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EDITORIAL

Genetic Determinants of Breast Cancer Risk in Childhood Cancer Survivors

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In recent decades, survival from childhood cancer has increased dramatically (1) because of improving treatments. In Europe, the five-year relative survival rate presently exceeds 75%, with comparable rates reported in the United States (2,3). These successes come at the cost of a well-documented increased risk of treatment-related second cancers in later life. The US Childhood Cancer Survivor Study (CCSS) reports a 30-year cumulative incidence of second cancers that exceeds 20% in this group of patients (4). Hodgkin’s lymphoma and Ewing sarcoma survivors have the greatest risk of developing a second cancer, of which breast, brain, bone, thyroid, soft tissue, melanoma, and acute myeloid leukemia are most frequently observed (5).

Despite occurring almost exclusively in women, breast cancer is one of the most common subsequent neoplasms in survivors of childhood or adolescent cancer. It is particularly prevalent in patients with a primary diagnosis of Hodgkin’s lymphoma who received supradiaphragmatic radiotherapy as part of their treatment. In a recent UK study of more than 5000 such women, Swerdlow et al. reported a sixfold increased relative risk of subsequent breast cancer when radiotherapy was the sole treatment (6). More dramatic increases were observed when breast cancer risks were estimated by age at first radiotherapy treatment; the relative risk of breast cancer increased more than 20-fold when radiotherapy was first administered at age 10 to 14 years, with a peak of almost 50-fold at age 14 years that presumably is coincident with breast development (6). Critically, elevated breast cancer risks may persist for up to 30 years post-radiotherapy treatment before diminishing (6), and consequently intensive long-term surveillance is warranted for this group of patients.

At present, apart from variables pertaining to treatment, little is known about the effects of either genetic or environmental exposures in modifying the risk of radiation-induced second malignancies in childhood cancer survivors. Predicated on the hypothesis that common or rare germline variants influence the risk of radiation-induced breast cancer in this group of individuals, Morton and colleagues (7) performed a genome-wide association study (GWAS), which they report in this issue of the Journal. They analyzed 178 case subjects and 2200 control subjects from the CCSS and 29 case subjects and 574 control subjects from the St. Jude Lifetime Cohort (SJLIFE), among whom a total of 131 case subjects (63%) and 493 control subjects (18%) received at least 10 Gy radiation exposure to the breast during treatment for their first malignancy. Genome-wide genotyping and imputation of unobserved genotypes was used to generate a data set of almost 17 million single nucleotide polymorphisms (SNPs) that were represented in both cohorts and that were of sufficient quality for statistical analysis.

SNP rs4342822, localizing to 1q41, was statistically significantly associated with an almost twofold raised risk of breast cancer in patients who received at least 10 Gy radiation to the breast in both CCSS and SJLIFE and surpassed the widely applied threshold for genome-wide statistical significance (P < 5 × 10⁻⁸) upon joint analysis. The same locus showed no evidence for an association with breast cancer risk in individuals who had received lower doses of radiotherapy. Despite the relatively small sample size, Morton and colleagues also explored the contributions of rare variants upon radiation-induced breast cancer risk and detected independent associations of two SNPs, rs74949440 and rs17020562, mapping respectively to 11q23 and 1q32.3. The associations attributable to these SNPs differed with respect to the magnitude of radiation exposure; the former was associated with risk in women receiving at least 10 Gy, while the latter was associated with risk only in women who received less than 10 Gy.

Though potentially exciting, a liberal measure of caution is warranted until these associations are replicated in independent cohorts; discoveries from small GWAS, particularly rare variant associations, often suffer from a high rate of attrition because hits that are initially promising subsequently fail to replicate. Indeed, the data from Morton et al. (7) cast an element of uncertainty upon joint analysis.

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identified and robustly validated hundreds of variants that influence the risk of most common types of the disease (10), these successes have depended on global collaborative efforts to pool vast numbers of samples for analysis (11–13). Clearly there are fewer analogous studies with sufficient epidemiological and treatment data that also have genetic material for germline analysis with which to prosecute susceptibility to breast cancer in childhood cancer survivors. Nonetheless, this field must now move to consolidate available resources in order to expedite validation of promising leads.

Much effort is currently being expended on the development of breast cancer predictive models for population-based risk stratification that incorporate germline predisposition SNPs. A challenge is that because the breast cancer relative risk effects of SNPs are small, they are of little use individually and instead must be combined to identify different levels of risk (14). The data reported by Morton et al. (7) indicate that larger effects might be conferred by a subset of loci involved in mediating the risk of radiation-induced cancer, suggesting that meaningful risk stratification might be attainable with a small number of independent loci. If confirmed, the associations presented by Morton et al. (7) represent an important step toward delivering on the promise of second cancer risk prediction in childhood cancer survivors.

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