Testicular cancers are rare, but they are also the most prevalent tumors in young men and their incidence continues to rise, with a projected 20.4% increase in new cases and 39.8% increase in mortality by 2040. Testicular cancers, even with germ cell metastases, if treated aggressively are often eradicated with 5-year survival rates reaching 90%. Despite effective chemotherapy, 15% to 20% of patients with disseminated disease experience recurrence and most relapses occur within the first 5 years. Currently, there are no randomized trials informing decisions related to metastectomy for germ cell tumors, leading to interinstitutional management variability. General consensus supports germ cell metastatectomy, especially with pulmonary nodules because it can provide high cure rates. However, many resected nodules are later determined to be benign, causing some to question the necessity of every operation.

Donahoe and colleagues performed a single-institution retrospective review of 89 patients over the course of 13 years to determine whether concordance rates between metastatic sites could be used to guide surgical treatment algorithms. Ninety percent of patients (9 out of 10) with benign histology found during their initial pulmonary nodule resection also had benign disease identified on subsequent pulmonary resections. However, 1 subsequently had a malignant nodule. Similarly, 3 out of 11 patients with pulmonary and mediastinal lesions had malignant histology on subsequent resections after an initial resection of benign disease. Although the histological concordance rates for germ cell metastatic sites was 81%, the authors’ conclusion that 19% discordance rate mandates resection of subsequent lesions rather than trust that resection of a previous benign nodule reliably predicts that subsequent nodules are also benign.

Serum tumor markers (STMs) also had inadequate sensitivity and specificity to warrant making absolute decisions against surgical resection of pulmonary nodules. STM remains an important element in the management of mediastinal germ cell tumors and they help guide whether residual masses should be treated by resection, chemotherapy, or surveillance. In this series, STMs before resection correlated with malignant histology. However, 9 patients with normal STMs actually had viable malignancies. Furthermore, normal STMs at the time of resection related to lower survival rates owing to somatic transformation. This corroborates previous findings by Schekel and colleagues that showed that germ cell tumors with malignant somatic transformation portends a worse prognosis, with survival rates of 50% to 60%. STMs are vital; however, they should not be the sole information upon which clinicians rely to guide management of metastatic nonseminomatous germ cell cancer. Donahoe and colleagues demonstrated histological correlations between serial resections of mediastinal and pulmonary lesions in the settings of germ cell tumors and relations of STMs and malignancies that are insufficient to direct surgical interventions for new lesions. As a result, these data imply that new thoracic lesions should be managed aggressively with surgical resection that contributes to long-term cancer survival. This approach is analogous to assuming that there may be a snake hiding under every rock.
Commentary: Good news travels fast

Kenneth A. Kesler, MD

Before the results of a phase II trial demonstrating the effectiveness of cisplatin-based chemotherapy for disseminated testicular cancer were published in 1976, the word was out and survival rates began to exponentially increase for a malignancy, which at the time was rarely cured.1,2 The treatment algorithm of combination chemotherapy followed by aggressive surgery to remove residual disease for metastatic nonseminomatous germ cell tumors is now considered one of most successful models for multimodality cancer therapy, which our thoracic oncology community emulates for other locally advanced neoplasms with improved but currently lesser success.

Although cure rates for testicular cancer are very high, there still remain frontiers for improvement, such as the quandary of chemorefractory nonseminomatous cancer and malignant (somatic) transformation (fortunately representing only a minority of cases) and the variety of postchemotherapy pathology, which has implications for the indications to remove residual disease. For 2 examples, avoidance of pulmonary metastasectomy would seem prudent where there is a high likelihood of complete tumor necrosis. In contrast, pulmonary metastasectomy for numerous areas of chemorefractory disease may be futile.

Donahoe and coworkers3 have taken a deep dive into their institutional experience, examining the pathology and survival outcomes after surgery for nonseminomatous testicular cancer metastatic to the lung and mediastinum. The authors’ messages are important. First, the ability to predict postchemotherapy pathology is overall good, taking under consideration serum tumor marker levels, computed tomography appearance, and orchiectomy pathology, but the predictive accuracy as well as the concordance between lung, mediastinum, and retroperitoneal pathology remains imperfect. Second, while “benign” teratoma is the most common pathology, other benign and malignant elements can be present.