Transient diabetes insipidus in a post-partum woman with pre-eclampsia associated with residual placental vasopressinase activity

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Summary

This case illustrates the exceedingly rare phenomenon of transient diabetes insipidus, in association with pre-eclampsia, occurring in the post-partum period following an in vitro fertilisation pregnancy, in an otherwise well 48-year-old lady. Diabetes insipidus can manifest during pregnancy, induced by increased vasopressinase activity secreted by placental trophoblasts and usually manifests in the third trimester. This presentation elucidates not only the intricate balance between the physiology of pregnancy and hormonal homeostasis, but also the importance of post-partum care as the physiological changes of pregnancy still hold pathological potential in the weeks immediately following delivery.

Learning points:

- Diabetes insipidus (DI) is a rare complication of pregnancy occurring in 1 in 30 000 pregnancies.
- It is associated with excessive vasopressinase activity, secreted by placental trophoblasts, which increases the rate of degradation of anti-diuretic hormone.
- It is responsive to synthetic desmopressin 1-deanimo-8-α-arginine vasopressin as this form is not degraded by placental vasopressinase.
- Vasopressinase is proportional to placental weight, which is increased in pregnancies conceived with assisted reproductive techniques including in vitro fertilisation.
- Vasopressinase-induced DI is associated with pre-eclampsia.

Background

Pregnancy offers unique challenges to the diagnosis and management of endocrinopathies, as the physiological changes of pregnancy introduce diagnostic complexity into the interpretation of investigative results. Diabetes insipidus (DI) is a rare complication of pregnancy, occurring in 1 in 30 000 pregnancies and can be difficult to recognise in late-pregnancy and the post-partum period as increased thirst, increased water intake and increased urine output are commonly reported. Furthermore, pregnancy is a state of water retention with increased maternal blood volume and an associated decrease in plasma osmolality. This case exemplifies these diagnostic intricacies in an exceptionally rare presentation of concomitant vasopressinase-induced DI and pre-eclampsia manifesting in the post-partum period.

Case presentation

A 48-year-old lady presented to the emergency department with a 3-day history of a progressively worsening, global
headache. It was sudden in onset, initially intermittent in nature, now persistent. There were no aggravating or relieving factors. She denied any nausea or vomiting, visual disturbance, speech disturbance or focal weakness. She noted swelling of her legs, emergent over the last 3 days. Interestingly, she also complained of an insidious onset of polydipsia with increased oral intake to 5 L in the last 24 h prior to presentation. This was associated with polyuria and nocturia. Of note, she was 8 days post-partum, having delivered a 3.4 kg male infant by elective lower segment caesarean section (LSCS) at 38-week and 3-day gestation. The delivery was complicated by 500 mL blood loss. The current pregnancy was achieved through in vitro fertilisation (IVF) utilising a donor egg and was preceded by two prior pregnancies resulting in miscarriage. Gestational diabetes was diagnosed in the second trimester and managed by dietary modification alone, with no hypertension or proteinuria documented during the pregnancy. She had no other significant past medical or family history of disease and was not on any regular medications. She was an ex-smoker, with a 10 pack-year history. She denied any alcohol intake.

On examination, she was hypertensive to 183/87 mmHg, not tachycardic or tachypnoeic and was afebrile. Neurological examination revealed no focal deficits and reflexes that were globally brisk but not hyper-reflexic. Cranial nerve examination was normal with no visual field defect. She was noted to have pitting oedema bilaterally to the knees and dry mucous membranes. Her cardiovascular examination was otherwise normal. Her abdominal examination was normal, and her surgical incision site appeared to be healing well.

Investigation

She underwent an urgent CT brain, which revealed an enlarged pituitary and queried lymphocytic hypophysitis or haemorrhage. Urine analysis demonstrated new-onset proteinuria with protein to creatinine ratio 110 mg/mmol. Laboratory data were as follows: glucose 5.3 mmol/L, sodium 146 mmol/L, creatinine 150 µmol/L, eGFR 35 mL/min/1.73 m² and urate 0.58 mmol/L (0.15–0.40). She had an isolated elevated alkaline phosphatase (ALP) of 143 U/L (20–105) with otherwise normal liver function tests. She had a normal platelet count of 258 (150–400). She was diagnosed with post-partum pre-eclampsia, however, did not meet the criteria for HELLP syndrome.

She remained polydipsic and polyuric, passing urine every 1–2 h. Urine osmolality was 175 mmol/kg with a serum osmolality of 308 mmol/kg. Pituitary MRI revealed a pituitary at the upper limit of normal considering her recent pregnancy, with no evidence of infarct or inflammation (Fig. 1). Anterior pituitary hormone profile was normal. She underwent a water deprivation test, which terminated after 4 h, with sodium rising to 146 mmol/L with serum osmolality 302 mmol/kg, urine osmolality 175 mmol/kg and anti-diuretic hormone (ADH) level <0.8 ng/L. She was diagnosed with DI. In the context of her pre-eclampsia, this was thought to be due to the residual action of placental vasopressinase.

Treatment

Following the diagnosis of pre-eclampsia, she was commenced on oxprenalol, resulting in the gradual resolution of her headache with normalisation of her blood pressure. Following confirmation of DI with the water deprivation test, she was commenced on desmopressin nasal spray, which resulted in decreased urine output.

Outcome and follow-up

On discharge creatinine had improved to 73 mmol/L and sodium was 142 mmol/L. She returned to clinic 1 week post discharge. She had stopped taking desmopressin due to an upper respiratory tract infection. Post cessation, she did not have any nocturia and passed urine 5–6 times per day. She continued to drink to thirst. Sodium
was 142 mmol/L, serum osmolality 303 mmol/kg, urine sodium <20 mmol/L, urine osmolality 223 mmol/kg and ADH 0.9 ng/L. (Table 1)

By 1 month post discharge, her symptoms had resolved with resolution of both her pre-eclampsia and DI, such that she no longer required any medications.

Discussion

Gestational DI is a rare complication of pregnancy, occurring in 1 in 30 000 pregnancies (1). It is characterised by hypotonic polyuria and polydipsia, manifesting as a transient phenomenon specific to the pregnant milieu, secondary to increased placental vasopressinase activity (1). It is associated with absent ADH levels, as demonstrated in our case (2). Placental vasopressinase, expressed in placental trophoblasts, degrades endogenous ADH but not 1-deanimo-8-arginine vasopressin (dDAVP), the synthetic form with a modified N-terminal, thereby allowing the resolution of symptoms with dDAVP administration (3).

Pregnancy denotes a change in maternal blood volume, osmoregulation and ADH secretion by the pituitary, whereby water retention increases and plasma osmolality decreases by 10 mmol/kg (4). There is a three-fold increase in the metabolic clearance of ADH in pregnancy, due to the excretion of vasopressinase from placental trophoblasts (4). Vasopressinase is a cystine aminopeptidase that inactivates ADH and oxytocin. It is produced from the seventh gestational week and concentrations rise coincidentally with the rise of trophoblastic mass, to peak in the third trimester. Following delivery, levels fall, becoming undetectable by 5–6 weeks post-partum (5). This reflects the natural history of vasopressinase-induced transient DI, which typically has its onset in the third trimester and remits spontaneously by 4–6 weeks after delivery (6).

Post-partum presentations of isolated DI are exceedingly unusual (1). Post-partum DI has previously been described in relation to placental abruption, due to large volume release of placental vasopressinase into the blood stream (7). There are no reports of post-partum DI following placental manipulation through LSCS, or following post-partum haemorrhage with retained placenta.

Vasopressinase concentrations are commensurate with placental mass (5), reflecting the known increased incidence of vasopressinase-induced DI in multiple gestations, which are associated with increased trophoblastic mass (7). Another fascinating facet to our case is the associated IVF pregnancy. Pregnancies conceived with artificial reproductive technology (ART), including IVF and intra cytoplasmic sperm injection, are associated with larger placental weights, and a higher placental weight/birth weight ratio, independent of length of gestation at delivery or method of ART (8). Extrapolating from this, it can be conjectured that IVF pregnancies are at increased risk of transient DI through this mechanism. There are not data yet supporting this theory.

Post-partum pre-eclampsia is atypical but described (9). The pathophysiology of pre-eclampsia is yet to be fully elucidated, but placental dysfunction is emerging as critical to its pathophysiology. While commonly attributed risk factors include pre-existing hypertension, nulliparity, obesity and maternal age greater than 40 years (10), there is emerging recognition of the risk in IVF pregnancies. Egg donation is a highly successful form of ART, however, is emerging as a demonstrable risk for the development of pre-eclampsia with a recent meta-analysis denoting a odds ratio of 4.5, compared to autologous egg IVF pregnancies (11). The postulated mechanism centres on discordant allogenicity of the foetus in relation to the mother (12). The use of a donor egg in this case, coupled with her advanced maternal age and nulliparous status, provide multiple risk factors for pre-eclampsia to develop.

Pre-eclamptic placental and end-organ dysfunction manifests in a myriad of ways, including maternal
hypertension, proteinuria, thrombocytopenia, renal insufficiency, intrauterine growth restriction and neurological disturbance (13). Vasopressinase is metabolised in the liver, with vasopressinase-induced DI described in patients with eclampsia and pre-eclampsia, especially HELLP syndrome, as the associated liver dysfunction impairs clearance of vasopressinase (14). It has also been described in other forms of liver dysfunction complicating pregnancy, leading to deranged liver function, including hepatic steatosis (15). This is especially important in our patient as she had concomitant deranged liver function, providing a constellation of circumstances for gestational DI to manifest.

This case illuminates the diagnostic complexities of gestational DI, especially in the rare situation of postpartum emergence of disease. A high index of suspicion for vasopressinase-induced DI should thus be maintained in patients presenting with polyuria, especially in the presence of other risk factors such as pre-eclampsia and hepatic dysfunction, multiple gestation and IVF pregnancies.

Declaration of interest
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