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Primate Erythrocyte-Immune Complex-clearing Mechanism

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University, Columbus, Ohio 43210

ABSTRACT Previous in vitro studies have shown that immune complexes (IC) that fix complement can bind to the C3b receptor on primate erythrocytes. The in vivo function of this erythrocyte receptor, however, is unknown. This study was undertaken to determine whether the binding of IC to erythrocytes in vivo might play a role in the removal of IC from the circulation. Baboons and rhesus monkeys were prepared with a catheter in the ascending aorta to infuse IC and in the abdominal aorta, renal, hepatic, and portal veins to monitor changes in binding and clearance of IC across kidney, liver, and spleen + gut, respectively. Autologous 51Cr-labeled erythrocytes were infused intravenously and allowed to equilibrate. Preformed IC (125I-labeled bovine serum albumin [BSA] rabbit anti-BSA) were then infused into the ascending aorta at a constant rate for 120 s. Blood samples were drawn at frequent intervals for 30 min from all catheters below the IC injection site. Each blood sample was then centrifuged on percoll to separate IC bound to erythrocytes from IC in plasma or bound touffy coat cells. This resulted in an "erythrocyte fraction" beneath the percoll that contained the IC bound to erythrocytes, and a "plasma/buffy coat fraction" above the percoll that contained the IC in plasma and IC bound to buffy coat cells. Analysis of these data showed that the majority of the IC infused into the circulation rapidly became bound to erythrocytes. However, by 5 min after beginning the IC infusion, most of this IC load had been removed from the erythrocytes as they traversed the liver. In contrast, IC on erythrocytes did not deposit in kidney. The IC-bearing erythrocytes themselves were not trapped or detained by any organ. IC in the plasma/buffy coat fraction of blood were removed from the circulation but at a relatively low rate and almost entirely by the liver. These findings suggest that primate erythrocytes intercept large complement-fixing IC in the circulation causing the IC to adhere to the erythrocyte until the IC-bearing erythrocyte traverses liver where the IC is deposited, and the erythrocyte is returned to the circulation. This primate erythrocyte-IC-clearing mechanism may be important in the protection against diseases mediated by deposition of circulating IC.

INTRODUCTION

Complement receptor type 1 (CR1),1 which has receptor specificity for the C3b and C4b components of the complement system, is present on the surface of human and other primate erythrocytes (1, 2). The average number of CR1 receptors per erythrocyte is small (mean < 2,000/cell) compared with that of circulating B lymphocytes, monocytes, and neutrophils where the average number of CR1 receptors per cell ranges from 21,000 to 148,000 (1–3). However, because circulating erythrocytes greatly outnumber circulating leukocytes, the vast majority of all CR1 receptors present in the circulation of humans and other primates is located on the erythrocyte (4).

The CR1 receptors on lymphocytes, macrophages, and neutrophils have been shown to play important roles in immune regulation and in phagocytosis (5, 6). By contrast, to date no important in vivo function has been found for the vast number of CR1 receptors in the circulating erythrocyte pool. However, it is well established that, in vitro, immune complexes (IC) that fix complement can bind to primate erythrocytes via the CR1 receptor (6–9). It seems possible that the erythrocyte, by virtue of its ability to bind IC and its ubiq

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1 Abbreviations used in this paper: BSA, bovine serum albumin; CR1, complement receptor type 1; IC, immune complexes.
uity in the vascular space, could play a role in the removal of circulating IC from the blood stream. The present study was undertaken to test that hypothesis. The results indicate that primate erythrocytes participate in a rapid and efficient mechanism for removal of large complement-fixing IC from the circulation.

METHODS

Preparation of IC

Crystallized bovine serum albumin (BSA, Sigma Chemical Co., St. Louis, MO) was labeled with 125I as previously described (10). Labeling efficiency was ~70% and specific activity ranged from 20 to 30 μCi/μg. Anti-BSA antibody was produced by immunizing New Zealand rabbits, as previously described (10). The serum antibody concentration was measured by the quantitative precipitin method. IC were formed by combining at equivalence heat-inactivated rabbit antisera with 125I-labeled BSA (125I-BSA), incubating for 1 h at 37°C and then for 4°C for 48 h. The resulting precipitate was centrifuged (1,200 g for 10 min), washed, and then incubated with 0.01 M phosphate-buffered saline (PBS, pH 7.2) to which five times excess unlabeled BSA was added. The solution was agitated for 4 h at 37°C, centrifuged (100 g for 20 min), and the precipitate discarded. Approximately 5–30% of the precipitate was solubilized by this method, as assessed by loss of 125I counts per minute from the precipitate. All IC preparations were made 24–48 h before use and were stored at 4°C. This IC preparation was chosen for study because it showed much higher levels of complement-dependent binding to human and simian erythrocytes compared with IC preparations subjected to centrifugation at 500 or 1,000 g (8).

To assess the extent to which the 125I-BSA was incorporated into IC, 1 ml of 20% polyethylene glycol 6000 (PEG) was added to 10 μl of IC solution. This concentration of PEG precipitates >90% of the 125I-BSA that is incorporated into IC, but <10% of the 125I-labeled BSA that is not incorporated into IC (12). By this procedure ~90% of 125I-BSA was in the form of IC.

Surgical preparation

Seven primates (three female rhesus monkeys, one male, and three female baboons) were used. The animals weighed 5.4–30.2 kg. Anesthesia was induced by intramuscular administration of Ketamine and atropine. An endotracheal tube was then inserted and anesthesia maintained with inhalation of halothane and nitrous oxide. Electrocardiographic monitoring was maintained throughout the experiment. Intravenous Ringer’s lactate was infused continuously at 1–3 ml/min, depending on body size. Under sterile conditions (four experiments) or nonsterile conditions (three experiments) the common femoral artery and vein of both lower extremities were isolated to facilitate angiographic catheter placement. An upper midline abdominal incision was then made to provide exposure of the portal vein in the porta hepatitis. A No. 5 French polyethylene feeding tube was then placed in the portal vein through a stab incision and stabilized with a 6.0 polypropylene suture. The animal was then transferred to the fluoroscopy room where an angiographic catheter (5 or 6 French with torque control) was placed, under fluoroscopic guidance and with the aid of infusion of contrast media, into each of the following locations: the ascending aorta just above the aortic valve, the abdominal aorta at the level of the renal arteries, the renal vein (right or left), and a right hepatic vein. The aortic arch catheter was used for periodic measurement of arterial blood pressure, for infusion of the IC and, in three experiments for infusion of 85Sr-labeled microspheres (15±0.8 μm, 3 M Company, St Paul, MN) to assess tissue blood flow. The remaining catheters made it possible to monitor changes in blood composition across kidney (arterial vs. renal vein blood), across liver (arterial + portal vein vs. hepatic vein blood), and across the portal circulation (arterial vs. portal vein blood). The portal vein provides the venous drainage of the spleen, stomach, small bowel, large bowel, and pancreas. Hereafter, these organs will be simply referred to as spleen + gut.

Experimental protocol

Animals were infused with autologous 51Cr-labeled erythrocytes as follows: 10–20 ml of blood were removed, the erythrocytes isolated, washed in Hanks’ balanced salt solution, labeled with 20–40 μCi of 51Cr (10), washed, infused, and allowed to circulate for a minimum of 20 min. 10–20 ml of IC (0.3–2.2 mg) were then infused at a steady rate with a hand-held syringe over 120 s. Blood samples (~1 ml) were drawn simultaneously from the catheters in the abdominal aorta, hepatic portal, and renal veins into tubes containing 10 U of heparin. The blood samples were drawn starting at time 0, at 20-s intervals until 120 s had elapsed, and then at 3, 4, 5, 15, and 30 min. Each blood sample was transferred to an iced tube immediately after it was drawn. The tubes remained iced until processed as described below. All processing of these blood samples was done within 2 h of completing the experiment. Arterial blood pressure was monitored immediately before starting the IC infusion and at 5 and 30 min into the experiment. After obtaining the last (30 min) blood specimen, the position of the catheters in the aortic arch, renal vein, and hepatic vein was reassessed angiographically. In no experiment did catheter displacement occur. In the four experiments performed under sterile surgical conditions, the arteriotomy and venotomy incisions were then sutured and the animals allowed to recover from the general anesthesia. In the three experiments performed under nonsterile conditions, 1–2 μCi of 85Sr-labeled microspheres were infused into the aortic arch catheter and, after 2–3 min had elapsed, the animals were sacrificed by infusion of pentobarbital. To determine the content of each isolate in each organ, an aliquot (≤10%) was taken from liver, kidney, spleen, stomach, large bowel, small bowel, pancreas, and lung and the isolate counts per minute determined as described previously (11).

Processing of blood samples

Each blood sample obtained during the experiment was immediately transferred to an iced tube. Under these conditions, additional binding of IC to erythrocytes ceases, and erythrocyte-bound IC do not dissociate (see below). Thus, iceding the blood samples helps stabilize the binding of IC to erythrocytes at the same level as that present in vivo. To determine the extent to which the IC had become bound to erythrocytes, in vivo, it was necessary to devise a technique for separating IC bound to erythrocytes from IC not bound to erythrocytes. This could not be done simply by centrifugation of the whole blood specimens because the centrifugation force necessary to efficiently separate erythrocytes from leukocytes results in the downward migration of some of the IC population with the erythrocytes, even though the IC are not bound to the erythrocyte. To obviate
this problem, the following technique was devised: whole blood (0.5 ml) was placed on top of 3 ml of iced percoll (Sigma Chemical Co.) diluted in 0.15 M NaCl to a density of 1.100 g/ml. This preparation was then centrifuged at 350 g for 15 min. Under these conditions, >97% of the erythrocytes and IC bound to erythrocytes migrate to the bottom of the centrifuge tube and >95% of free 125I-BSA and IC bound to the buffy coat cells or free in the plasma, remain at the top of the percoll. The separation of buffy coat cells from erythrocytes was highly efficient as shown by the fact that, after centrifugation of whole blood on percoll, leukocytes were undetectable in the erythrocyte fraction by Coulter counter techniques (Coulter Electronics Inc., Hialeah, FL). In addition, microscopic examination of Wright-stained specimens of the erythrocyte fraction revealed no evidence of leukocytes or platelets. When IC alone were placed on top of the percoll (with or without fresh serum) and centrifuged as described above, <4% of the IC preparation migrated to the bottom of the tube during centrifugation (8). The protocol for these experiments is described below.

In vitro binding of IC to erythrocytes

Equal volumes of IC (prepared as described above), fresh undiluted baboon serum, and packed baboon erythrocytes (washed three times in PBS; buffy coat cells removed following each wash) were added together in duplicate iced tubes (12 x 75 mm). The contents were mixed and incubated at 37°C for 10 min in a shaker water bath. After incubation the reaction mixtures were removed, placed on iced percoll and centrifuged at 350 g for 15 min at 4°C. After centrifugation, the unbound IC (remaining on top of the percoll) were removed along with the percoll, leaving the erythrocytes with bound IC as a pellet at the bottom. The 125I counts per minute of both fractions were measured in a gamma scintillation spectrometer (model 5230, Packard Instrument Co., Inc., Downers Grove, IL). The percent of IC bound to erythrocytes was calculated as follows: % bound = (125I cpm erythrocyte fraction)/0.9(125I cpm erythrocyte fraction + percoll fraction) x 100, where 0.9 = fraction of 125I cpm that represent IC. Reaction mixtures containing equal volumes of IC, heat-inactivated serum, and packed baboon erythrocytes were included during the 37°C incubation period as negative controls.

To assess whether the immediate cooling to 4°C of the primate blood samples stabilize the IC binding to erythrocytes that has already occurred but prevents further IC binding to erythrocytes, the following in vitro experiments were performed. IC + erythrocytes + fresh plasma were incubated at 37°C for 10 min and then immediately transferred to a 4°C bath. Another incubate, consisting of IC + erythrocytes + fresh plasma, was incubated at 4°C continuously. Aliquots of each reaction mixture were removed periodically to measure binding of IC to erythrocytes, as described above. The results of a representative experiment are shown in Fig. 1. As can be seen, at 4°C, IC binding to erythrocytes does not occur but IC already bound to erythrocytes do not dissociate. Thus, cooling the blood samples to 4°C maintains the level of IC binding to erythrocytes at the same levels as those present in vivo.

Analysis of IC size by isokinetic sucrose gradients

Isokinetic sucrose gradients were constructed as described by Johns and Stanworth (13). Briefly, exponential gradients were formed of 15-24.25% sucrose (wt/wt, isotonic at 20°C) using a constant volume mixing chamber (kindly provided by Dr. Ronald Taylor, Department of Biochemistry, University of Virginia School of Medicine). The gradients (total gradient volume = 4.8 ml) were placed in a temperature-regulated ultracentrifuge (model L8-55, Beckman Instruments, Inc., Palo Alto, CA) and equilibrated at 20°C. To determine the population of IC that bind to baboon erythrocytes the following three reaction mixtures were prepared in duplicate: (a) fresh serum + IC + erythrocytes; (b) fresh serum + IC + PBS; and (c) heat-inactivated serum + IC + erythrocytes. After 10 min at 37°C, each reaction mixture was placed on percoll and centrifuged at 350 g for 15 min. After centrifugation, the material above the percoll was removed, the 125I counts per minute determined and then placed on the 15-24.25% isokinetic sucrose gradients. The gradients were centrifuged at 1,500 rpm for 30 min (w2 = 4.39 x 107 rad2/s) at 20°C. The gradients were then removed and fractionated via bottom displacement with 33% sucrose. 0.2-m1 fractions were collected (total of 25 fractions) and 125I counts per minute determined. The 125I counts per minute of each fraction was then plotted against the sedimentation coefficient (S20, w) corresponding to each sucrose fraction.

Calculations

Peak in vivo binding of IC to erythrocytes. This was taken as ICc/(0.9 x ICf) x 100, where ICc = 125I counts per minute in the erythrocyte fraction at the time of the peak ICc; ICf = the total 125I counts per minute in the blood sample; 0.9 = fraction of total 125I counts per minute in blood that represents IC.

Fractional removal of IC from erythrocytes traversing liver or kidney or spleen + gut. This was determined as follows: the 125I counts per minute of the erythrocyte fractions of the blood entering and leaving the given organ were plotted as a function of time. The time course of IC removal from the erythrocyte fraction was divided into an early phase (0-5 min) and a late phase (5-30 min). 5 min was chosen as the dividing line since the period of 0-5 min corresponded to the period of the greatest rate of removal of IC from the blood (Results). The area under the early phase curve of the 125I counts per minute on erythrocytes entering (A) and leaving (B) the given organ was then determined. The fraction of the 125I-labeled IC removed during the early phase was then calculated as (A - B)/A. The late phase fractional removal of IC from erythrocytes was calculated as the ratio: 1 - (mean 125I cpm on erythrocytes leaving the organ at the 5, 15, and 30 min time points)/(mean 125I cpm on erythrocytes entering the given organ at the 5, 15, and 30 min time points). This simpler formula was used for the late phase calculation because, in general, the late phase values represented low and nearly steady-state values with only small differences between the 125I counts per minute on the erythrocytes entering and leaving the given organ.

The IC on erythrocytes entering and leaving the kidney were taken as the 125I counts per minute in the erythrocyte fractions of the arterial and renal venous blood samples, respectively. The IC on erythrocytes entering and leaving spleen + gut were taken as the 125I counts per minute in the erythrocyte fractions of the arterial and portal venous blood samples, respectively. To determine the IC on erythrocytes entering liver (via hepatic artery and portal vein), the following corrections were applied to the 125I counts per minute in the erythrocyte fraction of each hepatic artery and portal
vein blood specimen. In the experiments in which microspheres were not used, the relative contribution of hepatic artery and portal vein blood flow to total hepatic blood flow was apportioned as follows: 0.32 (I25I cpm in the arterial blood erythrocyte fraction) + 0.68 (I25I cpm in the portal vein erythrocyte fraction). This apportionment of total hepatic flow into hepatic artery and portal vein flow is based on our previously reported studies in dogs, using the microsphere method (described above) to determine hepatic artery and portal vein blood flow (11). In the 24 experiments in dogs, hepatic artery flow averaged 32% (range 5 to 60%) of total hepatic blood flow. It should be noted that the wide variation between values of individual experiments is inherent in this method. In the three experiments in primates in which microspheres were infused, the actual distribution of the microspheres was used to determine the relative contribution of hepatic artery and portal vein blood flow to total hepatic blood flow. The data in these three experiments were in general agreement with the data obtained in the dog. That is, hepatic artery flow in the three primate experiments was 46.2, 40.1, and 8.6% of total liver blood flow, for a mean value of 31.6%. The IC on erythrocytes leaving liver were taken as I25I counts per minute in the erythrocyte fraction of the hepatic vein blood samples.

Fractional removal of IC from the nonerythrocyte fraction of blood (plasma/buffy coat fraction). To determine the extent to which IC were removed from the blood by mechanisms other than the erythrocyte-IC-clearing mechanism, the I25I content of the entire percoll fraction of each blood specimen was measured and plotted vs. time. These data were then analyzed exactly as was done for the erythrocyte fractions of blood, as discussed above. The percoll fractions contained I25I-labeled IC free in the plasma, I25I-labeled IC bound to leukocytes and platelets, and the I25I-BSA free in the plasma. Hereafter, this fraction of blood will be referred to as the plasma/buffy coat fraction.

Fraction of total hepatic blood flow via the hepatic artery. This fraction was taken as the ratio: (total microsphere cpm in liver)/(total microsphere cpm in liver + spleen + stomach + large bowel + small bowel + pancreas). The rationale for this calculation is that the microspheres in liver represent hepatic artery blood flow, while the microspheres in spleen + gut + pancreas represent blood flow to the liver via the portal vein (11).

Fraction of total hepatic blood flow via the portal vein. This fraction was calculated as (1 - the fraction of hepatic blood flow via the hepatic artery).

Tissue vascular volume. The fraction of tissue volume

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**FIGURE 1** Absence of IC binding to or dissociation from erythrocytes at 4°C. In one experiment (panel A), equal volumes of IC, fresh baboon plasma (10 U heparin/ml), and packed baboon erythrocytes were mixed together in a series of tubes and placed at 37°C for 10 min, and then at 4°C thereafter. In another experiment (panel B), equal volumes of IC, fresh baboon plasma, and baboon erythrocytes were mixed together in a series of tubes and incubated from the outset at 4°C. At 0, 20, 40, 60, and 180 min duplicate tubes were removed and the reaction mixtures were placed on iced percoll for separation of erythrocyte-bound IC from unbound IC. For each experiment, mean percent IC bound to erythrocytes was plotted against incubation time at 4°C.
that represented blood volume was taken as the ratio: \((^{51}\text{Cr} \text{ cpm/g of given tissue})/(^{51}\text{Cr} \text{ cpm/ml whole blood})\) (11).

Vascular space correction. To determine the extent to which the tissue \(^{125}\text{I} \text{ counts represent} ^{125}\text{I}-\text{labeled IC actually deposited in tissues vs.} ^{125}\text{I}-\text{labeled material, which is simply passively retained in the vascular space at the time the tissues were excised, a correction factor based on the} ^{51}\text{Cr counts in the vascular space and} ^{125}\text{I} \text{ counts in whole blood was applied as follows:} ^{125}\text{I} \text{ cpm deposited in tissue} = \left(\frac{^{51}\text{Cr cpm/g of tissue}}{^{51}\text{Cr cpm/g of whole blood}}\right) \times \left(\frac{^{125}\text{I} \text{ cpm/ml of whole blood}}{^{125}\text{I} \text{ cpm/ml of whole blood}}\right), \text{as described previously (14).}

**Hemodynamic data.** Mean systolic blood pressure at the start of the experiment was 82±6 mmHg and was stable during the experiment. Mean hematocrit at the start of the experiment was 33±4% and decreased by ~5% during the experiment due to the volume of blood samples removed and saline infused. In no instance was hemolysis noted.

**RESULTS**

**Clearance of preformed IC**

The results of a representative experiment are shown in Figs. 2 and 3. As can be seen from comparison of the \(^{125}\text{I} \text{ counts per minute in the arterial blood erythrocyte fraction}\) (Fig. 2) to the \(^{125}\text{I} \text{ counts per minute in the arterial blood plasma/buffy coat fraction}\) (Fig. 3), the majority of the \(^{125}\text{I} \text{labeled IC infused into the arterial blood become bound to erythrocytes. Furthermore, as the IC-bearing erythrocytes traverse liver the great majority of their IC burden is deposited. This is shown in Fig. 2 by the low \(^{125}\text{I} \text{ counts per minute in the erythrocytes exiting liver (hepatic vein erythrocyte fraction) compared with the high} ^{125}\text{I} \text{ counts per minute on erythrocytes entering liver (hepatic artery + portal vein erythrocyte fractions). The erythrocytes themselves were not trapped in liver, as shown in Fig. 4.} ^{51}\text{Cr}-\text{Labeled erythrocytes bind IC in vitro to the same extent as do erythrocytes not subjected to the process of} ^{51}\text{Cr-labeling (Fig. 5). Thus, under these conditions, the behavior of the} ^{51}\text{Cr}-\text{labeled erythrocytes is representative of the general erythrocyte population.}

The spleen appears to function in the same manner as the liver as shown by the fact that the \(^{125}\text{I} \text{labeled IC are efficiently removed from the erythrocytes traversing the organs served by the portal vein (spleen + gut) and, as shown below, most of the IC deposited in spleen + gut are deposited in spleen. The} ^{51}\text{Cr-labeled erythrocytes are not detained by the spleen.}

The handling of IC-bearing erythrocytes by kidney stands in striking contrast to that of the liver. By comparing the \(^{125}\text{I} \text{ counts per minute in arterial blood to that of renal vein, it is evident that the IC-bearing erythrocytes exit kidney with IC burden virtually intact (Fig. 2).}

The handling of the plasma/buffy coat fraction of blood by liver, kidney, and spleen + gut is shown in

![Figure 2](image-url)
**Figure 3** Representative study. Sequential change in $^{125}$I counts per minute in percoll fractions of blood samples obtained from arterial (●), renal vein (○), portal vein (□), and hepatic vein (■) catheters as in the experiment depicted in Fig. 1. Percoll fraction refers to the plasma/buffy coat fraction.

Hepatic Vein Blood $^{51}$Cr cpm/ml/Arterial Blood $^{51}$Cr cpm/ml

Renal Vein Blood $^{51}$Cr cpm/ml/Arterial Blood $^{51}$Cr cpm/ml

Portal Vein Blood $^{51}$Cr cpm/ml/Arterial Blood $^{51}$Cr cpm/ml

**Figure 4** Fractional removal of $^{51}$Cr-labeled erythrocytes traversing liver, kidney, or spleen + gut, before, during, and after infusion of IC over the time period from 0 to 120 s. The fractional rate of removal of $^{51}$Cr-labeled erythrocytes by the respective organ systems was estimated from the ratios shown above each graph.
As can be seen, the $^{125}$I-labeled material in the plasma/buffy coat fraction is cleared at a minimal rate by liver and spleen + gut and not at all by kidney. Thus, IC free in the plasma phase or bound to leukocytes or platelets are cleared far less efficiently from the blood than are the IC bound to erythrocytes.

The IC clearance data from all seven experiments are summarized in Tables I and II. As can be seen from Table I, at 60–180 s the majority of the IC in blood were bound to erythrocytes. There was no correlation between the dose of IC administered and the observed level of IC binding to erythrocytes. However, by the end of the early phase (5 min) most (73.1%) of the IC had been removed from the erythrocytes. In contrast, at this point only 17% of the $^{125}$I counts per minute in the plasma/buffy coat fraction had been removed. By 15 and 30 min, a substantial increase was seen in the calculated fractional removal of IC from the plasma/buffy coat fraction. This, however, overestimates the actual fractional removal of IC since some of the loss of $^{125}$I-labeled material from the plasma/buffy coat fraction represents diffusion of free $^{125}$I-BSA into the interstitial space.

Table II shows the early phase clearance of IC by liver, kidney, and spleen + gut. As in the representative experiment (Figs. 2 and 3), there were high rates of IC clearance from the erythrocyte fraction by liver, no significant clearance by kidney, and intermediate rates of clearance by spleen + gut. Low but significant rates of IC clearance from the plasma/buffy coat fraction was observed for liver and spleen + gut.

The mean late phase clearance of IC from the erythrocyte fraction for all experiments was liver 50.1±8.4%, kidney 8.5±12%, and spleen + gut 2.4±6%. None of these mean values is significantly different from 0. The mean late phase clearance of $^{125}$I-labeled material from the plasma/buffy coat fraction of blood was liver 7.7±10%, kidney 8.5±12%, and spleen + gut 2.4±6%. None of these mean values is significantly different from 0. Thus, for the late phase data, only the hepatic removal of IC from the erythrocyte fraction showed values for fractional removal significantly different from 0. It should be noted, however, that the rate of fractional removal of IC from the erythrocyte fraction by liver during the
TABLE I

Binding of IC to Erythrocytes and their Subsequent Removal from the Erythrocyte Fraction and Plasma/Buffy Coat Fraction of Blood

Table: Early phase (0-5 min)

<table>
<thead>
<tr>
<th>Exp no.</th>
<th>IC bound to erythrocytes (Peak value)</th>
<th>IC in plasma/buffy coat fraction (Peak value)</th>
<th>Time of peak value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1‡</td>
<td>85.3</td>
<td>14.7</td>
<td>120</td>
</tr>
<tr>
<td>2</td>
<td>78.3</td>
<td>21.7</td>
<td>60</td>
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<td>3</td>
<td>47.6</td>
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</tr>
<tr>
<td>4</td>
<td>55.4</td>
<td>44.6</td>
<td>60</td>
</tr>
<tr>
<td>5</td>
<td>17.7</td>
<td>82.3</td>
<td>120</td>
</tr>
<tr>
<td>6</td>
<td>69.7</td>
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<td>80</td>
</tr>
<tr>
<td>7</td>
<td>74.6</td>
<td>25.4</td>
<td>120</td>
</tr>
</tbody>
</table>

Mean: 61.2±8.8 38.8±8.8

Table: Late phase (5-30 min)

<table>
<thead>
<tr>
<th>Exp no.</th>
<th>Percent reduction in IC from peak value of erythrocyte fraction (5 min)</th>
<th>Percent reduction in IC from peak value of plasma/buffy coat fraction (5 min)</th>
</tr>
</thead>
</table>

5 min: 36.4 34.8 34.5
15 min: 12.2 29.4 43.1
30 min: 3.6 17.0 18.8

‡ Mean Qₑ: 44.8±8.2 54.4±8.7 57.0±9.3
§ Mean Qₑ: 6.5±3.3 13.7±4.2 14.8±4.4

* As defined in Methods.
† Percent IC in plasma/buffy coat fraction (peak value) = 100% - % IC bound to erythrocytes (peak value).
‡ Experiment 1 is the representative study shown in Figs. 2 and 3.
§ Qₑ (percent of peak IC load removed from blood by erythrocyte fraction) = % IC bound to erythrocytes (peak value) × % reduction of IC in erythrocyte fraction at time t, where t = 5, 15, or 30 min. The mean value shown is the average of the individual experiments.
§ Qₑ (percent of peak IC load removed from blood by plasma/buffy coat fraction) = % IC in plasma/buffy coat fraction (peak value) × % reduction of IC in percoll fraction at time t, where t = 5, 15, or 30 min. The mean value shown is the average of the individual experiments.

The late phase was much lower than that observed during the early phase (late phase: 50.1%/25 min = 2.0%/min) vs. (early phase: 73.1%/5 min = 14.6%/min).

Tissue analysis

The following data were taken from the three experiments (1, 2, and 5) in which the animals were

TABLE II

Early Phase (0-5 min) Data on the Removal of IC from the Erythrocyte Fraction and the Plasma/Buffy Coat (Percoll) Fraction of Blood by Liver, Kidney, and Spleen + Gut

<table>
<thead>
<tr>
<th>Experiment no.</th>
<th>IC removed from erythrocyte fraction</th>
<th>¹³¹I-Labeled material removed from percoll fraction</th>
<th>IC removed from erythrocyte fraction</th>
<th>¹³¹I-Labeled material removed from percoll fraction</th>
<th>IC removed from erythrocyte fraction</th>
<th>¹³¹I-Labeled material removed from percoll fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1†</td>
<td>84.8</td>
<td>23.4</td>
<td>2.5</td>
<td>0.8</td>
<td>30.0</td>
<td>18.3</td>
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<td>2</td>
<td>59.6</td>
<td>-1.0</td>
<td>-0.8</td>
<td>11.3</td>
<td>28.1</td>
<td>24.3</td>
</tr>
<tr>
<td>3</td>
<td>47.4</td>
<td>32.5</td>
<td>-15.4</td>
<td>16.9</td>
<td>11.6</td>
<td>16.9</td>
</tr>
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<td>4</td>
<td>75.7</td>
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<td>23.5</td>
<td>12.9</td>
<td>40.7</td>
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</tr>
<tr>
<td>5§</td>
<td>64.9</td>
<td>33.3</td>
<td>16.8</td>
<td>2.7</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>92.0</td>
<td>14.6</td>
<td>6.3</td>
<td>-5.4</td>
<td>21.1</td>
<td>-2.6</td>
</tr>
<tr>
<td>7§</td>
<td>60.5</td>
<td>25.4</td>
<td>-2.5</td>
<td>-11.8</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Mean: 69.3±5.9 25.2±6.4

43±4.9 3.9±3.9 26.8±4.8 14.7±4.5

* As defined in Methods.
† Experiment 1 is the representative study shown in Figs. 2 and 3.
§ Portal vein blood samples were not taken in these experiments. The hepatic fractional removal of IC and ¹³¹I-labeled material was calculated using the values for arterial and hepatic vein blood samples.
killed immediately after the conclusion of the experimental protocol. The results of the individual experiments are given in the following sequence: experiment 1, then 2, then 5.

**Tissue uptake of IC.** The total $^{125}$I counts per minute injected into each animal varied. Thus, to permit comparison between experiments, the tissue uptake of $^{125}$I, after correction for vascular space $^{125}$I (Methods), is expressed as the ratio: ($^{125}$I cpm/g of a given organ)/($^{125}$I cpm/g of liver).

The following ratios were obtained: kidney (cortex)/liver: 0.012, 0.000, 0.000; spleen/liver: 1.99, 0.48, 0.76; lung/liver: 0.28, not done in No. 2, 0.88; gut (mean of stomach + large bowel + small bowel + pancreas)/liver 0.030, 0.01, 0.038. In these three experiments, the percent of total $^{125}$I counts per minute infused which were deposited in the various organs was liver 67, 32.8, 26.1; spleen 6.8, 0.7, 1.2; kidney 0.03, 0.00, 0.00; gut 0.4, 0.13, 0.5; and lung 7.9, not done in No. 2, 9.2. The tissue analysis data show that the liver traps by far the greatest amount of the IC. Lung is also a significant site of deposition. Kidney and gut trap relatively small amounts of IC.

**Tissue vascular volume.** The tissue vascular volume was calculated from the $^{51}$Cr in tissue and blood, as discussed in Methods. The values are as follows: liver 0.23, 0.23, 0.24; lung 0.94, not done in No. 2, 0.23; kidney 0.23, 0.19, 0.14; spleen 0.35, 0.50, 0.65. The calculated volumes were comparable to those we found in dog under similar conditions (11). Thus, there is no evidence of selective removal of $^{51}$Cr-labeled erythrocytes from the circulation. If this had occurred, even in a subtle fashion, the calculated tissue vascular allowance would have been inappropriately high (10).

**Isokinetic sucrose density gradient analysis of IC**

To determine the size of the IC used in these experiments and the size of IC which bind to erythrocytes, in vitro experiments were performed using erythrocyte suspensions depleted ofuffy coat cells before incubation with IC. These experiments are depicted and described in Fig. 6. As can be seen, IC ranging in size from 10,000 S to >288,000 S are present in the original IC preparation and this range of IC sizes was only slightly affected by incubation in fresh serum. The entire range of IC bound to erythrocytes. However, the percent of IC bound to erythrocytes increased markedly with increasing IC size.

**Search for evidence of the erythrocyte-IC-clearing mechanism under conditions in which IC can form in vivo**

Two additional experiments were performed to determine whether the operation of the erythrocyte-IC-clearing mechanism could be detected when IC are allowed to form in vivo by infusing antigen and antibody separately into the same animal. In one experiment antigen was infused first, followed 15 min later by antiserum. In the other experiment, the order of infusion was reversed. The proportion of antigen and antibody was that which approximated equivalence. In all other respects the protocol was the same as described in Methods. In both experiments <4% of the $^{125}$I counts per minute were found in the erythrocyte fraction. On the basis of the experiment in which $^{125}$I-BSA was infused first, it could be determined that this small amount of erythrocyte-associated $^{125}$I could be attributed to nonspecific binding. Throughout both experiments whole blood $^{125}$I counts per minute fell gradually, after the BSA infusion was completed, probably largely reflecting the diffusion of $^{125}$I-BSA into the interstitial spaces. There was negligible removal of $^{125}$I-labeled material from either the erythrocyte or plasma/buffy coat fractions traversing liver or kidney. Thus, the erythrocyte-IC-clearing mechanism, or any other IC-clearing mechanism, was not detected in either of these experiments. Although negative, these experiments are useful since they show that specific hepatic removal of albumin ($^{125}$I-BSA) from the circulation (15) cannot explain the high rate of hepatic removal from the circulation of the preformed IC used in this study. These experiments also suggest that future studies to elucidate the operation of the erythrocyte-IC-clearing mechanism will require the use of specific populations of preformed IC since the percent binding to erythrocytes of IC forming in vivo appears to be very low.

**DISCUSSION**

This study demonstrates that when large, complement-fixing IC are infused into the blood stream of primates, the great majority of these IC rapidly become bound to circulating erythrocytes. The mechanism of adherence of IC to erythrocytes in vivo is currently under investigation in our laboratory. However, in vitro, it has been shown that the binding of IC to primate erythrocytes is complement dependent (6-9). It is generally thought that C3b sites are generated on the IC, which then allow the IC to bind to erythrocyte CR1 receptors. C3b sites on IC are generated very rapidly. For example, previous studies have shown that, in vivo, C3b sites are generated on IgM-sensitized erythrocytes so rapidly that only liver blood flow limits the rate at which the sensitized erythrocytes are removed from the circulation by the C3b receptors of the hepatic mononuclear phagocyte system (16). It has also been shown that C3b-mediated binding of preformed IC to leukocytes in vivo can be detected within 30 s of in-
fusion of the IC (17). Thus, the speed of binding of IC to erythrocytes observed in this study is consistent with the hypothesis that the binding reaction is complement mediated.

IC bound to erythrocytes do not cause detectable hemolysis in vitro (8) or in vivo. It is tempting to speculate that this lack of hemolysis is due to the inhibitory effect of the CR1 receptor on activation of the complement cascade (18, 19). Thus, the erythrocyte is not damaged by attachment to the IC. Indeed, in vitro studies have shown that if IC are cleaved from the erythrocyte by prolonged exposure to the complement-dependent release activity of fresh serum (20), the erythrocyte is, once again, able to bind IC (21).

Although IC-bearing erythrocytes can be demonstrated to aggregate in vitro (immune adherence reaction [6]) there was no evidence in this study that the IC-bearing erythrocytes aggregate in vivo. If this had occurred, there should have been evidence of 51Cr-labeled erythrocytes being removed from the circulation as erythrocyte aggregates traversed liver, kidney, or spleen + gut. However, there were no 51Cr gradients across these organs and calculated tissue vascular volumes were normal. It is possible that the shearing forces on erythrocytes in the circulation are greater than the immune adherence forces that aggregate IC-bearing erythrocytes in vitro. In any event, there was no evidence that binding of IC to erythrocytes in vivo resulted in detectable erythrocyte aggregation.

This study further demonstrates that, after the IC have become bound to the erythrocyte in vivo, the IC do not deposit in organs such as kidney or gut as the IC-bearing erythrocytes traverse these organs. Instead, the IC are efficiently removed from the erythrocyte as the IC-bearing erythrocyte traverses liver or spleen. The erythrocyte then exits liver or spleen apparently able, once again, to participate in this "erythrocyte-IC-clearing mechanism."

The mechanism by which IC are removed from
erythrocytes as they traverse liver is not known. It is possible that, as the first step, the exposed Fc regions of the IC bound to the erythrocyte attach themselves to the Fc receptors of the tissue-bound hepatic mononuclear phagocyte. There is evidence that binding of IC to macrophages via the Fc receptor results in ingestion of the IC (5). Thus, the fact that the IC removed from the erythrocytes traversing liver are not released back into the hepatic circulation suggests that the IC have been ingested by the hepatic macrophages. This would be in contrast to the hepatic uptake of IgM-sensitized erythrocytes, which occurs via hepatic macrophage receptors for C3b. In that situation the erythrocytes are released back into the circulation within 30 min, probably through the enzymatic action of C3b inactivator (16). Although no evidence of release of IC back into the hepatic circulation was detected in the study presented here, the possibility that erythrocyte-bound IC bind to hepatic macrophages via both C3b and Fc receptors is not excluded.

If the first step in the hepatic uptake of erythrocyte-bound IC is the binding of the IC to hepatic macrophage C3b and/or Fc receptors, the second step is the breaking free of the erythrocyte from the macrophage-bound IC. This could occur mechanically, because of the force of blood flow on the relatively rigid erythrocyte causes the erythrocyte to tear free from its attachment to the hepatic macrophage. If the total force binding the IC to the hepatic macrophage is greater than the total force binding the IC to the erythrocyte, the erythrocyte will break free leaving the IC bound to the hepatic macrophage. Alternatively, C3b inactivator or some other enzyme could cleave the bond between IC and erythrocytes. Indeed, Medof et al. (22) have shown that erythrocyte CR1 receptor markedly enhance the activity of C3b inactivator. Thus, it seems possible that a synergistic interaction between mechanical and enzymatic factors in liver could be responsible for the extremely efficient uptake of erythrocyte-bound IC by liver.

In contrast to the rapid hepatic removal of IC from the erythrocyte fraction of blood, the hepatic removal of IC from the plasma/buffy coat fraction of blood was considerably slower. This could reflect strong binding or phagocytosis of IC by peripheral blood leukocytes rendering the IC unavailable to the hepatic mononuclear phagocytes. It is also possible that the IC in the plasma phase represent a population of smaller IC that are cleared by the liver less efficiently than are larger IC (23, 24).

The preformed IC used as the biological probe in this study are much larger than are those used traditionally and were chosen simply because they showed the highest level of binding to primate erythrocytes in vitro. The biological significance of such large IC is not clear. However, Cameron (25) has recently provided arguments that the nephritogenic IC are large, insoluble or poorly soluble IC formed near equivalence, and not the small, soluble IC that circulate in plasma. It is possible, nevertheless, that such large circulating IC do not occur in pathophysiologic states and that the erythrocyte-IC-clearing mechanism is directed mainly at clearing less large IC at a less efficient rate than the very large preformed IC used in this study. Indeed, it is clear that, in vitro, smaller IC can bind to primate erythrocytes. Waller et al. (26) have demonstrated low level binding of preformed DNA anti-DNA IC to human erythrocytes when IC size is <175 S but high level binding as IC size approaches 600 S. Medof et al. (27) have shown that preformed BSA-anti-BSA IC smaller (~200 S) than those used in the present study can bind to primate erythrocytes in vivo. However, high dilution of the complement source (1:8 to 1:16 dilution of serum) is needed for maximum binding of these smaller IC (9).

In this study, IC binding to erythrocytes was not detected in vivo when free antigen and antibody were infused separately. This, however, may simply indicate that IC of sufficient size to permit binding to erythrocytes form at a rate which is too low and/or are cleared too quickly to be detected under the present experimental conditions. Thus, studies to elucidate further the operation of the erythrocyte-IC-clearing mechanism will apparently also require the use of specific populations of preformed IC.

The erythrocyte-IC-clearing mechanism is, evidently, largely an adaptation of man and other primates. Lower forms of animals have some ability of their erythrocytes to bind to IC, possibly through Fc receptors (28) or antibody-aggregated C4 receptors (29). However, only primates have vast numbers of CR1 receptors on circulating erythrocytes. The present study demonstrates the existence in primates of a potentially important erythrocyte-IC-clearing mechanism and provides a teleologically satisfying explanation for the existence of the vast numbers of CR1 receptors on circulating erythrocytes. Failure or insufficiency of the erythrocyte-IC-clearing mechanism might increase the probability of deposition of IC in vulnerable organs, such as kidney, and concomitantly decrease the probability of safe disposal of IC in liver or spleen. Indeed, this may occur in patients with systemic lupus erythematosus who have deficient numbers of erythrocyte CR1 receptors (3, 30, 31) or in whom the erythrocyte CR1 receptors are occupied by IC (32, 33). Thus, inadequacy of the erythrocyte-IC-clearing mechanism could play an important role in the clinical expression of systemic lupus erythematosus and other diseases mediated by deposition of circulating IC.
ACKNOWLEDGMENTS
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REFERENCES