Erythropoiesis and renal transplant pregnancy


Abstract: Objective. To examine erythropoiesis in renal transplant pregnancies.

Methods. Retrospective cohort study of 30 renal transplant cases and 30 age, smoking and parity-matched healthy controls with normal index pregnancy. Retrospective chart review and assay of frozen antenatal serum (for serum erythropoietin concentration [serum EPO], transferrin receptor protein [TfR], ferritin, folate and B12) were performed. The linear regression equation for normal pregnancy controls was used to calculate predicted [serum EPO] and the observed/predicted (O/P) log [serum EPO] was plotted. The relationship between [serum EPO] and haemoglobin (Hb) among transplant cases was considered to be different from that among controls if the slope of the O/P log [serum EPO] versus Hb regression was significantly different from zero.

Results. The transplant (14 cadaveric) to conception interval was (median [range]) 33.5 [4, 189] months. Immunosuppressants were azathioprine (n = 25), cyclosporine (n = 22) and/or prednisone (n = 25). Cases were more often primiparous (20 vs. 7 [controls]; p = 0.01), had pre-existent hypertension (20 vs. 0 [controls]; p < 0.001), developed new/increased hypertension or pre-eclampsia (28 vs. 0 [controls]; p < 0.001) and an antenatal rise in creatinine (14 vs. 2 [controls]; p < 0.001). In early pregnancy, cases had similar EPO (15.2 [2.6, 84.6] vs. 15.7 [6.4, 41.0] U/L) but lower Hb (101 [65, 129] vs. 116 [106, 150] g/L; p < 0.001). Twenty-two (73%) cases had Hb < 100 g/L (vs. 4 [controls]; p < 0.001); Hb was comparable at 6 wk postpartum. With advancing gestational age (GA), Hb remained stable and serum EPO increased in both groups. The slope of the O/P log [serum EPO] versus Hb for transplant cases was significantly different from zero within both the 17–28 wk (slope ± SEM: 0.010 ± 0.002; p < 0.0001) and the 29–42 wk GA categories (0.006 ± 0.003; p = 0.02). Cases showed smaller rises in serum TfR (change 481 [−1471, 2780] vs. 1119 [−698, 4195] [controls] ng/mL; p = 0.005).

Conclusions. Anaemia frequently complicates renal transplant pregnancies, in which serum EPO is inappropriately low and the rate of erythropoiesis blunted.

The World Health Organization defines anaemia in pregnancy as a haemoglobin (Hb) concentration less than 110 g/L (1), although many clinicians use a cut-off of 100 g/L. Dilutional anaemia occurs in normal pregnancy because plasma volume increases to a greater degree than red blood cell mass (2).

As in non-pregnancy, anaemia in pregnancy may be caused by factors related to abnormal or decreased production and/or increased destruction.
of red blood cells. The oxygen sensor in the kidney controls production of erythropoietin (EPO), a glycoprotein growth factor that stimulates differentiation of erythroid progenitor cells in the bone marrow. There is an inverse relationship between serum EPO and Hb concentrations (3); in the presence of anaemia, EPO levels rise exponentially in both non-pregnancy (4) and normal pregnancy (5).

The John Radcliffe High Risk Obstetric Unit manages all renal transplant pregnancies within the Oxford Region. Progressive anaemia during pregnancy was observed in 3 women with stable creatinine concentrations. However, in a comprehensive literature review (6) of pregnancy outcome following renal transplantation (based on 2409 women and 3382 pregnancies from 1961 to 1994), anaemia was not mentioned as a problem.

The objective of this retrospective study was to examine, among gravidae with renal transplants, the prevalence of anaemia and the appropriateness of EPO production (as reflected by the relationship between antenatal serum EPO and Hb). Comparisons were made with normal pregnancy controls, all of whom were managed in the same High Risk Obstetric Unit.

Methods

All women with pregnancies associated with renal transplantation and followed by the High Risk Obstetric Unit at the John Radcliffe Hospital, Oxford, between 1986 and 1996, were identified by a computerised search of the Unit database. ‘High risk’ normal pregnancy controls, with a history of either idiopathic recurrent miscarriage or foetal loss or pre-eclampsia in previous pregnancies, were also identified by a search of the same database. Controls were selected based on the following criteria: no history of a haematological disorder, antenatal BP < 140/90 mmHg, no antenatal proteinuria or known renal disease, term delivery and birthweight appropriate for gestational age (GA). These ‘high risk’ normal pregnancy controls were matched to transplant cases according to age (± 4 yr), smoking (yes/no) and parity (± 1).

In order to examine the ‘normality’ of normal pregnancy controls who had been referred to a High Risk Obstetrics Unit, ‘general antenatal’ pregnancy controls were also sought (prospectively) from general antenatal clinics at the same institution. They were selected and matched to cases according to the same criteria as for ‘high risk’ normal pregnancy controls. Similar results would justify use of ‘high risk’ normal pregnancy controls who were recruited and managed in the same way as cases.

For transplant cases and ‘high risk’ normal pregnancy controls, standardised forms were used to collect the following background medical data by retrospective hospital chart review: maternal demographics (e.g. age, parity), history of renal disease, presence of pre-existent hypertension, baseline (pre-pregnancy) creatinine, other medical history, obstetrical history and immunosuppressant and other drug therapy. For ‘general antenatal’ normal pregnancy controls, the same information was collected by prospective chart review.

Collection and freezing of serial antenatal and postnatal serum have been routine in the Unit since the mid-1980s. Routine collection periods have been as follows: ≤ 17, 17–22, 23–28, 29–34 and 35–40 wk gestation, as well as both 2–5 days and 6 wk postpartum. In addition to the GA at which the frozen serum was taken, the following laboratory results (corresponding to the same serum sample) were abstracted from hospital records: Hb (g/L) (as Hct was not available for most patients), mean cell volume (MCV) of red cells, platelet count (× 10⁹/L), creatinine (µM), uric acid (µM), dipstick proteinuria greater than trace, mid-stream urine (MSU) culture result (growth/no growth; organism if relevant) and 24-h urinary protein (g). Blood film and reticulocyte counts were not available. There was insufficient information on creatinine clearance to be useful and cyclosporine A (CSA) measurements could not be used as they were not timed relative to ingestion of the drug.

Serum samples that had been stored at −20°C were allowed to defrost, for this first time, at room temperature. All assays, except serum transferrin receptor protein (TfR), were performed and the samples were refrozen at −20°C and shipped to Belgium where they were similarly defrosted, assayed for TfR and refrozen. No samples were lost. All samples were assayed for serum EPO, using a chemiluminescence immunoassay (Nichols Institute Diagnostic Ltd., Essex, UK) with results expressed in U/L; the intra- and inter-assay coefficients of variation (CV) were 5.5 and 6.7%, respectively. Only the first and last antenatal serum samples were assayed for ferritin, folate, vitamin B₁₂ and TfR. Soluble TfR are closely related to the number of red blood cell precursors in the bone marrow and provide a surrogate marker for the rate of erythropoiesis, which is reliable in the absence of severe iron deficiency (which elevates TfR values) (7); soluble TfR levels are particularly useful for detection of multiple mechanisms of anaemia as can exist among pregnant patients with renal transplantation. Serum ferritin levels were
assayed using a two-site incubation immunoradiometric assay (Bio-Rad Laboratories Ltd., Herts, UK) expressed in μg/L; the intra- and inter-assay CV were 4.2 and 11.1%, respectively. Serum folate levels were determined using a modification of the chloramphenicol-resistant Lactobacillus caseii method (8) and expressed in μg/L; the intra- and inter-assay CV were 4.0 and 11.5%, respectively. Vitamin B12 was assayed using a non-boil radioimmunoassay kit (Ortho-Clinical Diagnostics, Amersham, Buckinghamshire, UK), with the results expressed in ng/L; the intra- and inter-assay CV were 4.9 and 13.1%, respectively. In accordance with standard procedure, each assay included both internal and external control samples. TfR was assayed using an ELISA (9, 10) in which each sample was run in duplicate; the intra- and inter-assay CV were 4.3 and 8.1%, respectively.

The following outcomes were abstracted from hospital records: pregnancy outcome (i.e. miscarriage, elective termination, intrauterine foetal death, live birth, neonatal death), major malformations (defined as those having a major impact on structure or function (11)), development of pre-eclampsia (defined as BP ≥ 140/90 mmHg, ≥ 0.3 g of proteinuria by 24-h collection and/or oedema (12)), new antenatal (isolated) hypertension (i.e. BP ≥ 140/90 mmHg), worsening of pre-existent hypertension (to ≥ 140/90 mmHg) without proteinuria, maternal pregnancy complications (e.g. urinary tract infection, renal allograft rejection), rise in antenatal and postnatal creatinine by 15% (from the first antenatal sample, to a value > 80 μM (13)), GA at delivery (wk), prematurity (i.e. < 37 wk gestation), birthweight (BW) (g), small for gestational age (SGA) infants (i.e. < 5th percentile), method of delivery, neonatal health problems (defined as those which prolonged hospital stay) and long-term renal outcome (e.g. renal allograft rejection or graft loss).

The primary outcome of interest was the relationship between serum EPO and Hb. However, a two-way analysis of variance with repeated measures was first performed on one factor (GA) to assess the relationship of serum EPO or Hb, with both patient group and GA. Given that measurements of EPO and Hb were made in time intervals that were not equal for all women, an unequally spaced repeated measurements design was used in which the spatial power law modelled the time series covariance structure of the repeated measurements.

A detectable relationship between GA and either serum EPO or Hb prompted examination of the relationship between them within GA categories. Within each GA category, the equation from the linear regression of log [serum EPO] versus Hb (5) among normal pregnancy controls was used to calculate ‘predicted’ values of log [serum EPO] from the Hb values of transplant cases. For transplant cases, a linear regression was performed of the observed over predicted (O/P) values of log [serum EPO] versus Hb. A slope that was significantly different from zero was taken to reflect a different relationship between log [serum EPO] and Hb than that seen among normal pregnancy controls. p < 0.05 was considered statistically significant for all analyses. Analyses were performed using SAS for IBM-compatible computers or GraphPad Prism 2.01 (GraphPad Software, San Diego, CA).

Of secondary interest were changes, from first to last antenatal sample, in: i) platelets, as a marker of the adequacy of other cell lines, ii) MCV, iii) serum ferritin, iv) serum folate, v) serum B12, vi) serum TfR and vii) serum creatinine. Obstetric outcomes and maternal medical complications were also compared between groups. Between-group comparisons were made by Student’s t-test or Mann–Whitney U-test (for continuous data) and χ² or Fisher’s exact test (for categorical data).

Results

Baseline demographics

Twenty-four renal transplant cases, with 30 pregnancies, were identified and included in the study. The indications for renal transplantation were reflux nephropathy (n = 11, 1/11 nephrectomy of native kidneys), vasculitis including systemic lupus erythematosus (n = 5), chronic glomerulonephritis (n = 4), familial nephrosis (n = 3), congenital malformations (n = 2), focal segmental glomerulosclerosis (n = 2), polycystic kidney disease (n = 1) and unknown (n = 2). The transplant to conception interval was (median [range]) 33.5 [4, 189] months. Cadaveric (14 [47%]) and living related donor (15 [50%]) transplants were equally common; the nature of one transplant was unknown. Only 6 (20%) transplants had been followed by an episode of acute rejection within the first 3 months post-operatively. Immunosuppressants were most commonly azathioprine, CsA and prednisone (n = 13), followed in frequency by azathioprine and prednisone (n = 7), CsA and prednisone (n = 5), azathioprine and CsA (n = 4) and azathioprine alone (n = 1).

Thirty ‘high risk’ normal pregnancy controls, with a history of either pre-eclampsia in a previous pregnancy (n = 17) or unexplained recurrent miscarriage (n = 10), stillbirth (n = 2) or neonatal death (n = 1) were identified. Their Hb, serum EPO, platelet counts, serum ferritin, serum folate,
serum B12 and TfR were found not to differ from those of the 16 ‘general antenatal’ normal pregnancy controls (data not presented). Therefore, only ‘high risk’ normal pregnancy controls were compared with renal transplant cases for the remainder of the analyses.

Table 1 shows that there were no differences between renal transplant cases or ‘high risk’ normal pregnancy controls in terms of maternal age at delivery, ethnicity, smoking or GA at booking. Cases were more likely to be primiparous (whereas controls were more likely to have had one previous pregnancy) and have pre-existent hypertension. Most (17/20) with hypertension had been treated with antihypertensive medication before pregnancy, usually with a beta-blocker (13/17). One case was on captopril, which was discontinued when pregnancy was diagnosed. Cases also had higher baseline serum creatinine and serum uric acid and there was a trend toward higher 24-h urinary protein. Three women had baseline serum creatinine greater than 125 \( \mu \text{M} \) (i.e. 146, 167 and 185 \( \mu \text{M} \)).

Table 2 shows that at baseline, renal transplant cases also had lower Hb concentrations than normal pregnancy controls. The GA (mean ± SD) at which the first sample was stored was not significantly different between transplant cases and ‘high risk’ normal pregnancy controls. Cases had higher MCV and serum ferritin, but similar serum folate, B12, EPO and TfR values. Platelet counts did not differ between groups. Equivalent numbers of transplant cases and normal pregnancy controls took iron supplements during the pregnancy (i.e. 21/30 vs. 16/30 [controls]; \( p = 0.18 \)). No cases or controls reported bleeding complications.

### Table 1. Baseline characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Reference range(^a)</th>
<th>Renal transplant pregnancies ((n = 30))</th>
<th>‘High risk’ normal pregnancies ((n = 30))</th>
<th>( p ^b )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal age at EDD (yr) (mean ± SD)</td>
<td>n/a</td>
<td>26.7 ± 5.0</td>
<td>28.7 ± 5.0</td>
<td>0.12</td>
</tr>
<tr>
<td>Caucasian (%)</td>
<td>n/a</td>
<td>29 (97)</td>
<td>30 (100)</td>
<td>0.31</td>
</tr>
<tr>
<td>Primiparous (%)</td>
<td>n/a</td>
<td>20 (67)</td>
<td>7 (24)(^c)</td>
<td>1.00</td>
</tr>
<tr>
<td>Nonsmokers (%)</td>
<td>n/a</td>
<td>24 (80)</td>
<td>24 (80)</td>
<td>0.74</td>
</tr>
<tr>
<td>GA at booking (wk) (median [range])</td>
<td>n/a</td>
<td>10.5 [6, 34]</td>
<td>11.0 [5, 27]</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Pre-existent hypertension (%)</td>
<td>n/a</td>
<td>20 [67]</td>
<td>0 (0)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Serum creatinine (( \mu \text{M} )) (median [range])</td>
<td>50–100 ( \mu \text{M} )</td>
<td>100.5 [66.0, 185.0]</td>
<td>58.0 [43.0, 79.0]</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Serum uric acid (( \mu \text{M} )) (median [range])</td>
<td>120–420 ( \mu \text{M} )</td>
<td>337.0 [171.0, 609.0]</td>
<td>199.0 [122.0, 320.0]</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>24-h urinary protein (g) (median [range])</td>
<td>&lt;0.3 g/24 h(^d)</td>
<td>0.1 [0.1, 0.4]</td>
<td>0.1 [0.1, 0.2]</td>
<td>0.052</td>
</tr>
</tbody>
</table>

\( ^a \) Range for non-pregnancy unless otherwise stated.
\( ^b \) Values in bold are statistically significant at \( p < 0.05 \).
\( ^c \) Data missing for one pregnancy.
\( ^d \) Normal range for pregnancy; lower limit of the assay is 100 mg per 24 h.

EDD (expected date of delivery), GA (gestational age).

### Erythropoiesis

Fig. 1 shows that from baseline throughout pregnancy, the Hb of renal transplant cases remained significantly lower than that of ‘high risk’ normal pregnancy controls (\( p = 0.0001 \)). At some point in gestation, Hb was less than 100 g/L among 22 \((74\%)\) transplant cases (vs. 4 \([13\%]\) controls) (\( p < 0.0001 \)). There were few serum samples available before 17 wk GA; therefore, postnatal Hb values were examined as surrogates for antenatal values. Postnatal Hb values were available for 20 of the 22 transplant cases with a antenatal Hb less than 100 g/L; only 1 had a Hb less than 100 g/L, compared with no controls (\( p = 0.40 \)). No cases or controls were transfused postnatally.

Fig. 2 shows that among both transplant cases and ‘high risk’ normal pregnancy controls, serum EPO concentration increased with advancing GA (\( p = 0.0003 \)), but there was no difference between groups (\( p = 0.75 \)). In order to account for the GA effect on serum EPO, the relationship between serum EPO and Hb was examined within three GA categories: \(< 17, 17–28\) and \(> 29\) wk gestation to reflect early, mid- and late pregnancy.

Using data from ‘high risk’ normal pregnancy controls, regression equations were obtained for log [serum EPO] versus Hb for both the 17–28 wk (2.95–[0.14 × Hb]; \( n = 46 \) samples) and 29–42 wk categories (2.23–[–0.07 × Hb]; \( n = 52 \) samples). The analysis was not possible for early pregnancy given the few available samples. Fig. 3a and b shows that the O/P ratio of log [serum EPO] for transplant cases differed significantly from zero, for both the 17–28 wk (slope ± SEM: 0.010 ± 0.002; \( r^2 = 0.30; p < 0.0001 \)) and 29–42 wk categories (0.006 ± 0.003; \( r^2 = 0.08; p = 0.02 \)). This indicated a relationship between log [serum
EPO] and Hb that was different from that seen among normal pregnancy controls, for both GA categories.

One renal transplant case received two units of packed red blood cells for a Hb of 59 g/L and 4 wk later, went on to receive exogenous EPO with good effect. Prior to EPO administration at 19 wk gestation, her serum EPO and Hb were 7.8 U/L and 60 g/L, respectively. After exogenous EPO administration, her serum EPO and Hb peaked at 21.4 U/L and 92 g/L by 28 wk gestation. Her data were not influential in the analyses. Administration of EPO was not associated with poorly controlled hypertension, although the patient did go on to develop increasing hypertension and proteinuria (maximum 0.98 g/d). She was induced at 36 wk gestation and had a vaginal delivery of a infant weighing 2724 g (50th percentile for GA). He had Apgars of 3, 6 and 9 at 1, 5 and 10 min, respectively and required intubation. There was no neonatal polycythemia.

Only the change (from the first to last antenatal samples) in TtR, platelets and creatinine differed between groups; no differences were seen in the change in serum ferritin, serum folate, serum B12 or MCV (data not presented). Cases showed a smaller rise in TtR levels (median change [range] of 481 [−1471, 2780] vs. 1119 [−698, 4195] controls U/L; p = 0.005), as well as larger rises in platelet count (21 [−96, 202] vs. −12 [−102, 152] [controls] × 10^9/L; p = 0.003) and creatinine concentration (12.38 [−7.07, 6.19] vs. 1.77 [−14.14, 30.94] [controls] μM; p = 0.0002).

Pregnancy and maternal outcomes

Table 3 shows that pregnancy outcome was favourable for both groups. Among renal transplant pregnancies, there were two intrauterine foetal deaths (at 29 and 35 wk gestation); prematurity, SGA infants and neonatal health problems (i.e. special care baby unit admission due to prematurity [n = 8] and sepsis [n = 1]) were also more frequent. Prematurity among transplant pregnancies was most commonly iatrogenic (n = 11) and most commonly for pre-eclampsia (n = 7). Four women with renal transplants had two or three pregnancies, which did not appear to be more complicated than the first.

Table 3 also shows that transplant cases suffered an excess of medical complications such as pregnancy hypertension, urinary tract infections or asymptomatic bacteruria, rise in antenatal serum creatinine concentration and increasing proteinuria. Twenty transplant cases developed new/worsening hypertension; overall, 19 (63%) developed hypertension and proteinuria that were thought to be indicative of pre-eclampsia. Just over half of renal transplant cases developed proteinuria, with a peak (median [range]) value of 500 [200, 5900] mg/24 h. Most transplant cases who developed antenatal urinary tract infections or asymptomatic bacteruria did so recurrently, i.e. on two (8 vs. 1 [control]), three (1 vs. 0 [control]), four (1 vs. 0

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Table 2. Baseline laboratory data

<table>
<thead>
<tr>
<th>Reference range for non-pregnancy</th>
<th>Renal transplant pregnancies (n = 30) at 16.6 ± 3.9 wk</th>
<th>‘High risk’ normal pregnancies (n = 30) at 18.9 ± 3.9 wk</th>
<th>p(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hb (g/L) (median [range])</td>
<td>101 [65, 129]</td>
<td>116 [106, 150]</td>
<td>0.004</td>
</tr>
<tr>
<td>MCV (μM) (median [range])</td>
<td>93.3 [85.4, 107.6]</td>
<td>89.7 [80.9, 99.4]</td>
<td>0.005</td>
</tr>
<tr>
<td>Platelet count (× 10^10/L) (median [range])</td>
<td>150–450×10^9/L</td>
<td>258.0 [106, 345]</td>
<td>0.79</td>
</tr>
<tr>
<td>Serum ferritin (μg/L) (median [range])</td>
<td>14–200 μg/L</td>
<td>38.0 [10.0, 287.0]</td>
<td>0.004</td>
</tr>
<tr>
<td>Serum folate (μg/L) (median [range])</td>
<td>2.1–28.0 μg/L</td>
<td>8.2 [4.4, 18.0]</td>
<td>0.89</td>
</tr>
<tr>
<td>Serum B12 (ng/mL) (median [range])</td>
<td>211.0 [76.0, 457.0]</td>
<td>204.0 [130.0, 472.0]</td>
<td>0.70</td>
</tr>
<tr>
<td>Serum erythropoietin (U/L) (median [range])</td>
<td>2–17.0 U/L</td>
<td>15.2 [2.6, 94.6]</td>
<td>1.00</td>
</tr>
<tr>
<td>TtR (ng/mL) (median [range])</td>
<td>2 900−7 100 ng/mL</td>
<td>3 439.0 [1 658.0, 5 912.0]</td>
<td>0.12</td>
</tr>
</tbody>
</table>

\(^a\)Values in bold are statistically significant at p < 0.05.
\(^b\)As per the World Health Organization.

Hb (haemoglobin), MCV (mean red cell volume), TtR (transferrin receptor protein).

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**Fig. 1.** Gestational age versus Hb (mean ± SD) for renal transplant cases (Tt) and normal pregnancy controls (NPC). Both antenatal, as well as postpartum day 3–5 and wk 6 are presented.
Impaired erythropoiesis among women with renal transplantation in pregnancy may result from a number of factors. There was no evidence that transplant cases were deficient in iron, folate or vitamin B12. In fact, serum ferritin was actually higher among transplant cases. Despite this, they went on (inappropriately) to receive supplemental iron in most cases. There was no global bone marrow dysfunction, as indicated by similar platelet counts at baseline and actually higher platelet counts among transplant cases before delivery; this occurred despite a higher incidence of proteinuric hypertension. A toxic effect of azathioprine therapy on erythropoiesis must be considered (15). Most transplant cases (25 [83%]) were on azathioprine and had higher MCV values throughout pregnancy, consistent with azathioprine-induced bone marrow toxicity. However, azathioprine should not impair renal release of EPO. CsA, which may decrease renal production of EPO, was taken by 22/30 transplant cases (16). Although renal transplant patients may be more plasma volume expanded during pregnancy, explaining both lower Hb and EPO values, the relationship between the two should not have changed. There was no evidence of haemolysis or excessive blood loss to account for anaemia among transplant cases.

Impaired renal function must be considered as an aetiological factor in the demonstrated impaired erythropoiesis among renal transplant cases. Support for the importance of renal graft function (17) and inadequate production of EPO in the pathogenesis of anaemia in renal transplant pregnancies comes from the following data: i) a slightly elevated but inappropriately low serum EPO in 1 such patient (18), ii) a gradual increase in EPO concentration during normal pregnancy (6), iii) higher exogenous EPO requirements by some haemodialysis patients during pregnancy (19) and iv) the response of anaemic transplant patients to exogenous EPO (5, 20). However, there are 3 reported cases of transplant patients with stable creatinine concentrations who required exogenous EPO during pregnancy (17, 20). Some but not all of these women were on azathioprine, which could contribute to anaemia through bone marrow suppression.

Discussion

The history of renal transplantation and pregnancy dates back 30 years. At present, about 1/50 women of child-bearing age with a functioning renal allograft become pregnant. These pregnancies are recognised to be more complicated, both obstetrically and medically (6). This single centre retrospective cohort of women, with predominantly mild renal impairment and hypertension, experienced rates of obstetric and medical complications comparable to those described in the literature (6). However, what was also demonstrated in this study, and not previously shown, was the fact that these pregnancies are also frequently complicated by anaemia.

Hb values less than 100 g/L occurred among most (74%) renal transplant pregnancies, but among few (13%) normal pregnancy controls. Postnatal anaemia was not a problem among either transplant cases or controls. When assayed between 17 and 42 wk gestation and compared with normal pregnancy controls, transplant cases were shown to have serum EPO levels that were inappropriately low for concomitant Hb values; although 1 transplant case received exogenous EPO, the latter would not change the relationship between serum EPO and Hb (14) and her data were not influential. The rate of erythropoiesis, as reflected by serum TIR levels, was also blunted among renal transplant cases given that TIR levels failed to rise as much as those among normal pregnancy controls, among whom this rise was consistent with the erythropoiesis of normal pregnancy.

Of note, haematuria developed among most renal transplant cases and more frequently than among 'high risk' normal pregnancy controls; none of the women had a concurrent urinary tract infection or identifiable renal stone disease and in no instance were red or white blood cell casts identified. Graft loss occurred among 4 renal transplant cases (13%), due to a failed transplant artery repair (at 5 months postnatally) or rejection (n = 3; at 6, 21 and 60 months postpartum).

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Not surprisingly, transplant cases in this study were shown to have higher serum creatinine concentrations (at baseline and before delivery), 24-h urinary protein excretion and a rise in antenatal serum creatinine by more than 15%. However, most cases had only a small rise in serum creatinine during pregnancy, with only 3 having a creatinine > 200 μM prior to delivery. Anaemia is not usually seen at the level of renal function seen among transplant cases in this study.

The finding of unexplained haematuria, which was confirmed by urine microscopy and seen among both (the majority of) renal transplant cases and (17% of) normal pregnancy controls, was an unexpected finding and remains unexplained. This was not associated with urinary tract infection, renal stone or evidence of rejection.

Although this study is the first to report on erythropoiesis among renal transplant recipients who become pregnant, it has many limitations. It is retrospective in design. Many data were abstracted from hospital records. Neither antenatal nor early pregnancy Hb values were available, such that postnatal values had to be used as surrogates. Serum folate, not RBC folate, could be assayed. Reticulocyte counts and Hct were not available from the hospital charts. Most cases were taking azathioprine, whereas many transplant patients today are on non-azathioprine-containing drug regimes. Most cases were also taking CsA, so it was not possible to examine whether this medication had an impact on erythropoiesis independent of renal function. Finally, we have no information on other growth factors, such as interleukin-3, which also regulate erythropoiesis and may be altered by immunosuppressant therapy.

In summary, this retrospective cohort study from a single tertiary care centre, demonstrates that anaemia frequently complicates the pregnancies of women with renal transplantation. Even in

![Graph](image)

Fig. 3. (a) O/P (observed over predicted) ratio of log {serum erythropoietin concentration [serum EPO]) versus Hb, for transplant cases, for 17–28 wk gestation. Predicted values of log [serum EPO] for transplant cases were calculated from the regression of log [serum EPO] versus Hb for normal pregnancy controls. The slope differs significantly from zero (p < 0.0001). (b) O/P (observed over predicted) ratio of log {serum erythropoietin concentration [serum EPO]) versus Hb, for transplant cases, for 29–42 weeks’ gestation. Predicted values of log [serum EPO] for transplant cases were calculated from the regression of log [serum EPO] versus Hb for normal pregnancy controls. The slope differs significantly from zero (p = 0.02).
Table 3. Pregnancy complications and outcome

<table>
<thead>
<tr>
<th>Obstetric outcomes</th>
<th>Renal transplant pregnancies (n = 30)</th>
<th>‘High risk’ normal pregnancies (n = 30)</th>
<th>p*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pregnancy outcome (%) live births</td>
<td>28 (93)</td>
<td>30 (100)</td>
<td>0.49</td>
</tr>
<tr>
<td>Major malformations (%)b</td>
<td>3 (10)</td>
<td>2 (7)</td>
<td>0.66</td>
</tr>
<tr>
<td>Gestational age at delivery (wk) [median [range]]</td>
<td>36.0 [29.0, 39.0]</td>
<td>39.0 [37.0, 41.0]</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Pre-eclampsia (%)</td>
<td>17 (57)</td>
<td>1 (3)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Birthweight (g) (median [range])</td>
<td>2 526 [550, 3 540]</td>
<td>3 174 [2 660, 4 047]</td>
<td>&lt;0.009</td>
</tr>
<tr>
<td>Gestational age at delivery (wk) [median [range]]</td>
<td>17 (57)</td>
<td>10 (33)</td>
<td>0.12</td>
</tr>
<tr>
<td>Caesarean section (%)</td>
<td>12 (40)</td>
<td>9 (30)</td>
<td>0.59</td>
</tr>
<tr>
<td>Neonatal health problems (%)</td>
<td>9 (30)</td>
<td>1 (3)</td>
<td>0.01</td>
</tr>
<tr>
<td>Maternal outcomes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pregnancy hypertensionc (%)</td>
<td>28 (93)</td>
<td>0 (0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>AN positive urine culture (%)</td>
<td>19 (63)</td>
<td>3 (10)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Rise in AN creatinine by 15%d (%)</td>
<td>14 (47)</td>
<td>2 (7)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Rise in PN creatinine by 15% (%)</td>
<td>16 (53)</td>
<td>17 (71)</td>
<td>1.0</td>
</tr>
<tr>
<td>24-h urinary protein &gt;0.3 g (%)</td>
<td>16 (53)</td>
<td>0 (0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Haematuria (%)</td>
<td>20 (67)</td>
<td>5 (17)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

a Values in bold are statistically significant at p < 0.05.
b Malformations among renal transplant cases were reflux nephropathy and hypospadias (n = 2) and among controls were bilateral hydroceles and polydactyly of the foot.
c Defined as either pre-eclampsia or worsened pre-existent or new hypertension.
d Rise in creatinine was defined by a 15% increase, from the first antenatal measurement, to >80 μM.

AN (antenatal), PN (postnatal).

the absence of serious deterioration in renal function, the transplant kidney may be unable to increase production of EPO and stimulate erythropoiesis sufficiently to meet the increased demands of pregnancy. If treatable causes of anaemia have been ruled out, then consideration should be given to treatment with exogenous EPO, which does not cross the term placenta in vitro (21).

Acknowledgements

Our sincere thanks to Davina Buckley whose diligence and careful handling of serum samples made this study possible. Thanks also to Tansy Cheston, June Jennings, Vibeke Mannion and the rest of the Silver Star Unit Team for their meticulous record-keeping. Special thanks, as always, to Professor CWG Redman for the foresight to prospectively collect serum samples and patient data which made this study possible and for his considered advice and academic support. LAM was supported by a Duncan Gordon Fellowship of the Hospital for Sick Children Research Foundation and a Detweiler Travelling Fellowship from the Royal College of Physicians and Surgeons of Canada. Funding was provided by the Oxford Regional Health Authority, Janssen-Cilag Ltd., UK and the FNRS, Belgium.

References


