Cushing’s disease secondary to a pancreatic Ewing Sarcoma in an 18-month-old child: a case report

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Abstract

Background: Extraskeletal paediatric Ewing Sarcoma with resultant Cushing’s syndrome due to ectopic ACTH (adrenocorticotrophic hormone) production is a very rare presentation with a poor prognostic outlook. This is the first pancreatic case ever reported.

Case presentation: An 18-month-old Caucasian female presented with rapid onset weight gain (2kg in two weeks), stridor and overnight apnoea, and irritability. She had a random paired plasma cortisol of 1115nmol/L [170-550] and ACTH of 6.5pmol/L [1-12 pmol/L], and undetectable corticotrophin releasing hormone. An abdomen CT (computerised tomography) revealed a 64 x 69 x 80mm mass in the left upper quadrant which displaced the pancreatic tail and involvement of the splenic vasculature. After surgical resection, histopathology confirmed a diagnosis of Ewing Sarcoma. She received Children’s Oncology Group (COG) protocol AEWS1031, followed by consolidation chemotherapy, and gradually weaned hydrocortisone replacement. There were no signs of recurrence until a surveillance abdominal ultrasound scan 2 years after diagnosis revealed a 10mm porta hepatis lesion. Multiple further intra-abdominal lesions developed, and were unresponsive to 4 cycles of high dose ifosfamide. She was transferred to palliative care 31 months after her initial diagnosis with abdominal pain, hypertension and respiratory discomfort. Her plasma ACTH was raised at 18.6 pmol/L [1-2] with a random cortisol of 1232 nmol/L [170-500], and she redeveloped Cushingoid symptoms at 36 months post initial diagnosis. She died peacefully 37 months after her initial diagnosis.

Conclusions: There have only been six cases of ACTH producing Ewing Sarcoma reported, with this case being only the second extraskeletal one ever reported in literature, and the first that is pancreatic in origin. Regardless of treatment modality, no patients have survived for more than seven years after their initial diagnosis.

Keywords: Extraskeletal Ewing Sarcoma; Cushing’s disease; paediatric; case report.

Abbreviations: ACTH: Adrenocorticotropic Hormone; CT: Computerised Tomography; USS: Ultrasound Scan; IV: Intravenous; ICU: Intensive Care Unit; MRI: Magnetic Resonance Imaging; COG: Children’s Oncology Group.

Background

Paediatric Ewing Sarcoma is a well reported condition with known treatment options and prognostic indicators. A very rare subset of this condition consists of ACTH producing tumours which cause a resultant Cushing’s syndrome. Existing literature describes five such cases ever reported, with only one extra-skeletal case. We present the presentation and clinical course of an ACTH producing extra-skeletal Ewing Sarcoma. The aim of this case presentation is to add a pancreatic case to the current literature and contribute to the limited knowledge of prognosis for these patients.
Case Presentation

An 18 month old Caucasian female presented with rapid onset weight gain (2 kg in two weeks), stridor and overnight apnoea, and irritability. On examination she weighed 14 kg (+2.45SD, 99th centile) with a height of 76.4 cm (-1.48SD, 5th centile) with a BMI of 23.99 (+4.68, >99th centile). She was markedly hypertensive (100/50 mmHg) (95th centile). She was cushingoid in appearance (central obesity, rounded face and dorsocervical fat pad) as seen in Figure 1. Although she had a tanned complexion she had recently been on holiday in a Pacific Island country, and her tanning did not include non-sun-exposed areas. Hirsutism over the lower back was noted, but there was no pubic or axillary hair. The impression was that she had features consistent with Cushing’s disease and went forward for further investigation.

Laboratory results demonstrated paired random plasma cortisol of 1115 nmol/L (170-550) and ACTH of 6.5 pmol/L (1-12 pmol/L), and undetectable corticotrophin releasing hormone. A random urinary cortisol was 2711 nmol/L and a 24 hour collect was 996 nmol/day (<380 nmol/day). Catecholamines were normal. Aldosterone was below detection limits, and renin was 49.2 mU/L (5.3-99.1 mU/L). Glucose levels were in the normal range. A contrast CT of the abdomen showed a 64 x 69 x 80 mm mass in the left upper quadrant with partial displacement of the pancreatic tail medially and splayed over the surface. The mass abutted the underside of the stomach without a clear fat plane. Although it closely abutted other organs in the area there was no suggestion of invasion. The splenic vasculature was intimately associated with the surface of the mass (Figure 2).

She went forward for surgical removal with a presumptive diagnosis of pancreatic neuroendocrine tumour with ectopic ACTH production. A transverse left upper quadrant incision was made. The mass was identified and dissection around the pancreas performed. Intra-operatively, as expected from pre-operative imaging, the mass was found to be closely associated with the splenic artery and splenic hilum requiring splenectomy to maximise the chance of adequate surgical margins. The tumour was removed en bloc with the spleen and tail of pancreas and the distal pancreatic tail oversewn. Histopathology is shown in Figure 3, and confirmed Ewing sarcoma.

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Haematoxylin and eosin stained sections showed a malignant epithelioid tumour with small round blue cell morphology (A). Immunohistochemistry was positive for synaptophysin (B), CD99 (C) and cytokeratin AE1/3 (D). NXX2.2 immunohistochemistry was also positive, and subsequent fluorescent in situ hybridisation testing demonstrated an EWSR1 rearrangement, in keeping with Ewing sarcoma.

Post-operatively she recovered from the surgery well. Her blood pressure rapidly returned to normal. Endocrine management consisted of intra and post-operative hydrocortisone replacement at physiological levels. This was weaned over the following weeks with subsequent normal adrenal axis testing. She received Children’s Oncology Group (COG) protocol AEWS1031- six cycles of alternating Vincristine/Doxorubicin/cyclophosphamide with Ifosfamide/etoposide. This was followed by consolidation chemotherapy of Vincristine, Doxorubicin, cyclophosphamide, Ifosfamide and etoposide. After completing the six cycles she was discharged with a haemoglobin of 71 g/L (105-140), and returned the following day for a red blood cell transfusion. Twenty minutes into this she developed tachypnoea, tachycardia, fever and hypoxia, and was admitted to ICU (intensive care unit) for four days. She was treated with steroids, IV (intravenous) tazocin (for an episode of febrile neutropenia), high-dose co-trimoxazole and oxygen for a probable chemotherapy and transfusion-related
acute lung injury of uncertain aetiology. Diffuse ground glass opacities were noted on chest CT. She subsequently developed pulmonary hypertension and right heart strain. She remained oxygen dependent for 14 months, and was gradually weaned. She was on prednisone for two years, and stopped sildenafil after 14 months. Chemotherapy was continued for another five months, during which time she had one ICU admission for an acute respiratory deterioration, but otherwise progressed well. Prior to starting prednisone, at 3 months post initial presentation, her morning basal cortisol levels were normal at 203 nmol/L [170-500]. They increased appropriately after Synacthen was administered to a peak of 1262 nmol/L [>400], demonstrating she had normal adrenocortical reserves.

Follow up radiology for the subsequent 14 months after completing chemotherapy showed no disease progression. She remained asymptomatic, but a surveillance abdominal USS (ultrasound scan) two years after her initial diagnosis found an indeterminate 10mm lesion in the porta hepatis region. A surveillance chest x-ray also found a possible new lung nodule, but this was later dismissed. A subsequent CT scan 25 months after her initial diagnosis found an ill-defined lesion in the portacaval region that extended into the porta hepatitis, with hyper-attenuation of adjacent liver parenchyma. There were no other significant radiological findings. Abdomen MRI (magnetic resonance imaging) at 26 months found a further soft tissue nodule between the left adrenal and pancreatic tail, and involvement of multiple portal lymph nodes. She received four cycles of high dose ifosfamide at 29 months post presentation, but further CT scans showed growth of all known disease sites, with the porta hepatitis region at 14x19x27mm, and threatened occlusion of the duodenum, left portal and renal veins. No new disease sites were identified. No further chemotherapy was given. After stopping chemotherapy, her plasma ACTH was raised at 18.6 pmol/L [1-2] with a random cortisol of 1232 nmol/L [170-500] consistent with the recurrence ACTH secreting endocrine neoplasia.

She was transferred to palliative care at 31 months post initial presentation with abdominal pain, hypertension (138/72) despite ongoing antihypertensives, and increased respiratory discomfort due to hepatomegaly and decreased lung volume. At 36 months she developed Cushingoid features again with rapid weight gain (Figure 4), hyperpigmentation on extensor surfaces of joints, irritability, headaches and hypertension. Ketoconazole was started to try and alleviate her symptoms, however she deteriorated further and died peacefully 37 months after her initial diagnosis.

**Table 1:** Summary of published cases of ACTH producing Ewing Sarcoma.

<table>
<thead>
<tr>
<th>Author</th>
<th>Demographic</th>
<th>Presenting Symptoms</th>
<th>Tumour Site</th>
<th>Treatment</th>
<th>Outcome</th>
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</thead>
<tbody>
<tr>
<td>Preeyasombat C et al(2)</td>
<td>12-year-old female, Thailand</td>
<td>Lower L leg swelling, acne, voice deepening, rapid weight gain, muscle weakness, increased appetite, skin hyperpigmentation</td>
<td>Left tibia</td>
<td>Amputation of lower L leg</td>
<td>No recurrence of Cushing’s syndrome after amputation. Died 3 years after diagnosis due to metastases.</td>
</tr>
<tr>
<td>Guran T et al(3)</td>
<td>9-year-old female, Turkey</td>
<td>Rapid weight gain, acne, hyperpigmentation on neck, bilateral pretibial oedema, abdominal mass</td>
<td>retroperitoneal</td>
<td>10 courses of chemotherapy vincristine/adriamycin/cyclophosphamide, radiotherapy, surgical resection</td>
<td>No recurrence after surgery reported. Cushing’s syndrome resolved. Died 1 year later from chemotherapy complications.</td>
</tr>
<tr>
<td>Galland-Decker C et al (4)</td>
<td>13-year-old male, Switzerland</td>
<td>On relapse: truncal obesity, leg/facial oedema, acute psychosis</td>
<td>8th rib</td>
<td>Surgical resection, chemotherapy (EURO-EWING 99 protocol with vincristine, actinomycin and ifosfamide), ketoconazole</td>
<td>Relapsed after 5 years, developed Cushing’s and psychosis after 7 years, died at age 20</td>
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<tr>
<td>Di Ruscio V et al (5)</td>
<td>10-year-old male, Italy</td>
<td>Rapid weight gain, depression, muscle weakness, widespread acne, hirsutism</td>
<td>Right ischiopubic &amp; ilio-pubic branches</td>
<td>One round of chemotherapy (ISG/AIEOP ISG EW-2 protocol with vincristine, adriamycin, ciclofosfamide, ifosfamide and etoposide). Followed by two courses of high dose ifosfamide when neurological status worsened.</td>
<td>Disease was widely metastatic on diagnosis. Died 6 months after diagnosis.</td>
</tr>
<tr>
<td>Varghese J et al (6)</td>
<td>9-year-old female, Nepal</td>
<td>Abdominal distension, moon face, hypertension, proximal muscle weakness, hyperglycaemia, hypertension</td>
<td>Left proximal humerus</td>
<td>Chemotherapy (cyclophosphamide/vincristine/adriamycin, ifosfamide/etoposide), ketoconazole</td>
<td>Died 7 days after starting chemotherapy</td>
</tr>
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</table>

Declarations

Ethics approval and consent to participate: Written consent for participation has been obtained from the patient’s parents. No ethics approval was required.

Consent for publication: Written consent for publication and the use of images was obtained from the patient’s parents.

Availability of data and materials: Not applicable.

Competing interests: The authors declare that they have no competing interests.

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Authors’ contributions: KC and RL wrote the manuscript and reviewed the literature. MD performed the histopathology and wrote that section of the manuscript. KM was the principal surgeon. SC was the principal oncologist. MDB was the principal endocrinologist. All authors read and approved the final manuscript.

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References


