Transformation of a microprolactinoma into a mixed growth hormone and prolactin-secreting pituitary adenoma

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INTRODUCTION

Approximately 10% of all clinically overt intracranial neoplasms arise from the pituitary gland and at autopsy, if histological examination of unselected autopsy material is to be believed, adenomas are present in as many as one in five cases (Levy and Lightman, 1993). Prolactin-secreting adenomas are the most common of them, representing about 40% of adenomas (Mindermann and Wilson, 1994). Prevalence has classically been estimated as 6–10 per 100,000 inhabitants, but more recent data suggest it could be as high as 62 per 100,000 inhabitants (Daly et al., 2006). Early diagnosis of mixed adenomas is rendered difficult by the fact that they generally present with signs or symptoms that can be related to overproduction of a single hormone. Clinical diagnosis of combined anterior pituitary hormone excess is thus a rare instance, and these adenomas are mostly identified after neurosurgery, at the time of pathological diagnosis.

The most frequent combination of pituitary hormone excess associates growth hormone (GH) and prolactin (PRL); the combination of GH, PRL, and the alpha subunit (α-SU) of the glycoprotein hormones is also possible, whereas other hormone combinations within a single tumor are extremely rare (Kovacs et al., 1998; Ma et al., 2002). Ultrastructural and immune electron microscopic data allow to classify bi-hormonal or pluri-hormonal adenomas in three separate classes: monomorphous, bimorphous, and plurimorphous tumors. Monomorphous adenomas consist of a single cell type with synthesis of two or more hormones in the same cell. Bimorphous and plurimorphous adenomas are composed of two or more cell types, and each hormone is produced by a different cell population. However, overlaps may exist, and some tumors can consist in a mix of cells, some of them producing only one hormone and others producing two or more hormones. In some adenomas, one cell type can predominate; in others, various areas contain groups of similar cells, or a gradual transformation seems to exist between the cell types. Despite these overlaps, the classification of tumors into monomorphous, bimorphous, and plurimorphous types for diagnostic purposes retains its value and remains a useful practical tool (Kovacs et al., 1998). In this paper, we describe a patient who presented initially with a pure PRL secreting pituitary adenoma, and who secondarily developed co-secretion of GH with time.

CASE REPORT

Here we present the case of female patient who delivered her first child in July 1996 at the age of 31 years, after spontaneous conception and an uneventful pregnancy. She did not breast feed her baby, and was seen again in May 1997, where she complained of persistent secondary amenorrhea and intermittent galactorrhea of 10 months’ duration. No specific investigations were performed at the time, and substitutive treatment with Estradiol and Norgestrel (Cyclacur®) was initiated by her treating gynecologist. This resulted in regular menstruations, but the treatment was stopped by the patient in September 1997 because of a desire of pregnancy, allowing to confirm the persistence of secondary amenorrhea.

Combined prolactin (PRL) and growth hormone (GH) secretion by a single pituitary tumor can occur in approximately 5% of cases. However, in all previously reported patients, combined secretion of both hormones was present at the time of diagnosis. Here we describe a patient initially diagnosed with a pure prolactin-secreting microadenoma, who experienced the progressive apparition of symptomatic autonomous GH secretion while on intermittent long term dopamine agonist therapy. She was operated on, and immunohistochemical analysis of tumor tissue confirmed the diagnosis of pituitary adenoma with uniform co-staining of all cells for both GH and PRL. This patient represents the first documented occurrence of asynchronous development of combined GH and PRL secretion in a pituitary adenoma. Although pathogenic mechanisms implicated remain largely speculative, it emphasizes the need for long term hormonal follow up of patients harboring prolactinomas.

Keywords: microprolactinoma, growth hormone, IGF-1, galactorrhea, amenorrhea, somatostatin analog
She was otherwise in good general health, had no particular complaint and was on no medication, only reporting having lost 10 kg after her first pregnancy. On physical exam, she was a healthy appearing young woman weighing 51 kg for 162 cm, the only positive finding being the presence of bilateral galactorrhea on stimulation. A pregnancy test was negative, and investigation of anterior pituitary function disclosed the following baseline hormone levels (November 1997): LH 1.6 U/l (N: 2–12), FSH 4.8 U/l (N: 4–12), oestradiol 0.2 nmol/l (N: 0.14–0.34); PRL 42 μg/l (N: 3.4–24.1); TSH 0.79 mU/l (N: 0.2–3.5), free T4 15 pmol/l (N: 8–22). GH and IGF-1 levels were reportedly within normal limits at the time. The patient refused to undergo a pituitary MRI because of claustrophobia, and a head CT scan disclosed the presence of a microadenoma (7 × 4 mm) in the left lobe of the pituitary gland.

The diagnosis of microprolactinoma was retained, based on the good correlation between the relatively small size of the adenoma and the moderate, albeit symptomatic, elevation of prolactin levels. Thus, dopamine agonist therapy with Quinagolide (Norprolac® 75 mg/day) was initiated in January 1998. A few weeks later, the patient became pregnant, treatment was stopped, and an uneventful pregnancy resulted in the delivery of a normal second child in October 1998. The patient was able to breast feed her baby. At the follow up visit in March 1999, the patient was still amenorrheic and she had persistent bilateral galactorrhea on stimulation. Her baseline prolactin level was 66 μg/l (N: 2–19), and IGF-1 was normal at 342 μg/l (N: 98–442) in July 1999. A repeat head CT scan confirmed the presence of a stable microadenoma. She then refused to take any medication, and also refused to take oral contraception.

Over the course of the following years, her baseline prolactin levels increased slowly but steadily, reaching 92 μg/l (N: 2–19) in 2002. Because of the risk of osteoporosis carried by long term amenorrhea, the patient then agreed upon taking dopamine agonist therapy and was started on oral Cabergoline (Dostinex® 0.5 mg/week) in June 2002. As anticipated, her serum prolactin levels declined, galactorrhea disappeared and by October 2002, regular ovulatory cycles had resumed. However, she experienced a minor depression on cabergoline that required transient treatment (August 2002–June 2003) with the selective serotonin re-uptake inhibitor nefazodone (Nefadar®). In May 2004, recurrence of amenorrhea under treatment motivated a repeat head CT scan which disclosed a small decrease in size of the adenoma. At the time, serum PRL levels were stable at 57 μg/l and notably did not exhibit any significant change despite the transient treatment with a selective serotonin re-uptake inhibitor that had occurred a few months before. Cabergoline was increased to 0.75 mg/week, with return of the menses. The first MRI was obtained in June 2005, showing an oval hypointense (T1) nodule in the left lobe of the pituitary gland, measuring 5 mm by 3 mm (i.e., smaller than on all previous imaging studies). In December of that year, the patient decided to stop taking cabergoline. She became amenorrheic again shortly thereafter, with Prolactin levels rising to 110 μg/l in August 2006, prompting re-introduction of the treatment. IGF-1 levels at that time were evaluated in a novel assay because of a change in the methodology used at our hospital and amounted to 356 μg/l (N: 106–276, see Figure 1).

Once again, therapy was stopped by the patient in April 2007, an interruption that was followed by another episode of amenorrhea.
accompanied by intermittent galactorrhea. At her next yearly follow-up visit in November 2007, she mentioned spontaneously an increase in size of her fingers. On specific questioning, she also reported disseminated arthralgias, and symptoms suggestive of carpal tunnel syndrome. Baseline hormone work up at the time was as follows: PRL was stable at 51 μg/l (N: 4–29), but IGF-1 was significantly higher than a few months before, at 480 μg/l (N: 98–261). A repeat pituitary MRI was therefore obtained in November 2007 (see Figure 2A) disclosing the existence of a mixed solid and cystic nodule in the left lobe of the anterior pituitary gland, measuring 6 by 10 mm. This nodule was larger than on previous imaging studies, and was located in contact with the left cavernous sinus without sign of invasion. This MRI exam also showed a second lesion localized at the root of the pituitary stalk, measuring 3 by 3 mm, corresponding to a small cyst. A GH suppression test with oral glucose showed a nadir GH level of 0.51 μg/l after 120 min. The control MRI realized in 2010 showed no residual tumor and the small cyst had also disappeared (see Figure 2C). Blood hormone levels were as follows (July 2010): PRL 16 μg/l (N: 4–29), IGF-1 120 μg/l (N: 98–261), and IGF-BP3 3.8 mg/l (N: 3.3–6–6).

DISCUSSION

Combined prolactin and GH secretion has been reported in 5% of all pituitary tumors, an co-secretion is usually diagnosed simultaneously (Kasantikul and Shuangshoti, 1990). Mixed prolactin and GH secreting pituitary adenomas are relatively common because somatotrophs and lactotrophs share the common somatomammotroph progenitor lineage. Conversely, the occurrence of a prolactinoma evolving into clinically and biochemically active acromegaly seems to be a rare phenomenon. In a recent report, Lania et al. (2010) describe secondary apparition of GH hypersecretion. However, in contrast to the patient described here, the primary tumor in their report was an aggressive prolactinoma, suggesting a different pathophysiological mechanism from the start. In this context, older papers describing the occurrence of combined elevations of GH and prolactin do not allow to differentiate between concomitant or sequential development of the individual hypersecretion syndromes (Tournaire et al., 1985; Goldman and Klinges, 1989; Pagesy et al., 1991). Moreover, it should also be reminded that elevated prolactin levels in the context of a somatotroph adenoma can also result from desinhibition of lactotroph cells by pituitary stalk compression.

In contrast to these previous reports, the patient discussed here is remarkable in the sense that symptoms of GH excess appeared progressively, several years after the diagnosis of microprolactinoma. Very importantly, clinical symptoms were well correlated with the biological findings. Indeed, an elevation of IGF-1 levels was not observed before 2006, and the first spontaneous complaint suggestive of GH excess was recorded in 2007. The fact that once documented, these elevated IGF-1 levels continued to increase steadily on further testing, together with the good correlation with clinical symptoms, argues very much in favor of secondary apparition of GH hypersecretion and against an artifact related to the change in the IGF-1 assay that occurred in 2005. This is further confirmed by the clearly abnormal results of the oral glucose suppression test, confirming the abnormal GH secretion.

Interestingly, this change in phenotype from a pure prolactin-secreting adenoma into a mixed somato-lactotroph adenoma was
accompanied by significant growth of the known pituitary tumor, after several years of stability in size. This unfortunate evolution, in parallel with the change in the secretory profile of the tumor, could therefore be interpreted as a sign of de-differentiation. Indeed, hormone-specific anterior pituitocytes are embryologically derived from a pluripotent precursor. The process of anterior pituitary cell development and differentiation arises as a consequence of concerted spatio-temporal expression or repression of a series of transcription factors to produce fully differentiated cells of the various lineages (Melmed, 2003). In this differentiation process, prolactin-expressing cells are just one step downstream from cells expressing both GH and prolactin.

At that point of the clinical evolution, and regardless of the pathogenic mechanisms implicated, the indication for surgical resection of the adenoma was retained. Given the close vicinity of the cavernous sinus, pre-treatment with a long-acting somatostatin analog was initiated in order to increase the chance of curative trans-sphenoidal surgery (Ferone et al., 2000). This option proved successful since medical treatment induced a significant reduction of the tumor size, ultimately allowing radiologically complete resection of all tumor tissue. Immunohistochemical exam of the removed tumor disclosed a pituitary adenoma with uniform co-staining for both GH and PRL, confirming the hypothesis of an evolution into a mixed somato-mammotroph pituitary adenoma.

The cytogenesis of pituitary adenomas that consist of two different cell populations is not known and remains to be elucidated (Kovacs et al., 1998), although different hypotheses have been proposed. Most, if not all, of these adenomas are monoclonal, as demonstrated by X-inactivation studies (Herman et al., 1990; Ma et al., 2002). Moreover, at the end of the eighties, an activating mutation of the α-SU of G proteins was identified in somatotroph cells of up to 40% of sporadic GH-secreting pituitary adenomas in Caucasians (Landis et al., 1990; Levy and Lightman, 1993). The resulting oncogene, gsp, was thought to induce tumorigenesis by virtue of persistent activation of adenyl cyclase, with subsequent GH hypersecretion (Landis et al., 1989; Eugster and Pescovitz, 1999). This report constitutes the demonstration that specific molecular abnormalities may form the basis of pituitary adenoma formation and hormone hypersecretion, at least in some specific cases.

Alternatively, it may be hypothesized that such mixed pituitary adenomas are not monoclonal, and that the insult that causes the neoplastic transformation may affect two different cell types. It is also conceivable that some pituitary tumors originate in an uncommitted stem cell which, because of unknown factors, can differentiate into two separate cell types. Such multidirectional differentiation could explain the development of some pluri-hormonal tumors (Kovacs et al., 1998).

Another hypothesis would be that one cell type can “transdifferentiate” to another cell type as a result of subsequent mutations during tumor progression. Such phenomenon involves reversible transformation of one cell type to another by phenotypic switches, without cell division (Senovilla et al., 2004). It takes place in a few pituitary cell types under physiological conditions. This concept of transdifferentiation was for example introduced to explain the

Table 1 | Results of the oral glucose tolerance tests (→oGTT) performed before and after successful trans-sphenoidal surgery, demonstrating the lack of inhibition of GH before surgery, and the inhibition of GH levels to a nadir of 0.51 ng/mL followed by physiological rebound at 180 min after surgery.

| Time (min) | t000 | t030 | t060 | t090 | t120 | t180 |
|------------|------|------|------|------|------|------|
| GH value (μg/l) |      |      |      |      |      |      |
| Before surgery (Nov. 2007) | 8.21 | 7.08 | 6.32 | 6.76 | 7.78 | 8.99 |
| After surgery (Sept. 2009) | 3.01 | NA   | NA   | 0.54 | 0.51 | 15.8 |

NA, non-available.
existence of “somato-mammotrope” cells, a cell type that stores and secretes both GH and PRL which is generated by the con-
version of somatotropes into mammotrophs during situations demanding large amounts of PRL such as lactation (Frawley and
Boocskor, 1991). Transdifferentiation has also been shown in the pituitaries of rats made hypothyroid by chemical thyroidectomy
(Horvath et al., 1990), and reversible phenotypic switching of GH and PRL gene expression has long been reported in experimental
rat pituitary tumor cells (Ivarie and Morris, 1983; Melmed, 2003).

Thus, pathogenic mechanisms of pituitary adenoma formation remain poorly understood, especially in sporadic occurrences such
as in the patient discussed here. It should also be noted that this change in the phenotype of the tumor was observed while the patient
was on long term dopamine agonist treatment, but also that therapy had been discontinued spontaneously by the patient
on several occasions over the years. One could therefore speculate that poor adherence to dopamine agonist therapy may have played
a role in this rather unusual evolution, by allowing emergence of a less differentiated adenomatous cell lineage.

In conclusion, we believe that this patient represents the first definitive documented evidence of the secondary apparition of
GH autonomous secretion within a known microprolactinoma. This phenotypic change of the adenoma was accompanied by
a change in its growth potential, suggesting some degree of de-
differentiation. Complements of investigations by immunohisto-
chemistry, DNA analyses, and clonality of tumor specimens would
have been interesting to obtain, in order to understand what was
responsible for this change. Pathogenic mechanisms implicated
in this rather unusual evolution for a microprolactinoma remain
largely unclear, and the potential role played by poor therapeutic
adherence is only speculative. However and regardless of the path-
ogenic mechanisms implicated, the clinical course of this unusual
patient emphasizes the need for long term hormonal follow up of
all pituitary adenomas.

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