HYPERPIGMENTATION AND ACTH – AN OVERVIEW OF LITERATURE

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ABSTRACT

Introduction. ACTH (adrenocorticotropic hormone) is a key regulator of adrenal production involving cortisol as an essential hormone for life. The melanin is a pigment which is produced by melanocytes at the level of melanosomes (the melanogenesis). Both MSH and ACTH are generated by the cleavage of POMC (proopiomelanocortin) after CRF (corticotropin-releasing factor) stimulation and then MSH acts on the skin causing hyperpigmentation.

Aim. To introduce clinical data of literature that link hyperpigmentation with ACTH excess. Three main topics are introduced: Addison’s disease, ectopic Cushing’s syndrome, and Nelson’s syndrome.

Method. This is a short overview of literature including papers that have been mostly published within last 5 years.

General data. Hyperpigmentation in relationship to ACTH includes its high levels in addition to low cortisol in Addison’s disease and Nelson’s syndrome and high non-suppressible cortisol in ectopic Cushing’s disease. ACTH has a pituitary origin in first two situations and malignancy in the third one. A pituitary tumour is found in cases with Nelson’s syndrome. An autoimmune background may be associated with Addison’s disease. An iatrogenic component is brought by Nelson’s syndrome. All three situations are severe and life threatening of different scenarios.

Conclusion. Hyperpigmentation may be the clue to connect dermatology to endocrine pathologies and ACTH massive release by a pituitary or a non-pituitary origin involves a complex panel of conditions.

Keywords: hyperpigmentation, ACTH, Nelson’s syndrome, Addison’s disease, ectopic Cushing’s syndrome

INTRODUCTION

ACTH (adrenocorticotropic hormone) is a key regulator of adrenal production involving cortisol as an essential hormone for life (1,2). The steroid secretion is normally the effect of a complex frame that also includes autocrine and paracrine factors (1,2). Even chromaffin cells from medullary part of the adrenal glands may produce ACTH as the pituitary gland does (1,2). ACTH acts not only as a hormone at the level of adrenal glands but also at the level of blood vessels and a local mediator at adrenals (1,2). ACTH also has a role as an acute regulator of aldosterone production as well as serotonin (2,3).

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Beyond endocrine and cardiovascular connections, ACTH is also linked to the skin and melanogenesis and this goes through MSH (melanocyte stimulating hormone) interplay (4,5). Alpha-MSH is a regulator of pigmentation in normal humans (5,6). This is done in association with other melanocortins whose roles are more or less understood until present time (5,6). Alpha-MSH as well as ASP (Agouti signalling protein) acts on melanocortin receptor type 1 (MC1r) of melanocytes displaying melanogenic effects, but also mitogenic actions (5,6). Skin melanocytes are activated by paracrin and autocrin factors, for instance, in case of prolonged sun exposure (5,6). The melanin is a pigment which is produced by melanocytes at the level of melanosomes (the melanogenesis) (6). Both MSH and ACTH are generated by the cleavage of POMC (proopiomelanocortin) after CRF (corticotropin-releasing factor) stimulation (6,7). The acquired hyperpigmentation it comes with an abnormal ACTH in parallel with high MSH as it happens in ACTH tumour-derivate secretion of pituitary and non-pituitary origin (5,6,7).

AIM

To introduce clinical data of literature that link hyperpigmentation with ACTH effects at the level of the skin. Three main topics are introduced: Addison’s disease, ectopic Cushing’s syndrome, and Nelson’s syndrome. All these conditions are associated with high levels of ACTH of either pituitary or non-pituitary tumour as cause, benign or malignant, and this goes with hyperpigmentation.

MATERIAL AND METHOD

This is a short overview of literature including papers that have been mostly published within last 5 years regarding medical data that involves high ACTH and skin hyperpigmentation. A selection of 30 papers is done.

GENERAL DATA

Addison’s disease

Addison’s disease is a rare, heterogeneous condition where the destruction of the adrenal glands is done mainly based on an autoimmune or tuberculous background (8). Low cortisol causes an ACTH increase based on a feedback mechanism between pituitary and adrenal glands while hyperpigmentation at the level of skin (including scars) and mucosa is one of the main clinical features (9). Others skin conditions like vitiligo may be associated in cases with polyglandular autoimmune syndrome (10). Primary adrenal insufficiency may be lethal if adequate substitution is not provided (9). More than 30 causes are described, including genetic types (9).

Ectopic Cushing’s syndrome

Cushing’s syndrome is the excess of glucocorticoid of either endogenous or exogenous cause (11,12). Endogenous sources includes the adrenal itself and ACTH derivate forms like corticotropinoma and ectopic Cushing’s syndrome (also called ACTH-dependent types) (11,12). This last mentioned type comprises between 10% and 20% of all ACTH-dependent cases (11,12). The cause of ACTH is a non-pituitary tumour despite the fact that a concomitant pituitary incidentaloma might be registered due to the high prevalence in general population (13,14,15,16). The most frequent forms of cancer causing ectopic Cushing’s syndrome are located in thorax-abdominal region (12,17). Among them we mention small cell lung carcinoma, medullary thyroid cancer, neuroendocrine tumours of the lung and gastro-intestinal tract (17,18,19,20). The typical endocrine assessment includes the confirmation of persistent cortisol secretion which is not suppressible at high-dose dexamethasone suppression test (18,19). The imaging tests are complex from computed tomography, magnetic resonance imaging up to PET (positron emission therapy) imaging (18,19). The therapy targets the specific profile of the causing tumour and, in cases with unknown/unidentified cause, bilateral adrenalectomy might be necessary to control the damage due to Cushing’s syndrome (21). In these cases the CRF but mostly the ACTH excess associates with hyperpigmentation even in a relative short period of time (19).

Nelson’s syndrome

Nelson’s syndrome represents another particular situation where ACTH increase is part of the pathogenic loop and its values can get extremely high (22,23,24). This happens after bilateral adrenalectomy for Cushing’s disease and it represents a rather atypical option nowadays due to others several medical and surgical therapeutical resources which are preferred (22,25,26). The complication is rare; it involves high ACTH with known complications including the skin, and corticotropinoma enlargement (26,27). Prior radiotherapy might be protec-
tive, yet the general data are inconclusive (26,27). There is not general guideline as approach but radiotherapy, neurosurgery of the pituitary gland and potential use of somatostatin analogues and/or dopamine agonists are the therapy lines (26,27). A study published in 2019 showed that 38% of the subjects with Nelson’s syndrome displayed some level of progression during a follow-up of 10 years (28). The risk of progression is higher if complex therapies are applied to Cushing’s disease but this may also be correlated with a more severe form of disease (28). Nelson-Salassa syndrome has been revealed in half of the persons with Cushing’s disease treated with bilateral adrenal removal associating a mild evolution (29). A few cases are described actually underlying a pituitary ACTH-producing carcinoma (29). This is suspected in rapidly growing tumours and ACTH increase with suggestive symptoms (29). Despite this potential severe complication, difficult cases of Cushing’s disease should still be treated with synchronous bilateral adrenalectomy (30).

DISCUSSION

Overall, hyperpigmentation in relationship to ACTH includes its high levels in addition to low cortisol in Addison’s disease and Nelson’s syndrome and high non-suppressible cortisol in ectopic Cushing’s disease. ACTH has a pituitary origin in first two situations and malignancy in the third one. A pituitary tumour is found in cases with Nelson’s syndrome. An autoimmune background may be associated with Addison’s disease. An iatrogenic component is brought by Nelson’s syndrome. All three situations are severe and life threatening of different scenarios (Figure 1).

CONCLUSION

Hyperpigmentation may be the clue to connect dermatology to endocrine pathologies and ACTH massive release by a pituitary or a non-pituitary origin involves a complex panel of conditions.

REFERENCES


