The Medical Treatment of the Hypersecreting Pituitary Gland

Bernard Corenblum

ABSTRACT: Pituitary adenomas may produce local endocrine and neurological effects, as well as systemic metabolic complications due to hormonal hypersecretion. Medical therapy with pharmacological agents has been developed and is based on the neurotransmitter regulation of normal pituitary hormonal secretion. 189 patients with secretory pituitary adenomas underwent medical therapy for the hypersecretory state. 156 of these were prolactin-secreting adenomas, 16 of which were in males. The response of bromocriptine was almost universal with lowering of serum prolactin and reversal of the clinical symptoms, as well as tumor shrinkage of most large adenomas with suprasellar extension. 23 patients with acromegaly were treated with bromocriptine, with only one noting clinical improvement, and decreased tumor size in two. Five patients with Cushing’s disease were treated with cyproheptadine, with only one showing a biochemical and clinical improvement. Two patients with Nelson’s syndrome each had progressive tumor growth stabilized with cyproheptadine and bromocriptine in one, and sodium valproate in the other. There appears to be a role for medical therapy in the majority of prolactin-secreting pituitary tumors, some growth hormone secreting pituitary tumors, and selected adrenocorticotropic secreting-pituitary tumors.

RÉSUMÉ: Traitement médical de la glande pituitaire hypersécrétrice Les adénomes pituitaires peuvent produire des effets locaux endocriniens et neurologiques ainsi que des complications métaboliques systémiques dues à l’hypersécrétion hormonale. Une approche thérapeutique a été développée, basée sur la régulation par les neurotransmetteurs de la sécrétion pituitaire hormonale normale. 189 patients avec adénomes pituitaires sécrétants ont reçu un traitement médical de leur état hypersécrétoire. Dans 156 cas, il s’agissait d’adénomes sécrétant la prolactine dont 16 chez des mâles. La réponse à la bromocriptine est presque uniforme avec chute de la prolactine sérique et renversement des symptômes cliniques. Il y eut aussi un rapetissement des tumeurs, surtout des gros adénomes avec extension suprasellaire. 23 patients avec acromégalie furent traités à la bromocriptine. Ils notèrent une amélioration clinique et chez 2, la tumeur a diminué de grosseur. Cinq cas de maladie de Cushing furent traités à la cyproheptadine, un seul montant une amélioration biochimique et clinique. Une tumeur progressive s’est vue stabilisée dans sa croissance chez 2 cas de maladie de Nelson, l’un avec la cyproheptadine et la bromocriptine, l’autre avec le valproate de sodium. Il semble donc y avoir une approche médicale possible dans le traitement de la majorité des tumeurs pituitaires sécrétant la prolactine, pour certaines tumeurs pituitaires sécrétant l’hormone de croissance et pour quelques rares tumeurs sécrétant l’adrénocorticotropine.
ACTH is primarily controlled by the recently described peptide corticotropin releasing hormone. The most important neurotransmitter effects appear to be by the catecholamines and serotonin. Serotonin facilitates ACTH release, possibly via a cholinergic interneuron (Jones et al 1981). The endogenous opioids also appear to be important, likely inhibitory in nature (Stubbs et al 1978). Other roles for control have been reported for acetylcholine, melatonin, and gamma aminobutyric acid (GABA). All stimulatory and inhibitory effects and their sites of action are not well understood. These observations have led to clinical trials for ACTH secreting pituitary adenomas with the antiserotonergic and anticholinergic drug cyproheptadine, the dopamine agonist bromocriptine, and the GABA agonist sodium valproate.

This is a report of the treatment of hypersecretory pituitary adenomas with various neuropharmacologic agents over a ten year period.

**MATERIALS AND METHODS**

A total of 189 patients with secretory pituitary adenomas underwent medical therapy for the hypersecretory state (Table I). Prolactin secreting pituitary adenomas made up 156 of the subjects. 151 were treated with bromocriptine in dosage varying from 1.25 - 7.5 mg per day, and 89 have had continuous therapy under medical supervision for 5 - 10 years, as previously reported in part (Corenblum and Taylor 1983). Five were treated with pergolide. Indices of response included normalization of serum prolactin, return of menses, fertility, loss of galactorrhea in the females, and improved libido and normalization of serum testosterone in males. Tumor size was evaluated initially with pneumoencephalography, but for the past seven years by computerized tomography scanning. Any visualized tumor had its size monitored by repeated radiological evaluation as well as indirectly by visual field testing and pituitary reserve testing. Pituitary reserve was tested in response to insulin-induced hypoglycemia, TRH, and GnRH (Corenblum 1979). None of these patients had surgical or irradiation therapy prior to the onset of bromocriptine and pergolide therapy.

Twenty-six patients presented with acromegaly, 16 females and 10 males. All were treated with bromocriptine in dosages varying from 10 - 20 mg/day in four divided doses. Growth hormone secretion was assessed following glucose administration, the clinical response was assessed depending on the presenting symptoms and hand volume measurement, and tumor size changes were followed with repeated CT scanning. Six of these patients had concomitant hypersecretion of prolactin. Seventeen of 26 patients had previously undergone surgical and/or irradiation therapy for the acromegaly without clinical or biochemical cure, while nine were treated initially with bromocriptine. In 13 patients demonstrating clinical improvement, nine of the 13 subsequently subsequently underwent conventional pituitary irradiation therapy with maintenance of bromocriptine therapy, while four have remained on bromocriptine alone. All bromocriptine failures underwent irradiation therapy.

Seven patients presented with ACTH secreting pituitary adenomas. Five had Cushing’s disease, and were initially treated with cyproheptadine 24 mg/day for a minimum of three months. None of these patients had previous surgical or irradiation therapy, but four of the five subsequently underwent transphenoidal resection of the underlying pituitary adenoma. Two
Table 1: 189 Patients With Demonstrated or Presumed Pituitary Adenomas Undergoing Medical Therapy

<table>
<thead>
<tr>
<th>Hypersecretory State</th>
<th>Number</th>
<th>Sex distribution (F/M)</th>
<th>Prior Therapy</th>
<th>Medical Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactinoma</td>
<td>156</td>
<td>140/16</td>
<td>None</td>
<td>151 Bromocriptine</td>
</tr>
<tr>
<td>Acromegaly</td>
<td>26</td>
<td>16/10</td>
<td>None 10 Surgery</td>
<td>Bromocriptine</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4 Surgery + Irradiation</td>
<td>Cyproheptadine</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3 Irradiation</td>
<td></td>
</tr>
<tr>
<td>Cushing's disease</td>
<td>5</td>
<td>4/1</td>
<td>None</td>
<td>Cyproheptadine</td>
</tr>
<tr>
<td>Nelson's syndrome</td>
<td>2</td>
<td>2/0</td>
<td>Surgery</td>
<td>Cyproheptadine + Bromocriptine</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Surgery + Irradiation</td>
<td>Sodium Valproate</td>
</tr>
</tbody>
</table>

patients presented with Nelson's syndrome, eight and 17 years following bilateral adrenalectomy. One had pituitary irradiation five years following the adrenalectomy, once the onset of hyperpigmentation was noted. This patient had one attempted surgical removal, and after this failed was treated with cyproheptadine, 24 mg/day, and bromocriptine 5 mg/day. The second patient had irradiation and then two surgical attempts for removal, yet continued to demonstrate an invasive ACTH secreting adenoma. She was then treated with sodium valproate, 400 mg/day.

RESULTS

The 156 patients with prolactin secreting pituitary adenomas consisted of 140 women and 16 men, varying from age 15 to 46. The 140 women were divided into three groups, depending on the size of the tumor (Table 2). Twenty-one were classified with macroadenomas according to gross enlargement of the sella turcica and the adenoma being at least 1 cm in size as assessed by radiological visualization. The 78 women with microadenomas had distortion of sellar shape or normal sella turcicas, but CT scan demonstrated an adenoma less than 1 cm in size. The 41 classified with normal sellas had no radiological abnormality detected, but the clinical diagnosis was of a prolactin secreting pituitary tumor on the basis of clinical impression, exclusion of other causes of hyperprolactinemia, and failure to demonstrate any release of prolactin in response to an injection of thyrotropin releasing hormone (Corenblum 1979). The patients with macroadenomas included three presenting with primary amenorrhea. Mass signs were clinically evident in eight patients with macroadenomas, all with visual field abnormalities, and seven of these with headaches. Of these eleven, six had normal pituitary reserve to stimulatory testing. Of the six with normal pituitary reserve, all six had return of menses, and of four desirous of pregnancy, two did conceive. Of the five with some degree of abnormal pituitary reserve, three of the five had return of menses, and none were desirous of pregnancy. Ten patients did not have any clinical signs of an underlying mass. Seven of these had normal pituitary reserve to stimulatory testing, all seven had return of menses, and of the five desirous of pregnancy, three did conceive. The three with abnormal pituitary reserve had two of the three return to normal menstrual activity, and none were desirous of pregnancy. In these 21 patients the tumor was clearly visualized, and tumor regression was noted in seventeen of the twenty-one. Eleven of them no longer had any suprasellar extension that was previously present. Abnormal visual fields were documented in eight, with seven returning to normal. The other did not fully return to normal, despite the fact that the tumor had regressed. Of the eight patients with abnormal pituitary reserve, six of the eight had normalization

Table 2: Response of 140 Women With Prolactinomas to Medical Therapy

| A. Macroadenomas (n = 21)    | Normal pituitary reserve 76 (74 return of menses, 34/39 pregnant) |
| B. Microadenomas (n = 78)   | Normal pituitary reserve 2 (1 return of menses, 1/2 pregnant) |
| C. Normal Sellas (n = 41)    | Normal pituitary reserve 40 (40 return of menses, 24/28 pregnant) |
| D. Total (n = 140)           | Normal pituitary reserve 129 (127 return of menses, 60/71 pregnant) |

* Figure in brackets indicates number of patients with this symptom.
of pituitary reserve within six months of therapy with bromocriptine.

The 78 women with microadenomas included 76 with normal pituitary reserve, and 74 of these 76 had normal return of menses. Of 39 desirous of pregnancy, 34 did conceive. Two patients with abnormal pituitary reserve included one who did have return of menses and became pregnant, and the other showed no return of gonadal function despite normalization of serum prolactin following one year of bromocriptine therapy, or after the addition of clomiphene citrate.

Of the 41 women with normal sella turcica, 40 demonstrated normal pituitary reserve and all 40 had return of menses, and 24 of 28 became pregnant. One patient with abnormal pituitary reserve failed to have return of menses, and demonstrated progressive hypothalamic insufficiency resulting from a granulomatous infiltrate which became evident after a follow-up over two years. A biopsy of the hypothalamus demonstrated histiocytosis.

Of the three patients with primary amenorrhea, two of the three showed marked regression in size of the mass, and all three spontaneous onset of menses (Fig. 1a and Fig. 1b).

Eighty-nine of these patients have been followed for at least five years without any long-term problems arising that were not obvious within the short-term institution of therapy.

Of the 16 males, eleven had macroadenomas, and five microadenomas (Table 3). The 11 men with macroadenomas included seven with suprasellar extension of the mass, six of whom had mass signs. Six demonstrated regression of the suprasellar extension. Of the five with abnormal visual fields, four were normalized. One patient, who did not demonstrate any regression of the mass or normalization of the visual fields, underwent surgery for removal of a cystic adenoma. Headaches improved in all nine with complaints of this. There was an increase of serum testosterone and improvement of decreased libido in 11 of the 16 men, while five required testosterone supplementation to continued bromocriptine therapy.

The group as a whole had normalization of the serum prolactin in 138 of 156 patients. Those that did not completely normalize, generally had an improvement in their gonadal status. All women with symptomatic galactorrhea had it markedly lessen or disappear so that it was no longer clinically significant. Four patients requested discontinuation of bromocriptine therapy after successful therapy for a minimum of one year, all because of cost or personal preference. No long-term complications of note occurred, although minor chronic nasal stuffiness was volunteered in 21 patients after direct questioning.

The twenty-six patients with acromegaly varied in age from 19 to 67 and included 16 females, six of whom had concomitant hyperprolactinemia, and two patients had the multiple endocrine adenomatosis syndrome. Nine were treated with bromocriptine as the primary form of therapy, ten following unsuccessful surgery, four following unsuccessful surgery and irradiation, Table 3: Response of 16 Men With Prolactinomas to Medical Therapy

<table>
<thead>
<tr>
<th>A. Macroadenomas (n = 11)</th>
<th>B. Microadenomas (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normalize visual field abnormalities 4(5)*</td>
<td>Loss of headaches 2(3)</td>
</tr>
<tr>
<td>Decrease tumor size 10</td>
<td>Improved libido 3(5)</td>
</tr>
<tr>
<td>Loss of headaches 7(8)</td>
<td></td>
</tr>
<tr>
<td>Improved libido 8(11)</td>
<td></td>
</tr>
</tbody>
</table>

* Figure in brackets indicates number of patients with this symptom.
and three following unsuccessful irradiation at least five years
prior to the institution of bromocriptine therapy (Table 4). Clinical
improvement was noted in 13 patients, which included
decreased greasiness of the skin, decreased sweating, fewer
headaches, more energy, onset of menses in amenorrheic women,
regression of soft tissue swelling and symptoms of carpal tunnel
syndrome, and improvement of diabetes mellitus. Of seven
insulin-requiring diabetics, four clearly demonstrated decreased
insulin requirements, but none discontinued insulin. Biochemical
normalization of growth hormone was not noted in any patient,
with partial normalization demonstrated in six, all measured
after glucose administration. Clear decreases in tumor size
were noted with CT scanning in two patients, both with
concomitant hyperprolactinemia. None of the patients had any
signs of an underlying mass at the time of onset of medical
therapy. The bromocriptine failures subsequently underwent
irradiation treatment, if this was not previously administered.

Five patients with Cushing's disease had primary treatment
with cyproheptadine for a minimum of three months (Table 4).
A biochemical remission with lessening of the clinical signs was
noted in only one patient. This included normalization of the
urinary free cortisol, and a normal response to insulin-induced
hypoglycemia. This patient has remained on cyproheptadine
and has undergone pituitary irradiation. The other four patients
without any clinical or biochemical remission have subsequently
undergone transphenoidal resection of the underlying pituitary
adenoma. Two of these four normalized their biochemical
parameters and the remaining two subsequently required bilateral
adrenalectomy. The two patients with Nelson's syndrome both
presented with invasive ACTH secreting pituitary adenomas.
The first patient had this diagnosis for seventeen years, and had
previously unsuccessful conventional irradiation and surgery.
It response to the combination of cyproheptadine and bromo-
criptine, there was a decrease of the serum ACTH concentration,
decreased clinical pigmentation, loss of headaches, less depres-
sion, and cessation of tumor growth that was previously
documented to be progressing with repeated CT scanning.
With discontinuation of the medication there was a return
of headaches and definite tumor growth was visualized. With
reintroduction of bromocriptine and cyproheptadine, she has
again stabilized. The second patient had two previous unsuccessful
surgical resections, and pituitary irradiation eight years prior to
therapy, with CT scan evidence of continued tumor growth. In
response to sodium valproate for one year there has been loss
of headaches and cessation of previously documented tumor
growth as shown by repeated CT scanning.

**DISCUSSION**

Pituitary adenomas may have local effects, but if there is
hypersecretion of a hormone, then there may be significant
metabolic and cosmetic effects, especially seen in Cushing's
disease and acromegaly. These metabolic effects may decrease
longevity, mainly through cardiovascular manifestations.
Hyperprolactinemia may result in hypogonadism with possible
long-term problems such as osteoporosis, but generally longevity
is not affected. It is for these differences in prognosis that
ablative therapy with surgery and/or irradiation is generally the
treatment of choice for Cushing's disease and acromegaly,
whereas invasive therapy or medical therapy may be individual-
ized for any given patient with a prolactin-secreting pituitary
adenoma. Because medical therapy is not curative for Cushing's
disease or acromegaly and tumor regression only occasionally
occurs, medical therapy remains as an adjunct. Medical therapy
may be used to prepare a patient for surgery, for cases of failed
ablative therapy, or for the elderly patient. Prolactin-secreting
adenomas appear to do well both with dopamine agonists and
with surgery, and even though medical therapy is not curative,
tumor regression is almost expected for the large adenomas.
There are reports of spontaneous "cure" following bromocriptine-
duced pregnancies and two such cases are in this reported
series. Because long-term follow-up of surgical results may
have a recurrence of 15% (Faglia et al 1983) to 55% (Serri et
al 1983), there may not be a long-term major advantage over
medical therapy, especially since the disorder does not have
major metabolic complications, but rather results in reversible
hypogonadism. Only infrequently are there problems with
growing tumor progression or apoplexy (Breidahl et al 1983).

The control of prolactin is unique in the pituitary gland, as it
is predominately under inhibitory control. Bromocriptine
was developed as a dopamine agonist, and will normalize prolactin
in 80 - 90% of patients with pituitary adenomas (Friesen and
Tolis 1977), with resulting reversal of associated hypogonadism
and galactorrhea. Bromocriptine is absorbed orally and reaches
a maximum blood level after 90 minutes, but the maximum
effect on suppression of prolactin secretion occurs several
hours later. The initial side effects of nausea and light-headedness
may be minimized by beginning with a small dose taken with
meals or at bedtime, and building up over two or three weeks.
Chronic symptoms may include nasal stuffiness, Raynaud's
syndrome, and constipation. Neuropsychiatric abnormalities
usually occur only with high doses such as seen with the treatment
of Parkinsonism, but may rarely occur with low doses. Indications
for therapy are infertility, arrested or delayed puberty, galactor-
hea, decreased libido, presence of a tumor mass, return of
estrogenization or androgenization status, and to inhibit the
progression of osteopenia. Bromocriptine does not appear to
be teratogenic (Turkal et al 1982). It appears that bromocriptine
decreases prolactin secretion initially by inhibiting exocytotic
events, and later by decreasing intracellular prolactin content,
which infers decreased synthesis. The regression of tumor size
was first noted in animals, and later in humans (Corenblum et

| Table 4: Response of GH and ACTH Secreting Adenomas to Medical Therapy |
|-----------------------------|-----------------------------|
| Previous Therapy | Clinical Improvement |
| A. Acromegaly |  |
| None | 6(9) * |
| Surgery | 4(10) |
| Surgery + irradiation | 1(4) |
| Irradiation | 2(3) |
| Total | 13(26) |
| B. Cushing's disease |  |
| None | 1(5) |
| C. Nelson's syndrome |  |
| Surgery | 0(1) |
| Surgery + irradiation | 0(1) |
| Stabilize adenoma growth | 2(2) |

* Figures in brackets indicate total number of patients in each category.
Decreased tumor size has also been seen with other dopaminergic therapy had any worsening of the success rate of surgery (Faglia et al. 1983). The decrease in size of the mass appears to result in part from decreased cell volume (Tindall et al. 1982) without any net cell loss, and appears to be reversible. There is a regularly observed decrease in the size of large adenomas with suprasellar extension (Corenblum and Hanley 1981) that are rarely cured with surgery. The long-term effects of bromocriptine on tumor size have yet to be reported over a large series, but this study appears to support that initial regression in size is maintained with continued bromocriptine therapy for up to ten years. It has been suggested that therapy for more than one year results in fibrosis of the tumor resulting in more difficult tumor removal at the time of surgery (Landolt et al. 1982), while short term therapy with bromocriptine may improve surgical results, presumably by producing regression of tumor size. Another report failed to find that previous bromocriptine therapy had any worsening of the success rate of surgery (Faglia et al. 1983).

Other dopamine agonists such as pergolide appear to be very similar to bromocriptine and have the advantage of being longer acting, or producing a response in patients who are otherwise resistant or intolerant to bromocriptine (Franks et al. 1983). Decreased tumor size has also been seen with other dopaminergic agonists (Kendall-Taylor et al. 1982). Tamoxifen, an estrogen receptor antagonist, has been used to decrease prolactin when added to bromocriptine with patients resistant to bromocriptine alone (Volker et al. 1982).

Metergoline has been used as a serotonin antagonist, although it has some intrinsic dopamine agonist activity. It may improve the clinical and biochemical abnormalities in hyperprolactinemic patients, but appears to have similar side effects as bromocriptine without the same predictable results, and thus does not have any real advantage.

It is the author’s opinion that large prolactin-secreting pituitary tumors generally have an unsatisfactory response to surgery, in that rarely is the patient rendered endocrinologically normal, the frequency of hypopituitarism is relatively high, and the success rate of total tumor removal is low. In such patients, pituitary irradiation is too slow to adequately treat extrasellar complications. For this reason, bromocriptine is probably the treatment of choice and if no response to medical therapy for six months is demonstrated, including regression of mass signs, surgical therapy may then be performed. Tumor regression is usually apparent in 2 - 4 weeks, so a short trial is probably informative, but in some cases it takes several months to see tumor regression. The fear that the pituitary tumor may continue to grow despite successful decrease of symptoms of hyperprolactinemia with bromocriptine is more theoretical than real, although there have been occasional reports of pituitary tumor growth despite continued bromocriptine therapy (Breidahl et al. 1983). The author has never observed such an occurrence, nor a microadenoma enlarging to be classified as a macroadenoma. Tumor growth may occur during a pregnancy if the bromocriptine has been discontinued, but regresses with reinstitution of the drug.

The treatment of choice for acromegaly remains as ablative therapy, either surgery or irradiation, with the use of bromocriptine as an adjuvant. Supervoltage external irradiation is usually successful, but may take many years, and has a problem of hypopituitarism and possibly some progressive visual failure (Atkinson et al. 1979). Neural damage appears to be related to the dose, and inversely related to the number of treatments, so presently 4,500 to 5,000 rads are administered in 20 - 25 fractions over 35 days.

The paradoxical response to dopamine in suppressing growth hormone and TRH in stimulating growth hormone, in patients with acromegaly, appears to be due to a direct effect of these drugs on the pituitary tumor, based on *in vitro* studies, as well as by the direct effect of a dopamine infusion in decreasing growth hormone. Furthermore, these qualitatively abnormal responses disappear after successful surgery. It is possible that there is a common stem cell for both prolactin and growth hormone cells, and growth hormone secreting adenomas may represent a dedifferentiation to a common cell type. The suppressive effects of bromocriptine on growth hormone secretion in patients with acromegaly are shorter in duration that that seen with prolactin adenomas. For this reason, bromocriptine should be administered four times a day for the treatment of acromegaly, as opposed to twice for prolactin adenomas. Furthermore, the dosage in acromegaly is higher than with prolactin adenomas, averaging about 20 mg per day, but occasionally reaching up to 60 mg per day. These doses give rise to more common side effects, including hypotension, nausea, constipation, dryness of the mouth, digital vasospasm, and nasal stuffiness. Such problems may be present in about one-third of patients and occasionally may require discontinuation of the medication. Bromocriptine restores serum growth hormone levels to normal in only a minority of patients, yet clinical improvement is frequently observed, one study showing an improvement in 97% (Wass et al. 1977). Clinical indices of improvement are a decrease in hand, foot, and ring size, decreased sweating, loss of headaches, improved libido, and improved glucose tolerance. In one study, glucose intolerance became normal in 65% of patients (Wass et al. 1977). The clinical improvement has been noted in repeated studies (Wass et al. 1983), despite the frequent failure to normalize serum growth hormone concentrations. It has been postulated that bromocriptine preferentially decreases the biologically active growth hormone, the monomeric fraction. It is also postulated that there is a peripheral action in decreasing somatomedin generation without any change in growth hormone levels (Wass et al. 1983). It was initially thought that only the acromegalic who acutely suppressed with bromocriptine or released hormone to TRH would respond to long-term bromocriptine therapy, but recent follow-up has shown that over 75% of all patients demonstrate a clinical response. Clinical observations have noted that patients are more likely to respond if there is concomitant hypersecretion of prolactin, indicating a mixed adenoma (Corenblum et al. 1976). There have been reports of regression of tumor size (Atkinson et al. 1979) usually in patients with concomitant hyperprolactinemia, and this study is consistent with those observations.

The role of bromocriptine for the treatment of acromegaly is to complement patients treated with external irradiation, or in patients who are surgical failures, but may be considered as primary therapy in elderly patients, or in other selected patients. For this reason with any patients that demonstrate a clinical response to bromocriptine used as the primary form of therapy, we recommend a course of external irradiation while the patient is maintained on the bromocriptine.

ACTH secreting pituitary tumors remain the most complicated to treat, as well as the most important, due to significant peripheral metabolic complications along with potential central neurological
treatment of choice. Should this be unsuccessful, or should the adenoma, then either bilateral adrenalectomy or medical therapy along with external irradiation should be considered. Several neuropharmacological agents have been utilized, all based on the multitude of neurotransmitters known to effect ACTH secretion. Agents which affect adrenal synthesis of cortisol, such as metyrapone, aminoglutethimide, and the adrenolytic drug o,p-DDD have all been utilized to prepare the patient for surgery, or to await results of irradiation. A complication of bilateral adrenalectomy is continued growth of an ACTH secreting tumor, Nelson’s syndrome. Prophylactic pituitary irradiation is usually recommended, and this may decrease the incidence, but may not entirely prevent the occurrence of Nelson’s syndrome (Moore et al 1976). Cushing’s disease may result from an ACTH secreting pituitary adenoma or corticotroph hyperplasia. This may result from either inappropriate secretion of corticotropin-releasing hormone or else autonomous secretion by the corticotrophs, or possibly some inter-relation between these two factors. There is experimental evidence to suggest that serotonin and acetylcholine facilitate the release of corticotropin-releasing hormone. Cyproheptadine has antiserotonergic and anti-cholinergic activities, and has been shown to blunt the enhanced ACTH secretion that occurs in response to hypoglycemia (Plonk et al 1974), and metyrapone (Plonk et al 1976). Cyproheptadine does not appear to have any direct action on ACTH secretion in dispersed pituitary cells, but may have this action in adenomatous cells, and this effect may be abolished with the addition of serotonin (Ishibashi and Yamaji 1981). In approximately 100 patients, remission of Cushing’s disease occurred in 35 - 50% (Krieger 1983), and the longest reported successful therapy is up to five years. Patients who respond generally show a clinical and laboratory remission in two to three months of therapy, and remission for up to three years after discontinuing cyproheptadine has been reported. Unfortunately, relapse invariably occurs, occasionally while still receiving cyproheptadine. The utilization of cyproheptadine and irradiation therapy has allowed cyproheptadine to be discontinued within four months, with continued clinical remission. Unfortunately, there are also many unsuccessful reports of therapy with cyproheptadine, and the author’s small experience finds the beneficial response is noted in a minority of patients. The side effects of therapy include somnolence and hyperphagia and appears to be most marked in children. Along with a clinical remission, there is usually a return of normal biochemical parameters. This includes a return of normal diurnal rhythm, normal serum and urinary cortisol concentrations, and a return of normal suppression with administration of dexamethasone. Women may notice return of normal menses, lessening of hirsutism, loss of weight, and decreased facial rubor. Psychiatric manifestations may lessen. Cortisol concentrations begin to decrease in about two months, with clinical remission occurring in four to six months, and return of normal cortisol secretory dynamics in about six months.

The utilization of bromocriptine has its origin in the observation that dopamine may be inhibitory to the secretion of ACTH. Such patients may possibly be selected by the administration of bromocriptine (Lamberts et al 1982), and it has been suggested that such responsive adenomas or hyperplasia arise in the intermediate area of the pituitary where direct innervation of corticotrophs may occur from dopaminergic fibres. A small series has found that bromocriptine will improve the clinical and biochemical manifestations of Cushing’s disease in six of 13 patients, but it may not be long maintained (Lamberts et al 1980). We have used bromocriptine for at least three months in four patients with Cushing’s disease without any clinical or biochemical response.

GABA appears to decrease corticotropin-releasing hormone secretion (Jones et al 1976), and may be involved in the cortisol feedback action (Avs and Stark 1978). Sodium valproate decreases GABA transaminase which will increase GABA biological activity. Furthermore, sodium valproate may also have an effect on the post-synaptic GABA receptor, and again enhance GABA biological activity. Published reports find sodium valproate improving some cases with Cushing’s disease (Koppezaar et al 1982), but not others (Alloio et al 1982). One case reported was of a response to both cyproheptadine and valproate (Koppezaar et al 1983). In ten patients with Nelson’s syndrome, ACTH concentrations have decreased in response to sodium valproate (Dornhorst et al 1983). Tumor regression may occur in that loss of headaches and diabetes insipidus have been reported (Elras et al 1981; Jones et al 1981). Most reports to date are preliminary and continue to have the problem of unpredictability of response.

The opioid antagonist naloxone has been shown to decrease ACTH secretion in Nelson’s syndrome (Tolos et al 1982), but long acting opioid antagonists have yet to be utilized in clinical trials.

It appears that only a minority of patients with ACTH secreting pituitary tumors respond to medical therapy, and these patients cannot be predicted. The response does not appear to be related to the size of the tumor, not to the severity of the clinical symptoms. Due to the side effects and inconsistency of response, neuropharmacological agents remain as the second line of therapy, as an adjunct to radiotherapy, in failed surgical therapy, or to prepare a patient for surgery.

We have not had any experience with primary TSH secreting adenomas, but one report indicates that dopaminergic agents such as L-dopa and bromocriptine may be useful (Horn et al 1976). In both TSH secreting tumors and FSH secreting tumors related to long-standing primary target gland failure, there has been regression of both size and secretion of these adenomas in response to replacement of the feedback hormone. Three such cases have been treated in this centre, but are not reported in this study. Non-secreting pituitary tumors have had either isolated reports (Johnston et al 1981), as well as a series of cases responding to bromocriptine (Wollesen et al 1982). These studies have demonstrated decreased size of non-secreting pituitary tumors, but the decrease does not appear to be as rapid as what is seen with prolactinomas. Furthermore, they may need to be treated for at least one year to see any radiological improvement. We do not have any experience in treating non-secreting pituitary tumors with neuropharmacological agents, and ablative therapy remains the treatment of choice, if clinically indicated. Because some of these tumors may be prolactinomas that are not releasing the synthesized hormone, a trial of bromocriptine when ablative therapy fails may be attempted. Immunostaining of any surgically removed tissue for prolactin is useful to predict such a response.

In summary, the utilization of neuropharmacological agents...
based on neurotransmitter physiology has been utilized to treat the hypersecretory state and tumor growth of pituitary adenomas. Present indications vary from the first line treatment of choice for both prolactin secreting microadenomas and probably microadenomas, and as adjuncts to ablative therapy for treatment of acromegaly and ACTH secreting pituitary adenomas.

REFERENCES


