td>
<td>0.784</td>
</tr>
</tbody>
</table>

Notes: *, vs LMV + ADV group, P<0.05; **, vs LdT + ADV group, P<0.05.
Abbreviations: ALT, alanine aminotransferase; HBV, hepatitis B virus; LMV, lamivudine; ADV, adefovir dipivoxil; LdT, telbivudine; ETV, entecavir.

Treatment of patients

The patients were randomly divided into three groups: LMV + ADV (n=88), LdT + ADV (n=84), ETV + ADV (n=80). Drug usage16,17: LMV (GlaxoSmithKline Pharmaceutical Company Limited, London, UK) 100 mg oral administration once a day; LdT (Beijing Novartis Pharma Ltd, Beijing, People’s Republic of China) 600 mg oral administration once a day; ADV (GlaxoSmithKline Pharmaceutical Company Limited) 10 mg oral administration once a day; ETV (Bristol-Myers Squibb Pharmaceutical Company Limited, New York, NY, USA) 0.5 mg oral administration once a day. This study was approved by the institutional review board at Gannan Medical University and was conducted in accordance with the 1975 Declaration of Helsinki.

Biochemical and serologic responses

A peripheral blood sample was taken from the patients in the morning, after fasting for at least 8 hours. Hepatitis B surface antigen (HBsAg), anti-HBsAg, hepatitis B e antigen (HBeAg), anti-HBe, anti-hepatitis B core (Hbc) (total), and anti-hepatitis B core (Hbc IgM were identified by using the third-generation microparticle enzyme immunoassays (Abbott Laboratories, Chicago, IL, USA). Serum alanine aminotransferase (ALT), aspartate aminotransferase levels, creatinine, and creatine kinase were measured with an automatic biochemistry analyzer (Olympus AU5400, Tokyo, Japan) according to standard laboratory procedures. The biochemical response was defined as normalization of ALT levels.

Virological analysis

HBV DNA was investigated by real-time polymerase chain reaction (RT-PCR) (COBAS TaqMan 48 analyzer; Roche DiagnosticGmbH, Shanghai, People’s Republic of China) according to manufacturer’s instructions. Virological response was defined as a decrease in serum HBV DNA to undetectable levels by PCR assays (<500 copies/mL). HBeAg response was defined as seroconversion of HBeAg. Virological breakthrough was defined as an increase in serum HBV DNA by 1 log10 IU/mL above the nadir, or to detectable level (≥500 copies/mL) after achieving virological response during retreatment.

Genotypic analysis

HBV mutations were analyzed by direct sequencing. Extracted from serum samples, the HBV DNA polymerase gene region was amplified by nested-PCR using specific primers. We employed the primers and PCR program of Osiowy et al.18 Briefly, the primers, forward primer: spr1F (5’-GTTCCAAGAAACAGTAAAGCCC-3’) and the reverse
primer: spr1R (5'- GAAAGGCCTTGAAGTGGCG-3'), were used in the first round PCR. The inner primers: sense, spr2F (5'- GGTTGCCATTCTCCTATTTTCTAGG-3') and antisense, spr2R (5'- ACTTTCCACATCAAAGGCC-3') were used for nested PCR. The following PCR thermal-cycling program was performed: 35 cycles at 94°C for 30 seconds, 56°C for 30 seconds (first round) or 50°C for 30 seconds (second round), and 72°C for 40 seconds. The intended fragments were amplified using 2x PCR master mix solution (Tiangen Biotech Co., Ltd, Beijing, People’s Republic of China) with 5 µL of DNA extract and 2 µL of the first round PCR product. After amplification of the polymerase gene, the amplicons (730 bp) were visualized by agarose gel electrophoresis and gel purified using High Pure PCR Product purification kit (Roche Diagnostic GmbH). The purified PCR products were bidirectionally sequenced commercially (SEQLAB, Gottingen, Germany) using inner primers.

**Cost calculation**

A cost calculation was performed by means of identification and subsequent quantification of resources used, and assigning unitary cost to each. The costs are presented in dollars, for the year 2008. Unitary cost of LMV, LdT, ADV, and ETV was US$1.8, 2.2, 2.7, and 4.9 respectively.

**Cost-effectiveness ratios (CER) and incremental cost-effectiveness ratio (ICER)**

Cost effectiveness ratio (CER), standing for the cost for each unit of effectiveness produced by each therapeutic plan, was computed by dividing the difference in the mean costs of the three therapies (referred to as costs) and was assessed by the following formulas: 

\[
\text{CER}_{\text{LMV} + \text{ADV}} = \frac{\text{Cost}_{\text{LMV} + \text{ADV}} - \text{Cost}_{\text{ETV} + \text{ADV}}}{\text{Effectiveness}_{\text{ETV} + \text{ADV}} - \text{Effectiveness}_{\text{LMV} + \text{ADV}}}
\]

With reference to LMV, the ICER, defined as the additional cost incurred to achieve an extra unit of effectiveness, was calculated by applying the following formula: 

\[
\text{ICER}_{\text{LMV} + \text{ADV}} = \frac{(\text{Cost}_{\text{LMV} + \text{ADV}} - \text{Cost}_{\text{ETV} + \text{ADV}})}{(\text{Effectiveness}_{\text{ETV} + \text{ADV}} - \text{Effectiveness}_{\text{LMV} + \text{ADV}})}
\]

**Sensitivity analyses**

The uncertainties were minimized with sensitivity analyses that were performed using univariate analysis, varying the parameters of cost and effectiveness by either increasing or decreasing the value. To evaluate the robustness of the method and reduce uncertainties, the following types of analyses were performed: varying 10% discount on the prices for each drug; increasing the dosage of LMV from 100 to 150 mg/day; and adjusting the ratios of LMV arm response rate in LMV + ADV at 96 weeks with the intention of achieving the “threshold values”.

**Statistical analysis**

Statistical testing was performed by using SPSS version 12 (SPSS Inc., Chicago, IL, USA). The data are reported as median (range). Data are expressed as mean ± SEM of the indicated number of separate experiments. Statistical comparison between experimental group and control was performed by using one-way ANOVA analysis and unpaired two-tailed Student’s t-tests (for measurement data) or Chi-square test (for percentage). *P<0.05 was considered significant.

**Results**

**Biochemical response and seroconversion of HBeAg/hepatitis Be antibody (HBeAb)**

As shown in Figure 1A, at week 4, 12, 24, 36, 48, and 96, the ratio of patients with biochemical response was
6.8%, 25%, 51.1%, 72%, and 93.1% in LMV + ADV group respectively; 7.1%, 27.4%, 51.2%, 72.6%, 83.3%, and 95.2% respectively in the LdT + ADV group; 7.5%, 27.5%, 52.5%, 75%, 87.5%, and 97.5% respectively in the ETV + ADV group. There were no differences in virological response between the three groups. As shown in Figure 1B, at weeks 36, 48, and 96 of treatment, ratio of seroconversion of HBeAg/HBeAb was 23.7%, 34.1%, and 47.7% respectively in the LMV + ADV group; 23.8%, 34.5%, and 48.8% respectively in the LdT + ADV group; and 25.0%, 35.0%, and 51.3% respectively in the ETV + ADV group. There were no differences between the ratios of seroconversion of HBeAg/HBeAb in the three groups.

Virological response
As shown in Figure 2A, at weeks 4, 12, 24, 36, 48, and 96, HBV DNA levels decreased to 4.18±1.13, 3.62±0.38, 2.76±0.36, 2.03±0.23, 1.76±0.12, and 0.31±0.42 respectively in the LMV + ADV group; to 4.13±1.03, 3.26±0.62, 2.63±0.28, 1.94±0.43, 1.65±0.16, and 0.28±0.09 respectively in the LdT + ADV group; and to 3.98±0.92, 3.14±0.70, 2.43±0.09, 1.89±0.34, 1.55±0.12, and 0.26±0.38 respectively in the ETV + ADV group. As shown in Figure 2B, at week 4, 12, 24, 36, 48, and 96, the ratio of patients with virological responses was 6.8%, 26.1%, 65.9%, 76.1%, 84.1%, and 90.9% in LMV + ADV group respectively; 7.1%, 26.2%, 66.7%, 77.4%, 85.7%, and 92.9% respectively in the LdT + ADV group; and 7.5%, 27.5%, 70%, 78.7%, 87.5%, and 93.8% respectively in the ETV + ADV group. There were no differences between the virological response between the three groups.

Viral breakthrough and resistance
At week 96 of treatment, three patients (3.4%) developed viral breakthrough and genotypic mutation, including two RtM204V and one RtL180M, in the LMV + ADV group; two patients (2.2%) developed viral breakthrough and genotypic mutation, including one Rta181V and one Rtn236T, in the LdT + ADV group; and two patients (2.5%) developed breakthrough and genotypic mutation, including one Rtn236T and one Rtl180M + RtI184G, in the ETV + ADV group. There were no statistical differences between the ratios of viral breakthrough in the three groups (Table 2).

Cost-effectiveness and incremental cost-effectiveness ratio analysis of the three groups
At week 96 of treatment, in the LMV + ADV, LdT + ADV, and ETV + ADV group, the cost was US$3,024, US$3,292.8 and US$5,107.2 respectively, the ratio of negative conversion of HBV DNA was 90.9, 92.9, and 93.8 respectively, the cost-effectiveness ratio of negative conversion of HBV DNA was 33.3, 35.4, and 544 respectively; Ratio of seroconversion of HBeAg/HBeAb was 47.7, 48.8, and 51.3 respectively, cost-effectiveness ratio of seroconversion of HBeAg/HBeAb was 63.4, 67.5, and 99.5 respectively; Ratio of non-genotypic mutation ratio was 96.6%, 97.6%, and 97.5% respectively, cost-effectiveness ratio of non-genotypic mutation was 31.3, 33.7, and 52.4 respectively. In reference to LMV + ADV, incremental cost-effectiveness ratio of negative conversion of HBV DNA of LdT + ADV and ETV + ADV group was 134.4 and 718.3 respectively, incremental cost-effectiveness ratio of seroconversion of HBeAg/HBeAb was 244.4 and
578.7 respectively, incremental cost-effectiveness ratio of nongenotypic mutation was 268.8 and 2,314.7 respectively. These results showed that LMV was the least economically costwise, LdT the middle, and ETV the most economically costwise (Table 3).

### Sensitivity analyses

In the three groups of patients, varying the discount rate to 10%, LMV + ADV was more cost-effective than LdT + ADV and ETV + ADV. If LMV dose increases to 150 mg or the ratio of virological response increases to 91.9% at 96 weeks, there is a favorable impact on the average cost-effectiveness analysis for LMV + ADV. The sensitivity analysis demonstrates the robustness of the results obtained with the base case scenario with the three variables significantly impacting the results (Table 3).

### Side effects

All the patients tolerated treatment well, and no patient discontinued the therapy. At week 36 of treatment, ratio of side effects happening in the LMV + ADV group was 1.1%, with one patient demonstrating blood urea nitrogen (BUN) elevation (14.2 mmol/L), ratio of side effects in LdT + ADV group was 1.2%, with one patient demonstrating BUN elevation (14.7 mmol/L) and one patient with creatine kinase elevation (138.6 µmol/L), ratio of side effects happening in the ETV + ADV group was 1.3%, with one patient demonstrating BUN elevation (14.6 mmol/L). At week 96 of treatment, ratio of side effects happening in the LMV + ADV group was 4.5%, with two patients having blood urea BUN elevation (mean 13.9 mmol/L), one patient having diarrhea, and one patient having nausea; ratio of side effects in LdT + ADV group was 7.1%, with two patients having BUN elevation (mean 14.5 mmol/L), one patient having creatine kinase elevation (142.3 µmol/L); ratio of side effects in ETV + ADV group was 5.0%, with two patients having BUN elevation (mean 13.9 mmol/L), one patient having headache, and one patient having dizziness. The results showed there were statistical differences between the ratios of side effects between LdT + ADV group and LMV + ADV or ETV + ADV group. No statistical differences were observed between the ratios of LMV + ADV group and ETV + ADV group (Table 5). Acute renal failure or myopathy was not observed in any patient during the rescue therapy. The patients suffering side effects recovered after undergoing symptomatic treatments.

### Table 2 Viral breakthrough and genotypic mutation of the three groups

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Genotypic mutation (n, %)</th>
<th>Nongenotypic mutation (n, %)</th>
<th>Mutation genotype (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LMV + ADV</td>
<td>88</td>
<td>3 (3.4)</td>
<td>85 (96.6)</td>
<td>Rm204V (2), Rl180M (1)</td>
</tr>
<tr>
<td>LdT + ADV</td>
<td>84</td>
<td>2 (2.4)*</td>
<td>82 (97.6)*</td>
<td>Rl181V (1), Rn236T (1)</td>
</tr>
<tr>
<td>ETV + ADV</td>
<td>80</td>
<td>2 (2.5)**</td>
<td>78 (97.5)**</td>
<td>Rl236T (1), Rl180M + RlT184G (1)</td>
</tr>
<tr>
<td>P-value</td>
<td></td>
<td>0.523</td>
<td>0.631</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:** *, vs LMV + ADV group, P > 0.05; **, vs LdT + ADV group, P > 0.05.

**Abbreviations:** LMV, lamivudine; ADV, adefovir dipivoxil; LdT, telbivudine; ETV, entecavir.

### Table 3 Cost-effectiveness and incremental cost-effectiveness analysis of the three groups

<table>
<thead>
<tr>
<th>Group</th>
<th>LMV + ADV</th>
<th>LdT + ADV</th>
<th>ETV + ADV</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cost (US$)</td>
<td>3,024</td>
<td>3,292.8</td>
<td>5,107.2</td>
<td></td>
</tr>
<tr>
<td>Ratio of negative conversion of HBV DNA (%)</td>
<td>90.9</td>
<td>92.9*</td>
<td>93.8**</td>
<td>0.418</td>
</tr>
<tr>
<td>Cost-effectiveness ratio of negative conversion of HBV DNA</td>
<td>33.3</td>
<td>35.4</td>
<td>544</td>
<td></td>
</tr>
<tr>
<td>Incremental cost-effectiveness ratio of negative conversion of HBV DNA</td>
<td>–</td>
<td>134.4</td>
<td>718.3</td>
<td></td>
</tr>
<tr>
<td>Ratio of seroconversion of HBeAg/HBeAb (%)</td>
<td>47.7</td>
<td>48.8*</td>
<td>51.3**</td>
<td>0.852</td>
</tr>
<tr>
<td>Cost-effectiveness ratio of seroconversion of HBeAg/HBeAb</td>
<td>63.4</td>
<td>67.5</td>
<td>99.5</td>
<td></td>
</tr>
<tr>
<td>Incremental cost-effectiveness ratio of seroconversion of HBeAg/HBeAb</td>
<td>–</td>
<td>244.4</td>
<td>578.7</td>
<td></td>
</tr>
<tr>
<td>Ratio of non-genotypic mutation (%)</td>
<td>96.6</td>
<td>97.6</td>
<td>97.5**</td>
<td>0.742</td>
</tr>
<tr>
<td>Cost-effectiveness ratio of non-genotypic mutation</td>
<td>31.3</td>
<td>33.7</td>
<td>52.4</td>
<td></td>
</tr>
<tr>
<td>Incremental cost-effectiveness ratio of non-genotypic mutation</td>
<td>–</td>
<td>268.8</td>
<td>2,314.7</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:** *, vs LMV + ADV group, P > 0.05; **, vs LdT + ADV group, P > 0.05.

**Abbreviations:** HBV, hepatitis B virus; LMV, lamivudine; ADV, adefovir dipivoxil; LdT, telbivudine; ETV, entecavir.
Discussion

Persistent HBV infection is an important risk factor for the development of hepatic cirrhosis or for the occurrence of HCC. Oral nucleos(t)ide analogs have been widely used for more than 10 years, and the sustained suppression of serum HBV DNA to very low or undetectable levels has been associated with the prevention of liver disease progression and inhibition of the development of long-term complications. However, drug resistance has presented a serious challenge to CHB treatment. Insufficient antiviral efficacy caused by drug resistance has resulted in attenuated viral suppression, which can lead to significant clinical deterioration. LMV, LdT, and ETV can all be used as rescue strategies for HBeAg-positive chronic hepatitis B patients with resistance to ADV because all of them have no resistant mutations of ADV in the HBV polymerase and can quickly suppress the replication of HBV. In the present study, in the LMV + ADV, LdT + ADV, and ETV + ADV groups, a sustainably increasing proportion of patients achieved undetectable HBV DNA levels from 4 to 96 weeks, and a sustainably increasing proportion of patients achieved seroconversion of HBeAg/HBeAb from 36 to 96 weeks, which indicates that all the three plans have good prognosis, because the negative conversion of HBV is a predictor for the decreasing impairment of liver function, and hepatitis B e antigen seroconversion is a predictor for lower ratios of cirrhosis and slower disease progression. At week 96 of treatment, the ratio of patients experiencing viral breakthrough was 3.4%, with three resistance mutation (two RtM204V and one RtL180M) in LAM + ADV group, the ratio of viral breakthrough was 2.4% in LdT + ADV group, with two resistance mutation (one RtA181V and one RtN236T), and the ratio of viral breakthrough was 2.5% in ETV + ADV group, with two resistance mutations (one RtN236T and one RtL180M + RtT184G), which showed that the ratios of viral breakthrough of the three plans are rare and can be controlled.

Table 5 Side effects of the three groups

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Ratio of side effects at 36 weeks</th>
<th>Ratio of side effects at 96 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>LMV + ADV</td>
<td>88</td>
<td>1 (1.1%)</td>
<td>4 (4.5%)</td>
</tr>
<tr>
<td>LdT + ADV</td>
<td>84</td>
<td>2 (1.2%)</td>
<td>6 (7.1%)</td>
</tr>
<tr>
<td>ETV + ADV</td>
<td>80</td>
<td>1 (1.3%)</td>
<td>4 (5.0%)</td>
</tr>
</tbody>
</table>

Notes: * vs LMV + ADV group, P<0.05; ** vs LdT + ADV group, P<0.05.
Abbreviations: LMV, lamivudine; ADV, adefovir dipivoxil; LdT, telbivudine; ETV, entecavir.
dipivoxil resistance if the cost can be borne by the patients. However, there are still many problems that need to be studied such as: Can the combination therapy be switched to monotherapy after a fixed time or after the HBV DNA decreases to undetectable level? How about the side effects and how to deal with them after a longer time? How to deal with the viral breakthrough and multidrug resistance after combination therapy?

We will carry out further randomized longer term investigations with larger sample sizes to solve these questions.

**Summary**

LMV + ADV, LdT + ADV, and ETV + ADV are all relatively safe and effective rescue strategies for treatment of chronic hepatitis B with adefovir dipivoxil resistance, with LMV + ADV as the optimal selection and LdT + ADV as the alternative selection in the economically less developed regions, while ETV + ADV can be used in the economically developed regions.

**Acknowledgments**

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**
