A remarkably simple genome underlies highly malignant pediatric rhabdoid cancers

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Cancer is principally considered a genetic disease, and numerous mutations are thought essential to drive its growth. However, the existence of genomically stable cancers and the emergence of mutations in genes that encode chromatin remodelers raise the possibility that perturbation of chromatin structure and epigenetic regulation are capable of driving cancer formation. Here we sequenced the exomes of 35 rhabdoid tumors, highly aggressive cancers of early childhood characterized by biallelic loss of SMARCB1, a subunit of the SWI/SNF chromatin remodeling complex. We identified an extremely low rate of mutation, with loss of SMARCB1 being essentially the sole recurrent event. Indeed, in 2 of the cancers there were no other identified mutations. Our results demonstrate that high mutation rates are dispensable for the genesis of cancers driven by mutation of a chromatin remodeling complex. Consequently, cancer can be a remarkably genetically simple disease.

Introduction
Cancer is generally considered to arise due to DNA mutations that alter the function of numerous genes (1). Indeed, most cancer genomes are aneuploid, contain amplifications and deletions, and typically have hundreds to thousands of DNA point mutations. However, not all cancers are necessarily so complex as some highly aggressive tumors are diploid. Large-scale sequencing projects have revealed that mutation rates display 10- to 100-fold differences among cancer types and even among different cancers of the same type (2, 3). Even in cancer types possessing the highest mutation rates, there are often individual cases that contain many fewer mutations. This raises a fundamental question about the nature of cancer: How simple can the genomes of highly aggressive cancers be?

Rhabdoid tumors (RTs) are a useful type of cancer with which to address these questions. These extremely aggressive pediatric cancers of the brain, kidney, and soft tissues are highly malignant, locally invasive, frequently metastatic, and particularly lethal (4), and yet, they are typically diploid and lack genomic aberrations detectable by SNP arrays (5). Early-onset cancers also offer the opportunity to examine cancer genomes prior to the acquisition of substantial numbers of age-related mutations, the vast majority of which are likely passenger mutations.

Near-uniform biallelic inactivating mutations in SMARCB1 (also known as NF53, INI1, and BAF47), a gene that encodes a core subunit of the SWI/SNF chromatin remodeling complex (6, 7), are a hallmark of RTs. Alterations in genes involved in chromatin remodeling, and particularly in genes encoding SWI/SNF subunits (8), are increasingly being identified in a wide variety of cancers, raising the possibility that epigenetic dysregulation may be a central mechanism of oncogenesis.

Results and Discussion
We obtained DNA from 32 diagnostic pretreatment SMARCB1 mutant RT samples, of which 20 were from brain, 3 were from kidney, and 9 were from other soft tissues (Supplemental Table 1; supplementary material available online with this article; doi:10.1172/JCI64400DS1). The median age of the patients was 12 months. Whole-exome sequencing and SNP array analysis was performed on all 32 sample pairs. Analysis of SNP arrays identified a single region with significant focal somatic copy number alterations (SCNAs): deletions at 22q11.23 that contained the SMARCB1 gene were identified in 25 out of the 32 samples (GISTIC2.0, ref. 9; q < 10–50), which comprised focal deletions in 16 cases, monosomy 22 in 15 cases, and both in 6 cases (Figure 1). One sample (08-262A) had a germ line focal deletion. Tumor purity ranged from 43% to 97%, so the lack of additional detected SCNAs was not likely due to stromal contamination (Supplemental Table 2).

We next performed exome sequencing of DNA to a mean coverage of 83-fold across 32.6 Mb of targeted coding regions for each sample (Supplemental Table 3). This level of coverage resulted in a “call-able” exome of 28.6 Mb. Detection of SCNAs by sequencing data was consistent with SNP array findings (Supplemental Figure 1). Analysis revealed a total of 172 somatic substitutions and insertions/deletions (indels) in the 32 tumors (Table 1 and Supplemental Table 12). Other than SMARCB1 loss, 2 tumors (08-114 and 09-223) had no detectable mutations, and 4 tumors (07-057, 07-221, 08-172, and 09-131) had only subclonal mutations (Figure 2A). The mean mutation rate was 0.19 mutations per Mb, with a minimum of 0 and a maximum of 0.45 mutations per Mb. This rate is, to our knowledge, the lowest of all cancers sequenced to date, particularly for such a high-grade and lethal type of cancer (Figure 2B). Consistent with our tumor selection process, all tumors had combinations of SMARCB1 mutations and/or deletions predicted to cause homozygous loss of function (Supplemental Table 4 and Supplemental Figures 3 and 6). Over-
all, 71.5% of the mutations were classified as clonal. All 7 of the point mutations in \textit{SMARCB1} were classified as clonal (Figure 2A and Supplemental Table 5).

We looked for recurrent mutations that may cooperate with \textit{SMARCB1} loss to drive RTs. The only other recurrently mutated gene was \textit{GABRB2}, a subunit of the GABA A receptor, which was found to be clonally mutated in 2 out of the 32 samples (10-330 and SJDOS006; Supplemental Figure 4). However, the Catalog of Somatic Mutations in Cancer (COSMIC v51) database contains only 2 other instances of this gene being mutated, neither of which matched the RT mutations (10, 11). Only 2 mutations found among the 32 sequenced RTs, aside from \textit{SMARCB1} and \textit{GABRB2}, were present in the COSMIC database: 1 in \textit{NF2} (10-213) and the other in \textit{TP53} (10-330). The \textit{NF2} nonsense mutation (Y144*) may have been contributory; although subclonal (26% of cells), the mutation was present on a background of hemizygous deletion, implying...
homozgyous loss in the mutant subclone. The relevance of the TP53 mutation (D49N) was unclear; although clonal, the variant did not occur in one of the canonical mutation domains and was predicted not to be detrimental.

In addition to the 32 primary samples, we analyzed 3 independent recurrent tumor/normal pairs after chemotherapy (Supplemental Table 6). Sample 09-044 was found to be aneuploid (Figure 3A). The range of purity and coverage was comparable to that in the primary samples (Supplemental Tables 7 and 8). While the largest number of mutations per sample found in the 32 primary tumors was 13, the recurrent cancers contained 38, 47, and 47 mutations, resulting in a rate of 1.53 mutations per Mb (Supplemental Table 9), 8 times higher than that in the primary tumors (Figure 3B; \( P < 0.005 \)). Other than SMARCB1, which was homozgyously lost in all 3 recurrent samples (Supplemental Table 10), none of the mutations matched any in the COSMIC database and no gene contained recurrent, clonal mutations (Supplemental Figure 5). Notably, the treated samples had significantly more subclonal mutations than the primary tumors (\( P = 7 \times 10^{-4} \); Figure 2A and Supplemental Table 11). Despite their absence from the COSMIC database, we could not exclude the possibility that these mutations could be conferring a growth advantage for subclones that could ultimately contribute to recurrent or refractory disease. Regardless, these mutations are unlikely to constitute effective therapeutic targets up front, given their absence from the predominant cancer population.

The mutational profile was also distinct in the recurrent samples, as they contained a significantly reduced proportion of C→T transitions (\( P < 10^{-5} \), 2-proportion \( z \) test) and increased proportions of A→T (\( P < 0.05 \)) and C→A (\( P < 0.005 \)) transversions (Figure 3C). Overall, the recurrent tumors had a greater percentage of transversions (\( P < 10^{-4} \); Figure 3D and Supplemental Figure 2).

In part based upon the large number of mutations commonly present in tumors, genetic alterations that affect numerous protein coding genes have typically been considered a fundamental requirement for cancer development. The finding that SMARCB1 is the sole gene recurrently mutated at high frequency in extremely aggressive and lethal RTs, and in some cases may be the only mutated gene, prompts essential questions: What accounts for the extreme paucity of mutations, and how can these data be reconciled with current models of cancer that estimate that 5 to 15 driving mutations are required for oncogenesis (12)?

We considered 4 possible explanations. First, as we have only sequenced exome DNA, we cannot exclude the existence of mutations in noncoding portions of the genome, such as in noncoding RNAs or regulatory elements or in mutations in low coverage areas. Further, we cannot exclude balanced translocations or inversions, although these are not characteristic of RTs by karyotype (13). Nonetheless, occult events could cooperate with SMARCB1 loss. Second, as mutations were largely identified based upon differences between tumor and normal DNA, contributions from germ line events are difficult to exclude. However, since genetically engineered models have demonstrated that inactivation of Smarcb1 drives extremely rapid formation of cancer in all mice and since this occurs on several genetic backgrounds (14, 15), it seems unlikely that germ line alterations are essential for cancer formation driven by SMARCB1 loss. Third, the developmental stage/epigenetic stage may serve a contributory role. During development, there is relative enrichment of minimally differentiated cell populations that have a high proliferative capacity. Consequently, it is possible that developmentally restricted or lineage-specific populations of cells characterized by a certain epigenetic state are particularly susceptible, such that mutation of a single chromatin remodeler can drive transformation. This is consistent with our mouse model in which Smarcb1 deletion in the T cell lineage leads to transformation of a highly specific cell type, CD8+CD44hiCD122hi memory T cells, a population that has a high intrinsic capacity for proliferation. Notably, this transformation arises due to aberrant responses to lineage-specific T cell receptor
It is interesting that a type of cancer that has extremely few gene mutations at the time of initial diagnosis is characterized by a much higher number of mutations in recurrent samples. The reason for this is unclear. While we do not have specific treatment data, these patients were treated with chemotheraphy and potentially radiation therapy. It is possible that genotoxic chemotherapy directly causes such damage. This possibility is supported by the presence in recurrent samples of a high percentage of transversions, a mutation type known to be associated with chemotheraphy and similarly seen in recurrent AML (22). This raises the possibility that chemotherapy can cause the conversion of a remarkably simple cancer genome into one with 8-fold more mutations, a possibility with substantial clinical implications, as such mutations could potentially contribute to resistance. Alternatively, the selective pressure of chemotherapy may result in the outgrowth of subclones with high rates of mutation, or, conceivably, the fundamental nature of the recurrent disease has changed such that these cancers have acquired genetic instability and a high mutation rate.

Finally, while there is increasing evidence that epigenetic regulators are mutated in a large variety of cancers, it has been unclear whether these alterations are selected for because they act to facilitate genomic instability, because they potentiate the effects of other mutations, or because they directly drive oncogenic transformation. Particularly in cancers in which mutations in chromatin regulators exist in a genetically complex background, it has been extremely difficult to determine the relative contribution of epigenetic alterations. Our findings from RTs demonstrate that mutations in the SWI/SNF chromatin remodeling complex can act as potent drivers of cancer. Understanding the contributions of mutations in these remodelers to oncogenesis has the potential to facilitate development of targeted therapies for the wide variety of SWI/SNF mutant cancers.

Methods

Samples. Tumor tissue and matched blood from 32 newly diagnosed patients with cancer and from 3 recurrent tumors were collected. Mutation and deletion analysis of the SMARCB1 gene was performed as previously described (6). Tumors were reviewed to confirm the diagnosis and to estimate tumor content (The Children’s Hospital of Philadelphia, Hospital Sant Joan de Déu). DNA was extracted using standard techniques.

SNP arrays. All of these samples were processed and hybridized to Affymetrix SNP 6.0 arrays for genotyping and copy number analysis (2). SNP array data were further analyzed using the ABSOLUTE tool (23) to infer the tumor purity and ploidy (11, 24, 25).

Whole-exome sequence data. Library construction followed the procedure of previous publications (11, 24, 25). Descriptions of sequencing and analysis methods are in the Supplemental Methods. Data were deposited in dbGaP (accession no. phs000508). The complete list of all detected mutations can be found in Supplemental Table 12.

Statistics. Comparison of mutation rates was performed using a 2-tailed Welch’s t test for samples with unequal variance. Two-proportion z test and Pearson’s χ² test with a Yates’s correction for continuity were used to analyze the different proportions of mutation type. Subclonal mutation frequency was analyzed using Fisher’s exact test. Data in figures are shown as mean ± SEM.
Study approval. Patients’ guardians provided informed consent prior to their participation. Local IRBs (The Children’s Hospital of Philadelphia, Hospital Sant Joan de Déu) approved collection and testing of each sample. Subsequently, the Broad Institute’s IRB approved consents.

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