Surgical treatment of an abnormally positioned right adrenal tumor on segmental caudal vena cava aplasia in two dogs

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ABSTRACT. Segmental caudal vena cava (CVC) aplasia is a rare congenital vascular anomaly in dogs. Two dogs were diagnosed by CT imaging to have right adrenal tumors with concomitant segmental CVC aplasia. During surgery, a firm connection between the right adrenal gland and CVC was observed in both cases. The adrenal glands were found ventral to the CVC and the adrenal tumor was resected including the vascular wall. CVC venectomy for tumor removal will be required if the right adrenal gland is displaced in dogs with segmental CVC aplasia, even if there is no intravascular invasion.

KEY WORDS: adrenal tumor, dog, segmental caudal vena cava aplasia

Segmental caudal vena cava (CVC) aplasia is a congenital vascular anomaly with sporadic incidence in dogs. This malformation is usually observed incidentally on diagnostic abdominal imaging or autopsy since most cases are asymptomatic [1, 2, 9]. In dogs, vascular invasion is relatively common with adrenal tumors. Among them, pheochromocytomas are more likely to develop tumor thrombus into the CVC via the phrenicoabdominal vein [7]. Several surgical procedures have been described for CVC, including portosystemic shunts (PSS) and thrombi formation [2, 9]. However, the influence of an anatomical abnormality in determining the choice of surgical technique for segmental CVC is not well known. Herein, we describe the diagnosis and successful surgical management of segmental CVC in two dogs with right-sided adrenal tumors.

Case 1: A 10-year-old castrated male, Papillon, weighing 8.7 kg showed elevated alkaline phosphatase (>3,500 IU/l, reference range: 47-254 IU/l) in a comprehensive medical examination. A referral veterinarian performed an abdominal ultrasound examination to inspect the cause of this abnormality, and an enlarged right adrenal gland (14-mm) was found. Furthermore, the size and location of the right adrenal gland were also abnormal. Initially, the right adrenal gland did not seem to be located dorsal to the caudal vena cava (CVC); however, the precise position could not be identified due to the restless nature of the dog. The dog underwent an adrenocorticotropic hormone (ACTH) stimulation test, and the results were within normal limits. Adrenalectomy was scheduled, and preoperative CT imaging was performed for surgical planning. The CT scan showed that the tumor was located ventral to the CVC, but abnormal vessel anatomy was not observed, and dog was diagnosed with segmental CVC aplasia immediately before surgery. There was no vascular invasion observed into the CVC (Fig. 1).

Intraoperatively, a firm connection was found between the right adrenal gland and CVC. Transient vascular occlusion was performed using Rummel tourniquets during the tumor resection. A portion of the vascular wall of the CVC was removed, followed by reconstruction of the CVC wall defect by continuous suture pattern using 5–0 polypropylene suture (PROLENE: Johnson and Johnson, New Brunswick, NJ, USA). During anesthesia monitoring during the surgery, no cardiovascular abnormalities were observed. The dog recovered uneventfully. An anticoagulant (heparin sulphate, 75 units/kg, SC: subcutaneously, tid) was administered before surgery and was maintained until the next morning. The tumor was histopathologically diagnosed as a concurrent adrenocortical adenocarcinoma and pheochromocytoma. The dog expired 1,005 days postoperatively due to an unrelated cause.

Case 2: An 11-year-old intact female, Miniature Schnauzer, weighing 11.6 kg, was diagnosed with a 13-mm non-functional right adrenal tumor during an ultrasonographic assessment for diarrhea and vaginal discharge that persisted for a week. The position of
the right adrenal gland was ventral to the CVC (Fig. 2). Preoperative abdominal CT imaging was performed for surgical planning. The dog was diagnosed to have an adrenal tumor with concomitant segmental CVC aplasia (Fig. 3) and pyometra. Similar to the first case, venous invasion of the adrenal tumor was not observed. During surgery, the right adrenal tumor was bluntly dissected. It was observed to be firmly attached to the CVC (Fig. 4). The tumor was removed with transient occlusion of the CVC using Satinsky forceps. Vascular wall resection and reconstruction were performed using the same procedures as in case 1. The dog recovered uneventfully. An anticoagulant (heparin sulphate, 75 units/kg, SC, tid) was administered before surgery and was maintained until the next morning. The histopathological diagnosis was adrenocortical adenocarcinoma. This dog died 737 days postoperatively due to an unrelated cause.

Segmental CVC aplasia, also known as azygos continuation of the CVC or CVC uniting with the azygos vein, is very rare congenital vascular anomaly in dogs [1, 2]. The incidence of CVC without other anomalies is approximately 1–3% in dogs. This is usually observed on diagnostic abdominal imaging or on autopsy [1, 2]. Segmental CVC aplasia is rare in dogs, but more cases are being diagnosed due to the increase in use of CT scan for abdominal diseases [2]. In fact, in the first case under consideration, segmental CVC was overlooked on preoperative CT imaging. Since most cases are asymptomatic, the true incidence of segmental CVC could be much higher than its reported incidence.

There are seven types of CVC aplasia, classified by Schwartz et al. as follows: type 1, right lateral cavo-right-azygos shunt; type 2, right medial cavo-right-azygos shunt and small blind ending CVC cranial to the left kidney; type 3, large aneurysmal right medial cavo-right-azygos shunt with an isthmic connection to the azygos vein; type 4, split CVC and right medioventral cavo-right-azygos shunt; type 5, dorsal cavo-right-azygos shunt; type 6, aneurysmal cavo-left-azygos shunt with connecting portal vein; and type 7, split CVC, aneurysmal cavo-left-azygos shunt, and connecting portal vein shunt vessel [9].

Nine of the 37 reported cases had concurrent PSS [9]. In this case report, CT examination was conducted for the two dogs who developed right adrenal tumor with segmental CVC aplasia and the cases were identified as right-lateral cavo-right-azygos shunts (Type 1) [9]. The two cases were thoroughly assessed for the presence of shunt vessels and no vascular abnormalities were seen other than segmental CVC aplasia.

Most cases are asymptomatic and are usually identified as an incidental finding as the cases in this report [9]. Besides the concurrent PSS, this vascular abnormality poses certain risks, such as thrombosis [3, 5, 9]. Among the three reported cases, one dog was successfully managed with multiple surgeries and administration of aspirin [3], one dog died intraoperatively due to poor condition [5], and in one dog, the large thrombi were left without treatment and it survived for more than 1.5 years with no clinical signs [5].

Surgery is uncommon in dogs with segmental CVC aplasia because most cases are asymptomatic [9]. It is not known whether the anatomical differences from normal dogs could affect the treatment of surgical diseases. However, it was expected to have some difficulty in surgical intervention for abdominal disease, especially in the area between the right kidney and the liver in right lateral cavo-right-azygos shunt cases. In fact, the two cases in this report suggested that this abnormality could make removal of the right adrenal gland difficult.
In normal dogs, the outflow of blood from the adrenal gland into the CVC is through the adrenal vein and then returns through the common trunk (formerly known as the phrenicoabdominal vein) [4]. Adrenal tumors often develop vascular invasion into the CVC via the common trunk [6, 7]. In these cases, vessel occlusion is required and is performed using Rummel tourniquets and Satinsky forceps [6, 7]. In our cases, the normal right common trunks were present, which branched and coursed in the normal position, and in the opposite side of the displaced right adrenal gland.

The prehepatic CVC is from the embryologic subcardinal veins, and the renal part of the CVC originates from the subcardinal and suprarecardinal veins [5, 8]. In the present cases, the subcardinal veins may have degenerated while a remnant of the suprarecardinal vein influenced the abnormal running of the right common trunk. It was strongly suggested that due to this abnormality in the vessel anatomy, the right adrenal glands were directly attached to the CVC. Because of this finding, further dissection can cause severe bleeding from a punctured CVC during surgical manipulation. Besides, considering the principle of
In normal dogs, the right adrenal gland is located dorsal to the CVC [4], but in our experience, the abnormal location of the right adrenal gland relative to the CVC suggests a congenital abnormality. The right common trunk receives the right caudal phrenic vein and the right cranial abdominal vein [4], and these vessels connect to the diaphragm and dorsal muscles, which contribute to the dorsally fixed right adrenal gland. In the present report, the adrenal glands were displaced ventrally to the CVC. Furthermore, the right adrenal gland was medially displaced, as confirmed by CT images. The ventral displacement of the right adrenal gland by abdominal ultrasound may predict the existence of type 1 segmental CVC aplasia.

In conclusion, if a right adrenal gland tumor is found to be displaced in dogs with concomitant segmental CVC aplasia, CVC venectomy is required, even if there is no intravascular invasion.

POTENTIAL CONFLICTS OF INTEREST. The authors have nothing to disclose.

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