Original Article

Biomarkers for early diagnosis of aseptic loosening after total hip replacement

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Abstract: Numbers of patients are suffering from aseptic loosening after total hip replacement, which can need further revision surgery. Particle-induced osteolysis has been considered as a major cause of aseptic loosening after hip joint replacement. This is a prospective study, and there are 82 patients of unilateral osseous arthritis and 10 normal physical examination people in our hospital for this study. The participants were divided into four groups, including control group, No arthroplasty group, joint replacement group and loosened THA group. The osteoclasts cell morphology and osteoclast activity were observed by using the microscopy. The mRNA levels of MCP-1, TNF and RANKL were detected by using the Quantitative reverse transcriptase-polymerase chain reaction (RT-PCR). The results indicated that the bone erosion rates of the joint replacement respectively reached 12.9% and 31.6% after cultured 14 d and 21 d, compare with control, which was remarkably increased (P<0.05). The bone erosion rates of loosened THA also reached 43.4% and 88.4%, respectively, after cultured 14 d and 21 d, compare with control, which remarkably increased (P<0.01). The loosened THA group of expression level of MCP-1 gene were much higher than joint replacement. Expression level of RANKL gene of loosened THA group was higher 7.4 times than joint replacement. In conclusion, the osteoclast morphology and activity in the peripheral blood and expression levels of MCP-1 can be used for early diagnosis of aseptic loosening after total hip replacement.

Keywords: Biomarkers, aseptic loosening, total hip replacement, osteoclast

Introduction
In recent years, there are more and more end stage arthritis patients treated with joint replacement [1]. And total hip replacement had achieved remarkable achievements in worldwide range [1, 2]. Total hip replacement is an effective surgical intervention for end-stage joint diseases such as osteoarthritis and inflammatory rheumatoid. However part of patients treated with total hip replacement, are still suffering from torment by complications which can need further revision surgery, it also is significantly reduce patients of quality of life [3]. Aseptic loosening of the total hip replacement has been considered to resulting from aseptic inflammatory reactions and mechanical stress. Implanted materials such as debris of high density Polyethylene, metals, bone cement, wear debris resulted in inflammatory reactions. However, the precise biological mechanisms have not been clarified elucidated [4-6]. Particle-induced osteolysis has been considered as a major cause of aseptic loosening after hip joint replacement.

Currently, clinic diagnostics of aseptic loosening of the total hip replacement is that a circle bright zone around of implants is found by X-rays, but it appear after a long time to joint pain [7]. Emergence of bright zone may be caused by the osteolysis that is osteoclast absorbed and dissolved bone of directly contact with implants [8, 9]. And, osteoclasts were activated by wear debris such as metal ions; polyethylene particles guide the series of bone dissolution reaction, eventually cause prosthesis loosening [10].
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If this aseptic loosening can be detected before the symptoms appear, then we can take measures to correct the continued exacerbation of aseptic loosening, and avoid subsequent revision surgery [11]. Consequently, there are new methods aimed at improving diagnostic accuracy and speed of detection for aseptic loosening after total hip replacement. For this purpose, we hope find some effective means to detect the signs of aseptic loosening. The biological indicators can help patient discover loose implants, allowing us to take measures to correct the ongoing osteolysis.

As is known to everyone, multinucleated giant cells do not exist in normal peripheral blood, then the blood of mononuclear cells or macrophages of could not become osteoclasts except early proliferative immature monocyte-macrophage (osteoclast precursor cells, OCP) [12]. Osteolysis phenomenon appeared; the OCP of circulating blood volume and the local osteoclasts of numbers significantly increased, and the activity of osteoclasts also were gradually increasing result in resulting in osteolysis [13]. If we could detect osteoclast morphology and activity in the peripheral blood, joint replacement patients may be considered to have started early loosening occurs [14-16]. It may be is important factors of aseptic loosening failure that macrophages and fibroblasts have gathered around the prosthesis organization [17, 18]. Expression of nuclear factor-κB (RANK)/RANK ligand (RANKL) and formation of inflammatory cytokine TNF alpha (TNFα) may promote osteoclast formation and bone resorption [19]. Loosened THA patients had higher levels of chemokines IP-10, MCP-1, and MIG than primary THA patients in the synovial fluid [19, 20]. In order to confirm this hypothesis, the factor of biologic response, such as expression of RANKL, TNFα and MCP-1 were detected. The number and activity of osteolysis in peripheral blood could be as biomarker, it is vital significance for early diagnosis of aseptic loosening.

Materials and methods

Collection of patients’ clinical information

A prospective study, there are 82 patients of unilateral osseous arthritis and 10 normal physical examination people in our hospital for this study. The participants were divided into four groups, including control group (normal physical examination people), No arthroplasty group (similar strength of orthopedic surgery, but no prosthesis implantation of artificial joints), joint replacement group (accepted unilateral osteoarthritis unilateral total hip arthroplasty, selects the type of bone cement prosthesis handle and acetabular cup. All surgery performed by the same group of doctors) and loosened THA group (accepted unilateral osteoarthritis unilateral total hip arthroplasty and diagnosed as loosened).

Cell culture

Extracted approximately 10~20 ml peripheral blood from all participants and placed to anticoagulant tube, mixed with the same amount of α-MEM which contains 10% fetal bovine serum, M-CSF (macrophage colony stimulating factor) and RANKL (NF-kB activator receptor ligands). Then sucked up 9 ml mixed liquid, carefully covered on 5 ml human lymphocyte separation liquid level, low temperature (under 20°C), 3000 r/min centrifuge for 30 minutes. After centrifugation, Dark yellow liquid surface layer containing lymphocytes and monocytes were collected, Added to a 96-well plate (24 holes placed pretreated coverslips, wells containing the remaining dentin slices), each well was added 5 × 10⁵ cells and incubated at 37°C for 90 min, then washed away the non-adherent cells.

The TRAP (tartrate resistant acid phosphatase staining) and observation of cell morphology

The same as above, after cultured 7 d, 14 d and 21 d, respectively take out the dyeing, then sealed piece by neutral gum, then read number of multinucleated cells using phase contrast microscopy, calculated occupancy of osteoclasts which were considered as multinucleated cells of containing more than three nucleus and acid acid phosphatase positive.

Osteoclasts activity

After cells were cultured on dentin slices 7 d, 14 d and 21 d, quantitatively measured dentin slices of area and volume of the absorption under inverted microscope to evaluate osteoclast activity.

Quantitative reverse transcriptase-polymerase chain reaction (RT-PCR)

Total RNA of joint replacement group and loosened THA group of blood was isolated by the
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Table 1. Nucleotide sequences of primers used to determine agents responsible for loosening of the acetabular components of total hip prostheses

<table>
<thead>
<tr>
<th>Targets</th>
<th>Forward primer (5’→3’)</th>
<th>Reverse primer (5’→3’)</th>
</tr>
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<tbody>
<tr>
<td>TNFa</td>
<td>CATCTTTCTCAAATTCGAGTGACA</td>
<td>CACCAATGGAACGATGAGGGT</td>
</tr>
<tr>
<td>MCP-1</td>
<td>CTTGCACTTTAGCGCTAGACA</td>
<td>CTTGACTAGGCGATACGTAC</td>
</tr>
<tr>
<td>RANKL</td>
<td>GACATCCCATCTGGTCCC</td>
<td>AATACCTGTTGTCCTCCC</td>
</tr>
<tr>
<td>GAPDH</td>
<td>TCGACAGTCAGCGCATCTTTTT</td>
<td>TTCCAGCCTCAGTTGCTAACC</td>
</tr>
</tbody>
</table>

Table 2. Clinical characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Control (10)</th>
<th>No arthroplasty (30)</th>
<th>Joint replacement (26)</th>
<th>Loosened THA (26)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Numbers</td>
<td>10</td>
<td>30</td>
<td>26</td>
<td>26</td>
</tr>
<tr>
<td>Age (years)</td>
<td>58.2±10.3a</td>
<td>57.9±8.4</td>
<td>59.3±9.5</td>
<td>59.4±10.5</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>5/5</td>
<td>17/13</td>
<td>16/10</td>
<td>14/12</td>
</tr>
<tr>
<td>BMI (Kg/m²)</td>
<td>27.6±6.1</td>
<td>26.9±6.9</td>
<td>27.1±6.5</td>
<td>27.3±7.5</td>
</tr>
</tbody>
</table>

*a mean ± SD; M male; F female.

Table 3. Osteoclasts rate of participant

<table>
<thead>
<tr>
<th>Groups</th>
<th>Osteoclasts rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7 d</td>
</tr>
<tr>
<td>Normal persons</td>
<td>0</td>
</tr>
<tr>
<td>No arthroplasty</td>
<td>0</td>
</tr>
<tr>
<td>Joint replacement</td>
<td>3.4±0.5</td>
</tr>
<tr>
<td>Loosened THA</td>
<td>23.4±5.3</td>
</tr>
</tbody>
</table>

use of RNAi so Plus Kit (Takara, Japan). Reverse transcription was operated by the manufacturer’s instructions (Takara, Japan). Quantitative RT-PCR was achieved on an ABI StepOnePlus real-time PCR system (Applied Biosystems, USA). Each reaction contains 10 μl 2×SYBR Green Master Mix Reagent (Takara, Japan), 2.0 μl cDNA sample, and 400 nM of gene-specific primers in a final volume of 20 μl. Primers were designed by primer premier 5.0 software and the sequences were listed in Table 1. Amplifications were performed at 95°C for 30 s, followed by 40 cycles of 95°C for 5 s, 60°C for 30 s. GAPDH gene was used as a control to normalize the relative expression of target genes. The change of each gene mRNA level was calculated as 2-ΔCT, where ΔCT = (CT treatment-CT control). The quantitative RT-PCR analysis for each cDNA sample was repeated for three times.

Statistical analysis

In this study, Statistical Package for Social Sciences software (SPSS Inc., Chicago, IL, USA), version 19.0 were used for statistical analysis. The clinical data were all presented as mean ± SD and compared between groups by the student’s t tests. The P<0.05 was considered to indicate a statistically significant difference.

Results

Clinical demographics and characteristics

In the study, there are 82 patients of unilateral osseous arthritis and 10 normal physical examination people. The study included four groups: control group, No arthroplasty group, joint replacement group and loosened THA group. All participants of study are between the ages of 52~65 years; male 52, female 40; between the BMI of 26.1~27.9 Kg/m². The four groups show no significant difference in age, sex ratio and BMI (Table 2).

Abnormalities of monocyte cells morphology and activity in joint replacement and loosened THA

To observe cellular morphology and measure osteoclast activity, collect 10 to 20 ml peripheral bloods of all participants, and isolation and culture monocytes of bloods. These fertile mononuclear macrophages continually were cultured and observed by after 7 d, 14 d and 21 d. randomly select five positions, repeat three times, count and calculate the number and occupancy of multinucleated cells in cultured cells by TRAP tartrate resistant acid phosphatase staining. After lartrate-resistant acid phosphatase staining, positive activity material (multinuclear giant cell) was located in the cytoplasm. After monocyte cells were cultured 7 d, Joint replacement and loosened THA groups, cultured cells initiate fusion. And with the prolonged time of culture (14 d and 21 d), the integration of the cell gradually increased. After cultured 21 d, multinuclear giant cell rate of the joint replacement’s group reached 32.1±9.3%; multinuclear
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Collecting peripheral blood samples from participants of the joint replacement and loosened THA groups, extracted mRNA of the samples, detected mRNA level of relevant loosened genes by quantitative RT-PCR. The gene primers were showed in Table 1. In the Figure 2, the loosened THA group of expression level of MCP-1 gene was much higher than joint replacement. Expression level of RANKL gene of loosened THA group was higher 7.4 times than joint replacement. However, expression level of the TNF gene there were little difference between the joint replacement group and loosened THA group (Figure 2).

Discussion

Total hip arthroplasty has become a successful surgery strategy for end stage arthritis patients, which significantly resume normal life [5, 7, 8]. However, there are still many patients suffering from complications of the failure of the THA which could need further revision surgery [9, 11, 14].

Currently, the precise biological mechanisms of aseptic loosening of the total hip replacement have not been clearly elucidated [21-23]. Aseptic loosening of the total hip replacement has been considered to trigger by multi-factors such as biological, microbiological, biomechanical factors [24, 25]. Mechanical stress and local host responses by the implanted materials is the most important two factors [14, 26-29]. And the implanted materials include high density polyethylene, bone cement, and metals etc. [15, 30-32]. The stimulated inflammatory response by the released wear debris of implants can induce the osteoclast accumulating, excessive resorption, bone loss, and even-

Expression level of MCP-1 gene in the peripheral blood of joint replacement and loosened THA

Giant cell rate of loosened THA's group reached 92.8±20.6% (Table 3).

At the same time, others of cells were cultured by dentin slices. To evaluate osteoclast activity, quantitatively measured dentin slices of area and volume of the absorption under inverted microscope. Same as multinuclear giant cell rate, in the normal persons and No arthroplasty’s groups, bone erosion rates were low quietly after cultured 7 d, 14 d and 21 d. Bone erosion rates of the joint replacement respectively reached 12.9% and 31.6% after cultured 14 d and 21 d, compare with control, it remarkably increased \( P<0.05 \); bone erosion rates of loosened THA reached, respectively 43.4% and 88.4% after cultured 14 d and 21 d, compare with control, it remarkably increased \( P<0.01 \) (Figure 1).
tually periprosthetic osteolysis [33, 34]. Therefore, the new biomarkers which is applied for early diagnosis of aseptic loosening of the total hip replacement, can accurately, efficiently, sensitively and rapidly reduce the loose of occurrence and prevent further development to avoid the subsequent revision surgery [35, 36].

In the normal peripheral blood, Giant cells are not present the blood of mononuclear or tissue of macrophages cell cannot be transformed into osteoclasts, only early immature hyperplastic mononuclear macrophage can become the precursor of osteoclast (osteoclast precursor cells, OCP) [25, 27, 37-39]. When bone dissolve is initiated, the number of OCP in blood and local osteoclasts activity significantly increase, eventually leading to bone dissolve and loose [40]. So the hypothesis was proposed, if the number and activity of osteoclast increase, the joint replacement patients may appear early stage of aseptic loosening. In this paper, culture and observe osteoclasts cellular morphology, meanwhile, detect the osteoclast activity to find osteolysis.

Under the circumstances, we can timely take action to intervene further development of aseptic loosening. After cells come from participant of blood were cultured 7 d, 14 d and 21 d, the normal persons and No arthroplasty’s groups, multinucleated cells were absent. When monocyte cells (osteoclast precursors) were cultured on 7 d, Joint replacement and loosened THA groups, cultured cells initiate fusion. And with the prolonged time of culture 14 d and 21 d), the integration of the cell gradually increased. After cultured 21 d, osteoclast rate of the joint replacement group reached 32.1±9.3%; osteoclast rate of loosened THA’s group had increased to 92.8±20.6% (Table 2).

Further to verify the correlation between osteoclasts’ number and activity, then detect the activity of osteoclast (bone erosion rates). Same as osteoclast rate, in the normal persons and No arthroplasty’s groups, bone erosion rates were low quietly after cultured 7 d, 14 d and 21 d. However, bone erosion rates of the joint replacement compare with control, respectively reached 12.9% and 31.6% after cultured 14 d and 21 d, it remarkably increased, P<0.05; bone erosion rates of loosened THA reached respectively 43.4% and 88.4% after cultured 14 d and 21 d, compare with control, it remarkably increased, P<0.01 (Figure 1). Our results suggest that osteoclasts’ number and activity in peripheral blood could remind that the patients of accepted prosthesis have been as aseptic loosening after total hip replacement in the early days.

Loosened THA patients had significantly higher RANKL expression on only osteoblastic stromal cells, than primary THA patients [41]. Osteoblastic stromal cells play an important role in the periprosthetic osteolysis induced by wear debris. Numbers of studies showed that positive correlation between the expression levels of IP-10, MCP-1, or MIG in the synovial fluid and RANKL levels of synovial fluid or osteoblastic stromal cells in the periprosthetic bone marrow [42]. In our study, the loosened THA group of expression level of MCP-1 gene was much higher than joint replacement. Expression level of RANKL gene of loosened THA group was higher 7.4 times than joint replacement. The release and expression of cytokines and chemokines from synovial fibroblasts such as IL-6, IL-8, MCP-1 and MCP-2 could be induced micro-particles [43]. This study showed that loosened THA patients had higher levels of MCP-1 in the synovial fluid than primary THA patients. This finding suggests that the effects of MCP-1 periprosthetic osteolysis induced by wear debris in loosened THA are mediated through RANKL expression of synovial fluid. The multinucleated giant cells of blood and MCP-1 in the synovial fluid all may make as biological marker for early failure of aseptic loosening THA in the future. Certainly, we also need to further verify the reliability of this biological marker to guide the early clinical diagnose for aseptic loosening THA.

In conclusion, the present study was considered to evaluate possible function of biological marker (the multinucleated giant cells of blood and MCP-1 in the synovial fluid) for risk of early failure of aseptic loosening THA. The results showed that Emergence and increasing activity of the multinucleated giant cells of blood and the expression levels of MCP-1 in the synovial fluid could prompt osteolysis occur, also further diagnose risk of early failure of aseptic loosening THA.

Disclosure of conflict of interest

None.
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