Blood group antigen studies using CdTe quantum dots and flow cytometry

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Abstract: New methods of analysis involving semiconductor nanocrystals (quantum dots [QDs]) as fluorescent probes have been highlighted in life science. QDs present some advantages when compared to organic dyes, such as size-tunable emission spectra, broad absorption bands, and principally exceptional resistance to photobleaching. Methods applying QDs can be simple, not laborious, and can present high sensibility, allowing biomolecule identification and quantification with high specificity. In this context, the aim of this work was to apply dual-color CdTe QDs to quantify red blood cell (RBC) antigen expression on cell surface by flow cytometric analysis. QDs were conjugated to anti-A or anti-B monoclonal antibodies, as well as to the anti-H (Ulex europaeus I) lectin, to investigate RBCs of A, B, A₁B, O, A₁, and A₂w donors. Bioconjugates were capable of distinguishing the different expressions of RBC antigens, both by labeling efficiency and by flow cytometry histogram profile. Furthermore, results showed that RBCs from A₂w donors present fewer amounts of A antigens and higher amounts of H, when compared to A, B, A₁, and A₂w. In the A group, the amount of A antigens decreased as A₁ > A > A₂ = A₂w, while H antigens were A = A₁ > A₂ = A₂w. Bioconjugates presented stability and remained active for at least 6 months. In conclusion, this methodology with high sensibility and specificity can be applied to study a variety of RBC antigens, and, as a quantitative tool, can help in achieving a better comprehension of the antigen expression patterns on RBC membranes.

Keywords: erythrocytes, nanoparticles, ABO antigens, carbohydrates, covalent binding

Introduction
In recent years, several studies in the immunohematology field have been performed, mainly to investigate blood group genes.¹ These immunohematological molecular analyses have been contributing to the study of genes related to antigens and also to the understanding of a great number of polymorphisms;²-⁵ nevertheless, they do not provide information regarding quantitative analyses, nor distribution, of the antigens on red blood cell (RBC) membrane surfaces.⁴ Analyses of antigen expressions are to this day performed by using methods based on hemagglutination, which can be performed in tube, microplate, and gel, with different potentiating reaction solutions.² These nonquantitative methods with some improvements, though not new, have advantages such as low cost, simplicity, and enough sensibility and specificity to be used in the transfusional routine. Recently, cytometric assays have been used to quantify antigens and to investigate more complex cases.⁶ Thus, we herein propose to apply dual-color fluorescent CdTe quantum dots (QDs) as a complementary and alternative method for immunohematological investigation by flow cytometry, using the ABO blood group as a model. The application of the QDs can contribute not only to the quantification, but also to the understanding of expression patterns of antigens on the RBC surface.
Fluorescence-based assays present high sensitivity, which can provide the identification and quantification of biomolecules with high specificity. Moreover, new fluorescent probes, such as QDs, allow researchers to take advantage of the full potential of fluorescence. QDs present exceptional resistance to photobleaching and a high reactive surface, which enables conjugations with a variety of biomolecules. We show here that QDs are capable of recognizing and evaluating A, B, and H RBC antigens quantitatively; in addition, QDs can be applied in a direct fluoroimmunoassay in living cells, with no fixation agents, features related to immunohematology.

To evaluate A, B, and H antigen expression, we applied CdTe QDs covalently bound to the antibodies anti-A and anti-B, as well as to the lectin Ulex europaeus I (UEA I), also called anti-H. UEA I lectin can recognize the H antigen by L-fucose detection. CdTe QDs exhibit not only a narrow emission and a size-tunable fluorescence in a broad wavelength range (from green to infrared), but also a higher specificity and reproducibility for biological applications after bioconjugation assays, when compared to CdS/Cd(OH)₂ QDs already used by some of us in previous work.

The passivation of CdS/Cd(OH)₂ QDs is labile and can be removed by bioconjugation procedures decreasing emission quality due to the exposure of surface defects. Moreover, to bioconjugate CdS/Cd(OH)₂ QDs, stabilized with polyphosphate ions, is also necessary for growing a polymeric shell and promoting covalent couplings with biomolecules; it is also necessary, turning this process more laborious. Effective bioconjugations are still considered a challenge as they have to preserve the characteristics of the complete set: the QDs’ fluorescence and biomolecule biochemical functions. In this work, we also associated electrophoresis, fluorescence microplate assay, fluorescence correlation spectroscopy (FCS), and inhibition assay experiments to assure effective QDs bioconjugations.

There are only a few previous reports that have investigated some blood antigens by flow cytometry using standard organic dyes. However, these studies were usually performed in fixed cells and by indirect fluoroimmunoasays, using secondary antibodies, which can make the experiments more laborious and less specific. Our methodology with CdTe QDs enabled us to quantitatively evaluate antigens on RBC membranes in A₁, A₂, B, A₁B, O, and in some A₁ weak groups both by profile and by labeling efficiency. Furthermore, we have also been able to correlate A and H antigen expression in A RBCs. We believe we show a simple method, not laborious and with high sensitivity, which allows antigen identification and quantification with high specificity and reproducibility. This approach can be used as a complementary tool for improving the comprehension of RBC biology in blood systems.

**Experimental procedures**

**Synthesis and characterization of CdTe QDs**

Aqueous colloidal dispersions of orange and green CdTe QDs were synthesized by adapting a previously established method reported by some of us. Briefly, QDs were prepared by addition of Te²⁺ (obtained from metallic tellurium at 10⁻⁴ mol) (Sigma Aldrich Co., St Louis, MO, USA) in a 0.01 M CdCl₂ or Cd(ClO₄)₂ (Sigma Aldrich Co.) solution with pH >10 in the presence of 3-mercaptopropionic acid (MSP) (Sigma Aldrich Co.) as stabilizing agent. We used a 2:1:2.4 molar ratio of Cd/Te/MSA for orange-emission QDs and 5:1:6.0 for green-emission QDs. The Te²⁺ aqueous solution was prepared by reducing metallic tellurium with NaBH₄ (Sigma Aldrich Co.) in a 1:30 molar ratio of Te/NaBH₄, respectively, at a high pH using NaOH and under nitrogen-saturated atmosphere. The growth of the QDs proceeded, in inert atmosphere, with stirring at 90°C for 2 hours or 8 hours for green or orange emission QDs, respectively. After being synthesized, QDs were characterized by absorption (Evolution 600 UV-Vis; Thermo Fisher Scientific, Waltham, MA, USA) and emission spectroscopy (LS 55 spectrometer; PerkinElmer Inc., Waltham, MA, USA). The fluorescence spectra were obtained at 365 nm excitation. Structural characterizations of QDs obtained from a very similar procedure have been previously described by some of us.

**Blood samples**

RBCs from A₁, A₂, B, A₁B, A₁ weak (A₁, A₂, and A₁), as well as O blood donors were obtained at the Hematology and Transfusion Center of UNICAMP (Campinas, São Paulo, Brazil). RBC samples (A₁, n=18; A₂, n=10; A₁, n=1; A₂, n=2; A₁, n=1; B, n=10; A₁B, n=6; and O, n=10) were collected in a 4 mL ethylenediaminetetraacetic acid (EDTA) anticoagulant tube from healthy donors who had had their blood previously serologically typed. RBC phenotyping was performed using monoclonal anti-A (clone: 9113D10, lot: 71EF01EA; Fresenius Kabi [Barueri, São Paulo, Brazil]) and anti-B antibodies (clone: 9621A8, lot: 08D01A21; Fresenius Kabi), as well as anti-A₁ (reference: 116005, lot: 11691-B3; Lorne Laboratories) and anti-H or UEA I lectin (reference 115002, lot: 11568-A1; Lorne Laboratories). A₁ phenotype was differentiated from A group by using both anti-A antibody and...
anti-A lectin \((Dolichos biflorus)\), as \(A_1\) presents agglutination for the anti-A antibody, though not for anti-\(A_1\) lectin.\(^{20}\)

**QD bioconjugation**

Orange QDs were conjugated to anti-A (clone: 9113D10, lot: 71EF01EA; Fresenius Kabi) or anti-B (clone: 9621A8, lot: 08D01A21; Fresenius Kabi) by using N-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) (Fluka) and N-hydroxysulfosuccinimide sodium salt (Sulfo-NHS) (Sigma Aldrich Co.) as coupling reagents. We first adjusted the pH of 2 mL of CdTe QDs (3 µM) to 5.5 by using MSA at 4.9% (w/v). We then added 1 mL of EDC (at 0.4 mg-mL\(^{-1}\)) and, after 5 minutes, 1 mL of Sulfo-NHS at 1.1 mg-mL\(^{-1}\) to the QDs sample.\(^{7,21}\) Fifteen minutes later, we added 40 µL of anti-A antibody or anti-B antibody to the system (the same antibodies described in the section “Blood samples”).

Green QDs were also covalently bonded to the UEA I lectin, also known as anti-H, obtained from Sigma Aldrich Co. (reference L5505) by using EDC and Sulfo-NHS. To this end, we used 1 mL of green QDs (4.5 µM), 1 mL of EDC (at 4.0 mg-mL\(^{-1}\)), and 1 mL of Sulfo-NHS (at 10 mg-mL\(^{-1}\)). At the end of the process, we inserted 200 µL of UEA I (at 0.5 mg-mL\(^{-1}\)) into the system.

Prior to RBC labeling, systems were incubated with 50 µL of Tris base (at 1 mM) for 2 hours under slow agitation. This procedure was used to quench the free carboxyl groups of nonconjugated QDs to minimize unspecific targets.

**QD bioconjugation characterization**

**FCS analysis**

FCS analysis was performed in a confocal microscopy (LSM 780 equipped with the software ZEN; Carl Zeiss Meditec AG, Jena, Germany) by using 40× water immersion objective (NA =1.0, WD =2.5 mm).\(^{22}\) To obtain the correlation curves, bare orange QDs and their bioconjugates with anti-A (or anti-B) as well as bare green QDs and their bioconjugates with anti-H were, respectively, excited by a green \((\lambda=514\,\text{nm})\) and a blue \((\lambda=488\,\text{nm})\) laser, both with 1 µW. The pinhole aperture, in both cases, was set as 35 µm.

Ten correlation curves were obtained for each sample. From the correlation curves, we obtained the hydrodynamic radius \((R)\) for each system using Equation 1:

\[
R = \frac{4k_BT\tau_D}{6\pi\eta\omega_x^2}
\]

where \(k_B\) is a Boltzmann constant, \(\eta\) is the medium viscosity, and \(T\) is the temperature. We adopted \(T=303\,\text{K}\) as the temperature of the sample under laser irradiation and \(\eta\) as the water viscosity at this temperature (\(\eta=7.98\times10^{-4}\,\text{Pa}\cdot\text{s}\)).

Diffusion time \(\tau_D\) was extracted from the correlation curves and the lateral radius \(\omega_x\) of the focal volume (300 nm for 514 nm excitation and 260 nm for 488 nm excitation), which was obtained by calibrating the system with a solution of rhodamine B (at 10 nM).\(^{23}\) As conjugated QDs show higher diffusion times when compared to bare QDs,\(^{22}\) it is possible to use the R measurements to confirm the bioconjugation. The concentrations of QDs and conjugates used for FCS experiments were of the order of 10–200 nM.

**Electrophoresis**

QD conjugations to anti-A and anti-B were further confirmed by polyacrylamide gel electrophoresis (PAGE) technique. For this, we used a native and discontinuous PAGE with stacking gel of 6% and main gel of 9%. After polymerization, aliquots of each anti-A, QD, and QDs-anti-A sample were mixed with glycerol at ratio 5:1 (v/v) and then loaded into the gel wells. The systems were immersed in electrophoresis running buffer and PAGE proceeded at constant voltage (100 V) for 90 minutes. We evaluated the conjugation by comparing the running profiles of the samples by observing fluorescence signals in an L-Pix EX transilluminator (Locus Biotecnologia), and, afterward, by using the Coomassie blue dye (R 250) (Sigma Aldrich Co.) that binds to proteins. The same procedure was performed for anti-B and their conjugates.\(^{24}\)

**Fluorescence microplate assay**

This method, developed by our group, is based on the presence or absence of fluorescence signals from samples placed in a fluorescence plate reader.\(^{25}\) For this, QDs with coupling agents (EDC and Sulfo-NHS), the proteins (anti-A and anti-B antibodies), as well as the bioconjugates (QDs-anti-A and QDs-anti-B) were placed in a polystyrene microplate (black 96-well Optiplate F HB microplates; PerkinElmer Inc.). All systems were added in triplicate and the plate was incubated for 2 hours in an incubator (water bath, humid chamber) at 37°C. After incubation, the plate was washed three times with phosphate-buffered saline (1×).

Fluorescence measurements were performed using a WALLAC 1420 plate reader with the Victor² (PerkinElmer Inc.) software. The excitation band pass filter used was F405 (405 nm/5 nm) and the emission band pass filter was F595 (595 nm/30 nm). The acquisition time was 1 second. The lamp was set to 20,000 and normal slits were used for excitation of samples and for collecting the emission.

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\(^{1}\)B. \(^{2}\)Dovepress. \(^{3}\)International Journal of Nanomedicine downloaded from https://www.dovepress.com/ by 54.70.40.11 on 31-Dec-2018

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In this method, as the nonconjugated QDs do not have affinity for polystyrene, they are removed after washing. The nonremoval of the proteins presented no interference as these have no significant fluorescence under the conditions used in the experiment; only bioconjugates are capable of showing a significantly detectable signal. For this reason, QDs with coupling agents and bare proteins were used as our controls and the QD–protein conjugates were the systems tested. The intensity of the detected signal is proportional to bioconjugation efficiency: the higher the signal, the more efficient the bioconjugation.

The relative fluorescence intensity (RF) of bioconjugates was calculated by using the following Equation:

\[
RF(\%) = \frac{\text{Bioconjugates FL} - \text{Control FL}}{\text{Control FL}} \times 100\%
\]

where Bioconjugates FL is the average fluorescence intensity of the QD–biomolecule conjugates and Control FL is the average controls signal. According to Carvalho et al\textsuperscript{23} the bioconjugation was considered efficient when the systems showed an RF higher than 100% compared to their controls.

**Inhibition of lectin assay**

In order to prove that labeling results were due to lectin carbohydrate specificity, lectin binding inhibition assays\textsuperscript{28} were accomplished by incubating QDs-anti-H with 0.3 M L-fucose for 1 hour at 25°C prior to their incubation with O RBCs.

**Flow cytometric analysis of RBC samples**

To obtain RBCs, the blood collected in EDTA tubes was centrifuged at 1,700×g for 5 minutes. The pellet was then further washed three times in saline at 450×g for 2 minutes. A sample containing RBCs in saline at 1% (v/v) was then prepared. This procedure was used for all analyses. RBCs were incubated at 37°C for 1 hour for all bioconjugates.

Type A\textsubscript{1} and B RBCs were incubated with QDs-anti-A and QDs-anti-B, respectively, using a proportion of 3:1 v/v (conjugated QDs:cells). Type O RBCs were incubated with both bioconjugates and used as the negative control, as they do not present A or B antigens in their membranes.

Type O, A\textsubscript{1}, and B RBCs were also incubated with QDs-anti-H using a proportion of 1:1 v/v (conjugated QDs:cells). In this case, types A\textsubscript{1} and B were used as controls as they show a very low H antigen expression.

Following ABO experiments, types A\textsubscript{2} and A\textsubscript{weak} were incubated with QDs-anti-A and QDs-anti-H, using the same procedure as aforementioned, to identify the N-acetylgalactosamine and L-fucose carbohydrates present on RBC membranes.

RBCs were analyzed by flow cytometry (FACSCalibur or Accuri C6; Becton Dickinson). The software programs used for data processing were CellQuest™ or BD Accuri C6 Software (Becton Dickinson) and FlowJo vX. Twenty thousand events (gated) were acquired. The fluorescence was excited at λ=488 nm and measured with an FL1 filter (530 nm/15 nm) or FL2 filter (585 nm/21 nm) for green and orange QDs, respectively.

**Results and discussion**

**QD optical characterization**

All QDs employed in the experiments were characterized by absorption and emission spectra, shown in Figure 1. Green (Figure 1A) and orange (Figure 1B) QDs presented a maximum emission peak at 548 nm (full width at half maximum =53 nm) and 610 nm (full width at half maximum =48 nm), respectively. By using the first maxima of the absorption spectra, at 496 nm and 559 nm approximately, the average diameters were estimated at 2.6 nm and 3.1 nm for green and orange QDs, respectively.\textsuperscript{13,27} We estimated the concentration of nanoparticles as 4.5 µM for green QDs and 3 µM for orange QDs using the average sizes, Beer–Lambert law, the absorption values at the first maxima, and the extinction coefficient obtained by Yu et al.\textsuperscript{28} Details of this type of calculation can be found in previous study reported by some of us.\textsuperscript{19}

Figure 1 shows that neither orange QDs-anti-A nor QDs-anti-B presented considerable changes in the optical properties when compared to bare QDs (Figure 1D). The emission profile was similar for both conjugates after the bioconjugation process. Green QDs-anti-H (Figure 1C) presented a red shift of approximately 20 nm and orange QDs with anti-A and anti-B showed a small blue shift of approximately 5 nm. This is probably due to some modifications on the QDs’ surfaces caused by the binding of these biomolecules. In addition, the higher shift present for QDs-anti-H can be justified by the excess of EDC used, more than 20×, when compared to the anti-A or anti-B antibodies’ conjugation.

**Bioconjugation analysis**

A thorough characterization of the bioconjugation is required to avoid artifacts and nonspecific results. For this reason, this section is devoted to the characterization of our bioconjugation procedure.

A previous analysis of the bioconjugates, performed by us in the beginning of this study, demonstrated that the
labeling of A RBCs presented more specificity when 15-day unblocked QDs-anti-A were used, indicating that bioconjugation was more efficient for this period of time. Therefore, we decided to use all conjugates at least after 15 days of preparation and also blocked by Tris.

**QDs bioconjugated to monoclonal antibodies**

In fluorescence microplate assay, control A: bare QDs (with coupling agents) and anti-A antibodies, showed an average signal of 296 arbitrary units and control B: bare QDs (with coupling agents) and anti-B antibodies presented an average signal of 311 arbitrary units. As QDs are removed after washing and antibodies do not present considerable autofluorescence at excitation and emission wavelengths used for fluorescence microplate assays ($\lambda_{\text{exc}}=405$ nm and $\lambda_{\text{em}}=595$ nm), we can consider this signal as practically being background detection. On the other hand, QDs-anti-A and QDs-anti-B showed an average signal of 9,261 and 7,836 arbitrary units, respectively. These results correspond to an RF of 3,028% for QDs-anti-A and 2,420% for QDs-anti-B, indicating efficient bioconjugation as shown in Table 1.

After the analysis by fluorescence microplate assay, we further confirmed bioconjugation by electrophoresis experiments (Figure 2). The proteins (Figure 2A) exhibited

**Table 1** Fluorescence microplate assay results of fluorescence intensities and RF percentage of controls (average signal of antibodies and bare QDs) and bioconjugates

<table>
<thead>
<tr>
<th>Systems</th>
<th>Average signal (arbitrary units)</th>
<th>RF (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control A</td>
<td>296</td>
<td>–</td>
</tr>
<tr>
<td>Control B</td>
<td>311</td>
<td>–</td>
</tr>
<tr>
<td>QDs-anti-A</td>
<td>9,261</td>
<td>3,023</td>
</tr>
<tr>
<td>QDs-anti-B</td>
<td>7,836</td>
<td>2,420</td>
</tr>
</tbody>
</table>

**Notes:** Control A: (average signal of bare QDs [with coupling agents] + average signal of anti-A antibody)/2. Control B: (average signal of bare QDs [with coupling agents] + average signal of anti-B antibody)/2.

**Abbreviations:** QDs, quantum dots; RF, relative fluorescence intensity.
differential migration from nonconjugated (bare) QDs (Figure 2B). Furthermore, QDs conjugated with anti-A antibody (Figure 2C) presented a slow migration when compared with nonconjugated QDs. The electrophoresis gel also demonstrated that QDs were successfully conjugated to anti-A antibody. The same profile was observed when orange QDs were conjugated to anti-B antibody (data not shown).

Fluorescence microplate assay and electrophoresis indicate a successful conjugation to QDs-anti-A and QDs-anti-B; however, bioconjugations were further analyzed by FCS measurements. From the correlation curves shown in Figure 3, the average diffusion time for bare orange QDs was calculated as $t = 152.3 \, \mu s$, while for the QDs-anti-A and QDs-anti-B this was $t = 824.6 \, \mu s$ and $t = 747.0 \, \mu s$, respectively. By using these diffusion times and Equation 1, we obtained the QDs’ $R$. Bare orange QDs showed an average hydrodynamic diameter of $D = 3.8 \, \text{nm}$, QDs-anti-A of $D = 20.4 \, \text{nm}$, and QDs-anti-B of approximately $D = 18.5 \, \text{nm}$.

As expected, bare QD sizes, obtained by FCS analysis, were higher than diameters obtained by absorption spectra. This occurs as the $R$ is also related to the charges and profiles of the QD surfaces, while the absorption is an optical property related to the QD core. All bioconjugates presented higher diameters than bare QDs. The results show that anti-A and anti-B antibodies were bound to QDs, a further confirmation of the efficacy of the bioconjugation. Typical original FCS curves and fittings are shown in Figure S1.

**QDs bioconjugated to anti-H lectin**

Bare green QDs showed an average diffusion time of $t = 70.8 \, \mu s$, while the QDs-anti-H diffusion time was $688 \, \mu s$. Bare green QDs had an average hydrodynamic diameter $D = 2.9 \, \text{nm}$ and QDs-anti-H of $D = 28.9 \, \text{nm}$, also confirming conjugation. Typical original FCS curves and fittings are shown in Figure S1.

The carbohydrate inhibition assay showed that QDs-anti-H were not able to label O RBCs, confirming the bioconjugation. The fluorescence microplate assay with QDs-anti-H was not capable of discriminating bare QDs from conjugates, probably as lectin has different interactions...
with polystyrene. Furthermore, in the electrophoresis with QDs-anti-H, the visualization of the bands of the QDs with coupling agents was not possible. However, both FCS analysis and the inhibition carbohydrates assay demonstrated that anti-H was successfully conjugated to QDs.

Flow cytometric analysis of RBC samples

ABO blood group incubated with QDs-anti-A and QDs-anti-B

Table 2 shows the percentage of RBCs labeled with their correspondent conjugates. Our results demonstrated that O RBCs did not present considerable labeling with any of the monoclonal antibodies. In both cases, using QDs-anti-A or QDs-anti-B, labeled O RBCs were less than 5%. These results suggest that conjugated QDs were successfully blocked and the labeling was specific and reproducible.

A, B RBCs presented in their membranes two types of carbohydrates, N-acetylgalactosamine (B antigen) and N-acetylglucosamine (A antigen), thus A, B showed 98.8% of cells labeled with QDs-anti-A and 92% with QDs-anti-B. The cytometry showed that an average of 96.7% (varying from 95% to 98.5%) of A, B RBCs were labeled with QDs-anti-A. For B RBCs incubated with QDs-anti-B, the analysis showed an average of 94.7% (varying from 91% to 98.8%) of cells labeled by these bioconjugates.

Bare QDs did not label cells nonspecifically. All results demonstrated that conjugates labeled RBCs of ABO blood groups specifically and efficiently.

Figure 4 shows typical flow cytometric profiles of A, A weak, B, A, B, and O RBCs when incubated with QDs-anti-A or QDs-anti-B. Type O RBCs incubated with QDs-anti-A or QDs-anti-B showed the same pattern presented by control RBCs (RBCs without QDs). Our results agree with those presented by Aki et al.15 and by Hult and Olsson.16 The percentage of labeled RBCs and the labeling profile were similar to ours in both studies, where the authors analyzed ABO antigens stained by a secondary antibody conjugated to organic dyes using an indirect immunoassay. The overall results demonstrated that a direct immunofluorescence assay using our nanotechnology methodology is capable of differentiating ABO RBC groups. The small deviations presented in Table 2 reflect the small differences of antigen expressions that can be found for the same blood group, due to the intrinsic variability of biological systems associated to the error presented by flow cytometry analysis of approximately 5%.

ABO blood group incubated with QDs-anti-H

Additionally, we investigated the presence or absence of H antigens by L-fucose analysis in ABO blood group. The average percentages of labeled cells are shown in Table 2. O type RBCs had the highest amount of L-fucose on the membrane; approximately 85% (varying from 80% to 95%) of the cells presented H antigens on their surfaces. O RBCs showed a heterogeneous distribution of L-fucose on cell membranes. In A, B, and A, B groups, all or almost all of the L-fucose molecules were converted into A or B antigens; in this way, a small average labeling of <10% (A, 8% [varying from 4% to 10%]; B, 7.9% [4% to 12%]; and A, B, <4%) was observed. Typical histogram profiles are shown in Figure 5. As A, B, and A, B showed similar cytometric results, we presented only the data for the A group (Figure 5B).

A, A weak, A weak, B, and A, weak groups incubated with QDs-anti-A

A, A weak, and A weak, phenotypes (phenotyped as A, A, weak, A, weak, and A, weak) were incubated with QDs-anti-A. Figure 6 shows typical histograms of A, A weak, and A, weak types. The A, subgroup presented an average percentage of labeled cells of 68% (varying from 66% to 70%) and A showed approximately 11% of labeled cells, while A showed 5% and A, presented similar quantities as the A, subgroup.

According to the histogram profile of A (Figure 6), two populations were observed after incubation with QDs-anti-A: 1) a set of unlabeled cells (approximately 35%), probably of RBCs with very little or without A antigens on the cell surface, and 2) a set of labeled cell population (approximately 65%) that presented A antigens on cell surface. The subgroup, serologically defined as A, with QDs-anti-A, presented a

Table 2 Average percentage of ABO blood group red blood cells labeled with conjugated QDs

<table>
<thead>
<tr>
<th>Blood samples</th>
<th>Bioconjugates</th>
<th>QDs-anti-A (%)</th>
<th>QDs-anti-B (%)</th>
<th>QDs-anti-H (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O type</td>
<td>&lt;5</td>
<td>&lt;5</td>
<td>85 (80–90)</td>
<td></td>
</tr>
<tr>
<td>A, type</td>
<td>96.7 (95–98.5)</td>
<td>94.7 (91–98.8)</td>
<td>7.9 (4–12)</td>
<td></td>
</tr>
<tr>
<td>B type</td>
<td>98.8 (96.0–99.7)</td>
<td>92 (80.5–98.2)</td>
<td>&lt;4</td>
<td></td>
</tr>
</tbody>
</table>

Note: ‘We did not incubate A, type with QDs-anti-B and B, type with QDs-anti-A. Data are represented as the mean percentage of labeled cells (range).

Abbreviation: QDs, quantum dots.
more expressive unlabeled cell population (approximately 88%), while a smaller amount of cells showed a discrete labeling (approximately 12%). The A<sub>weak</sub> samples, serologically defined as A<sub>x</sub> and A<sub>el</sub>, showed a similar profile of type O RBCs when incubated with QDs-anti-A. In this case, we could not differentiate them by flow cytometry analysis. However, we were able to differentiate them from the other phenotypes.

These results of labeling and profile obtained for A<sub>weak</sub> and A<sub>x</sub> agree with those presented by Hult and Olsson<sup>16</sup> and with the hemagglutination patterns of serological tests.<sup>5</sup> Our results for A<sub>x</sub> are also consistent with the mixed-field
pattern displayed after agglutination reaction of RBCs with anti-A monoclonal antibody.\(^4,5\)

A\(_2\) and A\(_{weak}\) groups incubated with QDs-anti-H
The flow cytometric patterns of A\(_{weak}\) phenotypes are depicted in Figure 7B. A\(_x\) and A\(_{el}\) presented only 30% of labeling, approximately. A\(_x\) showed a similar profile to A\(_{el}\), with a smaller amount of cells labeled by QDs-anti-H. These results are in accordance with the serological results. However, it is not possible to quantify the amount of H antigens in these cells using serological hemagglutination reactions.\(^5\)

Figure 7A shows the flow cytometric pattern of A\(_2\) phenotype. According to the flow cytometry analysis, A\(_2\) showed approximately 70% (varying from 60% to 80%) of RBCs targeted by the QDs-anti-H conjugates. These results can be explained by the fact that QDs-UEA I recognize the L-fucose of H antigens that have not yet been converted, as well as the L-fucose of the A\(_2\) antigen, which is exposed in the structure of this biomolecule.\(^29\) This observation can be supported by the similar labeling obtained by the QDs-anti-A and QDs-UEA I in A\(_2\). Further studies are needed to better elucidate this observation. Furthermore, in our work, we used the anti-H lectin from Sigma Aldrich Co. that can be more efficient in recognizing L-fucose when compared with the anti-H used in serological routine.

**Conclusion**
The experiments showed that anti-A, anti-B, and anti-H biomolecules conjugated to QDs did not lose their ability to efficiently recognize their respective targets on RBC surfaces. In this work, we used commercial and routinely applied monoclonal antibodies to label living RBCs, with no fixation procedures. Moreover, bioconjugates remained fluorescent, stable, and specifically labeling the cells for at least 6 months. This was confirmed by the results obtained after incubating A\(_1\), B, and O RBCs with their respective
bioconjugates, which showed approximately 100% of cells labeled by flow cytometry. The results are similar to those obtained with bioconjugates of 15 days. This gives rise to the possibility of using these bioconjugates in hematology centers with good versatility.

We have been able to differentiate the ABO groups by using QDs-anti-A, QDs-anti-B, and QDs-anti-H. Likewise, we were also able to differentiate some A phenotypes. The QDs conjugated to anti-H were important tools to correlate A and H antigens in A blood group. Type A2 was less labeled by QDs-anti-A when compared to type A1, while the opposite was shown for QDs-(UEA I) conjugate results. As discussed before, QDs-(UEA I) recognized in A1 the L-fucose of H antigens that were not yet converted, as well as the L-fucose of the A2 antigen, which is exposed in the structure of this biomolecule, thus justifying the labeling observed with these bioconjugates. Our results still suggest that RBCs from types A1 and A2 have fewer amounts of both H and A antigens. In the A group, the amount of A antigens decrease in the order A1 > A2 > A0. On the other hand, H antigens decrease in the order A1 > A2 > A0. Further studies are needed in this field, mainly those that correlate molecular biology with immunophenotyping techniques.

Taken together, these results demonstrate a simple, uncomplicated, specific, quantitative, and reproducible methodology that can be used as complementary and versatile diagnostic tools.

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Disclosure
The authors report no conflicts of interest in this work.

References


Figure S1 Typical fluorescence correlation spectroscopy curves of QDs and their bioconjugates.

Notes: Correlation curve (dashed line) and a fitting of the correlation curve (solid line) of the systems are represented for (A) orange QDs, (B) QDs-anti-A, (C) QDs-anti-B, (D) green QDs, and (E) QDs-anti-H.

Abbreviation: QDs, quantum dots.

Supplementary material

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