Neuroendocrine predictors of the evolution of depression

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The assertion that the clinical efficacy of antidepressants is comparable between—and within—the classes may be true from a statistical viewpoint, but is of limited value in practice. Indeed, depression is, clinically and biologically, a heterogeneous illness and several lines of evidence suggest that the response to a pharmacological treatment depends on the patient’s biological state.2 Despite advances in psychopharmacology, more than one-third of patients do not respond to the drug of first choice.3 Therefore, a major issue is not only to have efficacious drugs, but also to optimize their use.

Keywords: HPA axis; thyroid; TRH test; serotonin; dopamine; noradrenaline; antidepressant treatment

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Depression is both clinically and biologically a heterogeneous entity. Despite advances in psychopharmacology, a significant proportion of depressed patients either continue to have residual symptoms or do not respond to antidepressants. It has therefore become essential to determine parameters (or predictors) that would rationalize the therapeutic choice, taking into account not only the clinical features, but also the “biological state,” which is a major determinant in the antidepressant response. Such predictors can derive from bioclinical correlates and, in this context, the neuroendocrine strategy appears particularly suited. Numerous studies have investigated neuroendocrine parameters—derived mainly from dynamic challenge tests—in order to (i) determine the predictive profiles of good clinical responders to given antidepressants; (ii) monitor the progression of markers in parallel with the clinical outcome; and (iii) evaluate “in vivo” in humans the mechanisms of action of antidepressant compounds (before, during, and after treatment). This article does not attempt to be exhaustive, but rather uses selected examples to illustrate the usefulness of the investigation of the adrenal and thyroid axes and the assessment of central serotonergic, noradrenergic, and dopaminergic systems by means of neuroendocrine tests. Given methodological constraints, most of these investigations—except for baseline hormone values and the dexamethasone suppression test—cannot be used routinely in psychiatry. Despite these limitations, the neuroendocrine strategy still offers new insights in biology and the treatment of depression. Its possible expansion depends mainly on the development of specific agonists or antagonists for better investigation of the receptors supposedly involved in the pathophysiology of depression. These investigations will help define more homogeneous subgroups from a bioclinical and therapeutic viewpoint.
During the past years, there has been increasing interest in the identification of predictors of outcome in depression. However, there is little consensus regarding which clinical and biological variables influence the therapeutic response to antidepressants. Among the possible predictors, those derived from neuroendocrine investigations have been extensively studied. These predictors can be measured at baseline (ie, after a sufficient drug-withdrawal period) and/or during the course of treatment. It is beyond the scope of this article to detail the numerous endocrine indicators that can be used as potential biological predictors of outcome. Rather, this paper illustrates, through selected examples, the usefulness of some pertinent neuroendocrine investigations.

HPA axis

Considerable research findings have accumulated over the last four decades regarding the role of the hypothalamic-pituitary-adrenal (HPA) axis in the psychobiology of depression. Increased cortisol secretion and failure to suppress cortisol in response to dexamethasone, a glucocorticoid agonist, have been consistently associated with severe, melancholic, and psychotic depression. It has been hypothesized that this stress axis overdrive is primarily a reflection of abnormal limbic-hypothalamic activation, with increased secretion of hypothalamic corticotropin-releasing hormone (CRH) and consequent excessive adrenal cortisol secretion. However, it remains uncertain whether the hypercortisolism is an epiphenomenon or directly contributes to depressive symptomatology and to the biochemical alterations seen in major depression (Figure 1).

The dexamethasone suppression test

Although the exact pathophysiology underlying dexamethasone suppression test (DST) nonsuppression remains unclear, it has been suggested that abnormal cortisol response reflects impaired negative feedback at the level of the pituitary corticotroph (ie, decreased type II glucocorticoid receptor function) on endogenous HPA axis hyperactivity (ie, increase in hypothalamic CRH drive that overrides the action of dexamethasone). However, (i) cortisol nonsuppression following DST is not specific for the diagnosis of major depression; and (ii) the sensitivity of the DST in depression is low. Indeed, only 15% to 25% of major depressed patients are nonsuppressors, while the rate of positive DST increases in severe depression (about 40% to 70%). Despite these limitations, the use of DST in psychiatric research still has considerable merit. For example, serial DST monitoring...

Selected abbreviations and acronyms

| Abbreviation | Definition                                      |
|--------------|-------------------------------------------------|
| ACTH         | adrenocorticotropic hormone                     |
| CRH          | corticotropin-releasing hormone                  |
| DA           | dopamine                                        |
| DST          | dexamethasone suppression test                  |
| GH           | growth hormone                                  |
| HPA          | hypothalamic-pituitary-adrenal (axis)           |
| HPT          | hypothalamic-pituitary-thyroid (axis)           |
| NA           | noradrenaline                                   |
| PRL          | prolactin                                       |
| SSRI         | selective serotonin reuptake inhibitor          |
| T₃           | triiodothyronine                                |
| T₄           | thyroxine                                       |
| TRH          | thyrotropin-releasing hormone (protirelin)      |
| TSH          | thyroid-stimulating hormone (thyrotropin)      |

Figure 1. Overview of the relationships between the monoamine systems and the hypothalamic-pituitary-thyroid (HPT) and hypothalamic-pituitary-adrenal (HPA) axes. 5-HT, serotonin; NA, noradrenaline; DA, dopamine; monoamine receptors, 5-HT₁₆, α₂-adrenoreceptor, DA-D₂; ACTH, adrenocorticotropic hormone; CRH, corticotropin-releasing hormone; TRH, thyrotropin-releasing hormone; TSH, thyroid-stimulating hormone. + and − arrows indicate stimulation and inhibition, respectively, of the secretion of monoamines and hormones.
of depressed patients undergoing drug treatment showed that DST gradually turned into suppression in treatment responders.\(^{15,16}\) Patients whose DST remained abnormal or who were initially suppressors, but became nonsuppressors during an observation period, had a poorer prognosis.\(^{6,17}\) In addition, over a long-term follow-up, DST suppressors at baseline have a better outcome than non-suppressors.\(^{18}\) Although conflicting results on the predictive value of the DST have been reported, it is generally accepted that (i) the presence of an abnormal DST indicates the need for a biological treatment, while (ii) the initial DST status has no predictive value in the choice of prescription of antidepressants.\(^{19}\)

**The combined DEX/CRH test**

After CRH became available for clinical studies, a more sensitive test than the DST was developed: the combined dexamethasone/corticotropin-releasing hormone (DEX/CRH) test\(^ {20}\) in which dexamethasone-pre-treated subjects receive a single dose of CRH during the afternoon of the next day. In healthy control subjects, owing to the normal inhibiting activity of the glucocorticoid receptors at the pituitary level, CRH administration induces only a small amount of corticotropin (adenocorticotropic hormone [ACTH]) and cortisol secretion. In depressed patients, the ACTH/cortisol response to the combined DEX/CRH test is significantly increased compared with controls. This phenomenon suggests an altered glucocorticoid feedback regulation (ie, decreased glucocorticoid receptor sensitivity), possibly associated with hypothalamic CRH and vasopressin overdrive.\(^ {21}\) The combined DEX/CRH test identifies HPA axis dysfunction with high sensitivity in severe major depression (about 80%).\(^ {22}\) Furthermore, DEX/CRH test normalization typically precedes or coincides with—rather than follows—clinical recovery, and failure to normalize portends poorly for clinical outcome.\(^ {22}\) Patients with persistent severe HPA dysregulation are more prone to relapse within 6 months than those with low cortisol response to the DEX/CRH test at discharge.\(^ {23}\) Moreover, early improvement (after 1 or 2 weeks of therapy) and beneficial treatment outcome after 6 weeks are associated with a lower HPA system activity.\(^ {21}\)

Taken together, these studies suggest that lowering HPA activity and clinical response are related. However, in a recent study, Watson et al\(^ {24}\) have found that the DEX/CRH remains abnormal in remitted bipolar patients, suggesting that HPA axis dysfunction is a potential “trait” marker in bipolar disorder.

**Effects of antidepressants on the HPA axis**

Recent research suggests that antidepressants could exert their clinical action in depression via the restoration of type II glucocorticoid receptor function with a subsequent reestablishment of HPA axis negative feedback.\(^ {25}\) Indeed, animal studies\(^ {26}\) have consistently shown that antidepressants (ie, tricyclics, selective serotonin reuptake inhibitors [SSRIs], moclobemide, tianeptine) increase type II and type I (or mineralocorticoid) receptor expression and function (ie, increased efficiency of signal transduction by increasing mRNA levels and hormone-binding activities). This, in turn, is associated with enhanced negative feedback by endogenous glucocorticoids, and thus with reduced HPA axis activity. Downstream consequences of lowered cortisol secretion is a lessening of expression of genes that are under corticosteroid regulatory control, among them those related to biogenic amine neurotransmission. Additional evidence indicates that cortisol-lowering treatments (ie, “antiglucocorticoids”) may be of clinical benefit in depressed patients. Indeed, open and controlled trials suggest that blockers of cortisol synthesis (ie, metyrapone, ketoconazole, aminoglutethimide), or type II glucocorticoid receptor antagonists, including mifepristone (RU-486) and ORG 34517, may exert antidepressant effects.\(^ {27}\) Although clinical usage of the currently available antiglucocorticoids is limited by significant side effects, the development of drugs that specifically target the glucocorticoid receptor may lead to innovative strategies in the treatment of depressive states. In the same way, development of effective CRH blockers\(^ {28}\) will provide an important tool for further study of the role of CRH hypersecretion in severe depression and other stress-related illnesses.

**HPT axis**

It is well established that major depression may be accompanied by a dysfunction of the hypothalamic-pituitary-thyroid (HPT) axis, including a slight elevation of serum thyroxine (T\(_4\)), subnormal (or “blunted”) thyrotropin (thyroid-stimulating hormone [TSH]) response to morning injection of protirelin (thyrotropin-releasing
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hormone [TRH]), and a loss of the nocturnal rise in TSH. Usually, this phenomenon is not secondary to hypercortisolism. Investigation of the HPT axis by means of the TSH response to morning administration of TRH (at 8 AM or 9 AM) has limited clinical value because of modest diagnostic sensitivity (about 25%). Owing to the circadian activity of the thyrotroph, which is maximal between 11 PM and 1 AM pituitary TSH secretion is more sensitive to TRH stimulation in the evening than in the morning, both in normal controls and in depressed patients. In 1990, our group reported that the 11 PM TRH-TSH test was more sensitive than the 8 AM TRH-TSH test, and that the difference in TSH response between 11 PM and 8 AM TRH tests (ΔΔTSH) was an even more sensitive measure. Consistent with our original observation, this chronobiological index is reduced in about 80% of major depressed inpatients. It has been suggested that blunted TRH-induced TSH stimulation might reflect downregulation of the TRH receptors in the pituitary gland secondary to a prolonged increase in hypothalamic TRH stimulation. Furthermore, the shift to higher iodothyronine levels in euthyroid depressed inpatients, both in the morning and in the evening, may contribute to the blunting of TSH response to TRH.

Predictive value of the TRH test

So far, studies of the therapeutic predictive value of the TSH response to TRH test have yielded conflicting results. Some studies have found that the normalization of the test is associated with remission, in which case the blunted TSH response—and more obviously the reduced ΔΔTSH values—may be considered as a “state” marker of depression. Conversely, persistence of blunted responses during remission could represent a “vulnerability” marker to depression. Some investigators have found no link between the initial status of the TRH test and the subsequent response to a particular antidepressant treatment defined according to its biochemical mechanism of action. Others have found an association between the blunted TSH response and the response to desipramine (which shows a predominant “noradrenergic” action). On the other hand, it has been observed that patients with the lowest pretreatment evening TSH secretion (basal and after 11 PM TRH stimulation) have the lowest rate of antidepressant response, and this may contribute to antidepressant treatment resistance. In such cases, it has been speculated that adjunction of thyroid hormones, could be particularly beneficial to amplify antidepressant effects, since, by increasing the negative feedback on the hypothalamus, thyroid hormones may decrease TRH overproduction at this level.

To date, only one study has examined the relationship between morning and evening TSH and prolactin (PRL) response to TRH tests prior to antidepressant treatment and 12-month outcome. In this study, dissociation between 11 PM TRH-induced TSH and PRL stimulation (ie, blunted TSH response associated with normal PRL response) was indicative of poor clinical outcome. Thus, normal PRL response to TRH, despite decreased TRH receptor responsiveness, might reflect a decrease in dopaminergic tone, since dopamine (DA) inhibits PRL secretion. In such patients, one may hypothesize that adjunctive therapy with DA agonists might be useful to amplify antidepressant effects.

Effects of antidepressants on the HPT axis

A number of reports suggest that treatment with antidepressant drugs leads to changes in thyroid function tests: either decreased peripheral thyroid hormone levels and/or increased TSH levels (basal or post–TRH stimulation). However, these results have not always been confirmed, owing in part to methodological limitations, eg, small sample sizes, variable definitions of depression, hospitalization status, and technical factors, such as differences in the sensitivity of the assays used in the measurement of thyroid hormones and TSH. Furthermore, it remains unclear whether changes in thyroid function are a direct effect of an antidepressant on the thyroid axis or a correlate of clinical improvement. Animal studies suggest that chronic antidepressant treatment decreases thyroid function. However, data from healthy volunteers support the notion that tricyclic antidepressants have no consistent effect on TSH secretion. In depressed patients, most studies have shown that antidepressant treatment with tricyclics, SSRIs, or monoamine oxidase inhibitors (MAOIs) does not induce significant changes in TSH levels. Moreover, it has been reported, but not consistently, that response to tricyclic antidepressants is associated with (i) higher pretreatment T4 levels; and (ii) decreased measures (within the normal range) of T4 and free thyroxine (FT4) without changes in triiodothyronine (T3) or TSH levels. Thus,
although this is not supported by all studies, changes in thyroid function appear to be related to clinical recovery rather than to a direct effect of the antidepressant drug. This is further supported by the fact that normalization of the $\Delta\Delta$TSH test is related to clinical recovery, while, irrespective of outcome, $\Delta\Delta$TSH values are not significantly changed by 4 weeks of treatment with amitriptyline, fluoxetine, toloxatone, venlafaxine, or tianeptine.38,64

**Neuroendocrine investigations of the noradrenergic system**

The original catecholamine depletion hypothesis of depression has been reformulated into the “noradrenergic dysregulation hypothesis,”65 which emphasizes a primary subsensitivity or downregulation in nerve terminal $\alpha_2$-adrenoreceptors, leading to impaired negative feedback on presynaptic neurons, which in turn may induce a disinhibition of noradrenaline (NA) output and exaggerated NA release in response to any activation of the catecholaminergic system. One of the most consistently reported abnormal findings in depression is a blunted growth hormone (GH) response to acute administration of clonidine, a partial $\alpha_2$-adrenoreceptor agonist. This suggests subsensitive postsynaptic $\alpha_2$-adrenoreceptors at the hypothalamic level. A dysregulation of the NA system may lead to increased anxiety in depressive patients.66,67 More generally, blunted GH response to clonidine does not appear specific to depression, but rather to the “anxiety spectrum,” since this blunting has also been observed in generalized anxiety disorder,68 panic disorder,69,70 and social phobia.71 The link between anxiety and NA dysregulation in depressed patients is further supported by the negative correlation between GH response to clonidine and the severity of anxiety as evaluated by the Hamilton Anxiety Scale scores.72

**Predictive value of the clonidine test**

Some studies have shown that remitted depressive patients show reduced GH responses to clonidine,73,74 suggesting that decreased sensitivity of $\alpha_2$-adrenergic receptors may represent a vulnerability marker for depression. This is further supported by the fact that, despite having differing mechanisms of action, antidepressants such as desipramine, mianserin, clorgyline, amitriptyline, and fluoxetine do not restore clonidine’s effect on GH secretion in responders or nonresponders to treatment.75,76

It has also been argued that deficiencies in NA function could lead to differential response to noradrenaline and serotonin reuptake inhibitors.77 In a study by Coote et al.,78 the decreased GH response before treatment was correlated with subsequent good clinical response to desipramine (a “noradrenergic” antidepressant). In a recent study, Correa et al.79 reported that amitriptyline, which primarily increases NA function, was more efficacious than fluoxetine in depressed patients showing at baseline blunted GH to clonidine (amitriptyline is at least 100 times more potent than fluoxetine in the inhibition of the noradrenaline transporter69). Taken together, these results suggest that the NA function might influence response to antidepressant treatment.

**Neuroendocrine investigations of the DA system**

It is known that the mesolimbic DA system plays a key role in goal-directed and motivational behavior. In depression, it has been suggested that hypofunction in mesolimbic DA system may be involved in anhedonia and amotivational apathy.80 DA agonists can facilitate the action of antidepressant drugs in certain animal models of depression and in some depressed patients.82 According to the neuroendocrine challenge paradigm, hormone responses to DA agonists may provide an indirect assessment of central DA neurotransmission at the postsynaptic receptor level within the limbic-hypothalamic-pituitary axis in humans.83,84 Apomorphine, a direct-acting DA agonist with high affinities for $D_2$ and $D_3$ receptors85 and a partial agonist at the $D_1$ receptor,86 decreases PRL and stimulates GH,84 ACTH, and cortisol secretion.87-89 In major depression, discrepant findings have been reported: unaltered responses76,88-90 or decreased GH response77,91 to apomorphine have been found. Some of these divergences may be explained by the diversity of factors that influence the hormonal response to apomorphine, and by the heterogeneity of the populations studied. Indeed, when depressed patients are classified according to their clinical features, differences in the apomorphine response are observed between subtypes. For example, it has been found,80,92,93 but not by all,90 that apomorphine produces a lesser decrease in serum PRL levels in bipolar patients than in normals and in unipolar patients. On the other hand, unipolar patients with melancholic and psychotic features often show reduced ACTH/cortisol responses especially when hypercortisolism coexists.84
From a pathophysiological viewpoint (i) blunted PRL response to apomorphine may reflect decreased D₂ receptor function in the pituitary (ie, lactotrophs); and (ii) blunted ACTH/cortisol response may reflect decreased DA receptor function (ie, D₂ receptor–like or D₁ receptor, or both) connected with the regulation of the HPA axis at the hypothalamic level.

**Predictive value of the apomorphine test**

Some preclinical studies suggest that long-term antidepressants upregulate and/or hypersensitize postsynaptic DA receptors (ie, D₂ and D₃) as reflected by increased apomorphine responses in animals treated with several classes of antidepressants. However, in depressed patients, changes in DA function (ie, increased ACTH/cortisol, but not GH and PRL, responses to apomorphine) following antidepressants appear to be transient (ie, after 2 weeks’ treatment, but not after 4 weeks). These changes are not correlated with clinical efficacy and are independent of the compound administered (venlafaxine, tianeptine).

On the other hand, it has been found that greater DA postsynaptic sensitivity (assessed by greater GH response to apomorphine) is associated with greater resistance to paroxetine treatment. This finding has lead to the hypothesis that pretreatment low DA receptor responsivity could predict antidepressant response to SSRIs.

**Endocrine disorders**

Endocrine disorders are among the factors that should be routinely searched for in the management of depressed individuals. Rare cases of endocrine disorder–related depression can be identified through the systematic measurement of some parameters, eg, TSH/FT₃/FT₄, PRL, cortisol/ACTH, parathyroid hormone/calcium, and glucose. Moreover, it has been well documented that endocrine disorders are factors that may contribute to treatment resistance. The dexamethasone test is also used by endocrinologists and this test can be used routinely in psychiatry because it is simple and has decent sensitivity and predictive value in clinical evolution and response to treatment.

**Conclusions**

The findings reviewed in this article add further to the body of data pointing to the utility of neuroendocrine measurement in discriminating among subtypes of depressive disorders. Depression is characterized by a complex configuration of disturbances in a number of neurotransmitter and hormonal systems. Given the multiple reciprocal relationships between these systems, it is not appropriate at the present to consider one system as primary in an etiological sense. Moreover, the biological changes that can be studied (“biological states of depression”) not only result from the pathophysiological process involved in the etiology of depression, but also from adaptive processes that maintain the homeostasis of the systems. This is why, in basal conditions, it is rare to find significant biological abnormalities in depressive states. In contrast, dynamic challenges destabilize the homeostatic balance and may therefore be used to better characterize heterogeneous biological states. Moreover, this characterization may lead to different therapeutic strategies.

According to many studies, the presence of positive (or abnormal) neuroendocrine test suggests a need for antidepressant somatic therapy of depression, but the predictive value of these tests need further clarification. Unfortunately, for methodological, ethical, and economic reasons, the neuroendocrine tests are rarely performed in battery (ie, several tests for each patient) and this limits their application from a pathophysiological and therapeutic viewpoint. For instance, in depression, the absence of chronobiological dysfunction of the thyroid axis (ie, normal ∆∆TSH) is significantly associated with decreased serotonergic function, and vice versa. Therefore, the response to ∆∆TSH test may be of great value since a normal ∆∆TSH test could orientate the clinician towards antidepressants that increase “serotonergic” transmission; while a blunted ∆∆TSH test, which is often associated with a blunted clonidine test, could orientate the clinician toward antidepressants that increase “noradrenergic” transmission.

The relationship between neuroendocrine test results and clinical outcome has mostly been described in retrospective study protocols. On the basis of our observations, and those of others, one may propose the following strategies, which could be the theme of prospective clinical trials of antidepressants (strategies marked with an asterisk have not yet been evaluated in depressed patients):

- SSRI appear to be perfectly suitable for the first-line treatment for depression, especially when there is no evidence for chronobiological dysfunction of the thyroid axis (normal ∆∆TSH).
• “Noradrenergic” antidepressants appear to be suitable when the GH response to clonidine is blunted and/or when there is evidence for chronobiological dysfunction of the thyroid axis (blunted ΔΔTSH).

• “Dopaminergic” antidepressants appear to be suitable in case of normal TRH-PRL response associated with blunted TRH-TSH response (performed at 11 pm) and/or in case of blunted PRL response to apomorphine test (which is often observed in bipolar depression).†

• In case of a positive DST, frequently associated with severe depression, antidepressant treatment alone will probably not suffice and therefore calls for a different approach (ie, adjunction of “antiglucocorticoids” to antidepressants, or antipsychotics, in case of melancholic/psychotic depressed patients DST is associated with blunted ACTH/cortisol response to apomorphine, reflecting a possible presynaptic DA hypersecretion at the hypothalamic level).

• In case of nonresponse or partial response in patients with pretreatment 11 pm blunted TRH-TSH response, one may propose adjunctive thyroid hormone therapy. In this indication, T₃ seems to be more efficacious than T₄. Since TSH blunting could be secondary to hypersecretion of endogenous TRH, it may be that treatment with exogenous thyroid hormone increases the negative feedback and, in this way, tends to correct the hypersecretion of endogenous TRH. Speculating further, the return to normal levels of TRH is perhaps part of the physiological normalization, which is associated with resolution of a depressive episode. Finally, it would also be interesting to study the above strategies in conjunction with pharmacogenomic approaches.

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Predictores neuroendocrinos de la evolución de la depresión

La depresión es una entidad heterogénea tanto desde un punto de vista clínico como fisiopatológico. A pesar de los progresos de la psicofarmacología, una proporción significativa de pacientes deprimidos o continuán con síntomas residuales o no responden a los antidepresivos. El poder determinar predictores que permitan racionalizar la elección de los fármacos teniendo en cuenta no solamente el estado clínico, sino también el “estado biológico” –puesto que influye en la respuesta terapéutica– parece constituir una necesidad. Tales predictores pueden ser el resultado de correlatos biológico-clínicos y es en este contexto que las estrategias neuroendocrinas se pueden aplicar de manera adecuada. El estudio de parámetros neuroendocrinos esencialmente resultados de pruebas dinámicas ha permitido (1) establecer perfiles de predicción de buena respuesta a ciertos tratamientos antidepresivos, (2) seguir la evolución de los marcadores en paralelo con la clínica y (3) estudiar los mecanismos de acción de los antidepresivos “in vivo” en el hombre (mediante estudios antes, durante y después del tratamiento). Este artículo, que no intenta ser exhaustivo, considera principalmente el interés en psiquiatría de la exploración de los ejes corticotrópico y tirotídeo y de los sistemas centrales serotoninérgico, noradrenérgico y dopaminérgico mediante pruebas neuroendocrinas. Mientras tanto, teniendo en cuenta las restricciones metodológicas de la mayoría de estas investigaciones –con excepción de los niveles hormonales basales y la prueba de supresión con dexametasona– es ilusorio considerar las pruebas neuroendocrinas dentro del conjunto de los exámenes de rutina. A pesar de estas limitaciones, la estrategia neuroendocrina ofrece innegablemente nuevas posibilidades de modelos biológicos y terapéuticos en la depresión. Su expansión futura depende en gran medida del desarrollo de agonistas o antagonistas más específicos que permitan explorar de manera más precisa los diferentes receptores supuestamente implicados en la fisiopatología de la depresión. Es razonable pensar que al futuro este tipo de investigaciones permitirá delimitar subgrupos más homogéneos desde una perspectiva tanto biológico-clínica como terapéutica.

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Paramètres neuroendocriniens prédictifs de l'évolution des états dépressifs

La dépression est une entité hétérogène tant d'un point de vue clinique que physiopathologique. Malgré les progrès de la psychopharmacologie, environ un tiers des patients ne répondent pas au traitement antidépresseur de première intention. Déterminer des paramètres permettant de rationaliser le choix des chimiothérapies en tenant compte non seulement de l'état clinique mais aussi de « l'état biologique » — puisque celui-ci influe sur la réponse thérapeutique — apparaît par conséquent une nécessité. De tels paramètres (ou marqueurs de prédictivité) peuvent être issus de corrélats biologico-cliniques et c'est dans ce contexte que les stratégies neuroendocriniennes peuvent être appliquées de façon pertinente. De nombreuses études ont déjà permis (1) d'établir à partir de paramètres neuroendocriniens — essentiellement issus de tests dynamiques — des profils prédictifs de bonne réponse à certains traitements antidépresseurs, (2) de suivre l'évolution des marqueurs parallèlement à la clinique, et (3) d'étudier les mécanismes d'action « in vivo » chez l'homme d'antidépresseurs (par des études avant, pendant et après traitement). Cet article, qui ne vise pas à être exhaustif, envisage principalement l'intérêt en psychiatrie de l'exploration des axes corticotrope et thyroidien et des systèmes sérotoninergique, noradrénnergique et dopaminergique centraux à l'aide de réponses hormonales à des tests spécifiques. Cependant, compte tenu des contraintes méthodologiques de la plupart de ces investigations — hormis les prélèvements basaux et le test à la dexaméthasone — il est illusoire d'envisager les tests neuroendocriniens dans le cadre d'examens de routine. En dépit de ces limitations, la stratégie neuroendocrinienne offre indéniablement de nouvelles possibilités de modélisation biologique et thérapeutique. Son essor futur dépend pour une grande part du développement d'agonistes ou d'antagonistes plus spécifiques afin d'explorer de manière plus précise la fonctionnalité des différents récepteurs supposés impliqués dans la physiopathologie de la dépression. Il est raisonnable de penser qu'à terme ce type d'investigations permettra de délimiter des sous-groupes plus homogènes tant sur le plan biologico-clinique que thérapeutique.
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