Evaluation of Erythrocytes Magnetized Technology for Measurement of ABO Isoagglutinin Titers

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Abstract

Background  A variation in the measurement of ABO antibody titer has been seen among different laboratories due to lack of standardization. In our study, we aim to evaluate automated ABO isoagglutinin titer measurements by erythrocytes magnetized technology (EMT) and compare with conventional tube technique (TT).

Methods  We performed ABO isoagglutinin titration on samples received in a reference laboratory during a period of 2 months. A total of 134 tests for immunoglobulin G (IgG) titer and 116 for immunoglobulin M (IgM) for anti-A or anti-B were included in the study. Samples were processed for ABO isoagglutination titers by both TT and EMT by QWALYS-3 (Diagast, France). Microsoft Excel was used to compile data, for all calculations, and to draw graphs and plots. The number and percentage of cases within ±1, ±2, or ±3 titer difference (TT-EMT) were calculated.

Results  Median titers and their ranges obtained by EMT were higher or equal to those by TT for all IgM and IgG ABO-antibodies in all blood group (BGs), except anti-A IgM in (BG) O that was lower by EMT (32 [4:128]) than TT (48 [8:256]). One hundred one cases (107/116, 92.2%) for IgM titer were within one titer difference by both the methods.

Conclusion  Results of titration by EMT-based automated instrument QWALYS-3 and conventional TT may vary by one titer dilution in the majority of cases. Use of consistent method for patient management is, therefore, advised.

Keywords  ► isoagglutinin titer  ► erythrocytes magnetized technology  ► conventional tube technique  ► QWALYS-3

Introduction

ABO antibodies are naturally occurring antibodies. Individuals normally produce antibodies directed against the A and/or B antigen(s) absent from their red blood cells (RBC). The ABO antibodies are predominantly immunoglobulin M (IgM), activate complement, and react at room temperature or colder. However, there may be small quantities of immunoglobulin G (IgG) type of ABO antibodies present. The predominant immunoglobulin class of antibodies in group O serum is IgG.1 These antibodies are of clinical significance due to their ability to cause hemolytic transfusion reactions, hemolytic disease of the newborn, acute rejection in solid organ transplantation, and delay in engraftment of erythrocytes and megakaryocytes in ABO-incompatible stem cell transplantation.2,5 Therefore, ABO isoagglutinin titer

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measurements are useful for monitoring engraftment and predicting prognosis in ABO incompatible stem cell transplantation, identifying subgroups of ABO blood type, and monitoring transfusion reactions in ABO-mismatched transfusions.5,7 Even for management purpose, an isoagglutinin titer is especially useful for the administration of treatment like plasmapheresis, splenectomy, and immunosuppressives like rituximab. It has been shown that the ABO barrier in renal transplantation can be circumvented by pretreatment of the recipient with a combination of immunosuppression and apheresis to reduce the ABO antibody concentrations permitting successful engraftment of an ABO-incompatible kidney transplant.4,9

Performing an antibody titration can help determine antibody concentration levels. Serial twofold dilutions of serum that contains the required antibody are prepared and tested against an RBC suspension having the target antigen. The titer is considered to be the reciprocal of the greatest dilution in which agglutination is observed. The strength of reactivity is also noted by assigning it a score. When new serum samples from the same patients are tested for titer, the initial titer specimen is always tested in parallel to nullify the effect of subjective variation among technologists and the relative strength of the target antigen on the red cells being used. A change in titer level of two or more tubes (fourfold increase) is considered to be significant.

There are many methods for measuring ABO isoagglutinin titers such as conventional tube hemagglutination technique (TT), microcolumn agglutination technique (CT), microtiter plate method using solid phase red cell adherence (SPRCA) assay, and flow cytometric analysis. Hemagglutination at room temperature is used for measuring the IgM titer. Indirect antiglobulin test (IAT) is used to measure IgG antibodies. Dithiothreitol (DTT) is used to specifically differentiate IgG antibodies.10,11

TT for ABO antibody titers is limited by the subjective variation of the agglutination reading. It is time consuming and labor-intensive, prone to technical errors, requires experienced staff to read results, and the reaction can be read only once without disturbing the cell button. Furthermore, TT demonstrates low reproducibility and variable sensitivity and has been proven to be difficult to standardize between different laboratories. The CT is considered to be more effective in standardizing the titer readings as it eliminates the interreader variability and also produces more stable and reproducible result. Its other advantages are that it is less time-consuming and uses lower volumes of serum and RBCs, and can also be used as part of an automated system. It is being used despite the need for expensive reagents.

Another technique is SPRCA assays (fully automated systems available by IMMUCOR, United States). In this one of the components of the antigen antibody reaction is immobilized on a solid medium usually the microtiter plates which reacts with free antigen or antibody in the sample. The end point of the reaction is denoted by use of RBCs that may be used as indicator cells or may be a part of the antigen–antibody reaction. Erythrocytes magnetized technology (EMT; Diagast, France) is another technology using microtiter plates and magnetized erythrocytes. These RBC are reacted with patient’s serum on a microplate with magnets underneath and the results are read after magnetization and magnetized shaking. Other advanced and sensitive techniques like flow cytometry (FCM) have been claimed to detect both IgG and IgM ABO antibodies more accurately by the use of iso-type-specific Ab.

A variation in the measurement of the ABO antibody titer has been seen among different laboratories due to the absence of a standard method of titration.12 Because of the many limitations for the TT, automated instruments using microcolumn agglutination, SPRCA assay, and EMT are being commonly used these days.13 The advanced methods have improved the quality of testing but none of them has been found to be better than the other. Using automated techniques have improved the quality of results by reducing manual or clerical errors, decreasing subjectivity, reproducibility in testing, increasing throughput in laboratories with high workload, and keeping records of patient results in forms of images that can be retrieved later.

Not many studies have been done for the evaluation of isoagglutinin titers by EMT. Therefore, in this study, we evaluated automated ABO isoagglutinin titer measurements by EMT by QWALYS-3 (Diagast, Loos Cedex, France). It is used for ABO blood typing, antibody screening, identification, and cross matching.14 Here, we have compared the EMT-based ABO isoagglutinin titer measurements with the conventional TT in Indian scenario.

Materials and Methods

Study Population

We performed this cross-sectional study of ABO isoagglutinin titration on samples received in the Department of Hematology and Immunology at a standalone private national reference laboratory in India during a period of 2 months. A total of 134 tests for IgG titer and 116 for IgM titer for anti-A or anti-B were included in the study. The samples used were residual samples from blood grouping test done at our laboratory for whom both ethylenediaminetetraacetic acid (EDTA) samples and serum samples were available. The serum samples were processed for ABO isoagglutination titers by both manual conventional TT method and EMT by QWALYS-3 (Diagast, Loos Cedex, France). EDTA samples were used for blood grouping by QWALYS-3.

Methods for Isoagglutination Titer

Titration Using Conventional Test Tube Technique

Titers were obtained using the conventional TT as described previously.15,16 Twelve test tubes were taken and labeled (1:1, 1:2, 1:4, 1:8, 1:16, and so on). One hundred microliter normal saline was added to each of 12 test tubes except the first. One hundred microliter patient serum was added to the first two test tubes. After mixing the contents of the second test tube sufficiently with a clean pipette, 100 μL of this mixture was dispensed into the next tube. Same process of mixing and sequential dilution was repeated using a clean pipette to mix and transfer each dilution. One hundred microliter of
mixture in the last tube was saved in case further dilutions were required. The red cells used were pools of known A and B blood group (BG) red cells, diluted to 5% suspension, for the corresponding antibodies. For determining the IgM titer, 50 μL of pooled red cell was added to each of the tubes containing 100 μL of serially diluted serum, and tubes were centrifuged at 1000 × g for 1 minute. For determining the IgG titer, first neutralization of the IgM present in recipient’s serum was done by heat inactivation at 63°C for 10 minutes. A dilution series was prepared using the previously neutralized serum (1:1, 1:2, 1:4, 1:8, 1:16, and so on). To each of the tube containing 100 μL of serially diluted serum, 50 μL of pooled red cell was added and incubated at 37°C for 45 minutes followed by centrifugation and observation of hemolysis and agglutination. The cells were washed three times with saline and final wash was completely decanted. Anti-human globulin (AHG; Diagast) was added to the dry cell button according to the manufacturer’s directions followed by centrifugation and observation for agglutination. The maximum dilution at which macroscopic agglutination (1+) was observed was noted and the reciprocal of the value was designated as the titer (e.g., 32, 64). If agglutination was seen in the tube containing the most dilute serum, additional serial doubling dilutions were prepared and tested. The prozone phenomenon may cause reactions to be weaker in the more concentrated serum preparations than in higher dilutions. To avoid misinterpretation of results, the tube containing the most dilute serum was examined first and then proceeded through the more concentrated samples to the undiluted specimen.

**Titration Using Erythrocyte Magnetized Technology**

All steps in the procedure of antibody titration were performed by following the recommendations of the manufacturer as provided in the technical inserts. All the steps in the procedure of antibody titration were performed following the recommendations of the manufacturer. The detection of ABO IgG isotype antibody is based on the direct agglutination of RBCs that are already magnetized, combined with a magnetic field (E.M. Technology).

Serial twofold dilution of patient’s serum is done in the D-plates (empty microplates for dilution) using DiluentLyx as diluent. DiluentLyx (350 μL) was added into 11 wells of D-plates. Next, 700 μL of patient serum was dispensed in the first well and mixed sufficiently and 350 μL of mixture was dispensed in the next well and diluted stepwise automatically. Furthermore, 25 μL of mixture was dispensed in a microplate, and 25 μL of 1% type A1 blood cell suspension or type B blood suspension (Mag-plate for Heamlys A1, Mag-plate for Heamlys B: Diagast, Eurasante Parc, France) was added and left for incubation at room temperature for 10 minutes (EMT IgM). This was followed by magnetization for 4 minutes and agitation for 2 minutes at 900 rpm then for 45 seconds at 450 rpm. The maximum dilution factor at which 1+ agglutination (read with camera) was read and the reciprocal of the value was designated as the agglutinin titer.

**Statistical Analysis**

Microsoft Excel (Microsoft Corporation, Redmond, Washington, United States) was used to compile data, for all calculations and to draw graphs and plots. The median, maximum, and minimum values of the agglutinin titers of IgG and IgM for each blood type obtained by both the methods were calculated. The number and percentage of cases within ±1, ±2, or ±3 titer difference (TT-EMT) were calculated. The agreement between the two methods was expressed as the percentage of cases showing within one titer difference.

**Results**

The distribution of subjects into various BGs, their mean age, and male to female ratio is given in Table 1.

<table>
<thead>
<tr>
<th>BG</th>
<th>Number (%)</th>
<th>Age (Mean ± SD)</th>
<th>M:F</th>
<th>Number (%)</th>
<th>Age (Mean ± SD)</th>
<th>M:F</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>34 (25.4)</td>
<td>30.23 ± 10.87</td>
<td>11/23</td>
<td>36 (31)</td>
<td>32.05 ± 13.72</td>
<td>13/23</td>
</tr>
<tr>
<td>B</td>
<td>38 (28.4)</td>
<td>26.26 ± 6.71</td>
<td>11/27</td>
<td>44 (37.9)</td>
<td>27.84 ± 15.05</td>
<td>17/27</td>
</tr>
<tr>
<td>O</td>
<td>62 (46.3)</td>
<td>29.72 ± 12.33</td>
<td>20/42</td>
<td>36 (31.1)</td>
<td>28.02 ± 10.91</td>
<td>13/23</td>
</tr>
<tr>
<td>Total</td>
<td>134 (100)</td>
<td>28.87 ± 10.69</td>
<td>42/92</td>
<td>116 (100)</td>
<td>29.20 ± 13.49</td>
<td>43/93</td>
</tr>
</tbody>
</table>

Abbreviations: BG, blood group; IgG, immunoglobulin G; IgM, immunoglobulin M.
and anti-A for BG B, were same by both methods: 64 (4–512) by TT and 64 (2–2,048) by EMT for anti-A in BG B; 64 (2–256) by TT and 64 (2–512) by EMT for anti-B in BG A. IgG titer was higher for both anti-A and anti-B by EMT 256 (64–2,048), 256 (32–2,048), respectively, than TT method 128 (64–1,024), 128 (16–2,048), respectively.

For IgM, the median titer for anti-B in BG A and BG O was higher by EMT than TT: 32 (0–512) by EMT and 16 (1:256) by TT in BG A; 64 (8–512) by EMT and 48 (8–256) by TT in BG O. It was equal for anti-A in BG B by both methods 16 (1:2,048) by EMT and 16 (0:512) by TT method. For anti-A IgM titer in BG O was lower by EMT than TT method 32 (4:128) by EMT and 48 (8:256) by TT.

The agreements between the two methods are shown in Table 2. One twenty one (121/134, 90.3%) cases of IgG titer showed an agreement by both methods (within ± one titer difference). All the cases for IgG titers gave results within two titer differences by both methods. One hundred seven cases (107/116, 92.2%) and one hundred fourteen cases (114/116, 98.2%) for IgM titer were within one and two titer differences, respectively, by both the methods. Fifty cases (50/134, 37.3%) in case of IgG and forty-one (41/116, 35.3%) in case of IgM titer were giving same results by both TT and EMT. For IgG, 53.7% cases gave higher titers, while only 9% gave lower titers by EMT as compared with TT. For IgM, 36.2% cases gave higher and 28.5% cases lower titers by EMT as compared with that by TT.

Discussion
The main findings of this study showed that the median titers and their range obtained by EMT were higher or equal to that obtained by the TT for all IgM and IgG ABO-antibodies in all BGs except anti-A IgM in BG O that was lower by EMT than TT method: 32 (4:128) by EMT and 48 (8:256) by TT. Around one-third of the cases showed similar results for IgG and IgM titer of ABO antibodies in all BGs. However, for IgG
titer ABO antibodies by EMT around 50% samples gave higher titers than TT. In case of IgM ABO antibody samples, around one-third samples gave higher and one-third gave lower titer when EMT was compared with TT.

The significant interlaboratory variations in TT and availability of methods with higher sensitivity for ABO Ab titer measurement have led to increased use of techniques like EMT, CT, and FCM recently. Many clinicians, however, are unaware of these differences that arise due to use of different methods and take into consideration only the target titer obtained. There are many possible factors that may cause these differences to be produced, most importantly the interference of IgG and IgM measurement with each other. The anti-A and anti-B of BG B and A, respectively, are mostly IgM and small amounts of IgG. These antibodies are mostly IgG in persons with BG O. In a study comparing isoagglutinin titers by TT, EMT and microcolumn agglutination showed that EMT-based automated instrument showed higher titration level than the conventional tube hemagglutination technique. The median titer obtained by EMT in their study for IgG anti-B in BG A, anti-A in BG B, anti-A and anti-B in BG O were 16 (2–64), 16 (4–256), 2,048 (8–8,192), and 256 (16–8,192), respectively. The median titers obtained by EMT for IgM anti-B in BG A, anti-A in BG B, anti-A, and anti-B in BG O were 16 (2–64), 16 (4–256), 32 (2–256), and 16 (2–128), respectively. The titer for all antibodies were highest as measured by EMT except median titers of IgG, anti-A in BG B and anti-B in BG A, that were highest by microcolumn agglutination method. The agreements between two methods in their study were taken as the percentage of cases within one titer difference. The percentage of agreement for the conventional tube hemagglutination technique and EMT for IgM was 80% for anti-B of group A, 70% for anti-A of group B, 88% for anti-B of group O, and 60% for anti-A of group O, respectively. For IgG, the percentage of cases in agreement with the conventional tube hemagglutination technique and EMT for IgG was 83% for anti-B of BG A, 35% for anti-A of BG B, 0% for anti-B of BG O, and 3% for anti-A of BG O, respectively. Even on comparison of card method with TT and EMT, agreement results were comparable in case of IgM but low in IgG. Taking all BGs together, our study gave more than 90% results of IgG as well as IgM within one titer difference.

Another study comparing five methods for anti-ABO titration on 50 BG O healthy donors, the median titers of IgG anti-A and anti-B antibodies by SPRCA were 64 (8–2,048) and 64 (4–512), respectively. These values were lower than ours in BG O individuals.

The testing for isoagglutinin titers depends on the temperature differences. IgM can be measured at room temperature and IgG after an incubation at 37°C that causes complement activation. But both IgM and IgG may react at room temperature and activate complements at 37°C leading to interferences in assay methods. When plasma is not treated with DTT or heated at 63°C for 10 minutes to inactivate IgM molecules, indirect agglutination titers may be a mix of IgM and IgG antibodies, reacting at 37°C. However, both IgG and IgM antibodies are likely to be active in humoral rejection, and cold-reacting (30°C) antibodies are usually not of clinical significance. In a study by Nayak et al, reduction in titers by DTT treatment in nearly 50% samples tested for both anti-A and anti-B titers as compared with the gel card titers read without the use of DTT indicating that there is a good amount of IgM type of anti-A and anti-B antibodies in the serum samples, which should be avoided before reporting the titers of candidates of ABO-incompatible transplants. For the conventional TT tests in our study, we have not used DTT. We used heat inactivation as a method as well as monospecific anti-IgG antibodies to remove the nonspecific IgM antibodies. Other studies have also used heat inactivation as a reliable method for the estimation of IgG isoagglutinins.

Few studies have also compared the performances of other methods for anti-A and anti-B titers. There are some studies comparing TT with gel method, one of which stated that gel is more sensitive, column agglutination results being approximately two and half fold higher (one more dilution) than that of conventional TT method.

Another study comparing the TT, gel, card, and FCM for isoagglutinin titers showed that there were significant differences in the titers obtained by these methods and each method showed a different detection capacity for each ABO antibody depending on the BG tested. They showed that in BG O, the mean titer in gel IAT was significantly higher than that of tube IAT for anti-A. FCM with anti-IgM showed the highest titer compared with tube or gel method in all of the BGs.

<table>
<thead>
<tr>
<th>Difference in number of titer steps between TT and EMT</th>
<th>Frequency (IgG titer)</th>
<th>Percentage (IgG titer)</th>
<th>Frequency (IgM titer)</th>
<th>Percentage (IgM titer)</th>
</tr>
</thead>
<tbody>
<tr>
<td>−3.00</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0.9</td>
</tr>
<tr>
<td>−2.00</td>
<td>13</td>
<td>9.7</td>
<td>4</td>
<td>3.4</td>
</tr>
<tr>
<td>−1.00</td>
<td>59</td>
<td>44.0</td>
<td>37</td>
<td>31.9</td>
</tr>
<tr>
<td>0.00</td>
<td>50</td>
<td>37.3</td>
<td>41</td>
<td>35.3</td>
</tr>
<tr>
<td>1.00</td>
<td>12</td>
<td>9.0</td>
<td>29</td>
<td>25.0</td>
</tr>
<tr>
<td>2.00</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>2.6</td>
</tr>
<tr>
<td>3.00</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0.9</td>
</tr>
<tr>
<td>Total</td>
<td>134</td>
<td>100.0</td>
<td>116</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Abbreviations: EMT, erythrocytes magnetized technology; IgG, immunoglobulin G; IgM, immunoglobulin M; TT, tube technique.
Another study comparing five different methods for anti ABO titration concluded that due to poor agreements between the results obtained by different methods, application of consistent and uniform method for titration throughout treatment of patient was recommended.18

Since most of the methods other than TT are or can be automated, they reduce the risk of manual errors, using standardized protocols with higher precision and accuracy, and minimize manual work providing an increased throughput. Another advantage of using EMT above TT for titration is that pictorial result files are converted to quantitative results that can be compared with follow-up pictures of the patient results, thus helping to decide the course of management for the patient.

It has been established that the titers of IgG are clinically more significant.22 The titer of isoagglutinins helps in managing patients as well as determining their prognosis. The tolerable amount of isoagglutinin titer varies among hospitals or clinicians. Risk of antibody-mediated hyperacute rejection, successful transplant by reducing anti-A and anti-B levels by giving immunosuppressant, or procedures like immunoadsorption/plasma exchange can only be determined if an adequate and consistent method for measurement is used. Also, an increase in titer levels of more than two is considered to be significant while patient is on follow-up. Therefore, the standardization of test methods and target isoagglutinin titer can contribute to increasing success rates in cases of transplantation.

In conclusion, the EMT-based automated instrument, QWALYS-3, when compared with conventional test tube method does not give similar results and may vary by one titer dilution in majority (around 90%) of the cases. Therefore, the use of consistent method along with clinical correlation of the isoagglutinin titer is a must for patient management.

Authors’ Contribution
Parul Chopra was involved in definition of intellectual content, literature search, interpretation of results, data analysis, and drafting and editing of the manuscript. Sunanda Bhardwaj interpreted results and edited the manuscript. Ajay Samkaria carried out the tests in the laboratory, analyzed the data, and edited the manuscript. Asha Amoli carried out the tests in the laboratory, interpreted the results, did data entry, and edited the manuscript. Anil Arora conceptualized the study, defined intellectual content interpretation of results, and edited the manuscript.

Ethical Approval
Since the tests were done on residual samples obtained in the laboratory for testing, ethical clearance was not required.

Conflicts of Interest
None.

Acknowledgment
We would like to thank our laboratory staff for their cooperation in conducting the study.

References
19 Schweizer RT, Bartus SA, Perkins HA, Belzer FO. Renal allograft failure and cold red blood cell autoagglutinins. Transplantation 1982;33(1):77–79