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Non-reproductive triggers of postpartum psychosis

Ian Brockington

Abstract Bipolar disorders, and other psychoses, are known to be triggered by a number of agents apart from the reproductive process. In some women, pregnant or recently delivered, psychosis may be due to these alternative triggers. There are substantial numbers of mothers suffering from childbearing psychoses, who have been prescribed bromocriptine or steroids, have had surgical operations or developed thyrotoxicosis. It is best to eliminate these episodes and cases from study samples of puerperal psychosis.

Keywords Bromocriptine · Post-operative psychosis · Corticosteroid therapy · Thyrotoxicosis

Introduction

Many perhaps think of ‘puerperal psychosis’ as triggered by the cascade of hormones after childbirth. In What is Worth Knowing about ‘Puerperal Psychosis’ (Brockington 2014), it is proposed that the early postpartum trigger is only one of a group of triggers acting at various times in the reproductive process, including pregnancy, later in the postpartum period, after weaning, after abortion and at a stage in the menstrual cycle.

But in addition to all these triggers acting on the bipolar/cycloid diathesis, there are several non-reproductive triggers, which may result in psychoses that present in pregnancy or the puerperium.

Bromocriptine

The argolide derivative, 2-bromo-α-ergocriptine, a dopamine D2 agonist, was developed in 1968 to block the release of prolactin (Parkes 1979), and has been used since 1978 in a dose of 5–7.5 mg/day to inhibit puerperal lactation. Since then, 14 postpartum episodes have been reported, associated with its use (Brook and Cookson 1978; Vlissides et al. 1978; Charbonnier and Planche 1981; Canterbury et al. 1987; Kemperman and Zwanikken 1987; Daw 1988; Iffy et al. 1989; Durst et al. 1990; Fisher et al. 1991; Reeves and Pinkofsky 1997; Pinardo Zabala et al. 2003; Misdrahi et al. 2006). In a fifteenth (Lake et al. 1987), the mother became overactive before bromocriptine was prescribed. The onset, where known, was early in the puerperium. In eight, the clinical picture was typical of mania, and another ran a bipolar course, starting with depression (Reeves and Pinkofsky 1997). In three, it was atypical (Daw 1988; Pinardo Zabala et al. 2003), including one with a seizure (Iffy et al. 1989), and in one, depressive, with ‘voices’ telling her to destroy the baby (Canterbury et al. 1987). In six, duration was short—1 week or less in five, and 17 days in the sixth. Two mothers recovered after withdrawal of bromocriptine without other treatment (Charbonnier and Planche 1981; Canterbury et al. 1987); but other episodes have lasted up to 2 months, and required much treatment, in one case ECT. Two mothers suffered previous manic episodes, including steroid-triggered episodes (Fisher et al. 1991; Misdrahi et al. 2006). One had a family history of psychosis (Fisher et al. 1991).

In my series, there was one possible example:

A 24-year old was delivered of her 1st child by Caesarean section because of a suspected foetal facial cyst. She was treated with bromocriptine 5 mg/day. She felt excited because the baby was normal. On day 4 she...
began to behave strangely and became increasingly anxious, overactive, emotionally labile and disinhibited. She was talkative and cheerful, and gave a running commentary on her actions. Admitted to hospital, she was confused and perplexed; she did not know the day of the week. She was unable to think clearly, and spoke little, with slow and deliberate answers. The possibility of a toxic delirium secondary to bromocriptine was raised; it was stopped and she was treated with chlorpromazine. She rapidly recovered, but, a month later, was ‘over the top’ for two days. She remained well in the next 32 years.

It is clear that bromocriptine can trigger postpartum episodes, perhaps only in mothers with a bipolar diathesis. It can also trigger episodes in other circumstances, as described in the treatment of parkinsonism (Lipper, 1976), acromegaly (Le Feuvre et al. 1982; Valdes et al. 1989) and prolactinoma (Turner et al. 1984). Psychiatric complications required withdrawal of bromocriptine in 5/66 cases (Serby et al. 1978).

The frequency 5/46 (11 %) is about the same as post-abortion episodes (14 %).

It should be noted that many mothers were delivered by Caesarean section or forceps under general anaesthesia; in my bipolar/cycloid group, this was the mode of delivery in 30 episodes. Postpartum psychoses after Caesarean section could be post-operative rather than puerperal psychoses.

Corticosteroid therapy

This was introduced into therapeutics in 1950. There is a scattered literature on the precipitation of mania, severe depression, paranoid disorders, delirium and other psychoses by adrenal corticosteroids (for example, Sirois 2003; Kenna et al. 2011); about half of the reported cases are bipolar. One would, therefore, expect an association with postpartum bipolar disorders. In the literature, there are two cases of this association (Svoboda 1957; Johnson 1996), to which I can add six, which are briefly summarized below:

A 29-year old became pregnant for the 1st time. Because of foetal distress, she was delivered @ 35 weeks gestation by emergency Caesarean section. She had two doses of dexamethasone at the time of delivery. On day 2 she became sleepless and began writing copious notes. By day 7 she became overactive and aggressive, and said the television and radio referred to her, and her husband was trying to infect her and the baby with AIDS, which he had contracted during an affair. She later had a second episode of postpartum mania, with onset six weeks after the birth.

A 35-year old, after six years of infertility, conceived with gamete intra-fallopian transfer, and became pregnant with twins. At 30 weeks gestation, she was given dexamethasone to increase the maturity of the foetal lungs. A week later she was admitted to the maternity hospital with antepartum bleeding. She became excited, elated and sleepless and expressed persecutory ideas. She was talking constantly, giving a running commentary on everything that was happening. At 32 weeks gestation, she gave birth; one of the twins had Down’s syndrome. Three weeks later her psychosis recurred, with head-banging, restlessness, insomnia and racing thoughts. On admission to hospital she was depressed, and did not recover until five months later. She remained well for the next 20 years.

A 28-year old gave birth to her 1st child. For two weeks she was ‘on a high’, talking non-stop. Six weeks later she developed pityriasis rosea, treated with steroids. She became very high, ‘brilliant’, her mind racing. This
lasted a week until she stopped the steroids. She then became depressed for a year. Her baby also became high on steroids.

A 35-year old, who for several years had suffered from poly-arthritis and Crohn’s disease, developed pre-eclamptic toxaemia during her 1st pregnancy, and was delivered by Caesarean section. On day 12 she developed a cycloid psychosis – slow, confused and perplexed – from which she recovered within three weeks. When the baby was four months old, her arthritis recurred. Treated with ibuprofen, she developed a purpuric vasculitis with bullous lesions (Stevens-Johnson syndrome), together with laboratory evidence of systemic lupus erythematosus. She was treated with prednisolone 60 mg/day. Within four days she became withdrawn and mute, staring into the distance. She washed obsessively, complaining of sweating and halitosis. A CAT scan showed diffuse cerebral abnormalities, and an EEG low frequency activity. She was again treated with steroids, and recovered in three months.

A 24-year old, whose mother suffered from puerperal psychosis after her own birth, became pregnant for the 1st time. Two days before the onset of labour she developed a rash, which was treated with prednisone 20 mg/day. After the birth, on day 7, she became weepy, and disoriented, then elated and confused. She felt her brain was exploding. She thought her partner was trying to kill her and believed she was the Messiah. She telephoned a minister to say she had the solution to the Irish problem. She was writing reams of gibberish, which she believed was important to the future of humanity. After recovery she suffered severe bonding problems, which continued after the second baby was born. In 28 years she suffered one further manic episode.

A 28-year old, @ 35 weeks gestation, was treated with prednisolone for idiopathic thrombocytopenia. At 39 weeks she was delivered by forceps, and steroids were discontinued. On day 3 she became agitated, weepy and perplexed, with confusion and ‘paranoid’ ideation. On admission to hospital, prednisolone was

### Table 1 Association with thyrotoxicosis

| First author          | Clinical features                                      | Evidence of thyrotoxicosis                                      | Comment                      |
|-----------------------|--------------------------------------------------------|---------------------------------------------------------------|------------------------------|
| Johnstone 1884        | Onset of psychosis 6 months after the birth of her third child | Signs of exophthalmic goitre developed concurrently            | Late postpartum onset        |
| Knauer 1897 Case 65   | Onset of a chronic depressive psychosis after her sixth birth | She developed thyrotoxicosis                                  | No data on timing            |
| Sivadon 1933 Case 15  | Onset of psychosis on day 9 after her first birth       | She had an enlarged thyroid and tremor.                        | She died on day 22, infection possible |
| Schröder 1936 case 38 | She gave birth at 38, and on day 11 developed a psychosis | She had a goitre at the age of 10 and puerperal psychosis at 38; 7 months later, while still psychotic, she was noted to have slight exophthalmos and sweaty hands | Thyrotoxicosis was noticed 7 months later |
| Abély et al. 1947     | Onset of psychosis shortly after the second, third, fourth and fifth births | During two episodes, thyroid enlargement was noted. During the fourth episode, tests showed transitory hyperthyroidism, whose disappearance coincided with improvement in the clinical state | Only the fourth episode of postpartum psychosis was affected |
| Retzeanu et al. 1960  | Onset of psychosis in the ninth month of gestation      | She had a large soft thyroid and tachycardia; she refused surgery on her goitre and was treated with psychotropic drugs and radioactive iodine. Improvement in mental state and reduction in goiter were concurrent | Prepartum psychosis, with a possible response to anti-thyroid treatment. She also had pleurisy and galactorrhea |
| Butts 1968            | Psychosis of unknown onset after first birth, and with onset day 7 after the second | During the second episode, she had an enlarged thyroid, systolic murmur and tremor; but her protein-bound iodine was only 4.4 µg/100 ml and her basal metabolic rate minus 12% | Only the second episode of postpartum psychosis was affected. Evidence of thyrotoxicosis was equivocal |
| From my series        | Onset of depressive psychosis 7 months after the birth  | Goitre, loss of weight and tremor during her second pregnancy  | Late postpartum onset, response to ECT |
|                       | Psychosis on day 5 after both her first and second births | She had a goitre and clinical signs; the diagnosis was confirmed by laboratory tests. Treatment included radioactive iodine | Only the second episode of postpartum psychosis was affected |

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started again. She remained retarded, vague, staring into space, suspicious, speaking in a monotonous voice of being evil, hopeless and a failure. She denied the existence of her husband and the baby. She suddenly disrobed and started shouting, “Let me die!” Seven days later she abruptly improved. Two weeks later steroids were stopped, but soon started again because a diagnosis of systemic lupus erythematosus was made. She had no further episode in the course of nine years.

Only four other mothers were treated with steroids in any form. Furthermore, these six mothers, in a total of 88 years observation, had, apart from those associated with childbearing or steroid therapy, only one psychotic episode between them. It seems best to regard these episodes as steroid psychoses and not childbearing psychoses.

**Thyrotoxicosis**

Note that this patient had myxoedema disease is occasionally associated with acute psychosis, as shown by at least nine case reports and Brownlie’s survey in New Zealand (Brownlie et al. 2000); three patients have recovered after surgical treatment (Bursten 1961; Lazarus and Jaffe 1986; Abbasie et al. 2009), and others after treatment with propylthiouracil (Øestergaard Jensen 1950) or propranolol (Lee et al. 1991). Although occasional cases had manic features (Bursten 1961; Stowell and Barnhill 2005), a wide variety of other psychotic syndromes have been seen.

Of particular interest are two episodes of thyroid disease that developed concurrently with late onset postpartum psychoses (Bokhari et al. 1998; Stowell and Barnhill 2005):

Eleven weeks after her 3rd child was born, a 29-year-old developed insomnia, weight loss and fatigue intolerance. She appeared confused and was disoriented in time and place. She heard Jesus talking to her, and also had visual hallucinations. She believed she was pregnant with the Christ child, and would be killed by hospital staff. She had thyrotoxicosis, associated with thyroiditis. Her psychiatric symptoms improved concurrently with its treatment.

A 35-year old gave birth to twins. Seven months later she presented with increased energy, lack of the need for sleep, racing thoughts, preoccupation with bible reading, and the belief that God had fathered her babies. She collapsed and was admitted. She had myxoedema due to postpartum thyroiditis. With thyroxin and risperidone she recovered within six days.

Table 1 shows the details of eight other mothers with some evidence of this association.

In all, there are 11 mothers with concurrent childbearing psychosis and thyrotoxicosis; 1 had prepartum, 4 late postpartum, 5 early postpartum and 1 unknown onset. Three had other episodes of postpartum psychosis that were not accompanied by thyrotoxicosis. One mother may have responded to thyroid treatment alone. There is no consistency in these data. It is possible that thyrotoxicosis is a factor acting synergistically with childbearing factors, but more evidence is required.

**Discussion**

Unless Caesarean section is included among the surgical procedures, there are not many ‘non-reproductive’ triggers of childbearing psychoses, but any steps taken to improve the homogeneity of samples will be beneficial (Brockington 2016). Where these triggers are associated with episodes, they should be discounted, and where these are the sole episodes in the mother’s history, her case should be removed.

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