HYPERPROLACTINEMIA ASSOCIATED WITH PSYCHIATRIC DISORDERS

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Abstract  Hyperprolactinemia may be associated with psychiatric disorders in the context of two scenarios: antipsychotic-induced hyperprolactinemia and psychiatric disorders arising from the medical treatment of hyperprolactinemia. Both situations are particularly common in psychiatric and endocrine clinical practice, albeit generally underestimated or unrecognized. The aim of this article is to provide tools for the diagnosis and treatment of hyperprolactinemia associated with psychiatric disorders to raise awareness, especially among psychiatrists and endocrinologists, so that these professionals can jointly focus on the appropriate management of this clinical entity.

Key words: hyperprolactinemia, antipsychotics, dopamine agonists, impulse control disorders

KEY POINTS

• Hyperprolactinemia may be associated with psychiatric disorders in the context of two scenarios: antipsychotic-induced hyperprolactinemia and psychiatric disorders arising from the medical treatment of hyperprolactinemia.
• It is important that psychiatrists and endocrinologists be aware and informed to approach together the management of hyperprolactinemia associated with psychiatric disorders in its different forms in order to avoid serious and devastating complications for the patient.

This review will discuss the relationship between hyperprolactinemia and psychiatric disorders from two different approaches: hyperprolactinemia secondary to the treatment of psychiatric disorders and psychiatric disorders induced by the medical treatment of hyperprolactinemia. In both scenarios dopamine plays a leading role.

Dopamine is a neurotransmitter particularly important, involved in daily brain functioning (such as the control of motor function, motivation, and learning) and in several common disorders.

Three main dopaminergic pathways are described in the central nervous system1:

- The nigrostriatal pathway, involved in control of motor function
- The mesocorticolimbic pathway, related to the so-called reward system, which regulates behaviour, pleasure and addiction
- The tuberoinfundibular dopaminergic pathway (TIDA), responsible for prolactin secretion

Alterations in these pathways will result in a variety of entities ranging from motor deficits (Parkinson’s disease), addictive and impulse control disorders and hyperprolactinemia1.

Prolactin physiology

Prolactin (PRL) is a 199-aa polypeptide hormone, with a molecular weight of 23 kDa, synthesized by lactotrophs in the adenohypophysis. PRL belongs to the large so-
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The recognition of the PRL receptor expression at numerous extra-pituitary sites including endometrial decidua, breasts, brain, ovaries, prostate, endothelial cells, lymphocytes, skin, adipose tissue and cochlea has opened up a wide range of potential functions. In any case, the most recognized role of PRL in humans is to induce lactation. The 23 kDa monomeric form is the main circulating variant of PRL (85%) and responsible for mediating its physiological actions. This variant acts through a membrane receptor belonging to the superfamily of type I cytokine receptors that are characterized by having a single transmembrane domain that transduces signals after phosphorylation of cytoplasmic kinases.

Other circulating forms of PRL are the “big” PRL and the “big big” PRL (macroprolactin). Big PRL is the dimer of monomeric form (50 kDa) and the “big big” PRL comprises high molecular (>150 kDa) complexes formed by 23 kDa PRL and IgG autoantibodies that can be found in different degrees in immunoassays for PRL. It seems that these forms of PRL have minimal or any biological activity.

Secretion of PRL by the adenohypophysis has a circadian rhythm with higher levels during sleep and lower during wakefulness. Increased levels are also seen during ovulation. Synthesis and secretion of PRL are both regulated by releasing and inhibiting factors. PRL inhibiting factors include dopamine, gamma-aminobutyric acid (GABA), and somatostatin. Dopamine is the main regulator of PRL secretion through its D2 receptor, and unlike other pituitary hormones, the inhibiting component is dominant. Consequently, PRL level will increase if the blood flow of the pituitary stalk is altered or if dopamine synthesis is inhibited or in case of receptor blockade.

Among the PRL releasing factors can be mentioned TRH (thyrotropin-releasing hormone), VIP (vasoactive intestinal peptide), serotonin, opioid peptides, GHRH (growth hormone-releasing hormone), neurotensin, oxytocin, vasopressin, galanin, and estrogens. Dopamine is released through the axons of the TIDA pathway in the median eminence. These axons are projected from the dorsomedial portion of the arcuate nucleus and the lower portion of the ventromedial nucleus of the hypothalamus.

There are five types of dopamine receptors that belong to the family of G-protein-coupled receptors. D1 and D5 receptors couple to Gs protein and stimulate the synthesis of cyclic AMP. Meanwhile, D2, D3 and D4 receptors couple to Gi protein and inhibit the synthesis of cyclic AMP and inositol phosphate metabolism, decreasing the intracellular calcium mobilization and transportation through its channels.

The dopamine action on the adenohypophysis is exerted through the D2 and D4 receptors located in the lactotroph membrane, inhibiting - tonically and directly - PRL gene transcription, PRL synthesis and secretion and lactotroph proliferation.

Hyperprolactinemia

It is the most common endocrine disorder of the hypothalamic-pituitary unit. The causes of hyperprolactinemia can be physiological, analytical, pathological and pharmacological (Table 1)

Serum PRL levels are higher in women than in men and hyperprolactinemia is diagnosed when the PRL levels are higher than 25 ng/ml in females and higher than 20 ng/ml in males.

Hyperprolactinemia occurs at any age and its prevalence varies around 1% regarding adult population (0.2% in men and 1% in women), and 5% in cases of women with infertility. Hypogonadotropic hypogonadism and galactorrhea are the most characteristic signs and symptoms of PRL excess and will be discussed below.

Hyperprolactinemia secondary to the treatment of psychiatric disorders

Drug-induced hyperprolactinemia is the most frequent cause of non-physiological hyperprolactinemia. In fact, an epidemiological study in Scotland evaluated 1301 cases of hyperprolactinemia (excluding pregnancy) and the results showed that 45.9% of the cases were drug-induced.

A considerable number of drugs cause PRL elevations through different mechanisms that are listed below:

- Increased transcriptional activity of PRL gene: estrogens
- Dopamine receptor antagonism: risperidone, haloperidol, phenothiazines, metoclopramide, domperidone, sulpiride
- Dopamine depletion: reserpine, methyldopa
- Hypothalamic dopamine synthesis inhibition: verapamil, heroin, morphine
- Serotonin reuptake inhibition: opiates, fluoxetine, tricyclic antidepressants, monoamine-oxidase inhibitors

Usually the range of drug-induced hyperprolactinemia varies from 25 to 100 ng/mL. However, levels > 100 ng/mL can be observed, especially with the use of neuroleptics (risperidone, haloperidol, chlorpromazine) and antiemetics (metoclopramide and domperidone). Table 2 shows the comparative range of hyperprolactinemia due to different etiologies.

As regards in particular to antipsychotics, butyrophenones and phenothiazines may induce PRL excess in 50-90% of treated patients, while risperidone in 70-100% of cases. In relation with risperidone, PRL serum levels even greater than 300 ng/ml have been reported.
TABLE 1.– Causes of hyperprolactinemia

- Physiological causes
  - Pregnancy
  - Breast lactation
  - Breast stimulation
  - Stress
  - Exercise
- Analytical causes
  - Macroprolactinemia
- Pathological causes
  - Prolactinomas and mixed adenomas
  - Hypothalamic and pituitary stalk pathologies (macroadenoma, hypophysitis, granulomatous disease, Rathke’s cleft cyst, radiation and/or trauma, other tumors including craniopharyngiomas, germinomas and metastases)
  - Systemic diseases: hypothyroidism, adrenal insufficiency, chronic kidney disease, ovarian polycystosis, cirrhosis, pseudocyesis, seizures, cranial radiation, neurogenic or traumatic thoracic lesions, herpes zoster
  - Ectopic prolactin secretion: dermoid ovarian cysts, hypernephroma, bronchogenic carcinoma
  - Genetics: mutation with loss of function of the PRL receptor
  - Idiopathic diseases or conditions
- Pharmacological causes
  - Dopamine antagonists (neuroleptics, antiemetics)
  - Dopamine synthesis inhibitors (methyldopa)
  - Other medications (antidepressants, verapamil, estrogens, phenytoin and opiates)

TABLE 2.– Comparative range of hyperprolactinemia in relation to different etiologies

<table>
<thead>
<tr>
<th>Etiology</th>
<th>Typical prolactin levels (ng/ml)</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microprolactinoma</td>
<td>100-250</td>
<td>PRL &lt;100 up to 25%; PRL &gt; 250 in ~10% of cases</td>
</tr>
<tr>
<td>Macroprolactinoma (MAC)</td>
<td>200-1000</td>
<td>PRL &lt; 100 only in cases of cystic MAC or due to Hook Effect; PRL &gt; 1000 on giant MAC (&gt; 4 cm)</td>
</tr>
<tr>
<td>Non-functioning pituitary adenomas (NFPA)</td>
<td>25-100</td>
<td>PRL between 100 and 250 in up to 20% of cases; No cases of PRL &gt; 250 in patients with confirmed NFPAs without macroprolactinemia</td>
</tr>
<tr>
<td>Drugs</td>
<td>25-100</td>
<td>PRL between 25 and 100 in ~ 60-70% of cases; PRL between 100 and 250 in ~30% of cases; PRL &gt; 500 is very rare; PRL &gt; 250 in 5% of cases; PRL &gt; 300 in 1-2%</td>
</tr>
<tr>
<td>Primary hypothyroidism</td>
<td>25-250</td>
<td>PRL between 100 and 250 in up to 15% of cases; between 250 and 300 in 1-2%</td>
</tr>
<tr>
<td>Macroprolactinemia</td>
<td>25-250</td>
<td>PRL &gt; 250 in 5% of cases; PRL &gt; 500 is extremely rare, and usually restricted to concomitant monomeric HPRL</td>
</tr>
</tbody>
</table>

PRL: prolactin; MAC: macroprolactinoma; NFPs: non-functioning pituitary adenomas; HPRL: hyperprolactinemia

Different classes of antidepressants, including monoamine oxidase inhibitors, tricyclic antidepressants, and selective serotonin reuptake inhibitors induce hyperprolactinemia mainly via serotoninergic pathways. Hyperprolactinemia is typically mild and rarely symptomatic\(^2\),\(^4\).

Mechanisms of antipsychotic-induced hyperprolactinemia

Antipsychotics can be classified into those with great potential for PRL elevation, and those that modify PRL to a lesser degree or even decrease its level. Explana-
tions for these differences include variations in the D2 receptor dissociation rate, ability to cross the blood-brain barrier, presence of D2 receptor polymorphisms (Taq1A genotype)\textsuperscript{19}, and degree of serotoninergic inhibition\textsuperscript{4}. The effects of antipsychotics on prolactin levels are variable as can be seen in Table 3.

In addition, the tenor of antipsychotic-induced hyperprolactinemia also depends on the affinity rate and the duration of the D2 receptor blockade\textsuperscript{20, 21}. Haloperidol, for example, blocks D2 receptors for a period of more than 24 hours compared to quetiapine, which shows a milder and more transient effect. Besides, quetiapine and clozapine are rapidly dissociated from D2 receptors and both act as antagonists of 5-HT2A serotonin receptors, causing modest increases in PRL levels\textsuperscript{21, 22}.

In contrast to typical antipsychotics, the atypical ones induce milder PRL elevations due to less affinity for the D2 receptor and greater blockade on the 5-HT2A receptor. The exception to the rule includes risperidone, paliperidone and amisulpiride that usually cause significant PRL elevation\textsuperscript{17, 18}. In fact, risperidone, as expressed before, has the greatest impact due to its high antagonistic activity on D2 and 5HT2 receptors\textsuperscript{17, 18, 20, 22}. Both oral and depot forms of risperidone and paliperidone increase PRL in a dose-dependent manner.

The threshold for the D2 receptor blockade at which the increase of PRL level occurs would range between 65-80\%\textsuperscript{21}. PRL levels increase within a few days after initiation of antipsychotic medication, usually within the first 2 to 3 weeks with a wide range of 7 to 75 days\textsuperscript{6}.

Aripiprazole deserves a special mention for its particular mechanism of action. It behaves as a partial D2 and 5-HT1A receptors agonist Consequently, the drug does not cause a significant increase of PRL concentration and even reduces it when associated with another antipsychotic\textsuperscript{23, 24}.

Finally, estrogens enhance PRL secretion via different mechanisms including upregulation of PRL gene expression, raise sensitivity to TRH, downregulation of pituitary dopamine receptors expression, and stimulation of lactotroph hyperplasia\textsuperscript{4}. For this reason, antipsychotic-induced hyperprolactinemia is more common in adolescents and premenopausal women\textsuperscript{17, 18}.

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|l|}
\hline
Antipsychotics & Increased PRL & Antidepressants & Increased PRL \\
\hline
Typical & & & \\
- Chlorpromazine & +++ & - Clomipramine & +++ \\
- Haloperidol & +++ & - Amitriptyline & + \\
- Zuclopenthixol & +++ & - Desipramine & + \\
Atypical & & & \\
- Risperidone & +++ & - Nortriptyline & - \\
- Paliperidone & +++ & - Imipramine & CR \\
- Amisulpiride & +++ & - Maprotiline & CR \\
- Molindone & ++ & - Amoxapine & CR \\
- Quetiapine & + & - Pargyline & +++ \\
- Olanzapine & + & - Clorgyline & +++ \\
- Aripiprazole & 0 & - Tranylcypromine & ± \\
- Clozapine & 0 & - SSRI: & \\
- Ziprasidone & 0 & - Fluoxetine & CR \\
& & - Paroxetine & ± \\
& & - Citalopram & ± \\
& & - Fluvoxamine & ± \\
& & Other: & \\
& & - Venlafaxine & 0 \\
& & - Trazodone & 0 \\
& & - Nefazodone & 0 \\
& & - Bupropion & 0 \\
\hline
\end{tabular}
\caption{Effects of antipsychotics on prolactin levels}
\end{table}

\textsuperscript{MAOIs: monoamine oxidase inhibitors; SSRI: selective serotonin reuptake inhibitor.}
\textsuperscript{0} no effect; ± minimum increase but not abnormal levels; + increase to abnormal levels in small % of patients; ++ increase to abnormal levels in 25-50\% of patients; +++ increase to abnormal levels in > 50\% of patients; CR reports of isolated cases of hyperprolactinemia, but generally without increased prolactin levels.
Pathological consequences of hyperprolactinemia

The clinical impact does not differ from other causes of hyperprolactinemia but hypogonadotrophic hypogonadism deserves special consideration. Recent evidences indicate that hyperprolactinemia causes lower production of kisspeptin at the hypothalamic level reducing the hypothalamic secretion of GnRH (gonadotrophin-releasing hormone) and consequently the synthesis and secretion of pituitary gonadotrophins (LH and FSH) with a loss of gonadal stimulation and infertility (Fig. 1).

In model mice with hyperprolactinemia as well as in women with hypogonadotrophic amenorrhea caused by hyperprolactinemia, GnRH and gonadotropin secretion and ovarian cyclicity were restored by kisspeptin administration.

Thus, hypogonadism induced by PRL excess is clinically characterized in women by menstrual disorders such as oligoamenorrhea and galactorrhea; hirsutism, acne and vaginal dryness may also be present. Menstrual disorders usually occur with PRL levels greater than 50 ng/ml. PRL levels between 30 to 50 ng/ml can cause a deficient luteal phase also being a cause of infertility.

Galactorrhea (either spontaneous or deliberate) is a common manifestation in premenopause (up to 90%), but occurs less frequently in menopause owing to the lack of estrogenic effect on the breasts.

In men PRL excess can be accompanied by gynecomastia (very rarely by galactorrhea), sexual dysfunction and infertility. In both sexes, hyperprolactinemia may induce decreased libido, osteopenia or osteoporosis, and fracture risk.

It is important to keep in mind those patients whose first psychotic outbreak occurs in children or adolescents (< 18 years of age) and who will require antipsychotic treatment for several years. This population is at increased risk of experiencing adverse effects in the short (< 6 months) and medium-term (6-12 months), such as sedation, extrapyramidal symptoms, hypogonadism, galactorrhea, delayed puberty, weight gain and dyslipidemia.

These metabolic alterations should be explained because hyperprolactinemia has been associated with an increase in food intake, weight gain and obesity. Moreover, PRL excess has been shown to reduce glucose tolerance and hyperinsulinemia in humans.

The effects of hyperprolactinemia will be exacerbated on the long-term (> 12 months) with deleterious effects especially on bone tissue. Its consequences will be the lack of reaching peak bone mass and the resulting risk of osteoporosis and fractures later in life.

While osteoporosis is a known complication of hypogonadism in general and also as mentioned before, of hypogonadism secondary to hyperprolactinemia, many other factors are involved in the increased risk of osteoporosis in the psychotic patient. Table 4 shows the different factors involved in the development of osteoporosis in schizophrenia. These factors include hyponatremia, which can increase the risk of fractures through two mechanisms. The former arises from an increase in falls triggered by gait disorders resulting from a decrease in brain glutamate concentration, which is the neurotransmitter responsible for gait. Even asymptomatic chronic hyponatremia increases falls (21% vs. 5% in normal controls). This happens because chronic hyponatremia stimulates osteoclastogenesis and bone resorption.
TABLE 4.– Factors inducing osteoporosis in schizophrenia

- Hyperprolactinemia
- Smoking - Alcoholism
- Hyponatremia
- Vitamin D deficiency
- Malnutrition
- Insulin resistance and diabetes mellitus

A third of total body sodium is stored in bone, in response to hyponatremia, sodium is mobilized outside the bone tissue similar to what is observed in calcium deprivation.

Causes of hyponatremia in schizophrenia can be attributed to psychogenic polydipsia (dilutional hyponatremia) that affects 6-20% of patients; increased antiuretic hormone (ADH) secretion induced by psychosis per se or produced by drugs such as haloperidol, phenothiazines, MAOIs, selective serotonin reuptake inhibitors and amitriptyline.

Hyperprolactinemia would also be a risk factor for obesity-related metabolic syndrome, insulin resistance and diabetes mellitus through apparently by way of a direct effect on pancreatic beta cells.

Approach to the patient with antipsychotic-induced hyperprolactinemia

Determination of serum PRL

Serum PRL levels should be measured in patients receiving any drug that may cause serum PRL increases, and in particular in those patients with signs and symptoms of hyperprolactinemia. Once these serum PRL elevations are confirmed, it is advisable to temporarily discontinue the drug intake for at least 72 hours and repeat the measurement. Specifically, with oral antipsychotics, PRL levels tend to normalize within 48 to 96 hours after the last dose. However, elevated levels can persist as long as about 3 weeks depending on drug half-life and storage in fatty tissue.

This practice, however, is not commonly applied to psychotropic drugs as there is insufficient data on the consequences of discontinuing its use for a few days to measure PRL and the risk for exacerbation of the psychiatric condition.

Therefore, it would be advisable to have baseline PRL levels, as well as glucose and lipid profiles before prescribing an antipsychotic, given the potential of these drugs to induce insulin resistance and metabolic syndrome.

During treatment, it is suggested to monitor prolactin levels two to three weeks after psychotropic drug initiation, and then regularly every two to three months. It is also advisable to check PRL levels in case it is necessary to increase the dose or to replace the drug with an alternative psychotropic medication.

It is also important to consider the presence of macroprolactinemia (analytical hyperprolactinemia) in particular in asymptomatic patients with hyperprolactinemia who are receiving antipsychotics, as this condition may also occur in these cases and may induce clinical and diagnostic confusion. The incidence of macroprolactin varies among different series, but it is estimated to occur in about 10-25% of cases of hyperprolactinemia.

Macroprolactinemia is detected by the polyethylene glycol (PEG) precipitation test in particular and, if available, by gel chromatography. With the PEG test, the higher molecular weight forms of PRL are removed by precipitation leading to the residual monomeric form in the sample supernatant. If the recovery of monomeric PRL after precipitation with PEG is less than 40% of the initial total value, then macroprolactin is the predominant variant present of immunoreactive PRL.

Other causes of hyperprolactinemia to be excluded are: pregnancy (in all premenopausal women), primary hypothyroidism, polycystic ovary syndrome (especially in presence of hyperandrogenism) and liver and kidney failure.

Imaging studies

Pituitary magnetic resonance imaging (MRI) with gadolinium is the best method for visualizing the presence of abnormalities in the sellar region. There is no PRL level that can differentiate a lactotroph tumor from drug-induced hyperprolactinemia, although a value > 200 ng/ml is suggestive of macroprolactinoma.

It is relevant for decision-making to know the previous PRL level and the gynecological (regular cycles) or andrological (sexual function) records of the patient prior to the treatment with psychotropic medication.

If the drug cannot be discontinued or replaced by another psychotropic (concerning exclusively to the psychiatrist) and the presence of hyperprolactinemia was not coincident with the treatment initiation, it is advisable to perform MRI studies of the sellar region.

Besides, a prolactinoma or any sellar lesion may coexist in patients treated with antipsychotics, fact that may exacerbate the serum PRL level.

It is important to be aware of the presence of non-functioning adenomas or other clinically silent pituitary lesions reported in about 10% of MRI images that may not correspond to a prolactinoma, even in the presence of altered serum PRL levels.

Figure 2 proposes an algorithm for the evaluation of patients with hyperprolactinemia.
Therapeutic approach to the patient with antipsychotic-induced hyperprolactinemia

Management of this situation requires a joint therapeutic approach with the psychiatrist and the endocrinologist. Hyperprolactinemia caused by antipsychotics is an underestimated, unrecognized or ignored adverse effect in spite of the potentially serious complications it causes.\(^5,17,19\)

The approach to this situation is controversial. The first option would be drug discontinuation, difficult to apply in patients under treatment with antipsychotics. Patients with asymptomatic hyperprolactinemia simply require periodical clinical and laboratory monitoring to evaluate the appearance of symptoms related to hyperprolactinemia.\(^7\) This approach should be reconsidered in the case of women of childbearing age with a desire for pregnancy, since excess PRL, even when mild, may be a cause of infertility.

In symptomatic patients with drug-induced hyperprolactinemia, one could try to reduce the dose of the antipsychotic, although this can result in an aggravation of the psychiatric condition. It may also be ineffective for the control of hyperprolactinemia, given that it is not always possible to normalize PRL levels with the reduction of the psychotropic dose. Another proposed option is to switch to another antipsychotic with less impact on PRL levels, such as quetiapine, but difficult to accomplish if the patient is well controlled with the initial psychiatric treatment.\(^6,17,18\)

Adjuvant therapy with aripiprazole, a partial dopamine agonist, has proven efficacy in reducing hyperprolactinemia. However, two possibilities are worth considering: one that the primary anti-psychotic efficacy may be decreased and second a higher risk of developing side effects when adding another psychotropic.\(^23,41,42\)

For those patients whose psychiatric state is well controlled but present signs of hyperprolactinemia such as hypogonadism or low bone mineral density, the Endocrine Society recommends replacement therapy with estrogens for women and testosterone for males.\(^7\) If the impact on bone mass is significant or if hormone replacement therapy is contraindicated, vitamin D and calcium treatment...

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**Fig. 2.**– Algorithm for the evaluation of the patient with hyperprolactinemia (Adapted by Samperi I, Lithgow K, Karavitaki N. Hyperprolactinemia. J Clin Med 2019; 8. pii: E2203)
should be implemented together with bisphosphonates, if necessary. Adding a dopamine agonist such as cabergoline is not recommended given the specific risk of exacerbation of psychosis and rare occurrence.

Cabergoline at lower doses could only be considered in very well selected cases such as fertility desire or contraindicated hormonal replacement therapy, and if the aforementioned alternatives could not be undertaken or were ineffective. This therapeutic approach will require close monitoring by the psychiatrist.

**Psychiatric disorders induced by the medical treatment of hyperprolactinemia**

Once pregnancy, hypothyroidism and drugs have been ruled out, the most frequent cause of chronic hyperprolactinemia is a prolactin-secreting tumor. Prolactinomas are the most common pituitary adenomas, accounting for approximately 50% of all pituitary tumors. They are classified according to their size in microprolactinomas (< 1 cm) and macroprolactinomas ≥ 1 cm, and in giant prolactinomas when measuring > 4 cm. These tumors most commonly affect women, especially between the ages of 25 and 44 years old, with a female to male ratio of 20:1 for microprolactinomas. The ratio between men and women tends to be similar for macroprolactinomas, although these tumors are usually larger in males.

Dopamine agonists (DA) have become the first-line therapy for prolactinomas. They induce not only the reduction of PRL secretion but also of the tumor size. Commercially available DAs are bromocriptine, cabergoline and quinagolide. In Argentina, cabergoline is the only DA commercially available. Bromocriptine and cabergoline are ergot derivatives with agonist activity on the D2 receptor. Cabergoline is a more selective agonist with a longer half-life that allows one to two weekly doses, usually ranging from 0.5 to 3 mg/week. Quinagolide is a non-ergot derivative that also causes agonist action at D2 receptors. Current guidelines recommend cabergoline as the first-line DA given its superior efficacy to bromocriptine, its improved tolerance and convenient dosage.

DA therapy is often prolonged and if PRL levels remain normal for at least two years and the tumor is no longer visualized on MRI studies, the drug can be discontinued. Another strategy if the tumor reduced its size by more than 50% and is more than 5 mm away from the optic chiasm, is to gradually taper the DA dose to the minimum concentration required to maintain both a normal PRL level and control the tumor volume or even to discontinue the therapy.

The most frequent adverse effects are nausea and/or vomiting, postural hypotension that may be accompanied by dizziness and syncope, headaches, Raynaud’s phenomenon and more rarely nasal congestion, flushing and cramps. These adverse effects are related to activation of D1 and 5HT1 receptors. High dosages of DA, as used for Parkinson’s disease, have been associated with valvulopathy caused by activation of the 5HT2B receptors.

Until present, it seems that risk of valvulopathy would not be significant with dosages usually used in patients with prolactinomas, with a very low incidence of 0.11% according to some series. However, if the cabergoline dosage used is higher than 2 mg/week, an annual echocardiogram is recommended.

More rarely, DA can induce mood swings and psychosis in susceptible patients.

Other classically reported psychiatric symptoms in patients with Parkinson’s disease treated with high DA doses are impulse control disorders that have been reported most recently in patients with prolactinomas, treated with even low DA doses.

**Impulse control disorders**

Impulse control disorders (ICD) are defined as failure to resist an impulse or temptation to perform a harmful act on oneself or others. These disorders include addictions to gambling, sex, food, compulsive shopping, kleptomania, intermittent explosive disorder, vigorexia, and punding. The term punding was introduced by Swedish psychiatrist Carl Rylander in 1968 in connection with consumers of phenmetrazine (sympathomimetic amphetamine). Punding is defined as a complex, prolonged, unproductive, and stereotypical behavior that is often associated with ICDs. It is characterized by compulsion and fascination for performing non-purpose repetitive mechanical tasks such as assembling and disassembling devices - watches, computers, collecting or classifying several objects simultaneously, emptying and rearranging household drawers and shelves, among others.

The physiopathological mechanism of these disorders is caused by an interaction between dopamine agonist and D3 receptors at the level of the mesocorticolimbic system, responsible for the processes that regulate behavior, pleasure and addiction.

The prevalence of ICDs is set at about 8% of the general population. The percentage increases considerably when ICDs caused by DAs are considered as well. In a retrospective data analysis published by the Food and Drug Administration (FDA) on the Adverse Effects Reporting System between 2003 and 2012, the prevalence of ICD induced by DAs in patients with Parkinson’s disease was 61.7%, in those with restless leg syndrome 24%, and in those with hyperprolactinemia 3.5%. ICDs were more common in patients treated with selective D3 agonists such as pramipexole and ropinirole, which would explain a higher frequency in Parkinson’s disease.
Thus, the occurrence of ICD in patients with prolactinomas treated with DAs such as bromocriptine or cabergoline has been considered an unusual event until recently, with a prevalence of about 3-4% and has been preferably associated with the use of high doses of DA. The first case of gambling addiction was reported in 2007 by a 47-year-old patient treated with cabergoline, 0.25 mg/week.69

However, more recent case series have described a prevalence of up to 17% in patients with prolactinomas treated mostly with cabergoline.66 Among different case series, the most frequent ICD was hypersexuality with a greater incidence in males. It is worth noting that about 33% of affected patients may develop multiple types of ICDs while treated with DAs.57, 63, 64

Other infrequent psychiatric disorders associated with the use of DAs are major depressive disorder, hallucinations, sleep disorders, manic episodes and psychosis; which generally resolve with the discontinuation of treatment.57

Some risk factors have been established for developing ICD such as the male sex, an early onset of the disease (particularly observed in relation to Parkinson’s disease), personality traits (depression, aggressiveness, impulsiveness), history of psychiatric disorders, alcohol consumption or other addictions.1, 56, 57

Consequently, before prescribing DAs, it is advisable to enquire about the patient’s habits and personality traits that may relate to an increased possibility of developing these complications.55-67

Once the treatment has been initiated, during follow-up consultations, it will be of importance to inquire the patient, and if possible their relatives or caregivers, about the onset of symptoms compatible with ICDs since the patients themselves may either not be aware of the symptoms or hide the situation.57

ICDs may occur at any time after the DA treatment initiation (early or late) requiring periodic monitoring. Some questionnaires and psychometric tests can be used: Barratt Impulsiveness Scale and the Behavioural Inhibition/Activation System (BIS/BAS), Depression Anxiety Stress Scale (DASS21), Questionnaire of Impulsive Control Disorders in Parkinson’s disease (QUIP-S), Hypersexual Behaviour Inventory (HBI), among others.55-66

Upon occurrence, ICDs should be managed together with the psychiatrist and the initial treatment will be to either reduce the DA dose or discontinue the drug. It is no use substituting one DA for another.67

Burack describes the case of a 32-year-old patient with microprolactinoma who developed an acute manic episode with weekly doses of 0.5 mg of cabergoline. The manic episode quickly reversed after cabergoline was discontinued and replaced with aripiprazole 5 mg/day which was sufficient to keep the PRL level suppressed and to stabilize the psychiatric state.67 Tumor size remained stable over 15 months of follow-up. Therefore, aripiprazole would be a very good alternative for the control of hyperprolactinemia for those patients with microprolactinomas who develop DA-induced psychiatric disorders.57 Another option to be considered in these situations is transphenoidal pituitary surgery.63, 57

The complications of DA-induced ICDs can be serious and catastrophic for the life of the patient, with great impact on their family/social and professional life, financial decisions, and mental and physical health.

In a recent study, only 37% of the interviewed patients with hyperprolactinemia (n= 51) had been warned about the relationship between ICDs and DA therapy.66

Although to date there are no reported cases of ICDs in patients treated with DAs for other endocrinopathies such as acromegaly, Cushing’s disease and non-functioning pituitary tumours it is advisable to consider their possible development when prescribing CBG.

In conclusion, in patients treated with antipsychotics, prescribing dose, treatment duration, antipsychotic potency (D2 receptor blockade), age and sex are contributing factors to the level of hyperprolactinemia, occurring more frequently in adolescent and premenopausal women. It would be advisable to measure the PRL level before treatment initiation and then periodically, in every patient treated with antipsychotics. Antipsychotic-induced hyperprolactinemia is an underestimated, unrecognized or ignored side effect that requires treatment in many cases. The consequences of hyperprolactinemia can be serious and compromise adherence to treatment. Management of hyperprolactinemia requires a joint therapeutic approach between the psychiatrist and the endocrinologist.

Although, ICDs have been considered relatively uncommon adverse effects in the endocrinology-related field, in recent years, with the use of DAs, the number of reported cases has increased considerably. It is prudent and advisable that prior to treatment initiation, physicians warn and educate patients and/or their families or caregivers about the potential adverse effects of DAs. ICDs can occur unpredictably, at any age, affect both sexes, also with low DA doses, and usually disappear when ceasing treatment.

Finally, the aim of this review is to inform psychiatrists and endocrinologists of the need for joint management of hyperprolactinemia associated with psychiatric disorders in its different forms in order to avoid serious and devastating complications for the patient.

Conflict of interest: Leonardo F. L. Rizzo is the Medical Director of Química Montpellier S.A. Daniela Mana and Héctor Serra are Medical Consultants for Química Montpellier S.A.
References


51. Caputo C, Prior D, Inder WJ. The need for annual echocardio-
52. Steeds RP, Stiles CE, Sharma V, Chambers JB, Lloyd G, Drake W. Echocardiography and monitoring patients receiving dopamine agonist therapy for hyperprolactinae-
55. Barake M, Klibanski A, Tritos NA. Management of Endo-
58. Moore TJ, Glenmullen J, Mattson DR. Reports of pathologi-
59. Davie M. Pathological gambling associated with cabergo-
60. Dogansen SC, Cikirkici U, Oruk G, et al. Dopamine ag-
61. De Sousa SM, Chapman IM, Falhammar H, Torpy DJ. Dopa-testotoxicosis: disruptive hypersexuality in hypo-
63. Bancos I, Nannenga MR, Bostwick JM, Silber MH, Erickson D, Nippoldt TB. Impulse control disorders in patients with dopamine agonist-treated prolactinomas and nonfunc-
67. Burbac L. Management of microprolactinoma with aripip-