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Rare allelic forms of PRDM9 associated with childhood leukemogenesis

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ABSTRACT

One of the most rapidly evolving genes in humans, *PRDM9*, is a key determinant of the distribution of meiotic recombination events (Baudat et al. 2010; Myers et al. 2010; Parvanov et al. 2010). Mutations in this meiotic-specific gene have previously been associated with male infertility in humans (Irie et al. 2009) and recent studies suggest that *PRDM9* may be involved in pathological genomic rearrangements (Berg et al. 2010; Borel et al. 2012). In studying genomes from families with children affected by B-cell precursor acute lymphoblastic leukemia (B-ALL), we characterized meiotic recombination patterns within a family with two siblings having hyperdiploid childhood B-ALL and observed unusual localization of maternal recombination events. The mother of the family carries a rare *PRDM9* allele, potentially explaining the unusual patterns found. From exomes sequenced in 44 additional parents of children affected with B-ALL, we discovered a substantial and significant excess of rare allelic forms of *PRDM9*. The rare *PRDM9* alleles are transmitted to the affected children in half the cases, nonetheless there remains a significant excess of rare alleles among patients relative to controls. We successfully replicated this latter observation in an independent cohort of 50 children with B-ALL, where we found an excess of rare *PRDM9* alleles in aneuploid and infant B-ALL patients. *PRDM9* variability in humans is thought to influence genomic instability, and these data support a potential role for *PRDM9* variation in risk of acquiring aneuploidies or genomic rearrangements associated with childhood leukemogenesis.

INTRODUCTION

Most effort in cancer genomics has focused on capturing somatic mutations from the screening of tumor and normal somatic tissue genomes, to identify factors mutated somatically during tumor progression. Genetic approaches aim to find genomic regions predisposing individuals to cancer, to capture inherited predisposing mutations segregating in the population by using genetic linkage or association studies. For late-onset cancers, such as breast and colorectal cancers (Turnbull et al. 2010; Peters et al. 2012), many predisposing allelic variants have been described, supporting a polygenic model of susceptibility (Easton and Eeles 2008) but only few genetic risk factors for pediatric cancer have been established (Healy et al. 2007; Sherborne et al. 2010). Dominant mutations causing cancer early in life are likely to be rapidly eliminated from the population, and as a result, it is unlikely that affected children will share inherited mutations. Parental germline events may play a role in pediatric cancer development with early evidence for epigenetically marking of imprinted genes during meiosis (Joyce and Schofield 1998), that may be involved directly in tumorigenesis for cancers of embryonal origin, such as Wilms' tumours, rhabdomyosarcoma, adrenocortical carcinoma and hepatoblastoma. Little is known about the contribution of meiotic events to the genetic instability driving the early onset of childhood cancer. In particular, novel genomic changes that occur during meiosis will not be detectable using standard genetic mapping approaches. However, interrogating normal and tumor genomes from families of patients provides an ideal framework to study *de novo* genomic events potentially linked to childhood malignancies.

Recent genomic studies using family data have shown that many early onset diseases arise from defects caused by *de novo* genetic aberrations, be they point mutations (Awadalla et al. 2010), copy number variants (Greenway et al. 2009), structural rearrangements (Kloosterman et al. 2011) or aneuploidies (Hassold et al. 2007). Recombination rates in children correlate with maternal age at birth (Hussin et al. 2011), which may have implications for understanding aneuploid conceptions. Intriguingly, children born with constitutional aneuploidies and rearrangements are at an increased risk for various malignancies (Ganmore et al. 2009). For example, children with Down syndrome have nearly a 20-fold increased risk for acute leukemia (Ross et al. 2005), suggesting that carcinogenesis and congenital anomalies may have a common basis for some pediatric cancers (Bjorge et al. 2008). Known recombination associated factors, such as DNA repair and histone modifications, are associated with genomic instabilities and cancers (Fernandez-Capetillo et al. 2004; Helleday 2010), and congenital genomic rearrangements and aneuploidies have been associated with errors in meiotic recombination (Hassold and Hunt 2001; Sasaki et al. 2010). Such gross genomic events are frequent in pediatric cancers.

Cancer is the leading cause of death by disease among children in western countries, and the overall incidence rate continues to rise steadily. The most common pediatric cancer, acute lymphoblastic leukemia (ALL), is a hematological malignancy resulting from chromosomal alterations and mutations affecting molecular pathways that disrupt lymphoid progenitor cell differentiation (Greaves 1999). Childhood ALL is likely explained by a combination of genetic predisposition and environmental exposure during early development, in fetal life and in infancy. Furthermore, ALL is a heterogenous disease presenting many molecular subtypes, with different populations having different incidence rates, such that the power of stratified analyses will be limited due to small number of cases in each subgroup. Finally, there is well-established evidence for prenatal initiation of the leukemogenesis process in children (Wiemels et al. 1999; Greaves 2006), and focusing

exclusively on child genetic material in ALL association studies may be insufficient for understanding disease etiology.

To characterize the importance of parental germline events in susceptibility to childhood ALL, we first set out to determine whether meiotic recombination patterns can lead to factors associated with the development of childhood ALL. From exome sequencing and genotyping data, we characterized meiotic recombination patterns in a unique family (referred herein as the ALL quartet) with two siblings having hyperdiploid B-cell precursor ALL (B-ALL). We observed unusual localization of maternal meiotic recombination events, with a small number of crossovers taking place in previously well-characterized population recombination hotspots. Such hotspots are short segments (1-2Kb) identified to be highly recombinogenic in the human genome (Myers et al. 2005). The mother of the family carries a rare *PRDM9* allele, potentially explaining the unusual placement of recombination events observed (Berg et al. 2011). *PRDM9* is a meiosis-specific histone H3 methyltransferase that controls the activation of recombination hotspots via its zinc finger (ZnF) DNA binding domain recognizing a short sequence motif, which then triggers hotspot activity through modification of the chromatin state (Grey et al. 2011). Analyses of next-generation sequencing and Sanger re-sequencing read data from a cohort of parents with B-ALL affected children of French Canadian descent, revealed a substantial excess of rare allelic forms of *PRDM9*. This association was successfully replicated in an independent cohort of children with B-ALL diagnosed in Tennessee, USA, where the effect was found particularly in aneuploid and infant B-ALL patients. Not only has *PRDM9* variability been associated with hotspot activation, but in humans it has been suggested to influence genomic instability (Berg et al. 2010; McVean and Myers 2010). The results presented here point to rare *PRDM9* allelic forms involvement in the development of preleukemic clones in B-ALL patients and we propose that *PRDM9* histone H3K4 methyltransferase activity in the parental germline could lead to the genomic instability associated with childhood ALL, a plausible mechanism consistent with the current understanding of molecular pathways of leukemogenesis and the disease.

RESULTS

The ALL family quartet

To study germline processes such as recombination, data from families with at least two siblings is required. Within the Quebec Childhood ALL cohort, we identified a family with two siblings having hyperdiploid B-ALL diagnosed at Sainte-Justine University Hospital. Families with two cases of childhood ALL in a sibship are rare and it is not clear whether siblings of children with ALL have an increased risk of developing ALL themselves (Draper et al. 1996; Winther et al. 2001). From studies published between 1951 and 2009 and registry-based childhood ALL data, an international collaboration only identified three sibships that were concordant for hyperdiploid ALL (Schmiegelow et al. 2011). However, the high concordance rate in ALL subtype within sibships is somewhat incompatible with a scenario where all cases in sibships occur randomly through independent events.

Sampling from the family included six biological samples from four family members: the mother and father, sampled once, and their two sons, patients 383 and 610, sampled at diagnosis and in remission (Figure 1). The brothers were both diagnosed with B-ALL with FAB-L1 morphology, at the age of 2 for patient 383 and 3 years later, at 14 years of age, for patient 610. At diagnosis, both siblings showed hyperdiploid leukemia clones (>50 chromosomes, Supplementary Results), a childhood B-ALL subtype that is very likely to be prenatally initiated (Gruhn et al. 2008). However, chromosomal instabilities found in preleukemic clones are generated prenatally in the normal population at approximately 100-times the rate of overt ALL (Mori et al. 2002), thus a second hit is required to trigger ALL during childhood, and results from other genetic and/or environmental factors. Because the patients were both diagnosed within a 3 year time period, it is likely that the second hit is due to environmental exposure (Greaves 2006).

De novo mutation and recombination events

Entire exomes were sequenced at high coverage using the SOLiD platform (Table S1A) to allow the full interrogation of the mutational and recombinational landscape occurring in coding regions (Material and Methods). Among variable positions discovered in the patients' exome, we looked for *de novo* mutations (Supplementary Methods). Given the human mutation rate, no more than one *de novo* point mutation is expected in a normal exome (Conrad et al. 2011). We only identified a single coding *de novo* point mutation in one of the patients (patient 383, Figure S1), which suggests that the parental germline mutation rate in this family is not higher than expected. Nevertheless, the putative *de novo* mutation identified is predicted to affect the structure and function of SMAD6. SMAD6 functions as an inhibitor of TGF-beta family signaling and was found to be a ligand-specific inhibitor of growth arrest and apoptosis in mouse B cells (Ishisaki et al. 1999). Furthermore, SMAD6 is required for HL-60 myeloid leukemia cell line differentiation (Glesne and Huberman 2006) and is a key determinant of hematopoietic stem cell development (Pimanda et al. 2007).

Combining the exome sequencing with genotyping data obtained from Illumina Omni2.5 arrays, we identified over 816,000 high-confidence variable genomic positions within the ALL quartet exomes that showed no aberration in allele inheritance (Table S1B). We performed fine-scale dissection of meiotic recombination events on autosomes and located a total of 102 and 47 crossovers in maternal

and paternal meioses, respectively (Figure 2A, Supplementary Methods). We also identified nine short tracts flanked by crossover events, possibly indicating gene conversion events (Supplementary Results, Table S2). The maternal and paternal mean number of recombination per meiosis for the ALL quartet were compared to the distribution of maternal and paternal means in a control cohort of French-Canadian families (FC family cohort, Material and Methods). The mother of the ALL quartet exhibits a high recombination rate with respect to the FC family cohort, with her mean number of crossovers per gamete found to be at the end of the spectrum (Figure S2). She carries two copies of the haplotype at the RNF212 locus associated with high recombination rates in females (haplotype [T, C] at SNP rs3796619 and rs1670533) (Kong et al. 2008). The father carries one copy of the rs3796619 T allele, which was estimated to decrease male genome-wide recombination rate by 2.62% per copy. This is consistent with the lower recombination rate seen in the father of the ALL quartet (Figure S2).

We next evaluated hotspot usage by computing the proportion of high-resolution recombination events (localized between informative markers less than 30 Kb apart) overlapping known population hotspots inferred from HapMap2 data (Myers et al. 2005). This measure does not directly evaluate hotspot usage, since a fraction of crossovers is expected to map in population hotspots by chance alone, but it is a good proxy to compare relative hotspot usage. Among recombination events identified in the ALL quartet, 85 maternal events and 34 paternal events were localized between informative markers less than 30 Kb apart. Fifty-four percent (46/85) of maternal events and 79% (27/34) of paternal events overlapped with HapMap2 recombination hotspots. These proportions are significantly different between the parents ($p = 0.0124$, Fisher's exact test). Since it was reported that, on average, 70% of events are expected to overlap population hotspots (Coop et al. 2008; Hussin et al. 2011), this result suggests that the mother has a lack of recombination in population hotspots. To validate this result, we further derived a null distribution of the proportion of recombination events expected to overlap with HapMap2 recombination hotspots in the FC family cohort (Material and Methods). The paternal crossovers show the expected enrichment in population hotspots inferred from HapMap2 data, while the maternal recombination landscape is unusual with a particularly low proportion of meiotic recombination events occurring in HapMap2 hotspots (Figure 2B).

Characterising PRDM9 in the ALL family quartet

Variability in recombination hotspot usage correlates with variation in *PRDM9*, a gene identified as a major hotspot determinant in mammals (Baudat et al. 2010; Berg et al. 2010; Myers et al. 2010; Parvanov et al. 2010) and the only locus known to be involved in hotspot specification in humans (Kong et al. 2010; Hinch et al. 2011). Allelic variation at the *PRDM9* locus consists of variable repeating units, encoded by a minisatellite formed by tandem-repeat C2H2 zinc finger (ZnF), and has a strong effect on recombination hotspots positioning and activity (Berg et al. 2010; Berg et al. 2011). The reduced proportion of recombination events overlapping population hotspots in the mother of the ALL quartet lead us to investigate genetic variation at the *PRDM9* gene. The sequencing data revealed that the father of the two affected brothers is homozygote for the most common 13-repeat allele (allele A) whereas the mother carries an allele A and a 14-repeat allele C, inherited by one of the two brothers (Supplementary Results). We further validated the presence of the C allele, which encodes major changes in the *PRDM9* ZnF array (Baudat et al. 2010), by Sanger re-sequencing (Material and Methods).

Although this allele is rare in populations of European ancestry (~1%), it is more frequent in individuals of African descent (~13%) (Berg et al. 2010). Because the parents have Moroccan Arab ancestry (Supplementary Results, Figure S3), we studied *PRDM9* diversity in 27 Moroccan individuals to establish whether the presence of the C allele in the mother reflects a different distribution of Moroccan *PRDM9* alleles relative to populations of European descent. Among 54 Moroccan *PRDM9* alleles sequenced, no C allele was found (Figure S4), suggesting that the frequency of the C allele in the Moroccan population is similar to the observed frequency in populations of European descent (Supplementary Results).

Motifs overrepresented in recombination hotspots in European and African American individuals have been inferred from population studies (Myers et al. 2008; Hinch et al. 2011): a 13-mer motif is enriched in linkage disequilibrium-based hotspots inferred from HapMap2 data, whereas a 17-mer motif is overrepresented in African-enriched hotspots. The A allele has been shown to bind to the 13-mer CCNCCNTNNCCNC motif, whereas the C allele has demonstrated inability to activate recombination hotspots presenting this motif (Berg et al. 2010). The C allele is predicted to specifically bind to a 17-mer motif CCNCNNTNNNCNTNNCC (Berg et al. 2011), very close to the motif found to occur at an increased frequency in African-enriched hotspots (Hinch et al. 2011). Compared to the distribution of these motifs seen at recombination events in the FC family cohort, we observed that the 17-mer motif is highly represented in the ALL mother's recombination events whereas the 13-mer motif is under-represented (Figure S5, Material and Methods). These observations confirm that the presence of the C allele in the mother is likely to have caused the genome-wide shift from HapMap2 hotspots observed in the maternal recombination landscape (Berg et al. 2011; Hinch et al. 2011).

Association between *PRDM9* and ALL in parents

Recent studies suggest that *PRDM9* is implicated in genomic rearrangements leading to congenital diseases (Myers et al. 2008; Berg et al. 2010; Borel et al. 2012). Because large-scale genomic rearrangements are common events in childhood leukemia, we sought to type *PRDM9* alleles in the parents of the B-ALL cohort. We assayed *PRDM9* ZnF alleles in a panel of 44 additional parents from 22 French-Canadian families with children affected by B-ALL (FCALL cohort) by analyzing reads aligning to the *PRDM9* ZnF array, in the parental trios where exome sequencing was previously performed (Material and Methods). The read data allows for the detection of all zinc finger repeats present in the common allele A, along with two rarer repeats, *k* and *l*, present in the C allele (Figure S6). Around one fourth of the parents (12/46, Table S3 and S4) have alleles with *k* finger repeats (*k*-finger alleles), usually rare in populations of European descent (Berg et al. 2010), suggesting that *k*-finger alleles are in excess in the FCALL cohort ($p = 0.0181$, Supplementary Results). To validate this result, Sanger re-sequencing of the *PRDM9* ZnF alleles was further performed in 13 pairs of parents from the FCALL cohort and in 76 parents from the ethnically matched FC family cohort (Figure S7, Supplementary Results). Among the 26 B-ALL parents re-sequenced, evidence for the presence of rare alleles in the read data had been found in 9 parents from 8 families, and all of them were confirmed. Through re-sequencing, we discovered rare alleles in two additional families, which were not originally detected in the exome sequencing reads. In contrast, only 5 parents from the FC family cohort carried *k*-finger alleles (Table S5). This confirms the excess of rare alleles in the FCALL cohort with respect to controls ($p = 3.22 \times 10^{-3}$, Figure 3),

with 76.9% of B-ALL families with at least one parent carrying a rare *PRDM9* allele. The rare alleles were preferentially carried by the mother ($p = 0.0235$, Fisher's exact test), although this maternal effect is not observed for *k*-finger alleles alone. Furthermore, the observation is not restricted to families with children having hyperdiploid B-ALL, since alternative *PRDM9* ZnF alleles are also found in parents of children presenting translocations and as yet uncharacterized genetic defects (Table S6). Finally, we found no significant evidence for transmission distortion, as *k*-finger parental alleles were transmitted to the affected child in 6 out of 11 cases with carrier parents (Table S3 and S4), resulting in 25% of the children of the FCALL cohort (6/24) carrying a *k*-finger allele.

Replication in a B-ALL patient cohort

The association was also detectable in the patients themselves ($p = 0.0123$). We replicated this latter association in an independent cohort of 50 children, sequenced whole genome, from St. Jude Children's Research Hospital. The children were affected by B-ALL from four subtypes: ETV6-rearranged, Philadelphia chromosome-positive, hypodiploid and infant B-ALL. We observed an excess of B-ALL patients with rare alleles in the St. Jude ALL cohort, with read data showing evidence for the *k* and/or *l* fingers in 10 children (Table S7). This excess is significant with respect to the French-Canadian controls ($p = 0.0143$, Table 1) and to 1000 Genomes Project controls from the CEU population ($p = 0.0353$, Supplementary Results). No *k*-finger alleles were detected in B-ALL patients from Philadelphia chromosome-positive and ETV6-rearranged subtypes. In hypodiploid and infant B-ALL subtypes, the excess of *k*-finger alleles is significant (Table 1). The children have no African ancestry, and 39 of them ethnically cluster with the controls of European ancestry, whereas the other children have different levels of Hispanic, Asian and Native American ancestry (Figure S8, Supplementary Results). Although the frequencies of *PRDM9* *k*-finger alleles in Chinese and Mexican individuals are likely similar to the ones observed in Europeans (Parvanov et al. 2010), to be conservative we also tested for the association between *k*-finger alleles and B-ALL when only non-admixed white patients were included. The association remains significant overall and in the hypodiploid subtype, although the effect becomes marginal in the infant B-ALL subtype (Table 1).

PRDM9 binding motifs and ALL translocations

Chromosome-number abnormalities and chromosomal rearrangements have been associated with altered recombination (Hassold and Hunt 2001; Sasaki et al. 2010). Given that *PRDM9* may be responsible for causing recombination-associated pathological genomic rearrangements (Myers et al. 2008; Berg et al. 2010; Borel et al. 2012), the high frequency of translocations and aneuploidies in leukemia raises the question of whether *PRDM9* is implicated in the generation of preleukemic cells early in development. In particular, the C allele accounts for an important fraction of the rare alleles detected in the ALL cohorts, and its binding motif as been predicted and validated (Baudat et al. 2010; Berg et al. 2011). To assess whether the C *PRDM9* allele was more likely than the common A allele to bind to genes known to be involved in ALL translocations (ALL gene list, Table S8), we performed a motif search to identify putative A and C binding sequences in the human reference genome. We found an enrichment of sequences potentially recognized by allele C relative to allele A in the ALL gene list in comparison to the rest of the reference genome (OR=1.53 [1.15;2.04],

Table 2) and to other coding regions (OR=1.61 [1.21;2.15], Table 2). The excess of C binding motif relative to A binding motif within the ALL gene list was found to be significant in unique DNA (OR = 2.39 (1.50-3.81 CI 95%), Table S9) but not in repetitive DNA, except for segmental duplications. The C motif is highly overrepresented relative to the A motif in segmental duplications in the ALL gene list compared to segmental duplication in other genes. Indeed, after correcting for the higher proportion of sequences matching the C motif than the A motif in the genome, the C binding motif is more than 4 times more likely to be found in a segmental duplication within the ALL genes than the A motif (OR = 4.78 (1.83-12.46 CI 95%), Table S9).

Infant B-ALL patients show an excess of C *PRDM9* alleles in the St. Jude ALL cohort and harbor leukemic clones with translocations involving the *MLL* gene on chromosome band 11q23. We therefore scanned the nucleotide sequence of *MLL* and found a motif matching the C putative binding sequence occurring within the breakpoint cluster region (Figure 4A) whereas no A binding motif was present in *MLL*. However, 75–90% of genomic DNA sequences are packaged into nucleosome particles, blocking the DNA from interacting with DNA binding proteins (Segal et al. 2006). Studies suggest that the sequence itself is highly predictive of nucleosome positioning, we thus used an *in silico* approach (Xi et al. 2010) to predict nucleosome positioning within the *MLL* breakpoint cluster region (Material and Methods). The tool predicts a potential starting position of the nucleosome at the end of the motif identified (Figure 4B), suggesting that the motif might be accessible in a stretch of unwrapped linker DNA. It follows that *PRDM9* C allele can potentially bind the *MLL* breakpoint cluster region, although this needs to be demonstrated *in vitro* and *in vivo*. Additionally, we re-analysed translocation data from sperm cells in men with known *PRDM9* alleles (Berg et al. 2010). The t(11;22)(q23;q11) translocations, often resulting in a *MLL*-rearrangements, occur at significantly higher frequencies in European males with *k*-finger alleles compared to those without *k*-finger alleles ($p = 0.0436$, Kruskal-Wallis test). However, no significant difference in translocation frequencies was observed between individuals of African descent with and without *k*-finger allele ($p = 0.7998$, Kruskal-Wallis test).

DISCUSSION

In this study, we examined germline processes in families with children having childhood pre-B acute lymphoblastic leukemia (B-ALL). With exome sequencing and dense genotyping data, we were able to capture parental germline recombination events in a unique family with two affected siblings. We identified *PRDM9* as being associated with unusual recombination patterns and discovered a substantial excess of rare allelic forms of *PRDM9* in two independent ALL cohorts. In both the initial and replication ALL cohorts, care has been taken in controlling for population structure (Supplementary Results), the cause of many false-positive genetic associations. These data support the hypothesis that *PRDM9* rare allelic variants are associated with ALL in children, but represent a relatively small dataset and the findings require further support in independent cohorts. The association should also be investigated in other types of childhood leukemias, such as T-lineage ALL and acute myeloid leukemia, as well as in parents of children with constitutional aneuploidies (Ganmore et al. 2009) or in women experiencing molar pregnancies (Roman et al. 2006). The minisatellite alleles of *PRDM9* have to be carefully typed from sequencing read data and rare alleles should be validated through re-sequencing. These alleles are known to cause functional biological variation, as variants in the *PRDM9* gene influence recombination locations, although they have little effect on the total genome-wide recombination rate (Kong et al. 2010). If confirmed, this novel association suggests additional biological function for *PRDM9* allelic variation that might impact other processes than meiotic recombination.

These findings raise many important questions about both leukemogenesis and *PRDM9* function. First of all, *PRDM9* activity is likely exclusive to parental germ cells but it remains unclear if it acts in the patient somatic cells. The parents carrying rare *PRDM9* alleles only transmit the susceptibility allele to half of their affected children in the FCALL cohort, indicating that these alleles may act during meiosis, giving rise to gametes that predispose the offspring to B-ALL. Furthermore, *PRDM9* specific expression and function at early stages of meiosis (Hayashi et al. 2005) support the parental model and point to a germline mechanism of ALL development. However, *PRDM9* expression has been observed, albeit at low levels, in several hematopoietic tissues including leukemia cell lines (Johnson et al. 2003). Genomic data from additional family cohorts is needed in order to definitely resolve this question. Additional family data will also be informative with respect to the sex-specificity of the effect. Indeed, the higher frequency of maternal rather than paternal carriers of rare alleles among parents in the FCALL cohort suggests a strong sex-specific effect, at least for some alleles. Maternal-specific effects on recombination with implications for child health have been demonstrated in humans, such as maternal age effect on recombination (Hussin et al. 2011) and a maternal origin of most division errors leading to trisomies (Hassold and Hunt 2009). The sex-specific effect of *PRDM9* alleles on B-ALL risk could result from the differences in recombination patterns along chromosomes and in hotspot usage between males and females mammals (Paigen et al. 2008; Kong et al. 2010) or from sexual dimorphism in the regulation of the meiotic process (Cohen et al. 2006).

The mechanism underlying this novel association is not known, yet previous evidence suggests that *PRDM9* may be responsible for chromosomal translocations due to its ability to determine sites of genetic crossing over (Myers et al. 2008; Berg et al. 2010; Borel et al. 2012). Therefore, *PRDM9* could be implicated in the generation of some chromosomal rearrangements found in leukemia, a hypothesis supported by analysis of putative binding motifs of *PRDM9* alleles occurring in a subset

of genes involved in chromosomal rearrangements frequently found in ALL (Table S8). In these analyses, we used the *in silico* predicted binding motifs for the PRDM9 C and A alleles as predictors for the binding activity of PRDM9 ZnF array to DNA sequences. PRDM9 binding properties are still mysterious (Segurel et al. 2011), and PRDM9 *in silico*-derived binding sites are not necessarily reliable for predicting PRDM9 binding activity. This is because *in silico* predictions using zinc finger databases and the algorithm developed by Persikov and colleagues (Persikov et al. 2009) give a vast excess of sites compared to those actually bound by PRDM9. Nevertheless, the binding predictions for PRDM9 common allele A led to the discovery of the role of this gene in human recombination (Myers et al. 2010) and the predicted C binding motif matches almost perfectly the DNA motif found in excess in African-enriched hotspots (Hinch et al. 2011). Therefore, it appears that human alleles A and C are able to bind to at least a subset of these genomic motifs. It follows that the significant excess of sequences matching the C motif compared to A motif in the ALL translocated genes, although not a demonstration that C binds these sequences, suggests that the C allele is more likely than the A allele to bind in these fragile regions. In particular, we identified a motif matching the C allele binding motif occurring within the breakpoint cluster region of *MLL* (Figure 4), a gene translocated in infant B-ALL patients, however *in vitro* and *in vivo* binding of PRDM9 C allele in this region remains to be demonstrated.

Importantly, translocations in ALL patients generally occur somatically. In humans, perturbation of the H3K4me3 dynamics at early stage of development specifically leads to inappropriate differentiation of haematopoietic progenitor cells (Chi et al. 2010). Furthermore, the H3K4me3 mark, specifically placed by PRDM9 (Hayashi et al. 2005), is a histone methylation event mis-regulated in many pediatric cancers (Schwartzentruber et al. 2012; Wu et al. 2012; Zhang et al. 2012) and has been linked to leukemia initiation (Chi et al. 2010). In particular, deregulation of factors that mediate H3K4me3 interferes with RAG-mediated V(D)J recombination, crucial for B cell maturation, and affects hematopoietic cell populations. For example, local accumulation of H3K4me3 has recently been shown to occur within the breakpoint cluster region of *BTG1*, a driver gene affected by deletions that result from aberrant somatic recombination events in B-ALL (Waanders et al. 2012). Aberrant histone methylation in germline may therefore help establish tumor-initiating cell populations in early leukemogenesis. However, this hypothesis implies that H3K4me3 marks would be passed on to the child and maintained until early development. Transgenerational epigenetic inheritance has been recently reported for the H3K4me3 mark in *C. elegans* (Greer et al. 2011) and depends on chromatin modifiers but also on the H3K4me3 demethylase RBR-2 acting in germline, suggesting that other contributors, acting in concert with PRDM9, would be required to disrupt normal H3K4me3 patterns.

As these results potentially link allelic variation at *PRDM9* with childhood ALL risk, it is reasonable to expect that higher frequencies of alternative alleles in individuals of African descent (Berg et al. 2010; Hinch et al. 2011) would indicate a potentially higher incidence of childhood leukemia in African populations. Incidence of childhood leukemia in sub-saharian Africa is not well documented, however the incidence rate among admixed African-American children is approximately half the rate in children of European descent (Gurney et al. 1995). Therefore, PRDM9 potential role in ALL will likely involve multi-locus interactions arising in specific genomic backgrounds. The significant difference in frequency of t(11;22)(q23;q11) translocations between men with and without PRDM9 *k*-finger alleles found in Europeans but not in Africans suggests that different alleles, or combination of alleles in a heterozygote, may not have the same impact on different genetic backgrounds. These differences could arise due to the existence of

variation between European and African individuals in factors implicated in regulating H3K4me3 in meiosis after DBS repair. Moreover, since PRDM9 interacts with specific binding motifs to regulate histone methylation and recombination, these loci, if mutated, could modify downstream PRDM9 deleterious functions. Two studies (Jeffreys and Neumann 2002; Myers et al. 2010) have shown that self-destructive drive due to biased gene conversion disrupts the common-allele binding motifs and, in the African population, the same process appears to be eliminating binding motifs for alleles at higher frequencies (Berg et al. 2011). Furthermore, a recent study argued that genetic ancestry is critical to understand ALL risk and failure to go into remission, an indicator of relapse risk in ALL (Yang et al. 2011). In this context, it makes sense to consider that not only are *PRDM9* alleles critical, but also that the ancestral background of the patients is a key factor for the role of PRDM9 in leukemogenesis.

While PRDM9 is known to be involved in sterility in human (Irie et al. 2009) and mice (Hayashi et al. 2005), this is the first study to specifically implicate PRDM9 in human disease and this novel association will hopefully inspire further investigation. PRDM9 clearly interacts with multiple factors to facilitate proper histone methylation and recruit recombination, but its molecular partners remain largely unidentified (Segurel et al. 2011) and the biological importance of PRDM9 is not fully understood. Finally, if PRDM9 is implicated in a germline mechanism of ALL development, it would mean that risk factors for ALL could be established as soon as during meiosis in the parental germline. Therefore, the results reported here raise the intriguing possibility that germline events and recombination processes play a role in the susceptibility to pediatric cancer, which have considerable implications for mapping strategies as well as prognosis and treatment of childhood leukemia.

MATERIAL AND METHODS

Datasets.

The initial French-Canadian B-cell precursor acute lymphoblastic leukemia cohort (FCALL cohort) includes blood samples from 23 French-Canadian nuclear families, which include 22 parental trios (both parents and an affected child) and one family composed of both parents and two affected brothers, referred herein as the ALL quartet. DNA from each sample was extracted from peripheral blood cells or bone marrow as previously described (Baccichet et al. 1997). Exome sequencing using the Applied Biosystems SOLiD 4.0 System, read mapping and variant calling were performed as outlined below. Study subjects are from the established Quebec Childhood ALL cohort (Healy et al. 2010) diagnosed in the Hematology-Oncology Unit of Sainte-Justine University Hospital, Montreal, Canada, between October 1985 and November 2006.

The replication cohort (St Jude ALL cohort) is composed of 50 unrelated children affected with B-ALL treated at St-Jude Children's Research Hospital, Memphis, Tennessee, USA. Whole-genome DNA sequencing was performed using Illumina HiSeq paired-end sequencing at coverage of 30X, aligned using BWA (0.5.5) aligner to the human NCBI Build 36 reference sequence. The Institutional Review Board of the respective hospitals approved the research protocol and informed consent was obtained from all participants and/or their parents.

Exome sequencing data from two control cohorts were used in this study. The first one is a French-Canadian exome dataset (FCEXOME) consisting of 68 French-Canadian controls for whom exome sequencing reads aligning to PRDM9 ZnF array were available. The second exome data control cohort comprises 99 individuals of European descent from the CEU population sequenced in the 1000 Genomes Project (The 1000 Genomes Project Consortium 2010). The read data from these control cohorts are further described in Supplementary Results. PRDM9 re-sequencing data and genotyping data from two additional control cohorts were included: a cohort of families of self-declared French-Canadian origin (FC family cohort) (Hussin et al. 2011) and a cohort of unrelated Moroccan individuals (Moroccan cohort) (Idaghdour et al. 2010). Table S10 summarizes cohorts' composition, data generated and analyses performed on each dataset.

We also used SNP data from the publicly available HGDP (Cann et al. 2002) and HapMap3 (Consortium 2005) datasets as controls in some ancestry and association analyses.

Exome Sequencing in the FCALL cohort.

Exome capture was performed with the *SureSelect Target Enrichment System* from Agilent Technologies using the protocol optimized for Applied Biosystems' SOLiD sequencing. For each sample of the FCALL, single-end exome sequencing cohort was performed and generated approximately 5 Gb of mappable sequence data per sample. Color space reads were mapped to the NCBI Build 36 reference sequence with BioScope v1.2 (Ondov et al. 2008). Base quality recalibration was performed using GATK and BAQ (Li et al. 2009; McKenna et al. 2010). PCR duplicates were removed using Picard implemented in Samtools (Li et al. 2009). In the ALL quartet, SNP calling from exome data was performed as described in Supplementary Methods. De novo

mutation discovery was performed using the DND software (Cartwright et al. 2012). In the 22 trios, we called SNPs in parents using Samtools and only SNPs within targeted regions with Phred score above 30 were kept.

Genome-wide SNP Arrays

Genotyping data was available for all individual included in this study except for the 22 trios from the FCALL cohort. The ALL quartet samples were genotyped using the Illumina HumanOmni 2.5-quad BeadChips and the Affymetrix 6.0 arrays (Supplementary Methods). Normal samples from children in the St Jude ALL cohort were genotyped on Affymetrix SNP 6.0 (48 individuals) or 250k Nsp and 250k Sty (2 individuals). The Moroccan cohort comprises 163 unrelated individuals genotyped on the Illumina's Human 610-Quad SNP Beadchip (Idaghdour et al. 2010). The FC family cohort comprises 69 nuclear families with at least two offsprings, genotyped on the Affymetrix 6.0 array (Hussin et al. 2011).

Recombination Analyses

Recombination events were called in two datasets using the algorithm described previously (Hussin et al. 2011) available at www.iro.umontreal.ca/~hussinju/NucFamTools.html, on two datasets :

- (i) the ALL quartet recombination SNP dataset, which is obtained by combining the exome and the Illumina SNPs (Supplementary Methods). The recombination events were called separately for the pre-treatment and post-treatment samples. All double recombination events separated by 1 and 2 informative markers were ignored. The pre-treatment and post-treatment sets of recombination events were subsequently compared and only events seen in both were kept. All events excluded were double recombinants separated by less than 5 informative markers. Full description of the recombination landscape in the parents of the ALL quartet is provided in Supplementary Results.
- (ii) the Affymetrix SNP dataset for the 69 families from the FC family cohort and the ALL quartet. These samples were genotyped on the same chip, allowing direct comparison of the ALL quartet recombination landscape with recombination patterns observed in the FC family cohort. The 355,271 SNPs used in (Hussin et al. 2011) were selected and 20,118 SNPs were removed because of missing genotypes or mendelian errors in the ALL quartet data. Recombination events were located using two children in each family : for families with three offspring or more in the FC family cohort, two of the latter were chosen at random to match the ALL quartet family structure. We removed all double recombination events separated by less than 5 informative SNPs.

The congruence of HapMap2 recombination hotspots (Consortium 2005) was assessed using events localized between informative markers less than 30 kb apart. We estimated the proportion of recombination events that overlap a HapMap2 hotspot by chance using the approach described in (Coop et al. 2008). We also investigated the overlap between the inferred maternal and paternal recombination events and the putative *PRDM9* alleles A and C binding motif. Again, we only considered events localized between informative markers less than 30 kb apart. We also excluded recombination events overlapping a DNA sequence matching both motifs and, for each individual, we computed the proportion of events overlapping sequences matching exclusively A or C predicted binding motifs.

PRDM9 Zinc Fingers Typing in Short Read Data

To identify PRDM9 zinc fingers (ZnF) repeat types present in an individual, we analyzed sequencing reads that aligned to the PRDM9 ZnF array (exon 11) of the human NCBI Build 36 reference sequence (hg18), corresponding to the region chr5:23562605-23563612. The reads were extracted from BAM files before removing PCR duplicates i.e. multiple reads starting at the same reference coordinate. This is because applying this filter would cause the removal of reads sampling additional ZnF repeats, absent from the reference genome, that will align to the repeats present in the reference genome. Each read was aligned to the known human PRDM9 ZnF types identified in previous studies (zinc finger repeat types *a* to *t*, Figure S6). We computed the proportion of reads that aligned uniquely and without mismatch to each ZnF type. Given the proportion of reads aligning to types *b*, *c*, *d* and *f*, included in all PRDM9 ZnF alleles reported so far (Berg et al. 2010), we determine an inclusion criteria of 1% to infer the presence of a ZnF in a sample. Validation experiments by Sanger sequencing allowed us to confirm the accuracy of the 1% empirical criteria, since ZnF types predicted using this approach were present in the re-sequenced *PRDM9* alleles.

Sanger Sequencing of PRDM9 ZnF Alleles

We sequenced the ZnF array of *PRDM9* in 26 parents from the ALL cohort (including the ALL quartet parents), 76 parents from the FC family cohort and 27 unrelated individuals from the Moroccan cohort, resulting in a total of 258 alleles sequenced. *PRDM9* ZnF alleles were amplified from 5-20 ng of genomic DNA using the primers HsPrdm9-F3 and HsPrdm9-R1 (Baudat et al. 2010), designed to discriminate *PRDM9* from his paralogous copy *PRDM7*. Alleles were sequenced from diploid PCR products with primers 214F, 731F, 1742R and 1992R (Figure S9). Nonmixed sequence traces matching the A allele of *PRDM9*, indicating A/A homozygosity, were identified. We subsequently used the web-based tool Mutliple SeqDoc (Crowe 2005) to compare mixed traces with nonmixed A/A traces. This algorithm produces aligned images of a reference and a test chromatogram together with a subtracted trace showing differences between chromatograms. These difference profiles allow rapid visual identification of base substitutions, insertions and deletions in the test sequence. The differences highlighted by the algorithm are then visually checked and interpreted to avoid potential artifact calls often introduced by automatic base-calling software. This procedure allowed us to determine allele status for all individuals (Table S3 and S5, Figure S4). Most of the individuals were homozygotes A/A (64%), and all remaining individuals were heterozygotes. We identified 10 previously-characterized alleles (B, C, D, E, L1, L3, L9, L20, L24, L14) (Baudat et al. 2010; Berg et al. 2010) and 7 novel alleles found only in this study (L32-38). The novel alleles are described in Supplementary Results.

Association Testing and Ancestry Analyses.

Association between *PRDM9* alleles and ALL in the FCALL cohort was evaluated using randomization inference based on two-tailed Fisher's exact test with 10,000 permutations. For replication in the SJDALL cohort, one-tailed Fisher-based permutation tests were performed. To test whether rare alleles of *PRDM9* were over-represented in ALL subtypes, we performed a permutation test where we permuted the 50 children from the St Jude cohort across subtypes 10,000

times and computed how many times patients with *k*-finger alleles were only seen in hypodiploid and infant B-ALL subtypes. The FC family cohort was used as control cohort for association between PRDM9 ZnF alleles and disease and HapMap3 CEU individuals were considered as controls for association between SNP rs12153202 and disease (Supplementary Results). Ancestry was studied by Principal Component Analyses of genotyped genetic variation of subjects and controls using the smartpca module from the Eigensoft package (Price et al. 2006). Detailed ancestry analyses can be found in Supplementary Results.

Genomic Motif Search

A motif search was performed to identify PRDM9 ZnF allele A and C binding sequences in the human genome. Motif search was performed on the human reference genome (hg18) and on a custom-made degenerate reference genome, constructed using biallelic SNPs in dbSNP v134 that were validated (VLD flag, set if the SNP has at least 2 minor allele counts). At each position where a SNP was reported, the nucleotide in the reference genome was replaced by the IUPAC code corresponding to allele variation. We then scan degenerate or non-degenerate genomes to search for specific degenerate motifs, denoted A and C (Table 2), representing DNA binding motifs predicted by Berg and colleagues (Berg et al. 2010) for PRDM9 ZnF allele A and C, containing only nucleotides predicted with >95% accuracy, based on the algorithm by Persikov and colleagues (Persikov et al. 2009). The sequences found were then annotated based on their starting position using a list of coordinates for regions of interest. We counted the number of non-overlapping motifs occurring whole-genome, in genes and in segmental duplications within genes. Coordinates of all annotated human genes and segmental duplications were obtained from UCSC tables.

Mapping PRDM9 binding motifs within the ALL gene list

We built an ALL gene list using an unbiased strategy based on Mitelman and dbCRID databases (Kong et al. 2011; Mitelman et al. 2011) as of July 2011. Translocations were selected only if they were reported in more than 10 entries for ALL in these databases (Table S8A). We next retrieved fusion genes involved in these translocations from the databases and a final total of 38 genes were kept, following a literature search performed to verify that they have been implicated in ALL in peer-reviewed publications (Table S8B). We computed the number of sequences matching the A and C motifs occurring within and outside of selected ALL genes. Chi-square tests are sensitive to sample size and will tend to reject the null when the sample becomes sufficiently large. Because we have huge numbers for the motif counts, we used Odds Ratios (OR) to compare the frequencies between motifs:

$$OR = \frac{p_{m=A} / 1 - p_{m=A}}{p_{m=C} / 1 - p_{m=C}}$$

with p_m the ratio between the number of motifs m in the ALL gene list and the total number of motifs m in a particular genomic dataset, such as the whole genome, genic regions (repetitive and unique DNA, Table S9) and in segmental duplication occurring in genes. This provides a measure of the strength of non-independence between the motifs and their occurrence in the ALL gene list. Confidence intervals were calculated following the procedure described in (Morris and Gardner 1988). We generated 1000 lists of randomly chosen genes, with the inclusion criteria being a gene

length of 3Kb or more. The experiments described above were repeated with each random gene list in place of the ALL gene list and we computed the number of time significant OR were seen to obtain a two-tailed p -value. We used the program NuPoP (Xi et al. 2010) to predict nucleosome positioning within the *MLL* breakpoint cluster region (chr11:117857800-117858100, hg18).

Translocation Data.

We used t(11;22)(q23;q11) translocation frequencies previously published by Berg and colleagues (Berg et al. 2010), based on *de novo* detection of translocations in sperm from men with different *PRDM9* alleles. We separated African from European men and performed a Kruskal-Wallis test to compare men carrying k -finger alleles and men carrying other alleles in each population. For Africans, the k -finger alleles (C, L4, L6, L14-19) showed no significant influence on translocation frequency ($\chi^2 = 0.0643$, $p = 0.7998$). However, in Europeans, men with k -finger alleles (C, L20) showed significantly increased translocation frequencies ($\chi^2 = 4.0735$, $p = 0.04356$).

DATA ACCESS

Sequences of the *PRDM9* novel alleles have been deposited in GenBank under the accession number JQ044371-JQ044377. Genomic reads aligning to *PRDM9* exon 12 from all individuals in the FCALL and control cohorts are deposited in NCBI Sequence Read Archive (<http://www.ncbi.nlm.nih.gov/sra>) under the accession number SRA060797. Whole-exome sequencing reads and genotyping data for the ALL quartet will be made available through the Quebec childhood leukemia web portal (<http://childhoodleukemiagenomics.org>). Genomic sequence data from the SJALL cohort is available through the European Genome-Phenome Archive (<https://www.ebi.ac.uk/ega/>) under the accession number EGAC00001000044 and can be accessed by application to the Pediatric Cancer Genome Project (PCGP) Data Access Committee. More information can be found at <http://explore.pediatriccancergenomeproject.org>.

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AUTHOR'S CONTRIBUTION

JH1 and PA designed the study. DS YI JH2 GG AA LH GA JRD CGM and PA contributed reagents, patient materials and samples. JH1 FC JFS ML SB performed experimental work. JH1 VB VS JCG TdM and PA performed computational data analyses of the Sainte-Justine cohort data. JH1, JM, LW, JZ performed computational data analyses of the St. Jude cohort data. JH1 and PA wrote the paper. (JH1: J. Hussin, JH2: J. Healy)

DISCLOSURE DECLARATION

The authors declare no competing financial interests.

FIGURE LEGENDS

Figure 1. The ALL quartet family pedigree

The ALL Quartet is composed of the two parents and two brothers (patients 383 and 610) affected by hyperdiploid B-cell precursor childhood ALL, sampled prior to and after chemotherapy treatment. The brothers were diagnosed within a 3-year time period. The parents report Moroccan origins. Both maternal and paternal grandfathers are deceased from cancer. One of the father's sisters had children with poly-malformation syndromes, likely due to the high degree of consanguinity reported. Age at death is shown for deceased individuals.

Figure 2. Map of recombinations events and hotspot usage in the ALL quartet

(A) Single and double crossovers in the two meioses that give rise to the patients, determined from analyses of SNPs from exome sequencing and genotyping data. Analyses were performed using pre- and post-treatment samples and only kept crossovers inferred in both. Using two somatic tissues allowed us to remove genotyping errors and double recombination events resulting from errors. All crossovers displayed are supported by at least 3 informative markers and high-resolution events are localized between informative markers less than 30 Kb apart (B) Fraction of high-resolution crossover intervals overlapping population hotspots in the FC family cohort and in the ALL quartet. Mothers (triangles) and fathers (circles) are ordered according to their proportion of overlap. We estimate that 11.78% (10.56-13.24 CI 95%) of these crossover intervals are expected to overlap population hotspots by chance.

Figure 3. Excess of rare *PRDM9* alleles in parents from the FCALL cohort

(A) Pie charts showing frequencies of *PRDM9* zinc-finger (ZnF) alleles obtained through Sanger sequencing of 26 parents of patients with B-ALL and 76 parents from the FC family cohort (controls). Alleles labeled as 'Others' are population-specific alleles. Individuals' alleles are detailed in Table S3 and S5. (B) Differences in allele frequencies between parents of patients and controls. The *p*-value in parentheses was calculated by including alleles inferred from exome sequencing reads for the 20 ALL parents for which *PRDM9* ZnF arrays were not re-sequenced by Sanger (Table S4). Applying a Bonferroni correction for performing the same test in two subsets of alleles (non-A and *k*-fingers alleles) would yield $\alpha = 0.025$, although this correction is particularly conservative since subsets are correlated. (C) Allelic structure of *PRDM9* ZnF array for alleles found in these cohorts (population-specific alleles not shown).

Figure 4. *PRDM9* C binding motif in the MLL breakpoint cluster region.

(A) Logo plot of the C allele binding motif (Baudat et al. 2010), predicted based on the 3 indicated residues forming the binding unit of the ZnF repeats (positions -1, 3 and 6 of the ZnF alpha helices) and the consensus sequence motif simplified showing the most strongly predicted bases (in lowercase for

>80% consensus for a specific base and in uppercase for >95% consensus (Berg et al. 2010) (B) Presence of a motif at chr11:117857908-117857950 (hg18), within the breakpoint cluster region of *MLL*, matching the predicted PRDM9 C allele binding motif for the 7 strongly predicted bases shown in uppercase in the consensus sequence presented in (A), and 3 predicted bases shown in lowercase. Intronic regions are displayed in light blue. Black arrows show the positions of all occurrence of sequences matching the motif at uppercase characters. No *PRDM9* A allele binding motif was found in *MLL*. (C) Nucleosome starting positions predicted by NuPoP (Xi et al. 2010). The red line shows the position of the predicted C binding motif.

TABLES

Table 1. Replication of the association between *PRDM9* *k*-finger alleles and in patients from St. Jude ALL cohort

ALL subtypes	All Patients			Patients of European Ancestry		
	Individuals	<i>k</i> -finger alleles	<i>P</i> -value ^a	Individuals	<i>k</i> -finger alleles	<i>P</i> -value ^a
B-ALL	50	10	0.0143	39 ^b	7	0.0396
Hypodiploid B-ALL	16	5	7.50 × 10 ⁻³	12	4	8.85 × 10 ⁻³
Infant B-ALL	18	5	0.0122	14	3	0.0630
ETV6 B-ALL	9	0	-	9	0	-
Ph+ B-ALL	7	0	-	4	0	-

^a *P*-values from permutation tests based on one-tailed Fisher's exact tests on counts between cases and controls^b The 39 patients represent a subgroup of the cohort of 50 patients

Patient's ethnicities were verified by principal component analyses on genetic variation using Eigensoft package (Figure S8, Supplementary Results. Patients do not have African ancestry. Controls consist of 76 French-Canadians individuals sequenced at the *PRDM9* locus and showing a total of 5 *k*-finger alleles (Table S5). Association testing between cases and controls was performed for subgroups with sample size greater than 10 individuals. Applying a conservative Bonferroni correction to correct for testing in two independent B-ALL subgroups would yield $\alpha = 0.025$.

Table 2. PRDM9 alleles binding motifs in the Human Reference Genome.

Reference genome	PRDM9 allele	Motif	Number of Motifs		
			Genome	Genes	ALL gene list
Standard	A	A: G...C.....CC.CC....C.CC	12319	6953	34
	C	C: G...C.....CC.....C....CC	176826	95241	748
Degenerate	A	A: G...C.....CC.CC....C.CC	13335	7504	42
	C	C: G...C.....CC.....C....CC	186374	100041	789

Motif comparisons	ALL gene list vs Genome OR [CI]	ALL gene list vs Genes OR [CI]	Random gene lists
C vs A in Standard	1.53 [1.15;2.04] **	1.61 [1.21;2.15] **	43/1000
C vs A in Degenerate	1.53 [1.16;2.01] **	1.60 [1.22;2.12] **	48/1000

** significant based on 95% CI (one-tailed $p < 0.025$)

The motif search was performed on the Human Reference Genome (hg18). Motifs consist in DNA binding sequences predicted by Berg and colleagues (Berg et al. 2010) for each allele. The number of motifs is reported for whole genome, coding regions and the ALL gene list. The ALL gene list was built based on the most frequent ALL translocations reported in databases (Table S8). We compared counts using odds ratios (OR), to measure the association between motifs and their occurrence in the ALL gene list. For a given motif comparison, OR values were computed for 1000 random gene lists (13) and we computed the number of times significant ORs (two-tailed) are seen for both whole genome and genic regions. Results for unique and repetitive DNA are shown in Table S9.

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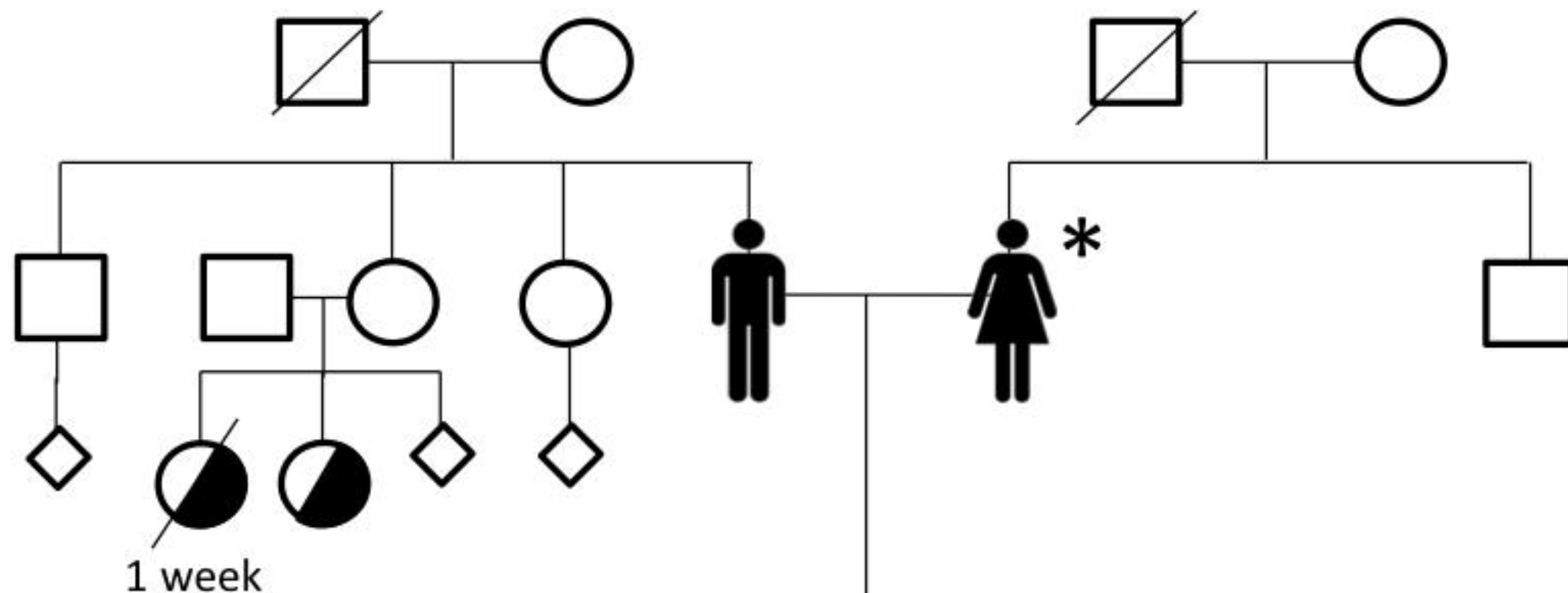
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Prostate cancer, 65yrs

Colon cancer, 69yrs

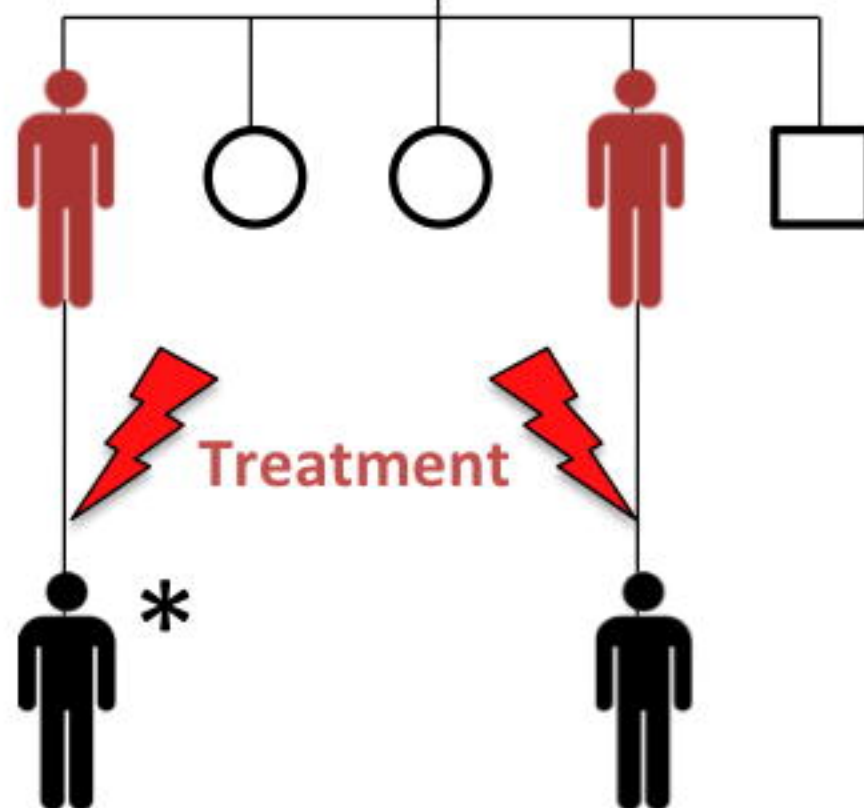


Sibling 383

Diagnosed at 2 years old
Hyperdiploid B-ALL

Sibling 610

Diagnosed at 14 years old
Hyperdiploid B-ALL



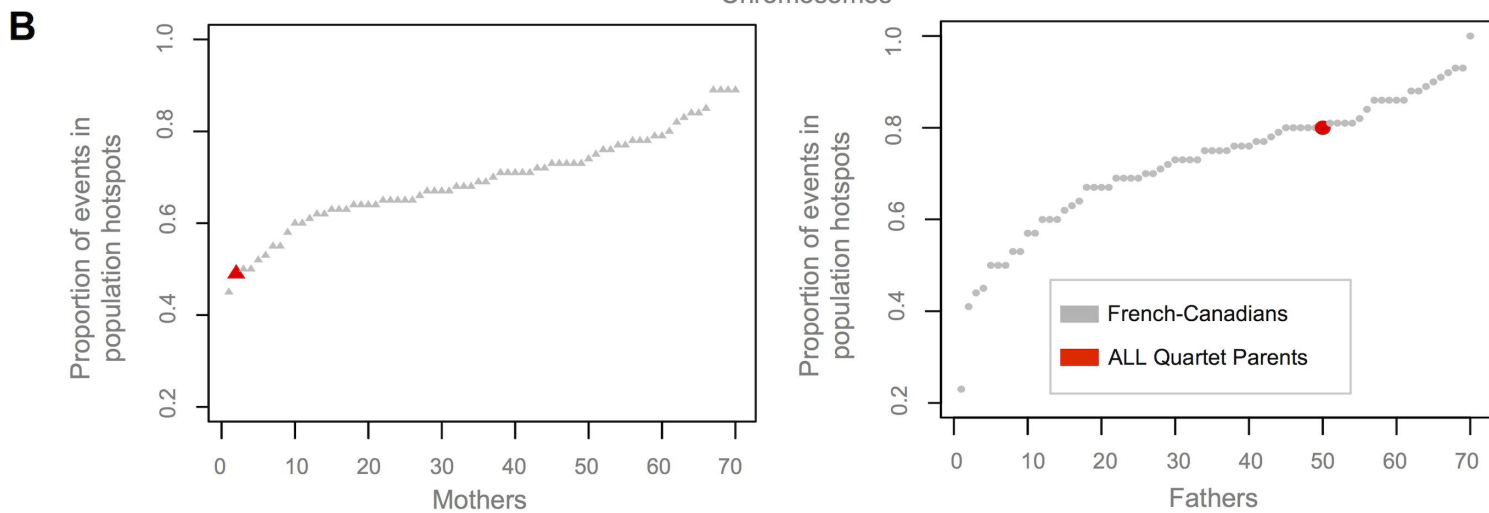
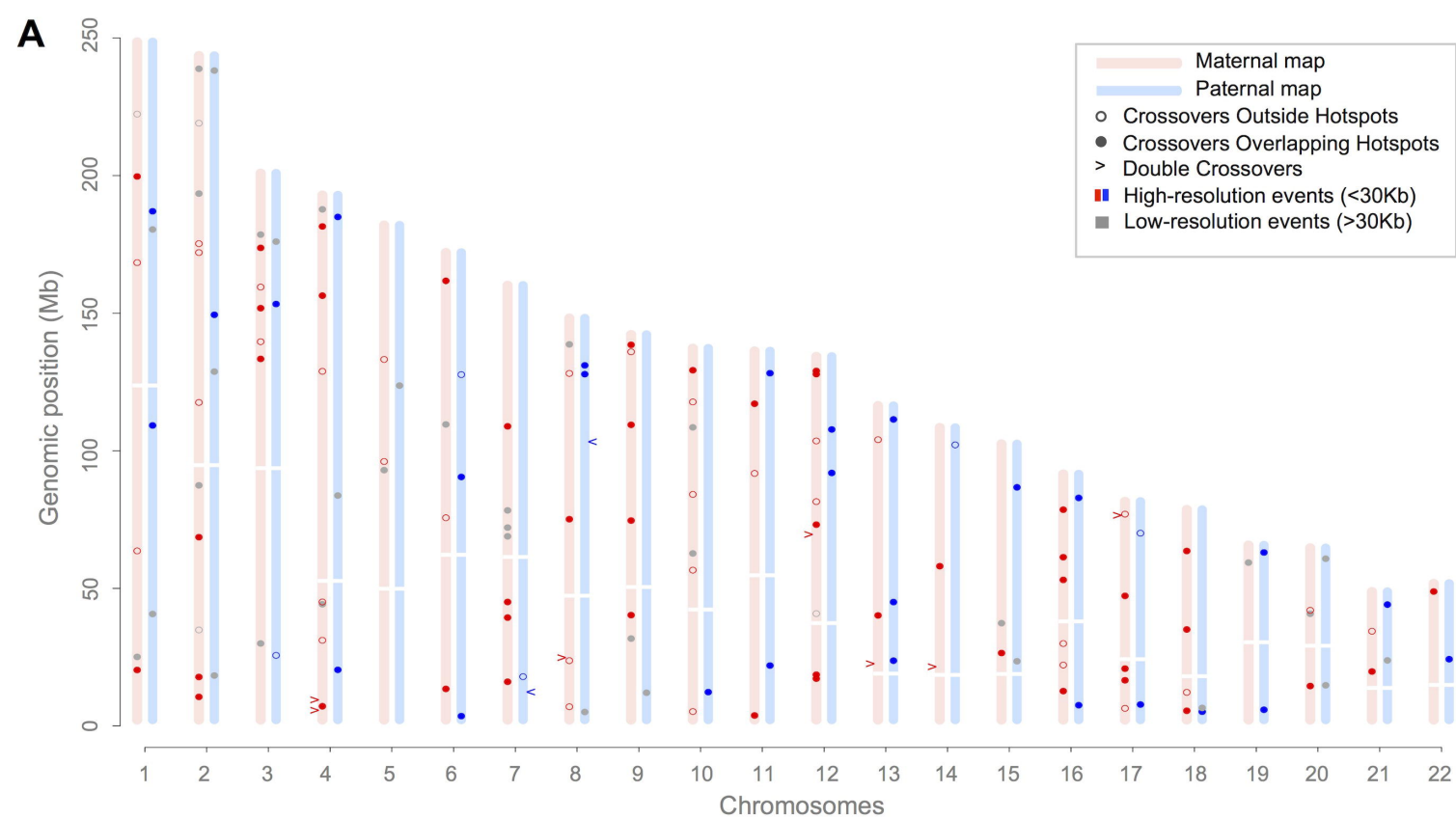
Blood samples sequenced



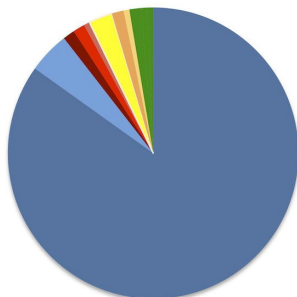
Poly-malformation syndrome



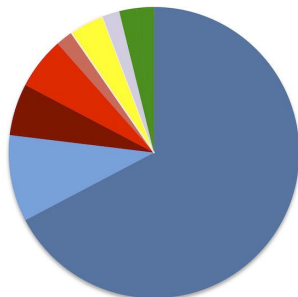
Carriers of $\text{inv}(2)(\text{p11q13})$



A French-Canadians



ALL Parents



■ A ■ B ■ C ■ D ■ L20
■ L24 ■ L3 ■ L9 ■ L1 ■ Others

B

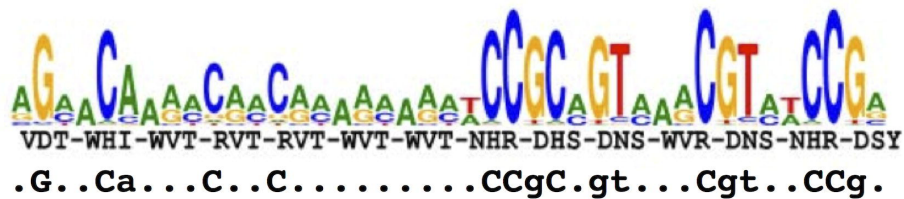
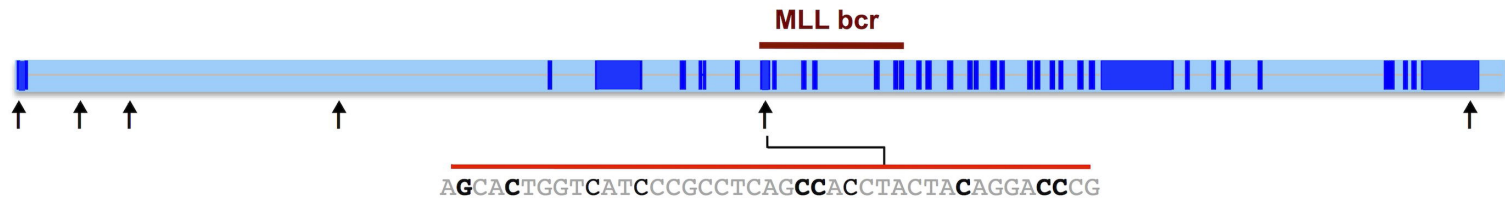
Statistic	ALL Parents	French-Canadians	<i>P</i> -value ^a
Non A alleles	32.7%	15.1%	3.22×10^{-3}
K-finger alleles (C, D, L20)	13.5%	3.3%	6.25×10^{-3} (2.10×10^{-3})
Couples with rare alleles (not A/B)	76.9%	33.3%	6.49×10^{-3}

^a *P*-values computed using two-tailed Fisher-based permutation tests

C

PRDM9 Zinc-Finger Array Alleles

A	a	b	c	d	d	e	c	f		g	h	f	i	j
B	a	b	c	d	d	c	c	f		g	h	f	i	j
C	a	b	c	d	d	c	c	f	k	h	l	h	i	j
D	a	b	c	d	d	e	c	f	k	g	h	f	i	j
L20	a	b	c	d	d	e	c	f		g	k	f	q	j
L24	a	b	c	d	d	e	c	f		t	p	f	q	j
L3	a	b	c	d		e	c	f		g	h	f	i	j
L9	a	b	c	d	d	e	c	f		g	p	f	q	j
L1	a	b	c	d						g	h	f	i	j

A**B****C**