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Castleman Disease in the Pediatric Neck: Case Report and Literature Review



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ABSTRACT

Objective: To investigate the common features of cervical pediatric Castleman disease (CD).

Study Design: Case report and literature review of pediatric patients with cervical CD.

Methods: Online medical journal databases were searched for patients aged 18 years or younger. Eighteen published papers were found, comprising 29 cases. One case from our institution was also included for a total of 30 patients.

Results: An asymptomatic mass in level V was the most common presentation. No gender differences were noted. Multiple forms of imaging were pursued, and no particular modality showed signs specific for CD. All cases were treated with complete surgical excision and diagnosed as hyaline-vascular type on histology, except for one case, where histologic type was not reported. No reports of multicentric disease, plasma cell, or mixed histology were found. No recurrences were reported.

Conclusions: This poster provides the largest known literature review of pediatric patients with cervical CD. In our analysis, there is a higher propensity for level V than previously reported in small studies. While CD is rare, it should be considered on the differential for a pediatric neck mass, particularly when presenting with an asymptomatic posterior neck mass and equivocal work-up. Fortunately, our study suggests that, if diagnosed as CD, the most likely diagnosis is hyaline-vascular type for which the long-term prognosis is good. Surgical excision is both diagnostic and therapeutic.

INTRODUCTION

Castleman disease (CD) was first described by Benjamin Castleman in 1954.^{1,2} Since that time, CD has become better known in literature as a lymphoproliferative disorder of unknown etiology.³⁻⁵ Castleman disease can occur anywhere throughout the lymphatic system. The most common sites include the mediastinum (60%), neck (14%), abdomen (11%), and axilla (4%).⁶⁻¹¹ While the underlying etiology is unknown, several hypotheses have been suggested. One theory postulates that the disease represents a reaction to a chronic viral antigenic stimulation. Some studies cite a role for interleukin-6 (IL-6).⁸

Castleman disease is rare in the pediatric population, although exact prevalence rates are not known.^{8,12} In children, CD has a more benign prognosis. It also has a different propensity for certain anatomic sites compared to adults, most commonly affecting the chest (33%), abdomen (30%), neck (23%), and axilla (7%).^{12,13} In contrast, the most commonly affected sites in adults are the chest (60%), neck (14%), abdomen (11%), and axilla (4%).⁷⁻¹¹

To better understand the clinical characteristics of this rare entity in children, in this study, we present a case study followed by an extensive literature review of pediatric cervical CD.

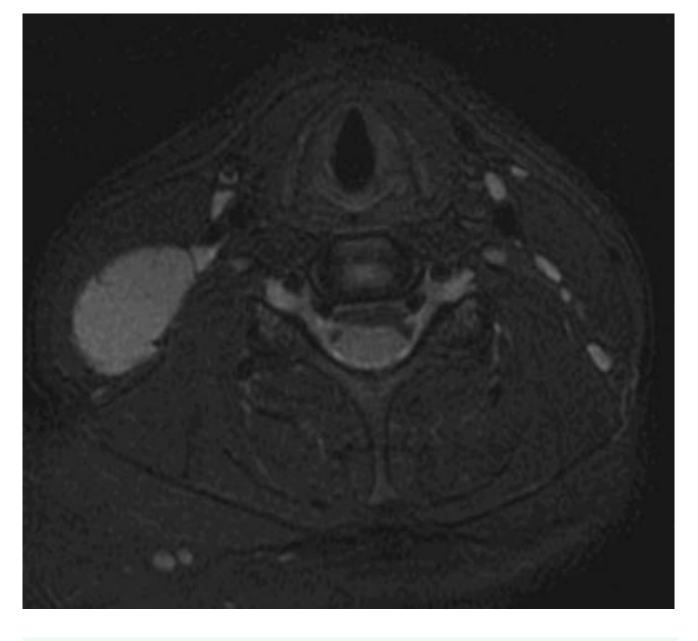


Figure 1. Neck magnetic resonance imaging of right-neck mass. Axial T2-weighted cut shows a hyperintense mass.

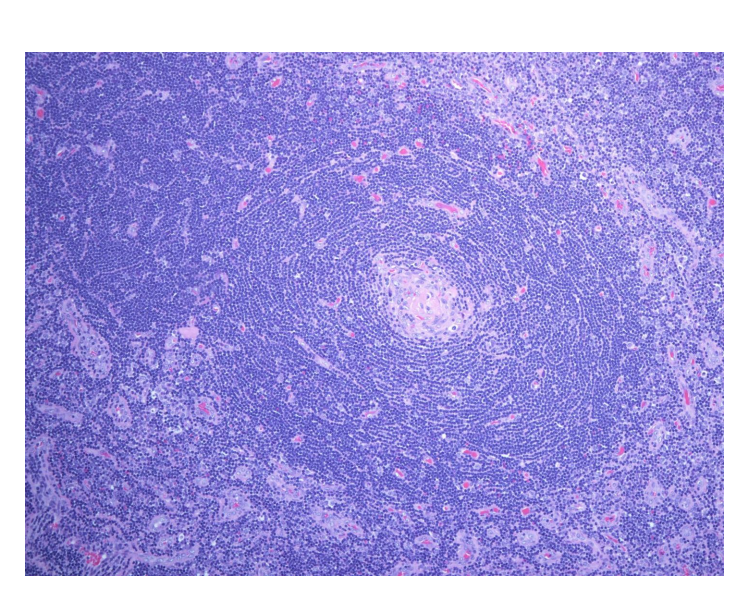


Figure 2. Castleman disease showing lymphoid follicle with onion-skinning of the mantle zone lymphocytes; 200x.

Nemours.

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CASE REPORT

A 13-year-old female presented to our institution with a tender right-neck mass that appeared suddenly. It had been present for six weeks, during which time the mass had not changed in size but did cause mild pain with head movement to the right. She had no significant previous medical history. Exam revealed a 4 x 6-cm mass deep to the lower half of the sternocleidomastoid muscle on the right. It was non-tender to palpation with no overlying skin changes. Routine laboratory tests were within normal limits. Titers for toxoplasma, cytomegalovirus (CMV), and Bartonella were negative. Epstein–Barr virus (EBV) IgG titers were elevated. Chest radiograph was within normal limits. Magnetic resonance imaging (MRI) revealed a well-defined, right-sided level III mass measuring 1.8 x 3.0 x 4.2 cm. It was bright on T2- and intermediately to slightly brighter than muscle on T1-weighted imaging (Figure 1). Several small vascular channels were apparent on the lesion by MRI. Several small lymph nodes along the inferior margin of the lesion extending down to the thoracic inlet were also noted. An additional 12 x 6 x 9-mm lesion was noted in the paraspinal musculature. A decision was made not to pursue this lesion given its small size and location, which would be unusual for CD. Fine needle aspiration (FNA) revealed atypical lymphoid proliferation. After discussion with the patient and family regarding the options, she was taken to the operating room, where a complete surgical excision of the mass was performed. Histologically, the nodal mantle zone showed concentric rings with an onion-skin appearance. Piercing blood vessels were frequently seen in these follicles ("lollipop" feature) (Figure 2). The interfollicular areas showed prominent hyalinized venules. Based on these findings, the postoperative histopathological diagnosis was HV-CD. At three months' follow-up, she was doing well with no signs of recurrence.

MATERIALS AND METHODS

Online medical journal databases were used for data collection. "Castleman's disease" in combination with "neck", "cervical," and "pediatric", were keywords used for searching the PubMed database. Only patients aged 18 and younger were included for analysis. After excluding reports on CD in other locations (ie, non-neck sites), 18 published papers were found, comprising 29 total reported cases of pediatric cervical CD (**Table 1**). The earliest case report published was in 1991 and the latest in 2012.^{5,19} In addition, one patient was diagnosed and treated at our institution. This patient was also included and brought our final patient count to 30 cases. This study was IRB exempt. All diagnoses of CD were based on histopathology.

Table 1. Castleman data

						Duration	n								Abnormal					
Reference	Year	Age	Gender	Location/Level	Signs/symptoms	(mo)	Size (cm)	CT Neck	MRI Neck	U/S Neck	CXR	CT C/A/P	Other Imaging	FNA?	Labs	DDx	Treatment	Path	F/U (mo)	Recurrence
in ¹⁴	2010	9	М	L level V	painless neck mass	6	1.5 x 2.0	well-defined neck mass	X	x	х	CT C/A- neg		x	х	X	excision	CD, HV-type	36	none
enfold⁵	1991	11	М	L level V ("post neck")	enlarging mass x 6mo	132	3.0	X	x	x	nl	x		х	x	x	excision	CD, HV-type	х	х
azer ¹⁵	1995	11	E			2	3.2 x 2.5 x 1.6	Ŷ	isointense w/ muscle, central		Y	MRI C/A- neg		Y	2020	×.	excision	CD, HV-type	Y	Y
azer	1995	11	F	R level V ("post neck")	slow growing mass	3	3.2 X 2.3 X 1.0	X	hypodensity, no enhance	appearance of large LN	X	-	Annia	X	none	carotid body chemodectoma,	excision	CD, HV-type	X	X
oslin ¹⁶	1985	17	М	R neck	recent growth, no pain, dysphagia, dyspnea, hoarseness, fatigue, wt loss	5	3.6 x 3.6	homogenously enhancing, displaced vascular bundle medially	x	x	x	х	Angio- mass w/ vasc blush	x	x	vagal neuroma, granulomatous d/o, lymphoma	excision	CD, HV-type	x	х
akabay ⁴	2009	17	F	R level V ("post neck")	neck swelling	12	4.0 x 2.0 x 2.5	irregular borders	Х	Х	х	х	х	Х	none	Х	excision	CD, HV-type	?	none
alisbury ¹⁷	1990	5	М	L cervical	slow growing mass	24	4.0 x 2.5 x 2.5	X	X	x heterogeneous mass w/	Х	x	X	X	X	x	excision	CD, HV-type	11	none
atel ¹⁸	1998	12.5	М	L level III ("ant tri, level thyroid cart")	asymptomatic	х	3.0 x 2.5 x 1.0	X	х	hypoechoic central portion, central flow	x	х	х	x	x	enlarged, necrotic LN	excision	CD, HV-type	х	х
140			_	, , , , , , , , , , , , , , , , , , , ,					intermediate T1 signal, elevated T2							-				
ond ¹⁰	2003	13	F	Level I (midline submental)	unsightly neck mass	24	2.0	x enhancing mass. Homogeneous,	signal	X	Х	X	Х	X	Х	X	excision	CD, HV-type	Х	X
ierlinckx13	1997	14	М	R lateral neck	asymptomatic	24	4.2 x 3.8 x 1.9	dense.	x	x	х	CT C/A- neg	Х	х	none	X	excision	CD, HV-type	8	none
									homogeneous, isointense T1 signal, hyperintense T2 signal, moderate			E	M aspiration- neg		?? Reference values not					
uerlinckx ¹³	1997	7	М	L cervical	increasing mass size	2	5.0 x 4.5 x 2.0	X	enhancement	Х	nl		Abd- neg	Х	given	Hodgkins	excision	CD, HV-type	5	none
hong ⁶	2010	14	F	R neck	none	12	2.0 x 2.0	X	X	X	х	x	Х	Х	none	X	excision	CD, HV-type	34	none
									uniform signal T1, nonuniform signa	oval shaped w/ clear boundary, uniform low- level echo, intact										
hong ⁶	2010	13	F	R neck	none	3	2.0 x 3.0	X	T2	envelope	x	x	x	х	none	X	excision	CD, HV-type	60	none
			_				3.4 x 0.9 (max4			performed- no characteristics				unspecific reactive	pos- lgG	lymphoma, reactive				
buza ¹¹	2008	12	F	L Level III	none	6	LN in total)	no description	X	discussed	X	CT C/A- neg	X	lymphoid hyperplasia	Rubella, CMV hypochromic	lymphadenitis	excision	CD, HV-type	36	none
										performed- no characteristics				unspecific lymphoid hyperplasia from	microcytic anemia, thrombocytosis , high ESR, rheumatoid	cystic hygroma removed from				
ouza ¹¹	2008	16	F	R Level II/III	slow growing mass	96	10.0	X	x	discussed elliptical hypoecogenic	x	x	x	previuos cytology many small lymphcytes	factor pos	same area 4 years prior	excision	CD, HV-type	34	none
	2009	11	F	R neck	increasing mass size	E	3.6	, and the second s		nodule, well-defined		v	v	and lymph-histiocytic	2020	lymphoid hyperplasia,	excision	CD, HV-type	25	2020
buza ¹¹ hen ¹⁹	2008 2012	13	F	L level II	x	24	4.0 x 4.0	x performed- no characteristics given	X	smooth limits x	x	x	x	aggregates x	none x	lymphangioma x	regional dissection	CD, HV-type	62	none x
hen ¹⁹	2012	13	M	R level II	x	12	4.0 x 3.0	performed- no characteristics given	x	x	x	x	X	X	x	X	excision	CD, HV-type	109	X
hen ¹⁹	2012	13	F	L level II	x	12	3.5 x 2.5	performed- no characteristics given	x	x	х	x	х	х	х	х	excision	CD, HV-type	83	х
nen ¹⁹	2012	13	М	L level lb	Х	12	2.5 x 2.5	performed- no characteristics given	х	x	х	x	х	x	х	x	level lb dissection	CD, HV-type	16	Х
hen ²⁰	2006	12	F	Level I*	asymptomatic	6	5.0 x 4.0 x 2.5	X	x	x	х	х	х	x	х	x	excision	CD, HV-type	х	none
hong ¹²	2004	6	F	R neck	asymptomatic mass	2	2.5 x 3.0 x 2.0	homogeneous mass	X	х	nl	х	MRA	х	none	х	excision	CD, HV-type	6	none
ong ²¹	2006	7	М	Level IV*	x	х	2.0 x 1.0	isodense to soft tissue, well circumscribed	x	x	х	x		? Inconclusive (doesn't say which pts got FNA)	none	x	excision	CD, HV-type	10	none
		I	IVI		Λ	^		isodense to soft tissue, well	^	^	~	~		? Inconclusive (doesn't		^			10	none
ong ²¹	2006	11	М	Level II*	Х	х	2.0 x 1.0	circumscribed isodense to soft tissue, well	Х	x	Х	Х		say which pts got FNA) ? Inconclusive (doesn't	none	Х	excision	CD, HV-type	11	none
ong ²¹	2006	17	F	Level III*	X	x	3.0 x 3.0	circumscribed	х	x	x	x		say which pts got FNA)	none	x	excision	CD, HV-type	13	none
n ²²	2010	16	F	Level II/III*	asymptomatic	2	4.2 x 3.5 x 2.0	x	Х	x	х	х	х	X	х	X	excision	CD, HV-type	х	none
1 ²²	2010	9	М	Level V*	asymptomatic	6	2.0 x 2.0	x	X	x performed- no characteristics	х	x	X	X	x	x	excision	CD, HV-type	X	none
a0 ²³	2008	5	М	L Level V	painless neck mass	11	1.5 x 1.5	Х	х	discussed	nl	x	abd u/s	х	none	reactive lymphadenopathy	excision	CD, type not mentioned	6	none
hen ²⁴	2007	9	F	L neck	X	24	X	X	x	X	x	х	X	x	X	X	excision	CD, HV-type	39	none
n ²⁵	2002	10	М		non tandar wahila wasa	2	4.0	ovoid enhancing node, low-attenuation	v	×.	Y		Y	, v		N.	Y			
an ²⁵ abinowitz not yet	2003	12	M	L level III	non-tender, mobile mass	3	4.0	stellate band in the center of mass	X bright T2, int to slightly bright to muscle on T1, slight uniform	X	X	X	X	x atypical lymphoid	X	x lymphoma, fibroma,	Х	CD, HV-type	X	X
INC YOL	2012			R Level III	painful neck mass		1.8 x 3.0 x 4.2									manphonia, noronia,				

x, unknown or not reported; *, unknown side; CD HV, Castleman disease hyaline-vascular type; abd, abdomen; u/s, ultrasound; CXR, chest x-ray; CT, computed tomography; MRI, magnetic resonance imaging; f/u, follow-up; C/A/P, CT chest/abdomen/pelvis; FNA, fine needle aspiration; DDx, differential diagnosis; mo, months

representing 25% of lesions.

- this conclusion definitively.
- Radiographic imaging is non-specific.⁴
- recurrences seen during a mean follow-up of 30 months.

Cervical pediatric CD is rare. It most commonly presents as an asymptomatic or slowly enlarging level V mass. Imaging characteristics are often non-specific and do not aid in the diagnosis. Imaging is important in excluding other diagnoses and to allow for preoperative planning. No specific lab abnormality is consistently seen in these patients, nor is FNA diagnostic. Excision is ultimately diagnostic and therapeutic, and when presenting in the pediatric neck, the diagnosis is likely HV-CD, which holds a favorable prognosis.

DISCUSSION

• In adults, the most common locations for CD include the chest (60%), neck (14%), abdomen (11%), and axilla (4%); in children, the chest (33%) remains the most common site of disease, followed by the abdomen (30%), neck (23%) and axilla (7%).⁶⁻¹³ Data are currently inconclusive as to the most common neck level for pediatric CD. In our literature analysis, level V was the most common location,

• All of the evaluated children had unicentric masses. While no multicentric disease was found, it is important to note that only 17% (5 of 30) of children in this literature review received full body work-ups to rule out this possibility. Therefore, it is difficult to make

• CT was the most common modality used for neck mass work-up in this analysis (47%); however, the results argue that CT is no more specific for diagnosing CD than any other modality. As such, it is reasonable to conclude that while CD should remain on the differential diagnosis for any pediatric neck mass, the imaging modality of choice should be whichever modality will evaluate for the etiology highest on the differential diagnosis for an individual patient. Additionally, since most of these patients will proceed to surgery, pursuing CT or MRI may serve a dual purpose of both attempted diagnosis and preoperative planning. Imaging to search for multicentric disease should be based on the patient's symptoms and on clinical suspicion.

• Definitive treatment of unicentric CD involves surgical excision, with excellent prognosis. Lin et al reported no recurrence after 109 months of follow-up in one patient, which represents the longest follow-up of pediatric neck CD to our knowledge.¹⁷ Another study showed a 100% five-year control rate after surgical excision of an isolated cervical mass.¹² Our analysis supports this data with no

CONCLUSIONS

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