-	Case Report
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3	Carotid Body Tumour a Challenging
4	Management: Rare Case Report in Baghdad
5	Radiation Oncology Center, Medical City,
6	Baghdad, Iraq
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10 11	ABSTRACT
	 Background: Paragangliomas are rare asymptomatic painless tumors, originating from paraganglionic bodies of autonomous nerve system. The carotid body tumors are relatively rare tumor but constitute majority of head and neck paragangliomas about 70%. These tumors are generally benign but possess aggressive local growth potential. Aim of study: The purpose of this case report article is to introducing the challenging we exposed to it during management such cases despite lack many conditions, facilities and circumstances found in other parts of world for treatment of CBT, which is the first time study in Iraq. Case presentation: We reported a case of 27 year-old female patient with a left neck swelling that had persisted for 5 years. She was first diagnosed and treated surgically by local excision in 2012. Histopathology revealed the histology of carotid body tumor. She was complaining of hypertension, headache, palpitation, and low grade fever. She was observed gradual onset of growing and pulsated lump on previous surgical scar site. At October 2017, the mass growing faster, and she felt pulsation on lying on left side resulting in discomfort on sleep and deglutition. On physical examination a pulsating firm painless mass measuring about 6 x 5 x 4 cm in size was found on the left side of her neck. Pulsations were felt on deep palpation and a faint bruit was heard on auscultation. Work up done for her included laboratory tests, neck US, neck CT scan, carotid angiography and slid review of histopathology. All suggested recurrent carotid body tumor. Conclusion: Throughout more than four decades working in this field, we faced 2-3 cases of CBTs, so it is very rare tumor. It is a challenging tumor whether decide to treat by surgery or radiotherapy. The main step in management of CBT is excluding of others tumour may be present in same region. Surgery is treatment of choice while radiotherapy is standard treatment for recurrent cases.
12 13 14 15 16	Keywords: Carotid body tumour; Paraganglioma; Chemodectoma; Glomus cells; Carotid Arteriography ABBREVIATIONS [[CBT: carotid body tumour, COPD: chronic obstructive pulmonary diseases, CCA: common

FNA: fine needle aspiration, US: ultra sound, CT: computed tomography, MRA: magnetic resonance arteriography, 3D: three-dimensional, RT: radiotherapy, CBC: complete blood pictures, RFT: renal function tests, LFT: liver function tests, Gy: gray, LAP: lymphadenopathy, SOL: space occupy lesion.]]

22 **1. INTRODUCTION**

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24 Carotid body tumour (CBT) or chemodectoma is a rare, highly vascular, mostly benign 25 tumour arising from the paraganglia of carotid body [1]. CBTs are nonchromaffin 26 paragangliomas arising from the chemoreceptor cells found at the carotid bifurcation. The 27 tumour is highly vascular; its blood supply is the richest per gram of tissue of any tumour [1]. 28 The carotid body, which originates in the neural crest, is important in the body's acute 29 adaptation to fluctuating concentrations of oxygen, carbon dioxide, and pH [2]. The carotid 30 body protects the organs from hypoxic damage by releasing neurotransmitters that increase 31 the ventilatory rate when stimulated [2, 3]. There are three different types of CBTs have 32 been described: familial, sporadic and hyperplastic. The sporadic form is the most common 33 type. The familial type (10-20%) is more common in younger patients. The hyperplastic form 34 is very common in patients with chronic hypoxia, which includes those patients living at a 35 high altitude and may be found in patients with COPD and cyanotic heart diseases. The first 36 anatomical description of carotid body was provided by Albrecht Von Haller in 1743 [2, 3, 4]. 37 Histologically, carotid body tumors have a characteristic growth pattern often referred to as a 38 zellballen [5, 6]. Carotid body tumors occur at any age but are typically diagnosed between 39 the third and sixth decades of life [5, 7, 8]. The usual presentation is a slow growing mass at 40 the angle of mandible [2, 4, 9]. It is must involve thorough evaluation for primary tumour of 41 the thyroid, the oropharynx, and the nasopharynx is essential, since metastases to a cervical 42 lymph node is a much more frequent cause of a neck mass than CBTs [9]. Carotid body 43 tumors can be a diagnostic challenge for the clinician and lack of pre-operative diagnosis 44 has been reported in up to 30% of the cases in different series [4]. The diagnostic work-up of 45 CBT may involve one or more of the following: Duplex US scanning, CT, MRI, MRA, carotid 46 arteriography, and serum and urinary catecholamine level assessment [2]. The treatment 47 modalities for CBTs are surgical excision and/or radiotherapy [5, 9, 10, 11, 12, 13]. 48

49 2. CASE PRESENTATION

A 27 year-old female patient presented with a left neck swelling that had persisted for 5 50 years. She was first treated by local excision at 29th July 2012 in a private hospital. 51 52 Pathology revealed the histology of carotid body paraganglioma. She was in good health for 53 two years before she developed her signs and symptoms of catecholamine excess such as 54 hypertension, headache, palpitation, and weight loss. She was consulted many doctors and 55 centers for her problem and she kept on medical treatment and follow up. In the last nine 56 months she observed gradual onset of growing and pulsated lump on previous surgical scar 57 site. Suddenly in two weeks exactly in October 2017, before she came to Baghdad Radiation 58 Oncology Center, the mass growing faster and she felt pulsation on is lying on left side resulting in discomfort on sleep and deglutition. Any information about past surgical data and 59 60 past history of patient were missed in Irag war against ISIS which included surgical file, IHC study and others investigations. 61

62 **1. Physical examination**

A pulsating firm painless and well circumscribed mass measuring about 6 x 5 x 4 cm in size was found on the left side of her neck, near the angle of the mandible. There were pressure symptom & more mobile transversely than vertically. Pulsations were felt on deep palpation and a faint bruit was heard on auscultation. She was complain of headache, tachycardia and hypertension.

- 68 **2. Work up**
- 69 2.1. Laboratory tests

70 CBC, ESR, RFT, LFT, electrolytes balance and urine analysis were done and all were 71 normal values. Catecholamine (vanilmandelic acid) levels in urine or serum was unavailable 72 in our city.

- 73 **2.2.** Imaging
- 74 CXR and Echo study also done.

75 **2.2.1.** Neck US

It revealed highly vascular ill-defined hypoechoic solid mass seen above level of Left carotid
 bifurcation between ICA and ECA measured 35 x 20 mm, reaching up to lower border of
 parotid gland suggestive of recurrent CBT. Normal other structures.

79 **2.2.2. CT** scan of neck

80 It demonstrated a well-circumscribed lobulated hypervascular, hypoechogenivic mass with 81 splaying of the carotid bifurcation. The mass was heterogeneously intense enhancing soft 82 tissue density mass of size 38 × 24 × 25 mm. the findings suggested recurrent CBT. There 83 were no infiltration into adjacent structures is seen. Both thyroid lobes were normal in size, 84 echogenicity, normal isthmus, no cervical LAP, normal both submandibular salivary glands, 85 normal neck vessels and no SOL.

86 **2.2.3.** Carotid angiography of neck

87 It showed a well-defined highly vascular blush mass (density is 250 HU while in the carotid artery is 340 HU), is about 33 x 27 x 25 mm, located at the lateral to both ICA and ECA of 88 left carotid arteries and about 2 cms above the carotid bifurcation. Mass is not intrinsic to 89 blood vessels & patency of the artery was preserved. This finding was considered to be 90 consistent with a diagnosis of recurrent CBT. In compares with previous carotid angiography 91 92 done in June 2012, there was irregular solid oval soft tissue density mass 63 x 39 x 27 mm 93 in size seen in left upper the neck in left retro mandibular region beneath left sternomastoid 94 muscle, extending from level of below left parotid gland down to below mandibular angle. It 95 was vascular mass and protruding between ICA and ECA. There was no cervical LAP. The 96 features were CBT.

97 **2.2.4. MRI, MRA and Octreotide Scan was unavailable and very cost to done.**

98 2.3. Histopathology

The previous histopathology was done at July 2012. Grossly there was a globular piece of tissue measured 5 x 3 x 2 cm, with a grey brown cut section, also there were three firm grey nodes. Histologically, the sections showed well defined nests of uniform cuboidal cells (Zellballen) with abundant granular basophilic cytoplasm, separated by highly vascularized fibrous septa. There was no mitoses or necrosis seen. The picture was consistent with carotid body tumour CBT. The three LNs showed reactive follicular hyperplasia. The slide review done and give the same result.

106 **3. Treatment**

At that time, the mass was quite large with size and attached to left common carotid artery.
Therefore, surgery was not suitable, because of very high risk for surgical injury to these
vessels and nerves and she was a recurrent case. Finally, she was referred to our

radiotherapy unit for definitive radiotherapy. The radiation technique was 3D-conformal
radiotherapy (3D-CRT) with 2 Gy/daily fraction to the total dose of 50 Gray in 25 Fractions.
The GTV (gross target volume), which was defined as gross tumor, was shown on contrast
CT scan. The CTV (clinical target volume) was the volume including GTV and 1 cm margin
around. The PTV (planning target volume) was the volume including CTV and 1 cm margin
around. The normal tissue constraint was limited within tolerance dose.

116 **4.** Follow up

There was not any complications regarding post radiation courses therapy. Symptoms like hypertension, palpitation, and flushing were disappeared. The tumor size was regressed regarding last neck U/S, and she will wait for the result of neck CT scan and CT angiography. She was planning for follow up every two months in this year.

121 **3. DISCUSSION**

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123 Carotid body tumors CBTs are rare and uncommon entities may be found in unilateral or 124 both sides of the neck and in both genders at the different rate of frequency, and they belong 125 mostly to a benign group of tumors and their surgical management is technically challenging. 126 The most common presentation in patients with CBT is a slowly enlarging painless mass in 127 the neck. Locally invasive growth of these tumors subsequently leads to cranial nerve 128 deficits along with compression symptoms like Horner's syndrome, syncope, hoarseness 129 and dysphagia since the carotid body functions as a chemoreceptor organ that is stimulated 130 by hypoxia, hypercapnea, and acidosis, it is involved in the control of blood pressure, heart rate, and respiration [1, 3, 5, 11, 13]. CBD are slow growing painless masses localized in the 131 132 neck, anterior to the sternocleidomastoid muscle at the level of the hyoid bone. As the 133 tumour grows, dysphagia, odynophagia, dysphonia, and symptoms due to compression of 134 cranial nerves IX to XII may be seen. The most commonly involved cranial nerve is the 135 vagus, up to one third of all cases will show cranial nerve palsies [3, 5, 8, 11].

US is the first non-invasive procedure which allows seeing that is vascular discrimination between the solid and cystic nature of the mass. Carotid arterial angiography is the most valuable diagnostic technique, it is the gold standard for diagnosis is carotid angiography, which serves both diagnostic and treatment purposes. MRI and CT angiography can demonstrate the extent of the tumor and its relationship to adjacent structures [2, 7, 8].

There are many choices of treatment for CBD including observation, surgery, external beam radiotherapy, and stereotactic radiotherapy. Surgery is the treatment of choice. The local control by surgery alone is approximately 85-100%. Most of the reports demonstrated local control with radiotherapy alone is approximately 80-90% comparable to surgery [10, 11, 13]. Depending on the size and location of the lesions, the indication for RT may be either:

- 146 1. Primary irradiation in the case of functional or other inoperability.
- 147 148
 - Adjuvant irradiation for R1 to R2 resections.
 Irradiation of recurrence if there is progression after surgery [2, 12].

149 Treatment with radiotherapy can achieve comparable local control and less morbidity than 150 surgical resection in paraganglioma. Regarding definitive radiation treatment of CBTs. There 151 are many techniques, protocols and radiation dose ranges of treatment. Although 152 stereotactic radiotherapy has been increasingly used and their results have been generally accepted, conventional radiotherapy and 3D radiotherapy are still commonly used in the 153 154 place where stereotactic radiotherapy is not available. Many reports used a radiation dose of 155 45 Gy in 25 fractions, with a daily dose of 1.8 Gy [3, 5, 6, 12]. Continued follow-up is 156 necessary, however, as recurrence and metastasis may occur years later [12].

157 **4. CONCLUSION**

CBT is a rare paraganglionic tumour affecting both sex of reproductive age. Treatment of choice is surgery. Radiotherapy is standard treatment in recurrent and inoperable cases. Since late recurrences are known, patients should be adequately need for periodic follow up so that recurrences can be identified early and treatments offered. The limitation of this study is that Immunohistochemistry was not done to support the histological diagnosis and was because the markers for this technique are currently out of stoke in country.

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Fig. 1. Head and Neck CT scan (axial and sagittal plan).

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170 Fig. 2. Carotid angiography

171 COMPETING INTERESTS

- 172 Authors have declared that no competing interests exist.
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175 CONSENT (WHERE EVER APPLICABLE)

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All authors declare that 'written informed consent was obtained from the patient (or other approved parties) for publication of this case report and accompanying images.

180 ETHICAL APPROVAL (WHERE EVER APPLICABLE)

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All authors hereby declare that all experiments have been examined and approved by the
appropriate ethics committee and have therefore been performed in accordance with the
ethical standards laid down in the 1964 Declaration of Helsinki.

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