Bilateral testicular germ cell tumors
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ABSTRACT

Testicular cancer represents 1% to 1.5% of neoplasias in males and 5% of urologic tumors in general. The incidence of bilateral testicular tumors is 1-5%. Approximately one third of the cases are diagnosed as synchronous, while the other two thirds are diagnosed as metachronous tumors. Additionally, 5% of all patients diagnosed with testicular cancer may have contralateral intratubular germ cell neoplasia and may develop a contralateral germ cell tumor. However, few data are available regarding bilateral testicular germ cell tumors (BTGCTs). In this review, we aim to provide an overview of the incidence, pathological features and clinical outcomes of BTGCTs.

Key words: Bilateral testicular cancer; bilateral testicular germ cell tumor; testicular germ cell tumor.

Though testicular cancers are rarely seen malignancies among male neoplasms, they are the most frequently encountered malignancies in men aged 20-40 years. Nonseminomatous tumors, and pure seminomas are most frequently seen in the third, and fourth decade of life, respectively. Besides, in developed countries, it has a rapidly increasing incidence within the last 30 years.[1] In the Western world, every year 3-6 new cases are seen among 100,000 men.[2] In Scandinavian countries testicular tumors are more frequently seen than Asian countries, and differ according to ethnicity, and socioeconomic level of the patients.

Recognized risk factors in the development of testicular tumors include past or current history of cryptorchidism or undescended testis, Klinefelter syndrome, testicular cancer in the first degree relatives, presence of tumor or intraepithelial neoplasia in the contralateral testis, and infertility.[3-5] In a relevant study, cryptorchidism was detected in 9.5% of patients with bilateral, and in 2.2% of unilateral germ cell tumors. [6] Besides, atrophic testis was associated with the development of testicular tumor.[8]

Pathological features, and incidence rates of BGCTs

Germ cell tumors constitute 90-95% of testicular tumors. Seminomas, and nonseminomatous tumors constitute 40, and 60% of these tumors. Bilateral testicular tumors generally have the same histological structure. Seminomas are the most frequently (48%) seen histological type, followed by nonseminomatous (15%), and nongerminal (22%) tumors.[10] In men over 50 years of age, most often testicular lymphomas are seen. They are bilateral in 35% of the cases with frequently bilateral involvement.[10]

The incidence of bilateral testicular tumor ranges between 1, and 5 percent.[11-14] One of the first investigations on bilateral testicular germ cell tumors (BGCT) was performed in the year 1941 by Gilbert, and Hamilton on 1466 patients with primary testicular tumors. In a study performed in M.D. Anderson Cancer Center on bilateral testis tumors, the investigators couldn’t reveal any history of cryptorchidism in any of their study population.[6] Patel et al.[7] published outcomes of a study on synchronous, and metachronous bilateral testicular tumors performed by Mayo Clinic, and detected past history of cryptorchidism in 10.5% of 19 patients (one patient with bilateral, and one with unilateral cryptorchidism). Besides, atrophic testis was associated with the development of testicular tumor.[8]
Bilateral testicular tumors are either synchronous (simultaneous) or metachronous (developed at different time points). Synchronous tumors become apparent at the time of diagnosis or within the first two months after establishment of the diagnosis. The incidence of synchronous tumors is 1-2.8%, while of metachronous tumors it is 2.4-5.2 percent. [9] Hentrich et al. [20] noticed a second tumor, and detected Stage I metachronous tumors in 30 out of 33 patients with metachronous tumors. In our country, Akdoğan et al. [19] investigated 987 patients, and detected bilateral testicular tumors in 3% of these patients.

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The terms testicular intraepithelial neoplasia (TIN), carcinoma in situ (CIS), intratesticular germ cell neoplasia (ITGCN) define the same histology. TIN is the precursor lesion of spermatocytic seminoma in the elderly, and yolk sac tumors in infants, and testicular germ cell tumors excluding mature teratoma. TIN is confined within seminifer tubuli, and therefore it is noninvasive. It was originally defined by Skakkebaek in 2 patients who were biopsied with the indication of infertility. [23]

The incidence of TIN in general population ranges between 1 and 5%, and various factors are effecting its incidence. [24-25] The reported incidence rates of TIN are as follows: cryptorchidism, 2-4%; infertility, 2.2%, and contralateral testicular tumor, 5%. [26-28]

Different biopsy criteria for the investigation of the presence of TIN which is the precursor lesion of testicular germ cell tumors have been suggested in the literature. In some European centers contralateral testicular biopsies were obtained because of detection of concomitant TIN in 4-6% of the cases. Von der Maase et al. [29] indicated risk of invasive growth of TIN as 40, and 50% within the first 3, and 5 years after establishment of diagnosis, respectively. However, in the USA, testicular biopsies are not routinely performed. Herr et al. [30] did not recommend routine contralateral testis biopsy in that it has a lower rates of positivity, and also induces emotional stress in patients. A consensus has been built in the literature concerning its use in high-risk patients, and those with a history of infertility, cryptorchidism, and atrophic testis. [31] European Germ Cell Cancer Consensus Group (EGCCCG) defined patients with smaller testicular volume (<12 mL), history of cryptorchidism or those younger than 30 years of age as high-risk patients for contralateral testicular biopsy. [32] In conclusion, despite lack of consensus concerning testicular biopsy, contralateral testicular biopsy should be included among alternatives which must be discussed with the patient before the final decision.

Clinical features, and treatment
Because of scarcity of bilateral testicular germ cell tumors, most of the publications are in the form of case reports or small-scale studies. This fact restricts the scope of debates on treatment strategies. [23] Bilateral testicular tumors are generally diagnosed at an early stage, and they have a good prognosis. Hentrich et al. [20] performed a study on 1180 patients with testicular tumors, and reported an average disease-free survival of 37 months in 14 patients with bilateral synchronous tumors. In metachronous tumors, after an average of 71 months, the investigators noticed a second tumor, and detected Stage I metachronous tumors in 30 out of 33 patients with metachronous tumors. Similarly Holzbeierlein et al. [21] shared their nearly 50 years of their experiences, and investigated 3984 patients with testicular tumors, and stated that generally low-stage metachronous
tumors became apparent an average of 50 months after their first diagnosis. In a review article by Zequi et al.,[22] 5-year survival rates were detected to be 88, and 95% for synchronous, and metachronous tumors, respectively. In the same review article, the authors indicated that nearly 50% of synchronous, and 74% of metachronous tumors were Stage 1 tumors. In our country, Akdoğan et al.[19] reported that metachronous tumors had developed after an average of 75 months, most of them being Stage 1 tumors. They also stated that seminomas carry 2-fold higher risk as for development of bilateral testicular tumors. Numerous studies suggested the effective role of platinum-based chemotherapeutics in the decreased incidence of contralateral testicular tumors. Van Basten et al.[33] detected 3-fold lower rates of contralateral tumors in patients receiving chemotherapeutic agents. In a SEER (Surveillance, Epidemiology, and End Results Program) study conducted by Fossa et al.[34], the authors attributed lower incidence rates of contralateral cancer to the use of platinum-based chemotherapeutic agents.

Current guidelines contain scarce information about bilateral disease.[35] Therapeutic approach of especially infertile patients with bilateral germ cell testicular tumors or solitary testicular cancer is organ preserving surgery. Management of secondary testicular tumors should be achieved in consideration of histology, and stage of the primary tumor which is not different from treatment principles of primary tumor. If adjuvant radiotherapy or retroperitoneal lymph node was applied for a primary tumor, then monitoring or chemotherapy can be implemented based on the stage of the secondary tumor.[20] Recently, organ preserving surgery is in the foreground, and partial orchidectomy which has been revived nowadays as an alternative to partial orchidectomy can be performed in experienced centers. Fundamental foundation of partial orchidectomy rests on the maintenance of physiological androgen production, and fertility. It was firstly performed by Richie in USA in the year 1984.[36] Weissbach[37] described the principles of partial orchidectomy. Partial orchidectomy can be applied in synchronous or metachronous organ-confined testicular tumors smaller than 2 cm localized away from testicular vasculature which involve less than 30% of the testicular volume with negative surgical margins without histopathologically detected TIN in intraoperative biopsy specimens.[38] Because of higher rates of TIN concomitancy (80-90%), the patients should receive postoperative radiotherapy at a dose of 18-20 Gy.[39] Potential development of infertility, and androgen insufficiency secondary to postoperative radiotherapy should be kept in mind, and before initiation of the radiotherapy the patients should be informed about this issue.

Patients with testicular tumors should be monitored for lifelong. In addition to examinations, and analytical tests required from the clinician, patient’s self-examination conveys importance. The patients should also receive appropriate education on this subject. Various studies have demonstrated that self-examination attempts were effective in the detection of smaller secondary tumors at an early stage.

Hormone replacement therapy should be provided to treat potential androgen insufficiency which might develop following bilateral or partial orchidectomy. Among various testosterone formulations, controlled-release transdermal patches which ensure more stable serum testosterone levels should be preferred.[39] Serum testosterone levels of these patients should be regularly monitored.

An important consideration in bilateral testicular tumors which are seen in the second, and fourth decade of life is that patients who wants to be a father should undergo preoperative sperm freezing procedure, and they should be informed about assisted reproductive technologies. Besides, the patients should be provided with detailed information about cosmetically important testicular prostheses.

In conclusion, despite its increasing incidence in recent years, still bilateral testicular tumors have lower incidence rates. Bilateral testicular tumors have generally the same histological structure, and most frequently testicular seminomas are seen. Since TIN is observed in 5% of the cases with contralateral testicular tumor, routine contralateral testicular biopsy is not recommended for the detailed evaluation of TIN in these patients. Partial orchiectomy can be an alternative for eligible patients in experienced centers. Self-examination of the patients is very important. Since these tumors are seen in youngsters, these patients should receive necessary treatments for infertility, and androgen insufficiency. With appropriate treatment modalities, excellent outcomes, and longer survival rates can be achieved in bilateral testicular tumors.

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**References**


17. Johnson DW, Morenau JE. Bilateral sequential germ cell tumors of the testis. Urology 1974;4:567-70. [CrossRef]


33. van Basten JP, Hoekstra HJ, van Driel MF, Sleijfer DT, Droste JH, Memorial Sloan Kettering Cancer Center experience 1950 to 2009;27:421-6. [CrossRef]


37. Richie JP. Simultaneous bilateral testis tumors with unorthodox management. World J Urol 1984;2:74. [CrossRef]

