Hypophosphatemia after right hepatectomy for living donor liver transplantation

Kelly W Burak MD1, Charles B Rosen MD1, Jeff L Fidler MD2, Gina K Hesley MD2, David Nagorney MD1, Michael R Charlton MD1, David J Brandhagen MD1

Hypophosphatemia has been described in patients undergoing right hepatectomy for liver cancer and in living donors for liver transplantation who also received total parenteral nutrition. At the study centre, significant hypophosphatemia (0.36 mmol/L or less) requiring intravenous replacement was seen in two of the first nine living donors for adult-to-adult liver transplantation. To determine the frequency of hypophosphatemia in living donors, the authors obtained phosphate levels on stored serum samples postoperatively days 0, 1, 3 and 7 in all nine patients, none of whom were on total parenteral nutrition. Within the first week, hypophosphatemia developed in 55.6% of patients and phosphate levels returned to normal by day 7 in all nine patients. One patient had normal phosphate levels during the first week, but had profound hypophosphatemia (0.32 mmol/L) on day 14 when he presented with a Staphylococcus aureus infection of a bile collection and significant hypoxemia. The extent of hepatectomy and the rate of liver regeneration, estimated by baseline and postoperative day 7 volumetric computed tomography scans, did not correlate with the development of hypophosphatemia. In conclusion, hypophosphatemia is common in living donors undergoing right hepatectomy and may be associated with complications. All living donors should be monitored for the development of hypophosphatemia during the first two postoperative weeks.

Key Words: Hepatectomy; Hypophosphatemia; Liver transplantation; Surgical complications

Hypophosphatemia has been described in patients undergoing major hepatic resections for liver cancer and has been associated with postoperative complications in these patients (1). In this initial report, serum phosphate levels fell in all 44 patients who underwent right hepatectomy for malignant tumours. Profound hypophosphatemia (less than 0.32 mmol/L) developed in eight patients and was significantly associated with the development of major postoperative complications including infections, cardiac arrhythmias, respiratory arrest and hepatic insufficiency (1). Because of the donor organ shortage, many liver transplant centres have now initiated living donor liver transplant programs (2,3). Pomposelli et al (4) reported the development of hypophosphatemia as a complication of adult-to-adult living donor liver transplantation. Of their first 18 donors, profound hypophosphatemia (less than 0.32 mmol/L) developed in 70% and severe hypophosphatemia (0.32 mmol/L to 0.48 mmol/L) developed in 30%. However, all donors were receiving total parenteral nutrition (TPN), which in itself is known to cause hypophosphatemia (5). In contrast, a recent report (6) of 95 right lobe donors found that hypophosphatemia was mild and was not associated with increased morbidity.

We began an adult-to-adult living donor liver transplantation program at our study institution in June 2000. Significant hypophosphatemia (0.36 mmol/L or less) was observed in two of nine living donors during the first year of our program. However, serum phosphate levels were not routinely tested in all donors. We therefore sought to determine the frequency and severity of hypophosphatemia in our liver donors by using...
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TABLE 1
Serum phosphate levels before (day 0) and during the first week following right hepatectomy in nine living liver donors

<table>
<thead>
<tr>
<th>Donor</th>
<th>Serum PO₄ levels (mmol/L)</th>
<th>% decrease in PO₄</th>
<th>Degree of hypophosphatemia*</th>
<th>Complications</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Day 0</td>
<td>Day 1</td>
<td>Day 3</td>
<td>Day 7</td>
</tr>
<tr>
<td>1</td>
<td>1.10</td>
<td>0.61</td>
<td>0.88</td>
<td>1.00</td>
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<td>2</td>
<td>0.94</td>
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<td>1.03</td>
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<td>0.87</td>
<td>0.90</td>
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<td>1.26</td>
<td>1.00</td>
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<td>1.23</td>
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<td>1.26</td>
<td>1.00</td>
<td>0.81</td>
<td>1.42</td>
</tr>
<tr>
<td>7</td>
<td>1.23</td>
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<td>0.45</td>
<td>1.03</td>
</tr>
<tr>
<td>8</td>
<td>1.42</td>
<td>0.36</td>
<td>0.61</td>
<td>1.03</td>
</tr>
<tr>
<td>9</td>
<td>1.10</td>
<td>1.16</td>
<td>0.87</td>
<td>1.07</td>
</tr>
</tbody>
</table>

Mean ± SEM 1.13±0.06 0.90±0.06 0.68±0.06 1.16±0.03 38.9±7.7

*Hypophosphatemia is defined as normal (>0.81 mmol/L), moderate (0.81 mmol/L to 0.52 mmol/L), severe (0.48 mmol/L to 0.32 mmol/L) and profound (<0.32 mmol/L). †Case 9 developed profound hypophosphatemia (0.32 mmol/L) on postoperative day 14 when presenting with an infected bile collection but had normal phosphate levels during the first week.

RESULTS
Nine living donors underwent right hepatectomy (five female, four male). The mean age of the donors was 35.8±10.9 years and their mean weight was 83.5±6.6 kg. Hypophosphatemia was prospectively identified in two of our nine living donors (donors 8 and 9). Donor 8, a 49-year-old woman who donated to her husband, was noted to have hypophosphatemia on the second postoperative day (0.36 mmol/L). At the time, she was found to have bile in surgical drains and endoscopic retrograde cholangiopancreatography confirmed a small bile leak that was treated successfully with endoscopic stenting. Intravenous phosphate replacement was started and phosphate levels returned to normal within one day. Donor 9, a 47-year-old man who donated to his uncle, presented on postoperative day 14 with fever (39.0°C) and was noted to be hypoxemic (oxygen saturation 80% on room air). A chest x-ray demonstrated a small right pleural effusion and a spiral CT scan excluded a pulmonary embolus. A CT scan of the abdomen demonstrated a subphrenic collection consistent with an abscess, and a CT-guided percutaneous drainage was performed. He was found to have profound hypophosphatemia (0.32 mmol/L) and intravenous phosphate was started. Cultures from aspirated fluid and blood grew Staphylococcus aureus and he improved on parenteral antibiotics.

Donor serum phosphate levels from the first week following surgery were measured from stored serum samples and are shown in Table 1. Serum phosphate levels were normal (greater than 0.81 mmol/L) in all donors before the operation (range 0.87 mmol/L to 1.42 mmol/L). Phosphate levels fell by the third postoperative day in eight of nine donors. The mean decrease was 39% (range 0% to 75%). Within the first week following surgery, hypophosphatemia developed in five of nine donors (55.6%) and phosphate levels returned to normal in all donors by day 7. Only one donor received replacement therapy within the first week (donor 8). As discussed above, donor 9 had normal phosphate levels during the first week but developed profound hypophosphatemia (0.32 mmol/L) on day 14 when he presented with an infected bile collection and 5 anerias bacteriums.

METHODS
The authors examined the first nine living donors in this adult-to-adult living donor liver transplantation program between June 2000 and July 2001. All donors were emotionally related to the recipients and all donors underwent an extensive medical, anatomical and psychosocial evaluation before surgery. All donors underwent right hepatectomy. Because serum phosphate levels were not routinely measured in all patients, stored serum samples from before the operation (day 0) and postoperative days 1, 3 and 7 were used to determine serum phosphate levels. Serum inorganic phosphate levels were measured by phosphammonium molybdate absorbance on a Hitachi 747-200 instrument (Roche Diagnostics, USA).

Helical CT scans with three-dimensional volumetric renderings were used preoperatively to estimate donor graft weight and degree of hepatectomy. All CT scans were performed on a multidetector helical scanner (LightSpeed, GE Medical Systems, USA). The three-dimensional volumes were calculated using a dedicated workstation (Vital Images, USA). Volumetric CT scans were repeated on postoperative day 7 to estimate the degree of liver regeneration. Estimated percentage liver regeneration was calculated by the following formula:

\[ \frac{\text{Day 7 liver volume} - \text{baseline residual volume following hepatectomy}}{\text{Day 7 liver volume}} \times 100 \]

Data are expressed as mean ± SEM. Correlation between liver resection, degree of liver regeneration and change in phosphate levels were calculated using Excel (Microsoft Corp, USA). Donors gave informed consent for liver donation and to have serum stored for research purposes. The Mayo Clinic Institutional Review Board approved the study.

stored serum samples. Phosphate supplements and TPN were not routinely used in our donors and, thus, the impact of these confounding variables was removed.

The reason for the development of hypophosphatemia after liver resection is unknown but is thought to be due to flux of phosphate into the liver for energy metabolism associated with liver regeneration. To examine this hypothesis, we used volumetric computed tomography (CT) at baseline to estimate the extent of liver resection and, on postoperative day 7, to estimate degree of liver regeneration. We attempted to correlate the extent of hepatectomy and degree of liver regeneration with the change in serum phosphate levels within the first postoperative week.
Complications were noted in six of the donors and are summarized in Table 1. Minor wound infections were noted in two donors. One donor developed a peptic ulcer that responded to therapy with a proton pump inhibitor. Three donors developed bile leaks, with one responding to endoscopic stenting and the other two requiring percutaneous drainage of bile collections.

The results of the volumetric CT scans are shown in Table 2. The estimated average hepatectomy was 63.8% (range 55.5% to 73.4%). Residual left lobe hypertrophy was seen in all donors by day 7 with an average increase in volume of 86.3% (range 17.6% to 209.7%). The percentage of hepatectomy strongly correlated with the degree of liver regeneration by day 7, with a correlation coefficient of 0.74 (Figure 1). The decrease in serum phosphate levels correlated poorly with the extent of hepatectomy (r = 0.25) and the percentage of liver regeneration was not predictive of the percentage decrease in phosphate levels (r = –0.006; Figure 2).

**DISCUSSION**

Hypophosphatemia occurred in two-thirds of our living donors after right hepatectomy. Within the first week following surgery, five of nine donors (55.6%) developed hypophosphatemia. The hypophosphatemia was moderate (0.81 mmol/L to 0.52 mmol/L) in two donors and severe (0.48 mmol/L to 0.32 mmol/L) in another three donors. One other donor developed hypophosphatemia two weeks after surgery but had normal phosphate levels during the first postoperative week.

Our results differ from two previously published studies. Pomposelli et al (4) reported that hypophosphatemia was universal in all 18 of their living donors following right hepatectomy. However, all donors in their series received TPN. TPN in itself has been reported to cause hypophosphatemia (5) and, therefore, TPN may have contributed to the hypophosphatemia in their living donors. Even when the last 12 donors in their series were given aggressive phosphate supplementation (60 mmol/day started on postoperative day 1 as part of their TPN), severe hypophosphatemia was still seen in 30% of donors and profound hypophosphatemia was seen in 8% (4). In contrast, a study by Tan et al (6) of 95 right lobe donors did not find hypophosphatemia to be a major problem. Most donors developed mild hypophosphatemia but profound hypophosphatemia (less than 0.32 mmol/L) was not observed.

**TABLE 2**

<table>
<thead>
<tr>
<th>Donor</th>
<th>Estimated total liver volume (g)</th>
<th>Estimated right graft volume (g)</th>
<th>Percentage hepatectomy</th>
<th>Estimated liver volume (g)</th>
<th>Percentage regeneration*</th>
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<tr>
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<td>1259</td>
<td>709</td>
<td>56.3</td>
<td>647</td>
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<td>209.7</td>
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<tr>
<td>Mean ± SEM</td>
<td>1674±123</td>
<td>1072±85</td>
<td>63.8±1.9</td>
<td>964±75</td>
<td>86.3±21.4</td>
</tr>
</tbody>
</table>

*Percentage regeneration was calculated by subtracting the estimated liver volume on postoperative day 7 from the baseline residual volume of liver following hepatectomy, dividing by the estimated liver volume on postoperative day 7 and multiplying by 100.

**Figure 1** Percentage of hepatectomy (estimated by baseline volumetric computed tomography scan) was strongly correlated with percentage of liver regeneration (estimated by volumetric computed tomography scan on postoperative day 7) in nine living donors undergoing right hepatectomy.

**Figure 2** Percentage of liver regeneration by the seventh postoperative day (estimated using volumetric computed tomography scan) was not predictive of the percentage decrease in serum phosphate levels from baseline within the first week following right hepatectomy in nine living donors.
and subsequent tissue hypoxia. Serum phosphate levels less than
phosphate diminishes levels of 2,3-diphosphoglycerate in ery-
through several potential mechanisms (12). Severe hypophos-
hepatic failure was seen in one patient with profound
(11) and, in the series by George and Shiu (1), death due to
neuropraxia in a live liver donor with associated hypophos-
plications include an altered level of consciousness, seizures,
(9). Pomposelli et al (4) described a live liver donor, with no
plications include arrhythmias (1) and myocardial suppression
in our series of cancer patients. They correlated profound hypophos-
phatemia with increased infectious, cardiac and respiratory
postsurgical complications. In our small series, there were six
postsurgical complications: two wound infections, one peptic
ulcer and three bile leaks. Severe or profound hypophos-
phatemia was seen in two of the three donors with bile leaks.
Overall, one-half of donors who developed severe or profound
hypophosphatemia had a complication. However, two of the
three patients who did not develop hypophosphatemia also
had postoperative complications. Therefore, based on our
small sample size, it is difficult to correlate the risk of compli-
cations with the presence of hypophosphatemia. We also can-
not answer the question of whether it is the partial
hepatectomy or the complication (bile leak and infection), or
the combination of the two, that led to the hypophos-
phatemia.

Hypophosphatemia has been associated with infectious,
cardiovascular, respiratory and neurological complications (7).
Hypophosphatemia may increase the risk of sepsis by impairing
leukocyte and macrophage function (8). Cardiovascular com-
plications include arrhythmias (1) and myocardial suppression
(9). Pomposelli et al (4) described a live liver donor, with no
previous history of cardiac disease, who developed pulmonary
edema during the phosphate nadir. Respiratory muscle dys-
function is a recognized complication of hypophosphatemia
(10), and respiratory arrest has complicated hypophosph-
phatemia following hepatic resection (1). Neurological com-
plications include an altered level of consciousness, seizures,
tremor and paresthesia, including a case of temporary radial
neuropaxia in a live liver donor with associated hypophos-
phatemia (4).

Hepatic failure has been attributed to hypophosphatemia
(11) and, in the series by George and Shiu (1), death due to
hepatic failure was seen in one patient with profound
hypophosphatemia.

Hypophosphatemia may lead to cell or organ dysfunction
through several potential mechanisms (12). Severe hypophos-
phatemia diminishes levels of 2,3-diphosphoglycerate in ery-
throcytes, resulting in increased oxygen affinity for hemoglobin
and subsequent tissue hypoxia. Serum phosphate levels less than
0.16 mmol/L are associated with impaired glycolysis. In addition,
severe hypophosphatemia is associated with depletion of cellular
ATP, the energy source for most cellular functions. Furthermore,
inorganic phosphate is an independent regulator of cardiac
adenosine formation and thus, potentially, cardiac contractility.
In skeletal muscle, hypophosphatemia is associated with
impaired membrane functional and structural integrity.

Hypophosphatemia is most likely to occur during the first
week following right hepatectomy. In our series, serum phos-
phate levels had returned to normal by day 7 in all donors. The
other two reports found that serum phosphate levels returned
to normal in all donors by day 5 (4,6). Interestingly, one of our
donors (donor 9) had normal serum phosphate levels during
the first week after surgery but presented with profound
hypophosphatemia two weeks following his liver resection. At
that time, he had an infected bile collection and Staphylococcus
septicemia. His significant hypoxemia (oxygen saturation
80%) was likely a result of his profound hypophosphatemia
(0.32 mmol/L). Because hypophosphatemia decreases produc-
tion of 2,3-diphosphoglycerate within erythrocytes, a shift in
the oxygen saturation curve may result in hypoxemia and tis-
tue hypoxia. In fact, in the series by George and Shiu (1), pro-
found hypophosphatemia was the cause of respiratory arrest in
one patient following liver resection. Our donor (donor 9)
demonstrates that hypophosphatemia can be a relatively late
development following liver resection, especially if there are
associated complications. Because hypophosphatemia is
thought to suppress immune function (1), one could argue that
he was at increased risk for septicemia because of low serum
phosphate levels. However, this donor had normal serum phos-
pate levels during the first week following surgery despite sig-
nificant liver regeneration (estimated at 209.7% on his day 7
CT scan). It is more likely that the additional metabolic stress
of septicemia, in the setting of rapid liver regeneration, resulted
in the late development of profound hypophosphatemia in this
donor.

The reasons for the development of hypophosphatemia
following right hepatectomy are poorly understood. The pro-
posed mechanism of hypophosphatemia is the movement of
phosphorous into a rapidly regenerating liver for the protein
synthesis and phosphorylation of nucleotides (ATP and
guanosine triphosphate). In our series, an average of 86.3%
liver regeneration was seen by the seventh day following sur-
gery. Interestingly, the degree of hepatectomy estimated on
the baseline CT scan was strongly correlated with the degree
of regeneration noted after the first week (r=0.74). In other
words, the larger the hepatectomy, the faster the liver regen-
eration. However, there was only a weak correlation between
the percentage hepatectomy and the percentage decrease in
serum phosphate levels within the first week. Furthermore,
the percentage of liver regeneration on day 7 was not corre-
lated with the percentage decrease in serum phosphate levels.
Therefore, reasons other than rapid liver regeneration may
account for the development of hypophosphatemia in these
donors.

Interestingly, hypophosphatemia was also seen in six of the
nine recipients of the right hepatic allografts within the first
week following liver transplantation. In the recipients,
hypophosphatemia was moderate in five patients and severe in
one patient. None of the liver transplantation recipients
received TPN during this time period. Significant liver regen-
eration was also seen in the recipient allograft after one week
(29.8±4.7%), suggesting that liver regeneration may con-
tribute to the hypophosphatemia in recipients of right liver
allografts.
CONCLUSIONS

Hypophosphatemia is very common but not universal in donors undergoing right hepatectomy for adult-to-adult liver transplantation. Hypophosphatemia may be associated with serious donor complications. Hypophosphatemia occurs most commonly during the first week following surgery. However, donors with later complications may also be at risk for hypophosphatemia. Although rapid liver regeneration is thought to be the cause of hypophosphatemia in these donors, we did not find a correlation between liver regeneration and a decrease in serum phosphate levels. All living donors should undergo monitoring for hypophosphatemia during their first postoperative week and the role of prophylactic supplementation of phosphorous in these patients deserves further study. Serum phosphorous levels should be checked in all donors with complications that occur within the first few weeks after donor hepatectomy.

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REFERENCES
