Inhibitor of prostacyclin production in sporadic haemolytic uraemic syndrome

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SUMMARY Prostacyclin (PGI₂) production was diminished when rat aortic rings were incubated with plasma from 5 of 6 patients with the sporadic form of haemolytic uraemic syndrome but was normal in the presence of plasma from 7 patients with the epidemic form of haemolytic uraemic syndrome or from patients with other renal diseases. The reduced PGI₂ production was caused by an unstable inhibitor, extractable into polar lipid solvents, in sporadic haemolytic uraemic plasma. These results suggest that there may be at least 2 different pathogenetic mechanisms in epidemic and sporadic haemolytic uraemic syndrome and that the reduced PGI₂ production observed in the sporadic type is due to an inhibitor of PGI₂ production rather than a deficiency of stimulating factors.

Prostacyclin (PGI₂) deficiency has been implicated in the pathogenesis of the haemolytic uraemic syndrome (HUS) and thrombotic thrombocytopenic purpura (TTP), but there are conflicting views on the pathogenetic mechanisms. Plasma from some patients with HUS shows a reduced capacity to support PGI₂ production by endothelial tissue and this has been ascribed to a deficiency of a factor present in normal plasma which stimulates PGI₁ production. This hypothesis is supported by the observation that fresh plasma infusions induce remission in some patients with HUS and TTP. Other patients, however, have responded instead to plasma exchange, suggesting that removal of an inhibitor of PGI₂ production may be implicated. More recently excessively rapid degradation of PGI₂ in HUS plasma has been observed and has been proposed as a third possible explanation for the PGI₂ deficiency. The interpretation of the importance of PGI₂ deficiency in HUS is further complicated by the fact that it is not a constant feature of the syndrome and that trials of PGI₂ infusions have had only moderate success in reversing the condition.

Although the heterogeneity of patients with HUS and TTP has been recognised, most reports on PGI₂ production have failed to distinguish the different subgroups within the HUS spectrum. Epidemiological studies on HUS in children, however, suggest the existence of at least 2 distinct subgroups in the United Kingdom. An epidemic form occurs mainly in summer, in younger children, usually with an explosive onset after diarrhoea, and has a good prognosis. In contrast a sporadic form occurs in older children, with no seasonal variation, developing insidiously without clear prodromal illness or after an upper respiratory infection, and the prognosis for renal function is poor.

In addition to the childhood epidemic and sporadic forms, HUS is seen in association with pregnancy, with collagen vascular disease, after some infections, and with drug exposure. Clearly, different pathogenetic mechanisms may underlie these different forms of HUS and may account for the disparate results obtained in studies on pathogenesis and in treatment trials in these disorders.

We show that plasma of patients with sporadic HUS (but not epidemic HUS) fails to support PGI₂ production by rat aortic rings and that this abnormality is due to the presence of an inhibitory substance. The results of preliminary studies on the nature of the inhibitor are reported.

Patients

HUS was diagnosed in 13 children with microangiopathic haemolytic anaemia, renal failure, and thrombocytopenia in whom other causes, particularly septicemia, had been excluded (Table). Seven younger children aged 11 months to 7 years (cases 1–7) in whom HUS developed acutely after a diarrhoeal prodrome during a summer epidemic were classified as having epidemic HUS. Five older
children aged 5 years to 13-5 years in whom HUS developed insidiously unassociated with a summer epidemic were considered to have sporadic HUS. Two of this latter group had recurrent disease (cases 8 and 13). One 9 year old girl (case 10) had a microangiopathic haemolytic anaemia and thrombocytopenia, but normal renal function.

To assess the specificity of PGI₂ abnormalities for HUS another group of 9 children aged 10 months to 13 years with various other renal diseases was studied. This group comprised 1 child with acute nephritis, 2 with chronic renal failure due to Henoch-Schonlein nephritis, 1 with steroid sensitive nephrotic syndrome, 1 with focal glomerulosclerosis, 2 with shock and renal failure, and 2 with hypertension due to renovascular disease.

Methods

Nine volumes of venous blood from patients and normal adult controls were each collected into 1 volume of 3-8% trisodium citrate. Platelet poor plasma (PPP) was obtained within 20 minutes of collection by centrifugation at 1300 g for 10 minutes at room temperature and was either tested immediately or stored at −20°C until required.

Platelet rich plasma (PRP) for aggregation studies was prepared from normal adults by centrifugation of citrated blood at 300 g for 15 minutes at room temperature. The final platelet count was adjusted to 250–300 × 10⁶/μl (250 000–300 000/mm³) by dilution with autologous PPP.

Ability to support PGI₂ production. The ability of plasma to support PGI₂ production by rat aortic rings was assessed by a modification of the method of Moncada et al.¹⁸—PGI₂ like activity produced being measured by the inhibitory effect on aggregation of normal human platelets challenged with 2·5 mmol/l adenosine diphosphate (ADP).

Aortic rings (1–2 mm width) were obtained from freshly sacrificed male Wistar rats. After rinsing 3 times in barbitone buffered saline (BBS), pH 7·4, the rings were kept in BBS at 0°C for up to 3 hours until used. In each experiment an aortic ring was incubated at 37°C for 5 minutes with 300 μl PPP from either a patient or a control subject, or with 300 μl BBS. A 100 μl aliquot was then removed and added to 300 μl normal PRP. The mixture was stirred for 1 minute in a Payton aggregometer and ADP was then added to a final concentration of 2·5 μmol/l. In the presence of PPP which had not been incubated with an aortic ring, full aggregation occurred. Inhibition of aggregation was taken to represent the effect of PGI₂ like material secreted from the aortic ring into the plasma tested. Each ring that had been incubated with test plasma was then rinsed in BBS for 30 seconds and reincubated in control plasma (and vice versa) for 5 minutes before retesting, so that each ring acted as its own control.

Detection of an inhibitor. To distinguish between inhibition of PGI₂ production and lack of PGI₂ stimulating factors a second set of cross over experiments were performed in which the production of PGI₂ like activity by each aortic ring incubated in test plasma was compared with that produced by the same ring following incubation in BBS. Diminished PGI₂ production after incubation of aortic rings in HUS plasma compared with PGI₂ production in BBS was taken to indicate the presence of an inhibitor.

Degradation of PGI₂ in HUS plasma. PGI₂ was added to 500 μl PPP from patients in cases 8, 11, and 12 or normal control PPP in final concentrations of 20–100 ng/l. After incubation for 5 minutes at
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37°C, 100 μl PPP was removed and added to 400 μl control PRP. After stirring for 1 minute in the aggregometer, ADP was added to 2.5 μmol/l final concentration and aggregation recorded.

Fractionation studies. Crude fractionation of plasma from the patient in case 8 into albumin rich and globulin rich fractions was performed using 45% saturated ammonium sulphate. Purification of the IgG fraction was achieved by ion exchange chromatography using DEAE cellulose. Fractionation of plasma proteins into low, medium, and high molecular weight ranges was performed by chromatography on a Sephadex G200 column. All fractions were concentrated to their original plasma concentrations by positive pressure filtration. All samples were prepared in phosphate buffered saline (PBS) pH 7.2 without azide.

Lipid extraction. Neutral lipids were extracted from 2 ml plasma from the patients in cases 8 and 11 by mixing with 8 ml hexane for 15 minutes, centrifuging at 1300 g at 4°C, removing the hexane layer, and evaporating the solvent under nitrogen. The residue was redissolved in 1 ml BBS.

Polar lipids were extracted into a 1:1 mixture of cyclohexane and ethyl acetate after acidification to pH 3–4. After mixing for 15 minutes the solvent was separated and evaporated under nitrogen. The residue was again dissolved in BBS and tested for ability to support or inhibit PGI2 production.

Results

Plasma from 5 of 6 children with sporadic HUS showed decreased ability to support PGI2 production by aortic rings (Fig. 1, Table). No abnormality of PGI2 production was detected on the incubation of aortic rings with plasma from any of the 7 children with epidemic HUS or of the 9 non- HUS patients when compared with the normal adult controls.

Less PGI2 production occurred when aortic rings were incubated in plasma from 3 of 4 sporadic HUS patients tested (no comparison with BBS made in the

![Fig. 1](image-url)  

**Fig. 1** Diminished PGI2 production on incubation of plasma from the patient in case 8 with aortic rings  
(a) Without prior incubation with an aortic ring, patient's plasma has no inhibitory effect on platelet aggregation induced by adenosine diphosphate.  
(b) After incubation with aortic rings, patients' plasma (ring A) shows much less inhibition of aggregation than control plasma (ring B).  
(c) Ring A now incubated with normal plasma and ring B with patient's plasma; only the normal plasma is inhibitory.  
HUS = haemolytic uraemic syndrome.
patients in cases 9 and 11) than when the same ring
was incubated in BBS (Fig. 2). This suggests
inhibition of PGI₂ production by HUS plasma rather
than lack of stimulation. No such inhibitory
activity was detected using plasma from the patient
in case 13, from any of the epidemic HUS patients,
the non HUS patients, or the controls.

The rate at which added PGI₂ lost its activity on
incubation with plasma from patients in cases 8, 11,
and 12, determined by inhibition of platelet aggrega-
tion, was no different from that in the presence of
normal plasma. This indicates that the effect of these
patients' plasma was on the production rather than
the degradation of PGI₂.

Stability and fractionation studies. Fresh plasma
from the patient in case 8 or her plasma stored at
−20°C for up to 3 months inhibited PGI₂ production,
but plasma stored at 4°C for 72 hours or dialysed
for 72 hours showed neither inhibitory activity nor a
failure to stimulate PGI₂ production compared with
normal plasma. Heating to 56°C for 30 minutes
resulted in only a partial loss of the inhibitory
activity. No inhibitory activity was detected in the
protein fractions, which supported normal PGI₂
production, but as these were prepared at 4°C or
dialysed, inhibitory activity may have been lost.

The cyclohexane/ethyl acetate extracted fraction
of plasma from patients in cases 8 and 9 inhibited
PGI₂ production whereas the same fraction of
normal plasma had no inhibitory activity (Fig. 3).
No inhibitory activity was detected in the hexane
extracted fraction of either HUS or normal plasma.

Discussion

We have shown that plasma from certain patients
with childhood HUS fails to support PGI₂ pro-
duction by aortic rings. This parallels previous
reports on adult patients with HUS. The abnormality
was found only in patients with sporadic HUS,
however, and no abnormality of PGI₂ production was
detected in patients with the commoner epidemic
form of HUS or in the other renal diseases studied.

Our findings suggest therefore that there may be
different pathogenetic mechanisms in the epidemic
and sporadic HUS subgroups and add weight to the epidemiological, genetic, and histological evidence suggesting that HUS comprises at least 2 distinct diseases that differ not only in their epidemiology, clinical features, and prognosis but also in their underlying pathogenesis. Therapeutic trials and laboratory investigations of HUS need to recognize the heterogeneity of patients with the syndrome.

Previous investigators have ascribed the abnormality of PGI2 production on incubation of endothelial tissue with HUS plasma to either lack of a stimulating factor or excessively rapid degradation of PGI2. Our observation, however, that less PGI2 production occurred on incubation of aortic rings in some samples of HUS plasma than occurred in BBS alone suggests the presence of an inhibitor of PGI2 production in them. Furthermore, prolonged storage or dialysis at 4°C resulted in loss of the inhibitory activity from HUS plasma which then showed normal ability to support PGI2 production. There was no evidence of excessive degradation of added PGI2 in our patients’ plasma. Although the cause of the abnormality of PGI2 production in our patients thus differs from that proposed by other workers, it is possible that more than 1 mechanism may induce PGI2 deficiency in different groups of patients.

Our preliminary studies to characterise the inhibitor of PGI2 production indicate that it is a polar lipid, extractable into a mixture of cyclohexane and ethyl acetate. It is of interest that 1 of the earliest known inhibitors of PGI2 synthesis was 15 hydroxyperoxyarachidonic acid and other lipid peroxides are also known to inhibit PGI2 synthesis. Several authors have reported reduced concentrations of antioxidants such as vitamin E, superoxide dismutase, and transferrin in patients with HUS. It is therefore possible that the inhibitor of PGI2 production in the sporadic HUS patients is a lipid peroxide (possibly an arachidonic acid metabolite) produced after an oxidizing insult or resulting from lack of antioxidants in HUS plasma. Further studies are required to define the nature of the inhibitor more precisely and to establish whether its presence is related to the reported defects in antioxidant potential in HUS plasma.

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