Endocrine paraneoplastic syndromes in lung cancer: a respiratory physician’s perspective

Abstract

Lung malignancy is known to be one of the leading causes of cancer-related mortality. Endocrine paraneoplastic syndromes in lung cancer are common. These are due to secretion of various substances and not because of direct tumour invasion or metastasis. These syndromes have also been associated with lung cancer prognosis. This review describes the many endocrine paraneoplastic syndromes seen in lung cancer and narrates their incidence, biology, clinical features, diagnosis, and management.

Key words: endocrine paraneoplastic syndromes, lung cancer, hypercalcaemia, hyponatraemia, small-cell lung carcinoma

Introduction

Lung cancer is considered to be the foremost cause of cancer-related mortality around the world [1]. Mortality rate in lung cancer is significant and is considered equal to that of prostrate and breast cancers combined. This is mainly because most of these patients present in advanced stages of cancer at the time of diagnosis [2]. The most important risk factor for lung cancer to date is tobacco smoke [3]. The late diagnosis of lung cancer in advanced stages is mainly due to the lack of clinical findings. Some patients may seek medical advice for symptoms not directly related to a malignancy because of the appearance of paraneoplastic syndromes, which in turn may lead to the diagnosis of cancer in the early stage, with early initiation of chemotherapy [4].

Paraneoplastic syndromes (PNS) are seen in malignant conditions with the clinical features caused by either production of hormones or functional peptides secreted by tumour itself. These should not be induced by direct infiltration and growth of the primary malignancy or metastases [5]. Improper immune cross-reaction of tumour cells with normal host cells can also be the rare cause of this syndrome [6]. Most common malignancy associated with PNS is lung cancer. Ten percentage of patients with lung cancer can have these syndromes [7]. In lung cancer, PNS are numerous (Table 1). Humoral hypercalcaemia of malignancy along with the syndrome of inappropriate antidiuretic hormone secretion seen respectively more in squamous-cell carcinoma and small-cell carcinoma are two of the most common endocrine PNS. The size of the primary tumour or stage of cancer has no relation with the symptom severity in these syndromes. These syndromes are diagnosed with specific criteria (Table 2), but all need not be fulfilled in clinical practice [5]. Especially, demonstration of hormones in tumour biopsy tissue is not practical in many cases.

The present review aims to present the incidence, tumour biology, clinical features, diagnos-
tic criteria, and treatment options for endocrine paraneoplastic syndromes in lung cancer. They are briefly described in Table 3.

**Humoral hypercalcaemia of malignancy (HHM)**

**Incidence**

Ten percent of all patients with advanced lung malignancy can have hypercalcaemia, and among

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**Table 2. Diagnostic criteria for paraneoplastic endocrine syndromes**

| Endocrine function abnormality with absent physiologic feedback regulation |
| Respective endocrine lung should not have any metastasis |
| Worsening not explained by increasing tumor burden |
| Cancer treatment improves endocrine function |
| Tumor biopsy sample showing evidence of either hormone substance or its production |

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**Table 3. Four most common endocrine paraneoplastic syndromes encountered in lung cancer and their features**

<table>
<thead>
<tr>
<th>HM</th>
<th>SIADH</th>
<th>Ectopic cushing syndrome</th>
<th>Carcinoid syndrome</th>
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<tbody>
<tr>
<td><strong>Incidence</strong></td>
<td>Baseline: 2–6%</td>
<td>10% of Cushing’s syndrome are paraneoplastic 50–50% of PNU-lung NET (SCLC, carcinoid)</td>
<td>1–5% in bronchopulmonary NET</td>
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<td>During course of cancer: 10%</td>
<td>In SCLC 7–16%</td>
<td>In NSCLC &lt; 1%</td>
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<tr>
<td><strong>Most common histology:</strong> squamous-cell carcinoma</td>
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<tr>
<td><strong>Mechanism</strong></td>
<td>PTHrP (most common): bind to PTH receptors in the bone, kidney and influence calcium, phosphorous regulation</td>
<td>Ectopic ADH secretion by cancer cells which inhibit free-water excretion in the distal tubule of the kidney</td>
<td>Cancer cells express POMC precursor gene which is translated into a prohormone later cleaved into ACTH</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Serotonin release by cancer cells. Can be precipitated by certain food, exercise, or alcohol</td>
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<tr>
<td><strong>Clinical features</strong></td>
<td>Altered sensorium, polydipsia, polyuria, renal failure, vomiting</td>
<td>1) Carcinoid: typical cushingoid features like centripetal fat distribution, systemic hypertension, proximal myopathy 2) SCLC: less cushingoid features, hyperglycaemia common, unexpected weight gain due to chronic water retention</td>
<td>1) Acute: prolonged flushing in upper torso anteriorly, bronchospasm, or diaphoresis 2) Chronic: fibrosis of right heart valves, retropertioneum 3) Rare: carcinoid crisis causing hypotension, cardiac arrest</td>
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<tr>
<td><strong>Diagnosis</strong></td>
<td>↑Ca</td>
<td>↑PTHrP</td>
<td>↓/N PTH</td>
</tr>
<tr>
<td>Clinically euvolemic: serum sodium &lt; 125 mEq/L, urinary sodium &gt; 40 mmol/L, urine osmolality &gt; 100 mOsm/kg</td>
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<tr>
<td>Rule out other causes of hyponatraemia — drug-induced, excess fluids, low intake due to cachexia</td>
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<td></td>
<td></td>
<td>1) Demonstration of hypercortisolism by: increased 24 UFC or salivary cortisol 2) 1 mg dexamethasone suppression test 3) High serum ACTH level with the absence of a pituitary tumour by CT or MRI brain</td>
<td>1) 24-hour urine 5-HIAA 2) Radiolabeled octreotide</td>
</tr>
<tr>
<td><strong>Treatment</strong></td>
<td>1) Tumour excision</td>
<td>1) Tumour excision 2) Ketoconazole, octreotide 3) Bilateral adrenalectomy in refractory cases 1) Surgical excision of tumour 2) During carcinoid crisis: octreotide 3) Pre lung surgery manipulation of tumour-octreotide not usually recommended</td>
<td></td>
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<tr>
<td>2) Restore intravascular volume</td>
<td>2) Firstline — fluid restriction (&lt; 1 L/day) 3) Demeclocycline, vaptans</td>
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<td></td>
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<tr>
<td>3) Bisphosphonates, calcitonin, hemodialysis</td>
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Conversion factors for SI: calcium values to mmol/L, multiply by 0.25; cortisol values to nmol/L, multiply by 27.588; osmolality values to mmol/kg, multiply by 1; PTH values to ng/L, multiply by 1; and sodium values to mmol/L, multiply by 1.

ACTH — adrenocorticotropic hormone; CT — computed tomography; HHM — humoral hypercalcaemia of malignancy; HIAA — 5-hydroxyindoleacetic acid; MRI — magnetic resonance imaging; NET — neuroendocrine tumour; POMC — proopiomelanocortin; PTH — parathyroid hormone; PTHrP — PTH-related protein; SIADH — syndrome of inappropriate antidiuretic hormone secretion
these, the thirty-day mortality rate can be as high as 50% [8, 9]. Bone metastases or altered parathyroid gland function are usually absent in these cases. In lung cancer, the incidence of hypercalcaemia is around 5% at baseline diagnosis and can be twice as this eventually (around 8–12%) during the disease course [5–7, 10]. Most common histology causing HHM is squamous-cell carcinoma, and in up to one-fourth of them, hypercalcaemia can be the presenting feature [5, 6, 10–12].

**Biology and mechanism**

In lung cancer patients, hypercalcaemia in the majority of the cases is usually caused by HHM [13, 14]. Four mechanisms have been observed in malignancy-related HHM (Table 4). The main and most important mechanism of action is the parathyroid hormone-related protein (PTHrP) secreted from cancer cells [12, 14, 15]. Ectopic parathyroid hormone production is one of the rare mechanisms [13]. Sometimes, post-chemotherapy, chronic G-CSF exposure can cause hypercalcaemia in few of them as it promotes osteoclastic bone resorption [16, 17].

Parathyroid hormone molecule has eighty-four amino acids while the PTHrP molecule has 139 to 173 amino acids. These two substances manifest C-terminal portions differently but have a similar first 13 amino acids N-terminal. PTHrP modifies itself into a configuration and binds to the PTH receptor. It can simultaneously bind to other PTH receptors and utilise different effects from PTH [18]. PTHrP binds to receptors of PTH in the kidney and bone. This subsequently influences the bone resorption and renal control of electrolytes, namely phosphate and calcium. PTHrP does not effect on the vitamin D3 1-alpha hydroxylase action unlike parathyroid hormone. In some animal model studies, in a squamous-cell carcinoma cell line, the amphiregulin-EGFR signalling system reconstitution caused HHM [19].

In patients with tumours originating from the lungs [8], including several other organ NETs, hypercalcaemia secondary to PTH secretion has been described [20].

Rarely, tumour cells can release interleukins that can activate the osteoclasts (like IL-1) which can be the cause of malignancy-related hypercalcaemia [21].

**Clinical features and diagnosis**

In lung cancer, HHM incidence is more frequent in those patients with significant disease burden (locally advanced or metastatic disease) [21]. The symptom level depends on the serum calcium concentration (calcium levels more than 14 mg/dL are considered severe), timing of the onset and the patient’s precancer neurologic and renal function [22, 23]. Moderate hypercalcaemia (serum level of 12–14 mg/dL) may result in neuromuscular symptoms (proximal myopathy, fatigue), neuropsychiatric symptoms (anxiety, confusion), circulatory disturbances (polydipsia, severe dehydration causing acute renal failure, polyuria), and gastrointestinal manifestations (abdominal pain, constipation, and rarely pancreatitis).

Hypercalcaemia, when severe, can cause cognitive impairment, confusion, and even coma. Cardiac conduction disturbance and hypotension causing death may also be seen [5, 6, 13, 22].

Many lung cancer patients can be emaciated due to cancer-related cachexia with low serum protein values. So, their serum calcium levels need correction depending on the serum albumin values. Patient’s calcium and albumin values estimation should be always made simultaneously [6].

The diagnosis is confirmed by the following laboratory tests: high serum levels of ionized and total calcium, low to normal parathyroid hormone (PTH) level and high PTH-related protein (PTHrP) concentration [8].

**Treatment**

Treatment of the underlying malignancy as radically as possible is the most successful therapy strategy [6]. The goal of medical care should be attaining electrolyte equilibrium and restoring intravascular volume to prevent the immediate acute complications of serum hypercalcaemia. Two to three liters of intravenous normal 0.9% saline solution will achieve this. Fluid substitution decreases calcium reabsorption in the kidney by increasing the glomerular filtration rate [7, 22]. Medications causing hypercalcaemia (like calcium-containing antacids, supplements of calcium, vitamin D, diuretics like thiazides) or that aggravating mental status changes should be stopped whenever possible [9].

In refractory persistent hypercalcaemia, treatment to reduce the elevated calcium levels should be done as per latest guidelines (including bisphosphonates and possibly denosumab) [6, 23]. Serum calcium level starts to decrease within a day.

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**Table 4. Humoral hypercalcaemia of malignancy causes**

<table>
<thead>
<tr>
<th>Parathyroid hormone-related protein</th>
<th>1.25 dihydroxy vitamin D</th>
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<tbody>
<tr>
<td>Parathyroid hormone</td>
<td></td>
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<tr>
<td>Granulocyte colony-stimulating factor</td>
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and normal values of calcium can be seen within a week following administration of intravenous bisphosphonates. Bisphosphonates have an influence on serum calcium levels for up to 3 weeks [5, 24]. In treating HHM, bisphosphonates are proven to be the safest and most effective agents [24]. They also enhance the survival of patients with bone metastasis [23]. The important complications of bisphosphonates are jaw osteonecrosis due to local vessel vasocnstriction and acute renal failure. Jaw osteonecrosis can cause severe pain locally, jaw swelling, tooth loss, and, rarely, soft tissue loss exposing the underlying bone [5, 6].

In cases of PTHrP-related hypercalcaemia, cinacalcet can be helpful [25]. A low calcium diet and steroids are usually less effective.

Dialysis is generally reserved for those with severe hypercalcaemia causing life-threatening symptoms or in patients with acute renal failure.

Prognosis
In lung cancer, those with significant disease burden (locally advanced or metastatic disease) have more chances of having HHM with mean survival of only around 2 months and poor prognosis [26].

In a study of 1,149 histologically proven lung cancer patients, hypercalcaemia was seen in 6%. Among those with hypercalcaemia, squamous-cell carcinoma was found in 51%, adenocarcinoma in 22% and small-cell carcinoma in 15%. Those with hypercalcaemia mostly had advanced disease (stage 3 or more) with a median survival of only a few months [11].

Serum PTHrP levels in patients with HHM can give information regarding prognosis. In some cases, it can be used to assess tumour response posttreatment. It may also predict the reaction to bisphosphonates. Studies have shown that concentrations above 12 pmol/L are frequently associated with both a smaller reduction in hypercalcaemia and with recurrence of hypercalcaemia within fourteen days of therapy [27, 28].

Prognosis of lung cancer-associated hypercalcaemia is generally poor. Those who become normocalcaemic after bisphosphonate therapy survive better (53 days vs 19 days) [27].

**Syndrome of inappropriate antidiuretic hormone secretion**

**Incidence**
A syndrome of inappropriate antidiuretic hormone secretion (SIADH) causing hyponatraemia with cancer background was noticed in 1957 for the first time. In 1963, from a lung tumour sample (small-cell carcinoma), an agent with ADH-like action was isolated [29, 30]. This syndrome manifests as hyponatraemia with euvolemia clinically, low serum osmolality and unsuitably high urine osmolality. Diagnosis has a specific criterion (Table 4). The tumour cell more commonly produces antidiuretic hormone and rarely atrial natriuretic peptide leading to paraneoplastic hyponatraemia. Antidiuretic hormone increases reabsorption of free water and atrial natriuretic peptide induces natriuresis [31].

SIADH is found in 1–2% of patients with lung cancer. SIADH causing clinical symptoms has been seen to occur in around 10% of lung cancer patients, mainly with small-cell carcinoma histology. Less than one percent of paraneoplastic hyponatraemia is caused by NSCLC. In SCLC, tumour stage does not correlate with the incidence of SIADH. Overall, around 70% of paraneoplastic SIADH cases are caused by SCLC [32–34].

**Biology/mechanism**
ADH is a 9-amino acid peptide usually produced by neurohypophysis. The peptide binds to receptors in the kidney to reduce the excretion of free water. When plasma osmolality exceeds 280 mOsm/kg, the pituitary increases ADH release, causing the kidney to retain more free water and maintain fluid and osmolar balance.

In patients with SCLC, ectopic ADH production causes hyponatraemia by inhibiting free-water excretion in the distal tubule of the kidney. ADH mRNA is expressed in SCLC cells and the peptide is translated and secreted. Measured levels of ADH in plasma are often increased in SCLC [35]. A subgroup of hyponatraemia and concurrent SCLC cases have no detectable levels of plasma ADH. The tumours in these patients express ANP mRNA, secrete the peptide, and have high plasma ANP levels [36].

**Clinical findings/diagnosis**
In SIADH, the level of sodium in serum and its time of onset determines the symptoms. Initially, the patients can have only headache and fatigue as presenting complaints. However, acute onset (< 48 hours) and severe hyponatraemia (serum sodium level of less than 120 meq/L) can cause general seizures, altered mental status and rarely death due to cerebral oedema. Usually, chronic hyponatraemia patients are asymptomatic, especially when it’s mild to a moderate degree.

In the background of hyponatraemia with clinical euvolemia status, > 40 mmol/L of sodium
in urine or > 100 mOsm/kg of osmolality in urine gives a clue to the diagnosis of SIADH [37]. The diagnosis of SIADH is confirmed by laboratory tests as given in Table 5.

Always, in patients having hyponatraemia in the setting of lung cancer, other causes of low serum sodium like inadequate sodium intake, drug-induced kidney injury or excess intravenous hypotonic solutions usage should be excluded.

Treatment

Surgical excision of the tumour is always the ideal treatment and when successful, can bring back the serum sodium level to normal in only some patients [33]. In SCLC, post-chemotherapy symptoms resolution can be seen in up to 80% of cases, but the syndrome would relapse along with the tumour (in 60–70% of patients) [33, 34]. In SCLC, rarely can chemotherapy result in tumour lysis syndrome contributing to acute SIADH [38].

As in any SIADH case, in asymptomatic milder hyponatraemia patients, reduced (1 L/day) intake of free water is the first step. In symptomatic hyponatraemia with serum sodium < 120 mEq, intravenous administration of 3% sodium chloride solution with an infusion speed of up to 1 mL/kg/hour is necessary for the first few hours at least. Correction of serum sodium level should be gradual as acute restoration can lead to irreversible demyelination.

Pharmacologic treatments can be used when conservative measures fail. Demeclocycline lowers renal response to ADH. Vasopressin receptor antagonists like conivaptan and tolvaptan enhance urine excretion of free water. These are effective in some cases.

Prognosis

SIADH per se carries an independent poor prognosis in malignancy [39].

In one study, SCLC patients with persistent hyponatraemia due to SIADH had worse survival. In this study, 61 patients had sodium level of less than 130 mEq/L and received at least two cycles of chemotherapy. Among these 61 subjects, compared to 46 patients whose sodium normalised (to 136mEq/L), the 15 patients in whom there was persistent hyponatraemia (< 136mEq/L) post-chemotherapy had worse survival [40].

Ectopic Cushing’s syndrome (ECS)

Incidence

In 1962, for the first time, connection between CS and ectopic production of adrenocorticotropic hormone (ACTH) was established in a patient with severe hyperadrenocorticism who was found to have SCLC [41]. In Cushing’s syndrome, in up to 10% of cases, the cause can be paraneoplastic [44]. In most of these cases, malignancy involved is lung NET (small-cell carcinoma or bronchial carcinoids) [41–44]. In non-neuroendocrine tumours, rarely are ECS reported [45].

Biology and mechanism

Ectopic production of ACTH happens to be the foremost cause of this endocrine syndrome in lung cancer patients [46]. Rarely, it is caused by corticotropin-releasing hormone (CRH) secretion from tumour cells [47].

The precursor gene, proopiomelanocortin (POMC), is expressed more in the cancer cells from which a 241-amino acid prohormone is translated and then cleaved into ACTH (39 amino acids), melanocyte-stimulating hormone, and opiate-like hormones. The ACTH binds to receptors in the adrenal gland, causing them to produce excessive glucocorticoid and mineralocorticoid hormones [48]. CRH is a 41-amino acid peptide produced and released in the hypothalamus paraventricular nuclei that stimulates the release of ACTH from the pituitary.

Clinical features and diagnosis

Systemic manifestations in this syndrome are mainly due to increased serum cortisol levels. The common clinical features of ECS are skin purple striae, moon like face, acne, proximal myopathy, oedema of the periphery, systemic hypertension, primary metabolic alkalosis, and persistent serum hypokalaemia. Weight gain (due to chronic water retention or centripetal fat distribution) may be one of the rare features in those with lung cancer-related ECS unlike in those without ECS where weight loss is seen because of cachexia [7]. Most of them also have hyperglycaemia [48, 49]. In ECS, due to SCLC, classical signs of Cushing’s syndrome are rare. An important reason for this finding could be the aggressive nature of SCLC causing only brief exposure to excessive ACTH [5, 50].
Those having SCLC with ECS getting chemotherapy are at increased risk for opportunistic infections [51]. ECS is also a risk factor for VTE (2%) [52]. This risk further may increase after surgery of the tumour (4%) [53].

Diagnostic laboratory findings include the following [54]:
1. Minimum of 2 increased measurements of 24-h urine free cortisol;
2. Salivary cortisol sample > 145 ng/dL between 23:00–24:00;
3. 1 mg of dexamethasone suppression test.

An ACTH level can differentiate ECS from Cushing’s in cases of proven serum or urine hypercortisolism. ECS suspicion is raised when an elevated morning ACTH is seen along with the absent pituitary tumour in brain CT or MRI. High-dose dexamethasone will not suppress an ectopic source (like in lung cancer) of ACTH. Bronchial carcinoids are an exception, because in some cases with this tumour type, serum ACTH and cortisol levels have been suppressed by high-dose dexamethasone [55].

To locate the primary tumour in the lungs, whole body somatostatin receptor scintigraphy or thorax imaging like CT can be used.

Treatment
Radical excision of the tumour is the best treatment [56]. When radical therapy of the tumour is unattainable, medications directed to cease the secretion of cortisol (ketoconazole, metyrapone and other drugs) or block (octreotide may block the release of ACTH) are required [7, 57]. With careful monitoring of serum potassium, antihypertensive agents and diuretics can also be used to control symptoms.

Bilateral adrenalectomy is the option used as a last resort in case of no response to medications [5].

Prognosis
Prognosis is affected by tumour type and degree of cortisol level as both these factors influence mortality and morbidity [56].

Most of the patients with SCLC and ECS present at an advanced stage. Even with chemotherapy, their mortality is significantly high since many of these tumours are chemoresistant [5, 7, 56].

Carcinoid syndrome

Incidence
Neuroendocrine tumours of the bronchopulmonary system account for around 20% of all lung malignancies and include typical carcinoid, atypical carcinoid, large-cell carcinoma and SCLC [58].

In neuroendocrine lung cancer, the release of serotonin by tumour cells might trigger the syndrome only in 1–5% of the cases [59]. A lower rate of carcinoid syndrome is seen in lung NET as they produce less serotonin than midgut NET [58]. In patients having localised disease (like in most of the cases of lung NET), carcinoid syndrome is seen most often with tumours of bigger size (> 5 cm) and those with concurrent liver metastasis [60].

Biology and mechanism
Carcinoid syndrome results from the release of vasoactive substances such as serotonin into the systemic circulation. As many as 40 substances (dopamine and many others) related to carcinoid syndrome have been identified as being the potential causes. The release of these substances can be triggered by increased adrenergic activities, such as physical exercise, or increased intake of foods rich in amines (chocolate, kiwi, avocado banana, and nuts) or alcohol.

Clinical features and diagnosis
Some may have acute symptoms like cutaneous flushing, secretory diarrhoea, and bronchospasm. The long-term results of persistently elevated hormone levels include telangiectasias of veins, valvular heart disease (right side more commonly involved), and retroperitoneum fibrosis. Flushing of the skin can be prolonged in the setting of carcinoid syndrome due to lung NET and it occurs more in the upper anterior part of the body [61, 62].

Carcinoid syndrome-associated bronchospasm is less typical and these patients usually have concurrent flushing, sneezing and dyspnoea [63]. In a retrospective study of 748 carcinoid syndrome patients, bronchospasm was seen in 15% [63].

In few cases, due to excessive production of serotonin and its release into the systemic circulation, an acute form of the syndrome can be seen. This is known as carcinoid crisis. These patients can have tachycardia, hypotension, bronchospasm, and even rarely sudden death. Carcinoid crisis is more common after stressful procedures such as anaesthesia, surgery or even radiologic interventions [64]. It can also happen spontaneously.

The evaluation of carcinoid syndrome is with a 24-hour urine collection for the most crucial metabolite of serotonin, 5-Hydroxyindoleacetic
acid (5-HIAA). This test has a specificity of approximately 90% [65].

Radiolabelled octreotide can be used to detect lung neuroendocrine tumours with ectopic hormone production as almost 80% of them demonstrate somatostatin receptors [66].

**Treatment**

Excision of the tumour is the best treatment [67]. Unlike other organ NETs (even with significant liver metastasis), carcinoid crisis risk with lung NETs is low, and hence prophylactic administration of octreotide before any tumour manipulation (biopsy or resection) is not recommended [62]. But still, when handling such tumours, clinicians should be aware of the possibility of the carcinoid crisis and the benefit of octreotide in such a scenario.

**Rare syndromes**

**Hypoglycaemia**

Lung cancer-associated paraneoplastic hypoglycaemia is rare. Non-islet cell tumours with insulin secretion and tumours releasing substances which can cause hypoglycaemia by non-insulin based mechanism are the main causes [68]. This condition is labelled as non-islet cell tumour hypoglycaemia (NICTH). Pulmonary tumours causing this condition are malignant mesothelioma, solitary fibrous tumour and adenocarcinomas [6].

In most of the cases, hypoglycaemia is caused by excess insulin release due to the secretion of peptides like precursors of insulin-like growth factor 1 (IGF-2), insulin-like growth factor 1, and sometimes glucagon-like peptide-1 which are capable of causing glucose utilisation by different mechanisms [68]. Rarely high tumour load having excess glucose utilisation, significant liver infiltration by tumour per se, or tumour metastasis to the endocrine gland (pituitary or adrenal) causing their destruction is the cause of hypoglycaemia in these patients [68].

NICTH is clinically characterised by recurrent hypoglycaemic episodes which affect elderly patients more often. In some, these hypoglycaemic episodes may direct towards the underlying undiagnosed cancer [69, 70].

Diagnosis depends on tumour type the patient is having. In NICTH acute phase, we can find decreased values of following substances in serum-insulin (normal range: 1.44–3.6 μIU/mL) and C-peptide (normal range: 0.3 ng/mL). They would also have increased levels of the following substances in serum: growth hormone, insulin-like growth factor 1, insulin-like growth factor 2 and IGF2:IGF1 ratio. In insulinoma acute phase, we can find increased value of both insulin and C-peptide levels in serum [6, 70].

Surgery of the tumour is the best management option in these patients. The essential goal in case of any hypoglycaemic emergencies is to bring back blood glucose to near expected values with 25% or 50% solution of dextrose. Oral glucose will help in few cases. In the long run, treatment of hypoglycaemia due to this syndrome may require glucagon, growth hormone and corticosteroids [6, 69–71].

**Acromegaly**

Only 1% of acromegaly is caused by growth hormone releasing hormone or growth hormone ectopic secretion by tumour cells. Of these, the majority are caused by carcinoid tumours of the lung and intestine [72]. In most cases, the GHRH gene is expressed by the lung cancer cells, and a 40–44 amino acid peptide is produced. Circulating GHRH peptide binds to receptors in the pituitary gland resulting in the production of excessive amounts of GH 73]. Rarely, lung carcinoid tumours express immunoreactive GHRH and result in abnormal GH secretion [74].

The earliest features of GH excess are hyper trophy of the extremities and face. The diagnosis of ectopic acromegaly is established by elevated serum levels of GHRH or GH, the absence of a pituitary tumour, complete recovery following lung tumour resection, positive GHRH immunostaining, detection of GHRH mRNA, positive bioassay (pituitary cells of rat on culture produce GH when subjected to tumour extract), or GHRH extraction from the tumour tissue [75]. However, coincidental pituitary tumours and lung solid tumours have also been described.

Management of ectopic acromegaly should be surgical resection of the tumour and is often curative in those with lung carcinoid. In those with unresectable or metastatic cancers, medical therapy with somatostatin analogues, such as octreotide and bromocriptine, have been shown to be effective [76].

**Ectopic secretion of chorionic gonadotropin**

This syndrome has been reported in large-cell lung carcinoma (LCLC). The syndrome causes Leydig cell hyperplasia which results in raised oestrogen levels and reduced testosterone production. This gives rise to atrophy of testicles with repression of spermatogenesis, and gynecomastia [77].
Conclusion

Despite the availability of new diagnostic techniques and biological and surgical treatment development, lung malignancy mortality is the foremost cause of cancer-related deaths since 1985. In lung cancer, the small-cell histology type is commonly associated with endocrine paraneoplastic syndromes. These syndromes are easy to diagnose and treat because they have a clear pathogenic pathway. Endocrine paraneoplastic syndromes have both a prognostic role as well as a predictive function in tumour treatment prediction. The prognosis also depends on the cure of the underlying tumour. Tumour progression can present along with syndrome recurrence. The most common syndrome is humoral hypercalcaemia of malignancy seen in squamous-cell histology and measurement of PTHrP can delineate it from primary hyperthyroidism.

Conflict of interest

None declared.

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28. Ling PJ, Allern RP, Hardy JR. Analysis of survival following treatment of tumour-induced hypercalcaemia with intrave-


