Massive reduction of tumour load and normalisation of hyperprolactinaemia after high dose cabergoline in metastasised prolactinoma causing thoracic syringomyelia

S H M van Uum, N van Alfen, P Wesseling, E van Lindert, G F F M Pieters, P Nooijen, A R M M Hermus

In 1970 a 20 year old woman presented with a pituitary chromophobe adenoma for which she underwent transfrontal pituitary surgery. In 1978 she had to be reoperated on because of local tumour recurrence, resulting in hypopituitarism. Bromocriptine (5 mg/day) was given for 15 years, but the plasma prolactin levels remained elevated. In 2000 the patient presented with signs and symptoms suggestive of a spinal cord lesion at the mid-thoracic level. A magnetic resonance imaging (MRI) scan showed an extensive leptomeningeal mass extending from the brainstem to L5, with a thoracic syringomyelia at the T7–T8 level. The plasma prolactin level was very high (5114 μg/l). A biopsy showed the presence of a metastasised prolactinoma.

On administration of high dose cabergoline, 0.5 mg twice a day orally, the plasma prolactin levels decreased within one month and then normalised within 26 months. Tumour load reduced considerably but unfortunately, her signs and symptoms did not improve. This case illustrates that a high dose dopamine agonist might be an important therapeutic option in patients with a metastasised prolactinoma.

In 1973, a magnetic resonance imaging (MRI) study of the brain and spinal cord did not show any abnormality in the sellar region, but did reveal multiple small areas of increased density around the medulla oblongata and the spinal cord, and between the cerebellum and cerebral peduncles. The lesions showed contrast enhancement with gadolinium. At that time the lesions were ascribed to Lipiodol used during previous myelographic studies. Since the patient had no complaints and there was no evidence of local recurrence, bromocriptine was discontinued. This resulted in an increase of plasma prolactin from 650 μg/l to 1280 μg/l. The prolactin level further increased to 7858 μg/l in 1999.

In 1998 the patient experienced numb, tight sensations in the fourth and fifth digits of the left foot. In April 1999 she developed pain in the left subscapular area and from October 1999 she noted icy and frozen sensations below the waist, with tingling sensations in both legs. These symptoms slowly progressed to a level where she had to visually guide herself when moving around. In addition, there were spontaneous kicking movements of the legs and micturition was impaired with urge incontinence. Her medication at that time consisted of cortisone acetate 37.5 mg/day, levethyroxine 0.05 mg/day and a combination of estradiol and norgestrel.

Physical examination at referral showed an alert, well oriented woman. There was increased muscle tone in the legs, and a partial drop foot on the left (MRC grade 4). She had a broad stance and gait and tandem walking was impossible. There was hypaesthesia below T9 level for both pinprick and position sense; Romberg’s sign was positive. There was hyperreflexia of the knee and ankle jerks. Abdominal cutaneous reflexes were absent at all levels. Plantar responses were normal. The symptomatology was most consistent with a spinal cord lesion at the mid-thoracic level.

Laboratory investigations revealed an extremely high level of plasma prolactin (5114 μg/l). On lumbar puncture the opening pressure was 14 cm H2O, and the viscous, yellow cerebrospinal fluid (CSF) flowed with difficulty. Fluid analysis showed 11 leucocytes/μl, 8 erythrocytes/μl, a hugely elevated protein content of 19 500 mg/l, glucose 1.9 mmol/l (serum value 4.2 mmol/l), and lactate 5645 μmol/l. This, and a negative Queckenstedt’s test, suggested a spinal subarachnoid block of CSF flow. Due to the limited amount of CSF obtained, no cytological studies were done and prolactin levels were not measured.

An MRI study of the whole cerebrospinal system was done (fig 1A, C). No tumour recurrence was found in the sellar region.

Abbreviations: CSF, cerebrospinal fluid; MRI, magnetic resonance imaging; MRC, Medical Research Council
region, but an extensive leptomeningeal mass surrounding and slightly compressing the intrathecal space down to the level of L5. Further, there was thoracic syringomyelia at the T7–T8 level.

Occipitocervical decompression was performed and a biopsy of the tumour mass surrounding the medulla oblongata was taken. Microscopic examination showed adenomatous tissue with closely packed monomorphic cells and a low mitotic index. The immunohistochemical staining for prolactin was strongly positive. A diagnosis of malignant prolactinoma with metastases to the leptomeninges was made.

In an attempt to improve or stabilize the patient’s symptoms a thoracic laminectomy at T7 with insertion of a syringosubarachnoidal CSF shunt was performed one week after the biopsy was taken. During the procedure extensive adhesions of the arachnoids and dura were found, with hardly any flow of CSF from the subarachnoid space. An arachnoid pseudocyst, which was compressing the spinal cord was laterally opened and drained. In the midline, a tumour, several millimetres thick extending over a bulging spinal cord, was removed, and following incision of the spinal cord the CSF discharged under pressure emptying the syrinx. Subsequently, the spinal cord tissue collapsed. A small catheter was inserted in the syringal cavity.

Unfortunately the patient’s difficulties with walking and her sensory disturbances did not improve after surgery. A week later an emergency ventriculoperitoneal shunt had to be placed because of acute obstructive hydrocephalus.

High dose cabergoline (0.5 mg twice a day orally) was started after the first operation. The plasma prolactin level decreased quickly and was below 300 μg/l within a month. Thereafter the prolactin level normalised and remained normal throughout the treatment which continued for 26 months. Treatment was well tolerated, and the patient did not show any further decline. She was discharged to a rehabilitation facility. MRI studies of the cerebrospinal system were repeated after four months. The tumour had decreased in size at the level of both the medulla oblongata and the spinal cord. Repeat MRI studies after treatment for 26 months showed a considerable, further decrease in tumour load (fig 1B, D).

Despite the impressive effect of cabergoline on plasma prolactin levels and tumour load, the patient continued to have disabling neuropathic pains in her legs and decreased sensibility below the T7 level, severely complicating her walking. Two years after the surgery she remains wheelchair dependent.

DISCUSSION

Our patient presented with a chromophobe pituitary adenoma in 1970, with local recurrence in 1978. Retrospectively, metastatic tumour deposits along the neuraxis were already present at least since 1993. When we first saw her 30 years after the first pituitary surgery, her neurological symptoms were not due to direct compression of the central nervous system by tumour metastases, but to blockage of CSF flow by arachnoid adhesions caused by metastatic tumour deposits and the subsequent formation of a thoracic syringomyelia.

Malignant prolactinomas are very rare. The first case was described in 1981 by Martin et al. Until now about 30 cases of metastasising prolactinoma have been described. The interval between the initial presentation of the prolactinoma, usually with a sellar tumour, and the diagnosis of the metastasised prolactinoma varies from two months to 27 years, with an average of seven years. Results of treatment of metastasised prolactinoma are generally poor. Surgery, radiation therapy, chemotherapy, and treatment with dopamine agonists either alone or, more commonly, in various combinations resulted in no, or only a partial, temporary response in almost all cases.

Our patient presented with metastases in spinal cord from a malignant prolactinoma resulting in a syrinx. Intraspinal metastases have been described previously in six patients with a malignant prolactinoma, but a syrinx, such as in our patient, was not present in any of them. Two patients died before the effect of therapy could be assessed. One patient had signs and symptoms due to a metastasis at L1 level. A combination of surgery, radiotherapy, bromocriptine, and anti-oestrogen therapy, resulted in stabilisation, but there was no improvement in the neurological situation and a decrease of prolactin of only about 50%. Another patient with a single intraspinal metastasis from a prolactinoma at the level of S1–S2 was successfully treated with surgery, radiation, and bromocriptine. Kuroki et al described a patient with multiple intradural masses on myelography, most probably metastases from a prolactinoma. Radiation resulted in improvement of severe low back pain, but the patient died within three months because of regrowth of intracranial lesions. The sixth patient presented with multiple prolactinoma lesions from the foramen magnum to L3. After biopsy of the largest nodule and radiation therapy the myelopathic symptoms considerably improved while on bromocriptine 100 mg daily.

Our patient had a large intraspinal tumour burden extending from the medulla oblongata to the lumbar region, which made curative surgery impossible. As we were concerned that radiotherapy would further damage the vulnerable neural tissue around the syrinx, we decided to treat the patient with a dopamine agonist only. In the past,

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**Figure 1** T1 axial, gadolinium enhanced, magnetic resonance imaging (MRI) scan of the brain and T1 sagittal, gadolinium enhanced, MRI scan of the spinal cord before (A, C) and after (B, D) high dose cabergoline for 26 months.
the plasma prolactin had increased during treatment with low dose bromocriptine (5 mg daily). We therefore selected another dopamine agonist, cabergoline, and because of the huge tumour load we chose a high dose, 0.5 mg twice a day orally (normal starting dose for a macroprolactinoma is 0.5 mg twice weekly). Our patient is the first reported patient with a metastasised prolactinoma in whom monotherapy with a dopamine agonist resulted in complete normalisation of the plasma levels of prolactin. This treatment also resulted in a decrease of tumour size, first documented after four months with further reduction seen after two years. The high dose cabergoline was very well tolerated and we conclude that this might be an important therapeutic option in patients with a metastasised prolactinoma.

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