Chapter 2
Clinical Presentation of Desmoid Tumors

Anastasia Constantinidou, Michelle Scurr, Ian Judson
and Charisse Litchman

Contents

2.1 Introduction.......................................................................................................................... 6
2.2 Incidence ............................................................................................................................ 6
2.3 FAP...................................................................................................................................... 7
2.4 Etiology............................................................................................................................... 8
2.5 Clinical Presentation ........................................................................................................ 8
2.6 Clinical Considerations ..................................................................................................... 9
  2.6.1 Risk Factors ............................................................................................................... 9
  2.6.2 Unique Tumor Locations ........................................................................................... 10
  2.6.3 FAP vs. Non-FAP ....................................................................................................... 11
  2.6.4 Multicentricity .......................................................................................................... 11
2.7 Clinical Course .................................................................................................................. 11
2.8 Conclusions ....................................................................................................................... 12
References.................................................................................................................................... 13

Abstract  Desmoid tumors (DT) constitute a rare fibroblastic proliferative disease. They present sporadically or as a manifestation of a hereditary syndrome such as Familial Adenomatous Polyposis (FAP). Despite the absence of metastatic potential, DT may cause debilitating symptoms and in some cases life-threatening organ damage because of their locally invasive nature. DT may range from small slow-growing masses to rapidly enlarging aggressive tumors. The clinical course of the disease is unpredictable but available data suggest an initial phase of growth may be followed by a long period of growth arrest with tumor stabilization or even regression. FAP-related DT are preferentially located in the abdomen whereas sporadic DT tend to involve mostly the extremities, although the abdomen and the thorax may also be affected. Antecedent trauma, pregnancy and estrogens play a role in the genesis of some desmoid tumors. Surgery is the favored current approach in the treatment of most desmoid tumors. Definitive protocols are not available as

C. Litchman (✉)
Department of Neurology, The Stamford Hospital, Stamford, CT 06904, USA
e-mail: cdlitchman@gmail.com

C. Litchman (ed.), Desmoid Tumors,
most studies have been retrospective, small and comprised of mixed populations of FAP and non-FAP as well as of mixed populations of extra-abdominal and intra-abdominal patients.

**Keywords**  FAP • Musculoaponeurotic • Sporadic • Primary tumor • β-catenin • Abdominal • Extra-abdominal • Intra-abdominal • Pregnancy • Head and neck • Trauma

### 2.1 Introduction

Desmoid tumors (DT) also known as aggressive fibromatosis (AF) constitute a rare fibroblastic proliferative disease. As suggested by their name (desmoid from the Greek word “δεσμος” meaning band-like) DT may occur in any musculoaponeurotic or fascial tissue [1]. Usually the masses are firm and fixed to surrounding tissue. It is uncommon to note lymphadenopathy, overlying skin changes, erythema, or dilated veins.

Desmoid tumors can occur anywhere in the body and are generally divided by anatomic designation as extra-abdominal, abdominal, or intra-abdominal (see Fig. 2.1). The behaviors of the tumors, including growth rates, age predilection and recurrence rates often vary with the location of the tumor [2, 3]. The most common locations are the extremities (around the limb girdles or the proximal extremities), the abdominal wall (most commonly in women during or after pregnancy), and intra-abdominal or mesenteric. Depending on their location, they tend to infiltrate adjacent organs, extend along fascial planes, compress blood vessels and nerves, erode bones or obstruct organs such as the bowel.

Though they have a benign histologic appearance, lacking the nuclear and cytoplasmic features of a malignancy and a metastatic potential, DT may cause debilitating symptoms such as pain, deformity and in some cases life-threatening organ damage because of their locally invasive nature. DT may range from small slow-growing masses to rapidly enlarging aggressive tumors. The clinical course of the disease is unpredictable but increasing information suggests that an initial phase of growth may be followed by a long period of growth arrest with tumor stabilization or even regression [4–6].

### 2.2 Incidence

Though the actual incidence is likely significantly higher due to misdiagnosis, multiple and confusing pathologic nomenclature and underreporting, the current estimate is an incidence of 2–4 per million per year. Desmoid tumors are undisputedly very rare, with only 900 new cases diagnosed each year in the US. These tumors constitute 0.03% of all biopsy-analyzed neoplasms and <3% of all biopsy-analyzed soft-tissue tumors [7]. These tumors have been documented in patients between 3 and 67 years [8], with a peak incidence of 25–35. The female to male ratio ranges from 1.4 to 1.8 [9–12]. Reitamo et al. noted that in females under the age of 15 an
extra-abdominal location was more common while in females aged 18–36 an abdominal location was more common. DT occur in the abdominal wall with a female to male ratio of 7:1 [13]. There was no association with race [14]. In one study, 16% of primary tumors were <5 cm, 28% were between 5 and 10 cm and 50% were greater than 10 cm [15]. While the majority of desmoid tumors are sporadic, approximately 5% are associated with Familial Adenomatous Polyposis (FAP).

### 2.3 FAP

Desmoid tumors may present sporadically or as a manifestation of a hereditary syndrome called Familial Adenomatous Polyposis (FAP). FAP is a familial cancer predisposition syndrome characterized by the development of hundreds to thousands of premalignant adenomatous polyps in the colon and rectum by the age of 40 years [16]. Unless treated at an early age, almost all patients with FAP will develop colorectal cancer [17]. In fact, FAP is responsible for 1% of all cases of colorectal cancer [18]. The treatment of choice is prophylactic surgery comprising colectomy with ileorectal anastomosis or restorative proctocolectomy [19].

A significant percentage (3.5–32%) of FAP patients will develop DT during their lifetime [20–22]. The risk of patients with FAP-developing DT is 800–1,000-fold
higher compared to the general population [23]. The peak incidence of DT in FAP is between the second and the third decade [24]. In the majority of cases DT occur following prophylactic surgery for FAP [25, 26] with surgical trauma identified as a trigger for the development of DT in FAP. However, in some cases, DT may be the first manifestation of FAP with about 4% of cases of DT appearing as an incidental finding at the time of primary surgery [27]. Family history is a predisposing factor for DT formation in FAP patients [28, 29], with an observed increased risk of 2.5 times in first-degree relatives [29].

2.4 Etiology

Desmoid tumors are the result of deregulation of connective tissue growth. Increased nuclear expression of β-catenin, a protein responsible for regulation of gene expression, proliferation and survival, is the characteristic feature in both sporadic and FAP-associated DT. Familial Adenomatous Polyposis is a hereditary (autosomal dominant) disease characterized by a germ-line mutation in the adenomatous polyposis coli gene (APC). In FAP-driven DT, inactivation of the APC gene leads to accumulation of β-catenin whereas in the sporadic setting, in approximately 85% of cases, mutations in the β-catenin gene CTNNBi lead to increased activity of β-catenin [30].

Desmoid tumors are viewed as a nonneoplastic process by some authors and as a well-differentiated low-grade sarcoma by others [31]. The characterization of desmoid tumors as a neoplastic process rather than as an inflammatory fibrous reaction has been bolstered by the molecular studies of X-chromosome inactivation that confirmed that DT are the result of a clonal process [31, 32]. Nonrandom X-chromosome inactivation, trisomy 8 and/or 20 was demonstrated in greater than 30% of sporadic DT [33]. DT behave aggressively as locally infiltrating mesenchymal monoclonal proliferations that lack metastatic potential [34].

2.5 Clinical Presentation

In sporadic desmoids, between 37 and 50% of DT arise in the abdominal region [35–37]. The most common extra-abdominal sites are the shoulder girdle, chest wall and inguinal regions [38] (see Fig. 2.2).

Patients with intra-abdominal desmoids may have asymptomatic masses which silently enlarge and infiltrate into adjacent structures [2] or may have symptoms of weight loss, cachexia, malaise, compression of ureters, renal failure, small bowel compression, perforation and peritonitis [35, 41, 42]. In sporadic DT, infiltration of intestinal or visceral structures is less common but muscle, nerve and vessel involvement may result in debilitating symptoms such as pain, restricted mobility or deformity. A characteristic example of such presentation is the infiltration of the brachial plexus by a shoulder girdle tumor which may result in pain in
the shoulder and arm and weakness of the upper limb. The management of such cases is challenging as surgical excision is often not a feasible option. Due to their aggressive infiltrating nature DT may cause impairment or loss of function of vital organs. DT of the upper chest wall may engulf organs in the mediastinum including the trachea or the esophagus. As a result patients may suffer from dyspnoea/asphyxiation and dysphagia, respectively. Weiss et al. reported a patient with quadriceps paralysis and neurogenic bladder from focal invasion of the lumbosacral plexus [43].

2.6 Clinical Considerations

2.6.1 Risk Factors

2.6.1.1 Trauma

Trauma has been theorized to increase the risk of DT occurrence. Antecedent trauma, often surgical, has been reported at the site of the DT in approximately 25% of cases [10, 29, 44]. Moreover, 68–86% of abdominal wall and intra-abdominal wall DT are noted after abdominal surgery, the majority within the first 5 postoperative years [21]. FAP patients appear to be at even greater risk for DT development following surgical trauma with a reported 84% of cases of FAP-associated desmoids occurring within 5 years of abdominal surgery [45]. There have been reports of DT
in laparoscopic port sites [46], following a total hip replacement [47], around silicone implants [48], at the site of an internal jugular catheter [49] and at the site of a previous rib fracture [50].

2.6.1.2 Estrogen and Pregnancy

There are several lines of evidence to support a role for estrogen in modulating the behavior of DT. Several studies have shown that DT in females of childbearing age have a greater growth rate than that of those in males or in pre- or postmenopausal women [3, 51]. Further, an increased frequency rate was demonstrated during pregnancy [9, 51] and in females taking oral contraceptives [28, 52]. Additionally, there have been reports of tumor regression during menarche and menopause [51, 53, 54] and with Tamoxifen treatment [55].

In the lab, fibrous tumors have been induced in animal models following the administration of exogenous estrogen [53] and estrogen was shown to exert a mitogenic influence on many cell types, including fibroblasts [56]. Additionally, in a study of human DT, estrogen receptors (ER) were observed in 33% of all DT examined, with an equal incidence in males and females and with antiestrogen binding sites found in 79% of samples, including some which were ER negative [57].

In pregnancy-associated DT, the mass is most frequently located within one of the two rectus muscles of the abdominal wall without involving the midline [58, 59]. Pregnancy-associated DT may develop during any trimester or postpartum.

While the history of antecedent trauma is 28% of sporadic DT [60], such a history is ostensibly missing in pregnant DT patients. It has been theorized that the combination of an altered hormonal milieu and the trauma of stretching of the abdominal aponeurosis during the advancement of pregnancy are contributing factors [61]. There has been one report of a DT that developed at the site of a prior cesarean section scar during a subsequent pregnancy [16].

A study of FAP patients revealed no association between the female gender or pregnancy and the risk of the development of DT [62]. After examining the divergent natural histories and behaviors of pregnancy-associated DT and FAP-associated DT, one group of investigators concluded that these two types of DT are separate entities [61].

2.6.2 Unique Tumor Locations

2.6.2.1 Head and Neck DT

Head and neck DT are a more aggressive disorders that affect a younger population. Twelve percent of extra-abdominal DT arise in the head and neck [63]. The mean age is 16.87 years, with 57.32% of cases under 11 years. Children with DT of the head and neck are younger at the time of diagnosis than children with DT at other
sites [64–66]. There is a 30% local recurrence (LR) with a male to female ratio of 1:1 [67]. One explanation for the often difficult clinical course is the restricted anatomy containing crucial neural and vascular structures [67].

2.6.2.2 Breast

Desmoid tumors are rarely seen in the breast and can simulate breast carcinoma [68].

2.6.3 FAP vs. Non-FAP

Anatomic locations differ between FAP and sporadic DT, with more intra-abdominal or abdominal than extra-abdominal wall tumors. In a Mayo clinic review from 1976 to 1999, 67% of FAP-associated DT were abdominal as compared to 11% sporadic. Limb DT accounted for 1.4% in FAP patients and 34.7% in non-FAP patients [69]. While one large study reported a female to male ratio of 3.0 in FAP patients with DT [28], some studies failed to show the female predominance in FAP-associated DT that has been shown in sporadic DT [29, 44]. Additionally, desmoid development occurred an average of 6 years earlier in FAP patients [22]. Eighty to 90% of FAP individuals will carry an alteration in the APC gene on chromosome number 5. The majority will have a family history of colorectal cancer and polyposis. But, up to 33% of FAP patients with DT will have a de novo mutation within the APC gene and therefore no family history of DT [69].

2.6.4 Multicentricity

There have been 10–20 reports of multicentric extra-abdominal DT, mostly in FAP patients [70–73]. These usually recur in the same limb in proximity to the site of the primary tumor. They do not grow simultaneously, with the second growth generally occurring years later [74].

2.7 Clinical Course

DT remains an enigmatic disease with a variable course that can range from an incidental small tumor that can remain small and stable or become large and grow rapidly, causing death in a matter of months or years. The morbidity and mortality is largely determined by the location of the tumor and therefore the adjacent structures the tumor may infiltrate or compress. According to Church, 10% of DT will resolve
spontaneously, 30% will undergo cycles of progression and resolution, 50% will remain stable after diagnosis and 10% will progress rapidly [75].

Some of the local recurrence (LR) rates are determined by tumor location. For example, extremity tumors are considered locally aggressive and have LR ranging from 24 to 77% [76–80]. LR rates for intra-abdominal tumors are higher than for extra-abdominal tumors, reported to be 57–86% [28, 81, 82]. One review found LR to be 24% for abdominal wall, 43% for extra-abdominal and 77% for intra-abdominal tumors [2]. In a study of 78 FAP patients that studied progression-free survival rates after surgery versus conservative care, it was determined that extra-abdominal and abdominal wall DT had better outcomes and more benefit overall from surgical intervention than intra-abdominal tumors [22].

Gender has been shown not to be a prognostic factor for LR [4, 83]. There is disagreement about whether age may play a role in recurrence. Some studies have shown that younger age was associated with increased local treatment failure [39, 84] while others did not [75, 85]. One study found the recurrence rate in children to be 88%, twice that of adults (38%) [10]. Also controversial is the role of age in LR risk. Some studies show increased risk of LR in female patients older than 30 [88] while others show increased risk in patients under 30 [9]. One larger study of 103 patients over 26 years found no correlation with recurrence to age, gender, or site [83]. There is some suggestion that size of the primary tumor is an important predictor for recurrence [40] but that a single recurrence did not significantly increase the likelihood of a subsequent recurrence [10].

There is ongoing controversy over the significance of margin status in predicting LR. In one series, response rates of 72% and 41% were reported for tumor-free and tumor-positive margins, respectively [86]. Other studies show no correlation with margin status. The MSKCC (Memorial Sloan-Kettering Cancer Center) and Instituto Nazionale Tumori experiences showed no significant difference (22% negative vs. 24% positive [76] and 21% positive vs. 18% negative) [87].

The limitations in the studies stem from the small subject numbers and the mix of intra- and extra-abdominal tumors as well as primary and recurrent lesions, leading to conflicting results about the biology of these elusive tumors [9, 70, 76–81, 88–90]. The difficulties of interpretation of the data are compounded by the unpredictable natural course of this tumor that can apparently regress even without treatment [75].

2.8 Conclusions

Desmoid tumors are an enigmatic, elusive disease that continue to defy definition. Due to their rarity and the practical limitations in their study, these tumors often evade accurate characterization. As they can arise in many locations throughout the body, thereby presenting unique challenges to physicians in many different fields, the most appropriate and fruitful approach to caring for any individual desmoid tumor patient is a multidisciplinary one.
References

Clinical Presentation of Desmoid Tumors


74. Barber HM, Galasko CSB, Woods CG (1973) Multicentric extra-abdominal desmoid tu-
Colon Rectal Surg 6:29–32
Am J Surg Pathol 7:477–482
153–157
79. Scott RJ, Taeschner W, Heinimann K et al (1997) Association of extracolonic manifesta-
tions of familial adenomatous polyposis with acetylation phenotype in a large FAP kindred. Eur J
Hum Genet 5:43–49
80. Thomas JA, Kothare SN (1972) Desmoid tumors of the abdominal wall. Indian J Cancer
9:66–69
patients with familial adenomatous polyposis. Cancer 74:1270–1274
adenomatous polyposis. Surg Gynecol Obstet 177:263–268
cal Orthop and Related Research 375:207–213
A retrospective study of 72 patients followed for 1–27 years. Acta Orthop Scan 73:213–219
tients with aggressive fibromatosis or desmoid tumors. A comparative review of 22 articles.
Cancer 88:1517–1523
dominal aggressive fibromatosis: a series of patients surgically treated at a single institution.
J Clin Oncol 21:190–197
88. Pritchard DJ, Nascimento AG, Petersen IA (1996) Local control of extra-abdominal desmoid
89. Miralbell R, Suit HD, Mankin HJ, Zucker berg LR, Stracher MA, Rosenberg AE (1990) Fi-
bromatoses: from postsurgical surveillance to combined surgery and radiation therapy. Int J
Radiat Oncol Biol Phys 18:535–540
Surg 182:369–377
Desmoid Tumors
Litchman, C. (Ed.)
2012, XI, 221 p., Hardcover
ISBN: 978-94-007-1684-1