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ABOUT COVER
Editorial Board Member of World Journal of Clinical Cases, Dr. Antonio Corvino is a PhD in the Motor Science and Wellness Department of University of Naples “Parthenope”. After obtaining his MD degree from the School of Medicine, Second University of Naples (2008), he completed a residency in Radiology at the University of Naples Federico II (2014). Following post-graduate training at the Catholic University of Rome, yielding a second level Master’s degree in “Internal Ultrasound Diagnostic and Echo-Guided Therapies” (2015), he served on the directive board of Young Directive of Italian Society of Ultrasound in Medicine and Biology (2016-2018). His ongoing research interests involve ultrasound and ultrasound contrast media in abdominal and non-abdominal applications, mainly in gastrointestinal, hepatic, vascular, and musculoskeletal imaging. (L-Editor: Filipodia)

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Late-onset non-islet cell tumor hypoglycemia: A case report

Shunichi Matsumoto, Eijiro Yamada, Yasuyo Nakajima, Naoki Yamaguchi, Takashi Okamura, Toshiki Yajima, Satoshi Yoshino, Kazuhiuko Horiguchi, Emi Ishida, Masashi Yoshikawa, Jun Nagaoka, Sho Sekiguchi, Mai Sue, Shuichi Okada, Izumi Fukuda, Ken Shirabe, Masanobu Yamada

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Author contributions: Yamada E contributed to the care of the patient, data collection, the literature review, and drafting the manuscript; Matsumoto S contributed to the care the patient, data collection and the literature review; Nakajima Y, Yamaguchi N, Okamura T and Yajima T contributed to the care of the patient; Yoshino S, Horiguchi K and Ishida E contributed to drafting the manuscript; Yoshikawa M, Nagaoka J, Sekiguchi S and Sue M contributed to the care of the patient; Okada S contributed to the drafting of the manuscript; Fukuda I contributed data on performing the procedure;

Abstract

BACKGROUND
Hypoglycemia due to non-insulin-producing tumors is referred to as non-islet cell tumor hypoglycemia (NICTH). As NICTH is a rare lesion, the natural course of NICTH is not well understood. We report a case of NICTH that was observed 30 years before the onset of hypoglycemia.

CASE SUMMARY
A 50-year-old man was diagnosed with an abnormal right chest shadow during a routine X-ray examination, but no further examination was undertaken because the lesion appeared benign. Thirty years after the tumor discovery, the patient was admitted to the hospital with symptoms of severe hypoglycemia, which was diagnosed as NICTH based on a complete examination. The tumor was resected and found to be a solitary fibrous mass (15.6 cm × 13.7 cm × 10.4 cm); thereafter, the patient’s blood glucose levels normalized and he completely recovered.

CONCLUSION
NICTH can have an acute onset, even if the tumor has been present and asymptomatic over a long time period.

Key Words: Hypoglycemia; Non-islet cell tumor hypoglycemia; Blood glucose; Solitary tumor; Late onset; Case report; Insulin-like growth factor II
Shirabe K and Yamada M did critical revisions to the manuscript; all authors issued final approval for the version to be submitted.

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CARE Checklist (2016) statement: The authors have read the CARE Checklist (2016), and the manuscript was prepared and revised according to the CARE Checklist (2016).

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Core Tip: Since hypoglycemia due to non-insulin-producing tumors is a rare lesion, so the natural course of non-islet cell tumor hypoglycemia (NICTH) is well understood. Here we describe a rare case of NICTH that was caused by a tumor that had been asymptomatic for 30 years. To our knowledge, this is the longest reported latency period before the onset of severe hypoglycemia. The sudden-onset of severe hypoglycemia in the patient described in this report indicates that NICTH can have an acute onset even when the tumor has been present for a longer time.

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INTRODUCTION

Hypoglycemia due to non-insulin-producing tumors is referred to as non-islet cell tumor hypoglycemia (NICTH)[1-3]. NICTH is a rare lesion, but constitutes the second commonest cause of hypoglycemia after insulinoma; approximately 25% of hypoglycemic cases occur due to NICTH[4]. Moreover, NICTH is typically induced due to the overconsumption of glucose or overexpression of insulin-like growth factor II (IGF-II) by tumors[5-7]. Moreover, high-molecular-weight IGF-II, known as big IGF-II, has potent insulin-like activity that leads to hypoglycemia[8]. However, big IGF-II fails to build a complex with both IGF-binding protein-3 (IGFBP-3) and its acid-labile subunit, whereas it binds easily to insulin receptors to induce glucose uptake in muscles and adipose tissues, thus leading to hypoglycemia[9]. Tumors of mesenchymal or hepatic origin are usually described as NICTH. Solitary fibrous tumor (SFT) is another rare mesenchymal tumor, wherein approximately 4%-6% of SFT cases develop NICTH[10].

CASE PRESENTATION

Chief complaints

During a routine chest X-ray examination, a 50-year-old man was diagnosed with an abnormal shadow in the right chest. As there were no signs to suggest the mass was malignant, the patient decided to forego further examination or treatment.

History of present illness

Ten years after the initial discovery, a routine chest X-ray revealed an increase in the size of the previously detected shadow, and the patient was advised a biopsy and tumor removal. However, the patient refused further examination or treatment in the absence of any symptoms. Thirty years after the initial discovery, the patient was hospitalized with symptoms of severe hypoglycemia. Prior to the acute episode necessitating hospitalization, the patient had no history of hypoglycemic symptoms, such as sweating, palpitations, and loss of consciousness. Moreover, there was no history of an increase in weight for the preceding 6 mo. As the patient’s hypoglycemia was resistant to treatment by frequent oral glucose supplementation, he was hospitalized for further management.

History of past illness

The patient had no medical records or family history of hypoglycemia.

Personal and family history

The patient was on treatment with azelnidipine for hypertension and rosvustatin for...
Physical examination
Clinical examination revealed no signs, except attenuation of respiratory sounds over the right chest.

Laboratory examinations
Laboratory test results (Table 1) after hospitalization showed a slight increase in C-reactive protein (CRP) levels to 2.11 mg/dL. The anti-insulin antibody test result was negative, and fasting blood glucose was 48 mg/dL, accompanied by a low immunoreactive insulin (IRI) level of < 5.0 µU/mL. Interestingly, there was no change in the levels of insulin-counterregulatory hormones, such as cortisol, growth hormone (GH), and norepinephrine (Table 2).

Imaging examinations
Chest X-ray and computed tomography (CT) scanning showed a giant solid tumor (11 cm × 14 cm × 15 cm) in the right lower chest region (Figure 1A and B). A fluorine-18 fluoro-2-deoxy-D-glucose (FDG) positron emission tomography showed uneven accumulation of the maximum standardized uptake value (SUVmax 3.1) indicating possible characteristics of a malignancy (Figure 1D). Interestingly, the mass showed a propensity for accumulation by octreotide scintigraphy, which was characteristic of a neuroendocrine tumor (Figure 1E).

Further diagnostic work-up
From the biochemical and histological findings, we suspected an NICTH. To confirm this diagnosis, we undertook a core biopsy of the tumor. Immunoblots of serum IGF-II and tumoral tissue IGF-II from tumor biopsies were conducted[10], and a high-molecular-weight form of IGF-II was identified in both types of samples on Western blotting (Figure 2A). Furthermore, immunohistochemical staining for IGF-II in tumoral tissue showed numerous immunopositive tumor cells (Figure 2B).

FINAL DIAGNOSIS
Based on the findings from the examination and investigations, the patient was diagnosed with NICTH.

TREATMENT
Because of problematic symptoms that were refractory to clinical treatment, we obtained written informed consent to carry out tumor resection. The resected mass was shown to be an SFT (15.6 cm × 13.7 cm × 10.4 cm; Figure 2C).

OUTCOME AND FOLLOW-UP
Postoperatively, the patient’s blood glucose levels quickly normalized, and the patient recovered completely.

DISCUSSION
Hypoglycemia is a usual feature that is observed during the treatment of NICTH. However, in the present case, NICTH was mainly diagnosed on the basis of a hypoglycemic attack[8]. Another symptom that implicated the tumor as an NICTH was its size, which exceeded 10 cm[2,8]. There are no reports in the literature whether an increase in the tumor size induces hypoglycemia; however, in our case, the size of a tumor could be related to the occurrence of hypoglycemia. Moreover, this assumption is supported by the theory that partial resection of the tumor could reduce the incidence of hypoglycemic episodes[11]. In the present case, we observed the progression of the tumor over 30 years, wherein the increase in tumor size eventually caused hypoglycemia. Interestingly, the initial phenomenon of a hypoglycemic attack
Table 1 Laboratory values at admission (post glucose infusion)

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<thead>
<tr>
<th>Clinical values (normal range)</th>
<th>Clinical values (normal range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC (/μL)</td>
<td>6800 (4000-9600)</td>
</tr>
<tr>
<td>Hb (g/dL)</td>
<td>14.7 (13.2-17.3)</td>
</tr>
<tr>
<td>Platelets × 10^4 (/μL)</td>
<td>21.9 (16-35)</td>
</tr>
<tr>
<td>Total protein (g/dL)</td>
<td>7.3 (6.3-7.9)</td>
</tr>
<tr>
<td>Albumin (g/dL)</td>
<td>3.7 (3.9-5.0)</td>
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<tr>
<td>Total bilirubin (mg/dL)</td>
<td>0.6 (0.3-1.2)</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>18 (13-33)</td>
</tr>
<tr>
<td>ALT (U/L)</td>
<td>10 (8-42)</td>
</tr>
<tr>
<td>LDH (U/L)</td>
<td>221 (119-229)</td>
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<tr>
<td>ALP (U/L)</td>
<td>229 (115-399)</td>
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<tr>
<td>γ-GTP (U/L)</td>
<td>16 (10-47)</td>
</tr>
<tr>
<td>ChE (U/L)</td>
<td>257 (213-501)</td>
</tr>
<tr>
<td>AMY (U/L)</td>
<td>78 (49-136)</td>
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<td>Blood urea nitrogen (mg/dL)</td>
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<td>Creatinine (mg/dL)</td>
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<tr>
<td>Na (mEq/L)</td>
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<tr>
<td>K (mEq/L)</td>
<td>4.0 (3.5-4.8)</td>
</tr>
<tr>
<td>Cl (mEq/L)</td>
<td>107 (100-107)</td>
</tr>
<tr>
<td>T-Cho (mg/dL)</td>
<td>206 (128-219)</td>
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<tr>
<td>TG (mg/dL)</td>
<td>44 (30-149)</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>5.4 (4.6-6.2)</td>
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<tr>
<td>Glycoalbumin (%)</td>
<td>15.8 (11.0-16.0)</td>
</tr>
<tr>
<td>Insulin antibody</td>
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<tr>
<td>CEA (ng/mL)</td>
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<td>SCC (ng/mL)</td>
<td>1.0 (0-1.5)</td>
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<td>NSE (ng/mL)</td>
<td>10.6 (0-12)</td>
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<td>CYFRA (ng/mL)</td>
<td>2.1 (&lt; 3.5)</td>
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<tr>
<td>ProGRP (pg/mL)</td>
<td>51.8 (&lt; 80)</td>
</tr>
<tr>
<td>SLX (U/mL)</td>
<td>19.4 (0-38.0)</td>
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<td>Blood glucose (mg/dL)</td>
<td>130 (80-110)</td>
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<tr>
<td>Immunoreactive insulin (μg/mL)</td>
<td>6.8 (1.0-21.74)</td>
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<tr>
<td>Serum C-peptide (ng/mL)</td>
<td>1.50 (1.1-3.3)</td>
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<td>Free T3 (pg/mL)</td>
<td>2.30 (1.88-3.18)</td>
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<tr>
<td>Free T4 (ng/dL)</td>
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<td>Adrenocorticotropic hormone (pg/mL)</td>
<td>77.0 (7.2-63.3)</td>
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<tr>
<td>Cortisol (μg/dL)</td>
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<tr>
<td>Human growth hormone (ng/mL)</td>
<td>&lt; 0.07 (&lt; 2.10)</td>
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<tr>
<td>IGF-1 (ng/mL)</td>
<td>92</td>
</tr>
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</table>

WBC: White blood cell; Hb: Hemoglobin; AST: Aspartate aminotransferase; ALT: Alanine aminotransferase; LDH: Lactate dehydrogenase; ALP: Alkaline phosphatase; γ-GTP: gamma-glutamyl transpeptidase; ChE: Cholinesterase; AMY: Alpha-amylase; T-Cho: Total cholesterol; TG: Total triglyceride; HbA1c: Hemoglobin A1c; SCC: Squamous cell carcinoma-related antigen; CEA: Carcinoembryonic antigen; NSE: Neuron-specific enolase; CYFRA: Cytokeratin-19
fragment; ProGRP: Progastrin releasing peptide; SLX: Sialyl Lewis X; T3: Triiodothyronine; T4: Tetraiodothyronine; IGF: Insulin-like growth factor.

<table>
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<tr>
<td>Blood glucose (mg/dL)</td>
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<tr>
<td>Immunoreactive insulin (μg/mL)</td>
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<td>Serum C-peptide (ng/mL)</td>
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<td>Free T3 (pg/mL)</td>
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<td>Glucagon (pg/mL)</td>
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<tr>
<td>Epinephrine (ng/mL)</td>
</tr>
<tr>
<td>Norepinephrine (ng/mL)</td>
</tr>
<tr>
<td>Dopamine (ng/mL)</td>
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</tbody>
</table>

T3: Triiodothyronine; T4: Tetraiodothyronine.

**Figure 1 Imaging findings.** A and B: Chest X-ray (A) and computed tomography (B) (left-hand side is transverse plane; right-hand side is coronal plane) showed a heterogeneous giant mass measuring 11 cm × 14 cm × 15 cm in size on the right lower chest; C: Computed tomography from ten years ago showed the tumor’s growth in the past decade; D and E: Positron emission tomography (D) and octreotide scintigraphy (E) showed the tumor’s accumulation. Without any increase of body weight possibly implies that sudden-onset NICTH could occur during the tumor progression.

Besides the overexpression of big IGF-II, NICTH of hepatic origin could be attributed to irregular of gluconeogenesis[2], or an increase of glucose consumption by
big tumors\(^{[12]}\). Notably, the tumor, in the present case, showed an increase in the FDG uptake, indicating glucose uptake by the tumor. The histopathological diagnosis confirmed the tumor as an SFT with no evidence of malignancy; therefore, the sign of FDG uptake could indicate the possible development of malignancy, and increased glucose uptake by the tumor may have induced hypoglycemia through big IGF-II. Indeed, insulin-like effects of big IGF-II are reported to lead to increased glucose uptake in insulin-sensitive tissues, especially muscle and fat\(^{[8]}\), although they may induce glucose uptake in the tumor itself\(^{[1]}\).

NICHTH is supposed to be induced by big IGF-II insulin-like activity; however, there are reports that IGF-II might regulate other insulin-counterregulatory hormones, such as GH and IGF-II, which could lead to hypoglycemia\(^{[2]}\). Furthermore, IGF-II could downregulate the expression of IGF-II\(^{[1]}\). Interestingly, the initial pathological phenomenon of a hypoglycemic attack without previous weight gain and potentially implies the sudden-onset occurrence of NICHTH, to immediately exceed the threshold during tumor progression.

The complete treatment of NICHTH is total resection of tumor, which could be difficult because of the characteristics of the tumor itself, metastasis, location, size, and so on\(^{[1-2,7]}\). In the present case, the total resection of the tumor could be undertaken despite the inconveniences of the patient’s age and treatment preference. Therefore, we considered other treatment options: Introduction of intravenous hyperalimentation, enteral tube feeds, local therapies (e.g., embolization, radiation), systemic therapies (e.g., chemotherapy, targeted antitumor therapy such as imatinib), glucocorticoids, rh GH, glucagon, octreotide, diazoxide, or bendrofluazide\(^{[2,3]}\). However, we found none of the options to be sustainable for self-management. This was especially because the patient showed an accumulation of octreotide, demonstrating tumor characteristics typical of a neuroendocrine tumor. Nonetheless, a high concentration of octreotide is not a definitive indicator for NICHTH even if octreotide is accumulated within the tumor\(^{[13]}\). Considering all of these factors, we finally obtained informed consent from the patient to undertake lung resection.

**CONCLUSION**

In conclusion, we describe a case of NICHTH that was observed for 30 years, which is most likely the longest reported duration up to the onset of severe hypoglycemia thus far. The sudden-onset severe hypoglycemia in the present case indicates that NICHTH could occur immediately with IGF-II levels above threshold during the tumor progression even over a longer time course.
REFERENCES


