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Somatic mutational landscape of hereditary hematopoietic malignancies caused by germline variants in *RUNX1*, *GATA2*, and *DDX41*

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Access to *RUNX1* genomics data is available through the *RUNX1* database (<https://runx1db.runx1-fpd.org/>). Original data may be obtained by email request to the

corresponding author, Anna L. Brown (anna.brown@sa.gov.au). Access to additional deidentified genomics data is available on request.

The full-text version of this article contains a data supplement.

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Key Points

- Germline *RUNX1*, *GATA2*, and *DDX41* HHMs are associated with driver somatic variants during leukemogenesis which are unique for each syndrome.
- Ongoing molecular monitoring of germline carriers without HM is needed to assess the risk profile and clinical actionability of somatic markers.

Individuals with germ line variants associated with hereditary hematopoietic malignancies (HHMs) have a highly variable risk for leukemogenesis. Gaps in our understanding of premalignant states in HHMs have hampered efforts to design effective clinical surveillance programs, provide personalized preemptive treatments, and inform appropriate counseling for patients. We used the largest known comparative international cohort of germline *RUNX1*, *GATA2*, or *DDX41* variant carriers without and with hematopoietic malignancies (HMs) to identify patterns of genetic drivers that are unique to each HHM syndrome before and after leukemogenesis. These patterns included striking heterogeneity in rates of early-onset clonal hematopoiesis (CH), with a high prevalence of CH in *RUNX1* and *GATA2* variant carriers who did not have malignancies (carriers-without HM). We observed a paucity of CH in *DDX41* carriers-without HM. In *RUNX1* carriers-without HM with CH, we detected variants in *TET2*, *PHF6*, and, most frequently, *BCOR*. These genes were recurrently mutated in *RUNX1*-driven malignancies, suggesting CH is a direct precursor to malignancy in *RUNX1*-driven HHMs. Leukemogenesis in *RUNX1* and *DDX41* carriers was often driven by second hits in *RUNX1* and *DDX41*, respectively. This study may inform the development of HHM-specific clinical trials and gene-specific approaches to clinical monitoring. For example, trials investigating the potential benefits of monitoring *DDX41* carriers-without HM for low-frequency second hits in *DDX41* may now be beneficial. Similarly, trials monitoring carriers-without HM with *RUNX1* germ line variants for the acquisition of somatic variants in *BCOR*, *PHF6*, and *TET2* and second hits in *RUNX1* are warranted.

Introduction

Hereditary hematopoietic malignancies (HHMs) are hematologic syndromes characterized by Mendelian inheritance patterns and an increased lifetime risk for hematopoietic malignancies (HMs).^{1,2} Individuals with HHM-associated germ line variants have a highly variable risk for leukemogenesis, and many HHM-variant carriers do not develop malignancies (carriers-without HM).³ Very little is understood about the premalignant states in carriers-without HM, the molecular and genetic factors that affect leukemogenic risk, or the environmental factors that drive leukemogenesis in HHMs. This knowledge gap has hampered efforts to refine the clinical surveillance of carriers-without HM, identify individuals with the highest risk for HMs, and develop interventions that delay or prevent leukemogenesis in high-risk carriers-without HM. Moreover, treatments used for malignancies in HHM-variant carriers (carriers-with HM) are not tailored to these syndromes aside from *DDX41* and *GATA2* carriers, for which there is a limited role for lenalidomide therapy or prophylactic hematopoietic stem cell transplant, respectively.⁴⁻⁶ Instead, carriers-with HM are treated with standard-of-care therapies for sporadic HMs, which may carry an uncharacterized gene mutation-specific risk of additional treatment effects, such as engraftment failure or secondary therapy-related neoplasms. Given the paucity of HHM families at individual institutions, a coordinated, multi-institutional effort is required to understand the natural history of HHMs, leukemogenic mechanisms, and the unique biologic factors that may be present in individual HHM syndromes.

HHMs have been recognized phenotypically for over 100 years. Autosomal dominant (AD) predisposition to myeloid malignancies

is the most well characterized, with more than 15 AD HHM-related genes identified to date.⁷ Pathogenic germ line variants in *RUNX1*, *GATA2*, and *DDX41* collectively represent the most common causes of AD HHMs and are primarily associated with myeloid malignancies. These HHMs are more common than previously recognized and may have highly penetrant leukemogenic phenotypes. Germ line *DDX41* carriers account for ~2% to 4% of all patients with seemingly sporadic HMs, and *GATA2* carriers have a 90% lifetime risk of developing HMs. *RUNX1*-driven HHMs were the first known HHM syndrome and have a high penetrance for HM (~44%).^{3,4,8-11} Identifying these syndromes can be challenging because of limited syndromic features, and recognition is often made based on a high-risk family history, an early-onset HM, or the identification of an HHM-associated variant on tumor-based molecular profiling.¹² Individuals harboring germ line variants in these genes often present with cytopenias: *RUNX1* most commonly with thrombocytopenia;¹³ *GATA2* with monocytopenia, dendritic cell, B, and natural killer cell (NK) lymphoid deficiency;¹⁴ and *DDX41* with variable cytopenias that can include leukopenia, neutropenia, and/or erythroid dysplasia.^{15,16} The age of myelodysplastic syndrome/acute myeloid leukemia (MDS/AML) diagnosis also differs between HHMs, with *GATA2* carriers developing MDS/AML at a mean age of 19 years, *RUNX1* carriers at 29 years, and *DDX41* carriers at 67 years.^{16,17}

The mechanisms driving leukemogenesis in these variant carriers are unclear. Most work to date has focused on germ line *RUNX1* variant carriers.⁸ *RUNX1* carriers have an increased risk for clonal hematopoiesis (CH) (67%-75% CH^{18,19}). However, because of the rarity of *RUNX1* HHM, single-center studies have limited numbers of patients available (9¹⁹ and 3¹⁸). Recent studies looking at CH in germ line *GATA2* carriers have shown an association between CH

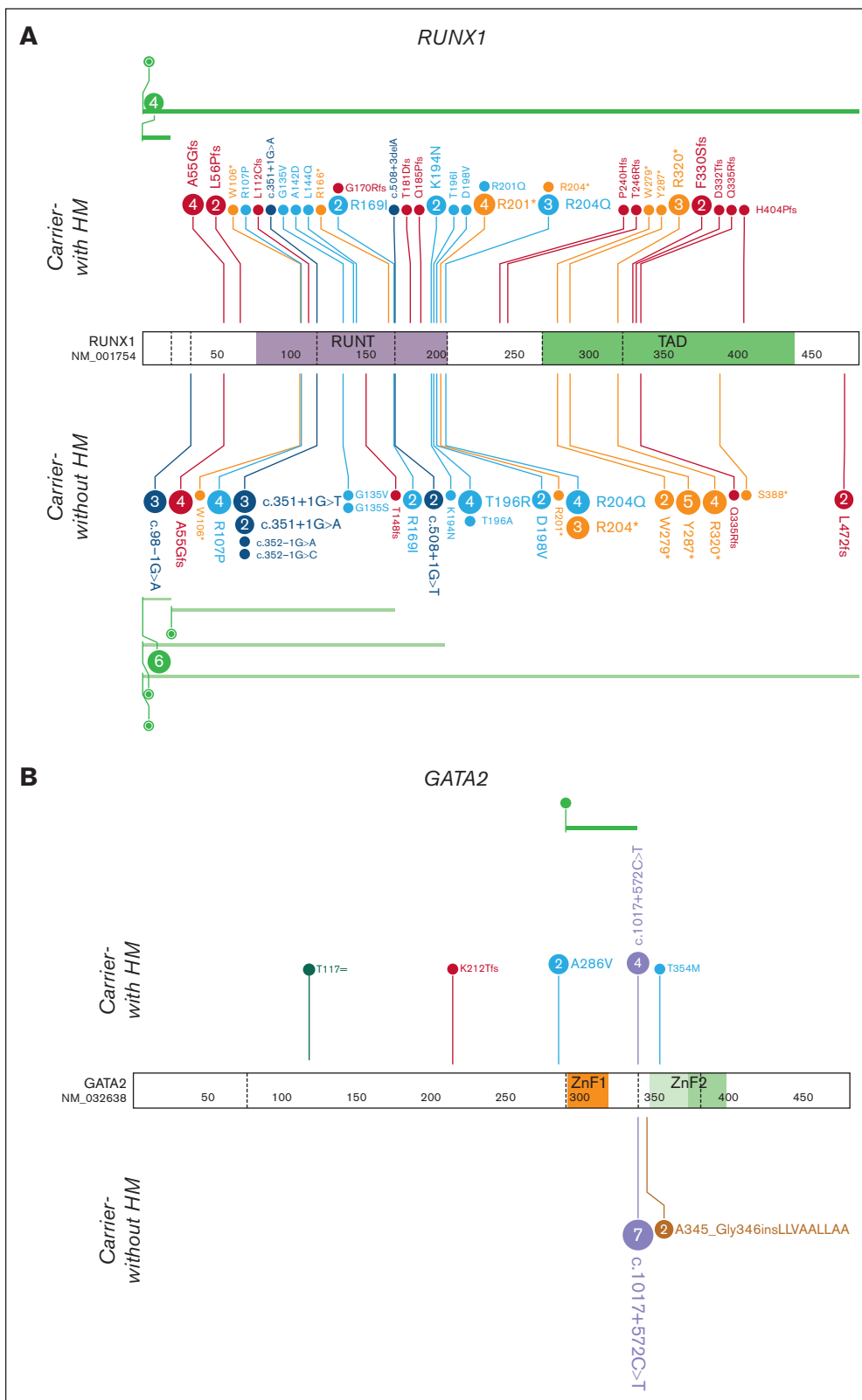


Figure 1. HHM genomics cohorts. Germ line variants in the HHM cohorts were visualized using the ProteinPaint web application.³⁰ Carriers-with HM cohorts (diagnosed with HM) are visualized above the protein. Carriers-without HM cohorts (no HM diagnosis) are below the protein. Variants (displayed as protein changes where possible) are color coded by variant type. The number of individuals with each variant is indicated within the circle when the number is greater than 1. (A) Germ line *RUNX1* (66 carriers-without HM and 52 carriers-with HM individuals); (B) Germ line *GATA2* (9 carriers-without HM and 13 carriers-with HM individuals); and (C) Germ line *DDX41* (22 carriers-without HM and 29 carriers-with HM individuals) cohorts.

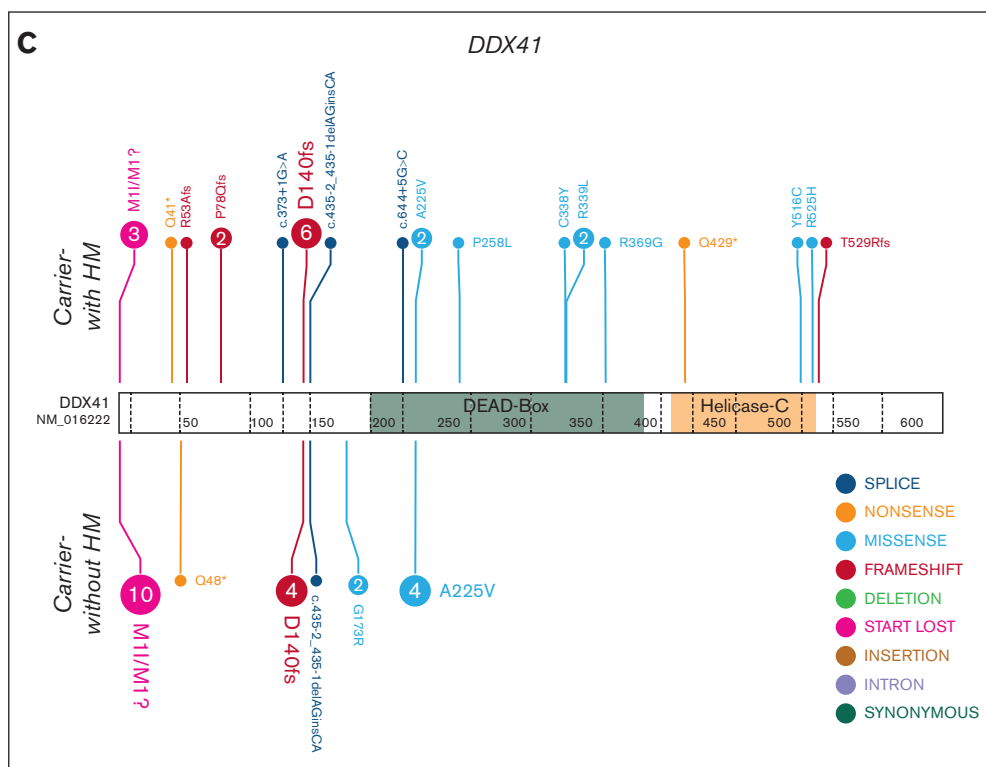


Figure 1 (continued)

and a hypocellular marrow while also linking specific somatic events with the likelihood of leukemic transformation.^{20,21} Similarly assessment of CH patterns in comparative cohorts of HHM-variant carriers may identify specific leukemogenic patterns for different HHMs and ultimately inform clinical trials to define guidelines for clinical surveillance of unaffected HHM-variant carriers.

To address this knowledge gap, we collected retrospective next-generation sequencing (NGS) data from hematopoietic tissue samples from an international cohort of patients with HHM driven by germ line *RUNX1*, *GATA2*, or *DDX41* variants. Our cohort is the largest comparative HHM-focused cross-sectional collection of its kind, with 240 patient samples evenly distributed between carriers-without HM (n = 120) and carriers-with HM (n = 120). We used a uniform variant calling and curation approach to identify driver somatic variants in each sample. This unique distribution of samples from carriers-without HM and carriers-with HM and across multiple HHMs, in conjunction with a uniform bioinformatic approach, enabled us to determine driver somatic variants that develop within hematopoietic tissue in *RUNX1*, *GATA2*, and *DDX41* variant carriers before and after diagnosis of a blood cancer in the HHM syndromes.

Patients and methods

Patient cohort

Clinical and genomics data from germ line *RUNX1*, *GATA2*, or *DDX41* variant carriers were collected from the *RUNX1* database (<https://runx1db.runx1-fpd.org/>),²² the Centre for Cancer Biology (Australia), the University of Chicago (USA), and the National

Institutes of Health (USA). In total, data from 195 patients who had undergone genomics profiling (whole-exome sequencing or panel-based sequencing) were retrospectively collated to form the *RUNX1*, *GATA2*, and *DDX41* cohorts. All procedures in this study involving human participants were performed in accordance with the Declaration of Helsinki. Studies were approved by institutional human research ethics committees and/or institutional research boards. All participants signed an informed consent form to share genomics and protected health information.

NGS reanalysis and variant calling pipeline

NGS data were collected and reanalyzed with the bioinformatics pipeline used for the *RUNX1* database.²² Original FASTQ (textfile format for sequencing data) or Binary Alignment Map (BAM) files were obtained. Sequence reads were aligned to the GRCh37 (hs37d5) human reference genome with BWA-MEM (ver 0.7.12).²³ Sambamba (ver 0.6.5)²⁴ was used for marking polymerase chain reaction duplicates, and GATK (ver 3.8-1) was used to recalibrate base-quality scores. Freebayes (ver 1.2)²⁵ was used to call single nucleotide variants (SNVs) and insertions/deletions (indels). Variant-, gene-, and protein-level annotations were performed using an in-house pipeline (<https://github.com/SACGF/variantgrid>). Somatic variant curation was performed as previously described²² (supplemental Methods). All data sets were independently curated by at least 3 variant curators.

Tumor mutation burden (TMB) analysis

SNVs and indels were identified with Seurat, Shimmer, Strelka, and SomaticSniper,²⁶⁻²⁹ using paired germ line samples (cultured skin fibroblasts or hair) from the same patient to remove germ line

Table 1. Driver somatic variants detected in germ line *RUNX1*, *GATA2*, or *DDX41* carriers-without HM

Individual	Phenotype			Gene	Somatic variant			
	Clinical presentation	Age	Sex		Genomic coordinates	c_HGVS	p_HGVS	VAF (%)
M09_1	Thrombocytopenia	16	F	BCOR	X:39933270 CA>C	NM_001123385.2:c.1328del	p.Leu443CysfsTer8	8.3
M01_3	Thrombocytopenia	17	M	ATP10A	15:25953381 G>A	NM_024490.3:c.2411C>T	p.Ala804Val	5.0
M03_2	Thrombocytopenia	17	M	PHF6	X:133527608 CTG>C	NM_032458.3:c.321_322del	p.Ala108IlefsTer3	39.4
				CUX1	7:101882720 G>A	NM_001202543.2:c.3776G>A	p.Arg1259Gln	42.3
				IDH2	15:90631934 C>T	NM_002168.3:c.419G>A	p.Arg140Gln	9.7
G01_2	Thrombocytopenia	19	F	NOTCH3	19:15276860 GC>G	NM_000435.3:c.5404del	p.Ala1802LeufsTer23	3.2
V01_1	Thrombocytopenia	23	M	EP300	22:41574723 TCCACACCACGTTTCC>T	NM_001429.3:c.7014_7028del15	p.His2338_Pro2342del	29.6
G07_3	Thrombocytopenia	33	M	BCOR	X:39932334 G>GT	NM_001123385.2:c.2264dup	p.Tyr755Ter	2.4
M06_2	Thrombocytopenia	37	F	TET2	4:106164068 G>A	NM_001127208.3:c.3578G>A	p.Cys1193Tyr	11.3
G07_2	Thrombocytopenia	40	M	TET2	4:106164787 C>T	NM_001127208.3:c.3655C>T	p.His1219Tyr	5.3
S_G_3	Thrombocytopenia	40	F	TET2	4:106196461 T>G	NM_001127208.3:c.4794T>G	p.Tyr1598Ter	5.0
W01_3	Thrombocytopenia	40	M	BCOR	X:39921617 CG>C	NM_001123385.2:c.4202del	p.Pro1401ArgfsTer83	20.8
S_D_2	Thrombocytopenia	43	M	DNMT3A	2:25463242 AG>A	NM_022552.5:c.2250del	p.Phe751SerfsTer28	3.1
		48	M	DNMT3A	2:25463242 AG>A	NM_022552.5:c.2250del	p.Phe751SerfsTer28	4.9
U02_1	Thrombocytopenia	49	M	TET2	4:106162587 G>A	NM_001127208.3:c.3500+1G>A	p.?	2.6
A01_1	Thrombocytopenia	53	F	TET2	4:106158510 T>C	NM_001127208.3:c.3409+2T>C	p.?	33.3
				DNMT3A	2:25497943 CG>C	NM_022552.5:c.505del	p.Arg169GlyfsTer56	29.6
				SRSF2	17:74732959 G>A	NM_003016.4:c.284C>T	p.Pro95Leu	22.9
				DNMT3A	2:25497943 CG>C	NM_022552.5:c.505del	p.Arg169GlyfsTer56	42.8
		56	F	TET2	4:106158510 T>C	NM_001127208.3:c.3409+2T>C	p.?	30.5
				SRSF2	17:74732959 G>A	NM_003016.4:c.284C>T	p.Pro95Leu	28.0
				BCOR	X:39932898 T>TG	NM_001123385.2:c.1700dup	p.Ala568SerfsTer43	6.7
				BCOR	X:39923055 C>T	NM_001123385.2:c.3653G>A	p.Trp1218Ter	4.3
G02_2	Thrombocytopenia	55	M	BCOR	X:39911577 GA>G	NM_001123385.2:c.5052del	p.Pro1685GlnfsTer40	2.5
				BCOR	X:39932898 T>TG	NM_001123385.2:c.1700dup	p.Ala568SerfsTer43	6.4
				BCOR	X:39923055 C>T	NM_001123385.2:c.3653G>A	p.Trp1218Ter	2.2
				BCOR	X:39933676 TG>T	NM_001123385.2:c.922del	p.Gln308ArgfsTer70	1.2
				BCOR	X:39933416 TG>T	NM_001123385.2:c.1182del	p.Lys395ArgfsTer47	0.7
				BCOR	X:39932109 ACT>A	NM_001123385.2:c.2488_2489del	p.Ser830CysfsTer6	17.6
W01_2	Thrombocytopenia	68	M	BCOR	X:39933593 A>AG	NM_001123385.2:c.1005dup	p.Ser336LeufsTer45	7.1
I03_2 (BM)	Thrombocytopenia	72	F	DNMT3A	2:25470029 T>C	NM_022552.5:c.1015-2A>G	p.?	13.3
				BCOR	X:39933492 TG>T	NM_001123385.2:c.1106del	p.Ser369Ter	4.0
F01_6	Thrombocytopenia	76	M	BCOR	X:39916476 C>T	NM_001123385.2:c.4527G>A	p.Trp1509Ter	13.6
				ATM	11:108213949 G>A	NM_000051.3:c.8269G>A	p.Val2757Met	17.0
				GRIN2A	16:9857831 G>C	NM_000833.4:c.3570C>G	p.His1190Gln	14.5

Table 1 (continued)

Individual	Phenotype			Gene	Genomic coordinates			Somatic variant		
	Clinical presentation	Age	Sex		Genomic coordinates	c_HGVS		p_HGVS		VAF (%)
						c_HGVS	p_HGVS			
F01_8	Thrombocytopenia	76	M	<i>BCOR</i>	X:39921490 TG>T	NM_001123385.2:c.4329del	p.Thr1444ProfsTer40	34.7		
				<i>TP53</i>	17:7578442 T>C	NM_000546.6:c.488A>G	p.Tyr163Cys	4.1		
U02_3	Asymptomatic	NA	M	<i>CCND3</i>	6:41903707 G>A	NM_0011760.4:c.850C>T	p.Pro284Ser	3.5		
S_E_4	Thrombocytopenia	NA	F	<i>SRSF2</i>	4:106162587 G>A	NM_001127208.3:c.3500+1G>A	p.?	1.2		
D01_2	Thrombocytopenia	NA	M	<i>ATR</i>	17:74732959 G>T	NM_003016.4:c.284C>A	p.Pro95His	13.9		
				<i>DNMT3A</i>	3:142281940 A>G	NM_001184.4:c.304T>C	p.Trp102Arg	3.9		
				<i>BCOR</i>	2:25466797 C>T	NM_022552.5:c.1906G>A	p.Val636Met	24.7		
				<i>DNMT3A</i>	X:39923092 TA>T	NM_001123385.2:c.3615del	p.Lys1207AsnfsTer31	3.6		
				<i>BCOR</i>	2:25464460 C>T	NM_022552.5:c.2053G>A	p.Gly685Arg	3.2		
D02_2	Thrombocytopenia	NA	F	<i>PTPN11</i>	X:39933373 TGCCCGG>TT	NM_001123385.2:c.1220_1225delCCGGGCGinsA	p.Pro407GlnfsTer31	3.1		
U02_4	NA	NA	F	<i>TET2</i>	12:112926887 G>A	NM_002834.4:c.1507G>A	p.Gly503Arg	25.2		
Family_53_8	Asymptomatic	16.5	F	<i>KDM5A</i>	4:106157215 C>T	NM_001127208.3:c.2116C>T	p.Gln706Ter	7.3		
Family_53_3	Asymptomatic	47	M	<i>DNMT3A</i>	12:416952 C>CT	NM_001042603.3:c.3597dup	p.Gly1200ArgfsTer7	3.2		
Family_0127.041	Asymptomatic	87	F	<i>ASXL1</i>	2:25457171 T>A	NM_022552.5:c.2716A>T	p.Lys906Ter	3.6		
				<i>DNMT3A</i>	20:31022576 TAC>T	NM_015338.5:c.2062_2063del	p.Thr688fs29	4.2		
					2:25467022 A>G	NM_022552.5:c.1851+2T>C	p.?	4.1		

variants. Somatic variants identified in 3 or more callers were included with high confidence. Variant calling thresholds were set at alternate allelic depth ≥ 3 and variant allele frequency (VAF) $\geq 5\%$. Somatic variants were filtered and annotated with the variant effect predictor package (hg19). The total number of somatic variants in the tumor exome was divided by the length of exome capture (38 Mb) to calculate the TMB.

Statistical analysis

GraphPad Prism 7.03 and RStudio Version 1.4.17 with tidyverse, ggplot2, ggrepel, caTools, and ROC packages were used for statistical calculations and figures. ProteinPaint was used to create lollipop plots.³⁰ Circos plots were created using ShinyCircos software.³¹ Unless otherwise stated, the *P* value was calculated using a one-way analysis of variance with Tukey multiple comparisons test using a single pooled variance. *P* value of sex differences were calculated using a two-sided Fisher exact test. The prop.test function/z-value was used as a 2-sample test for equality of proportions with continuity correction. Logistic regression modeling was used to determine the relationship between age and CH. The nonparametric Mann-Whitney *U* test was used to calculate the significance of the TMB between the *RUNX1* and *DDX41* cohorts.

Results

Genomic cohorts for germ line *RUNX1*, *GATA2*, or *DDX41* HHMs

Through international data sharing, we created cohorts of carriers-without HM (no HM) and carriers-with HM (diagnosed with an HM) with germ line *RUNX1*, *GATA2*, or *DDX41* variants (Figure 1; supplemental Methods). NGS data included samples from germ line controls, complete remission patients, carriers-without HM, and carriers-with HM. Multiple samples were collected from individuals when available, including longitudinal. The *RUNX1* cohort included 66 carriers-without HM and 52 carriers-with HM individuals (including 80 and 66 independent NGS samples, respectively). The *GATA2* cohort included 9 carriers-without HM and 13 carriers-with HM individuals (9 and 13 NGS samples, respectively). The *DDX41* cohort included 22 carriers-without HM and 29 carriers-without HM individuals (including 31 and 41 independent NGS samples, respectively). Each cohort is summarized in supplemental Table 1 and supplemental Figures 1A and 2. We used a standardized bioinformatics and variant curation approach²² to identify clinically relevant and potentially clinically relevant somatic variants (driver somatic variants, detailed in the supplemental Methods).

CH is prevalent in *RUNX1* and *GATA2* but not *DDX41* HHM carriers-without HM

Age-related CH is frequently observed in healthy populations, with the prevalence of CH in HHMs an area of active investigation.^{18-21,32-34} We evaluated our cross-sectional cohorts of *RUNX1*, *GATA2*, and *DDX41* carriers-without HM for CH-related variants at the time of sample collection (Table 1; Figure 2A). We identified CH in 35% of *RUNX1* carriers-without HM (23 of 66 individuals; Figure 2A,C) and 22% (2 of 9 individuals, Figure 2A; supplemental Figure 1B) of *GATA2* carriers-without HM, respectively. The prevalence of CH was significantly lower (3%, 1 of 31 individuals, *P* = .002; Figure 2A; supplemental Figure 1C) in *DDX41* carriers-without HM. The reduced prevalence of CH in the

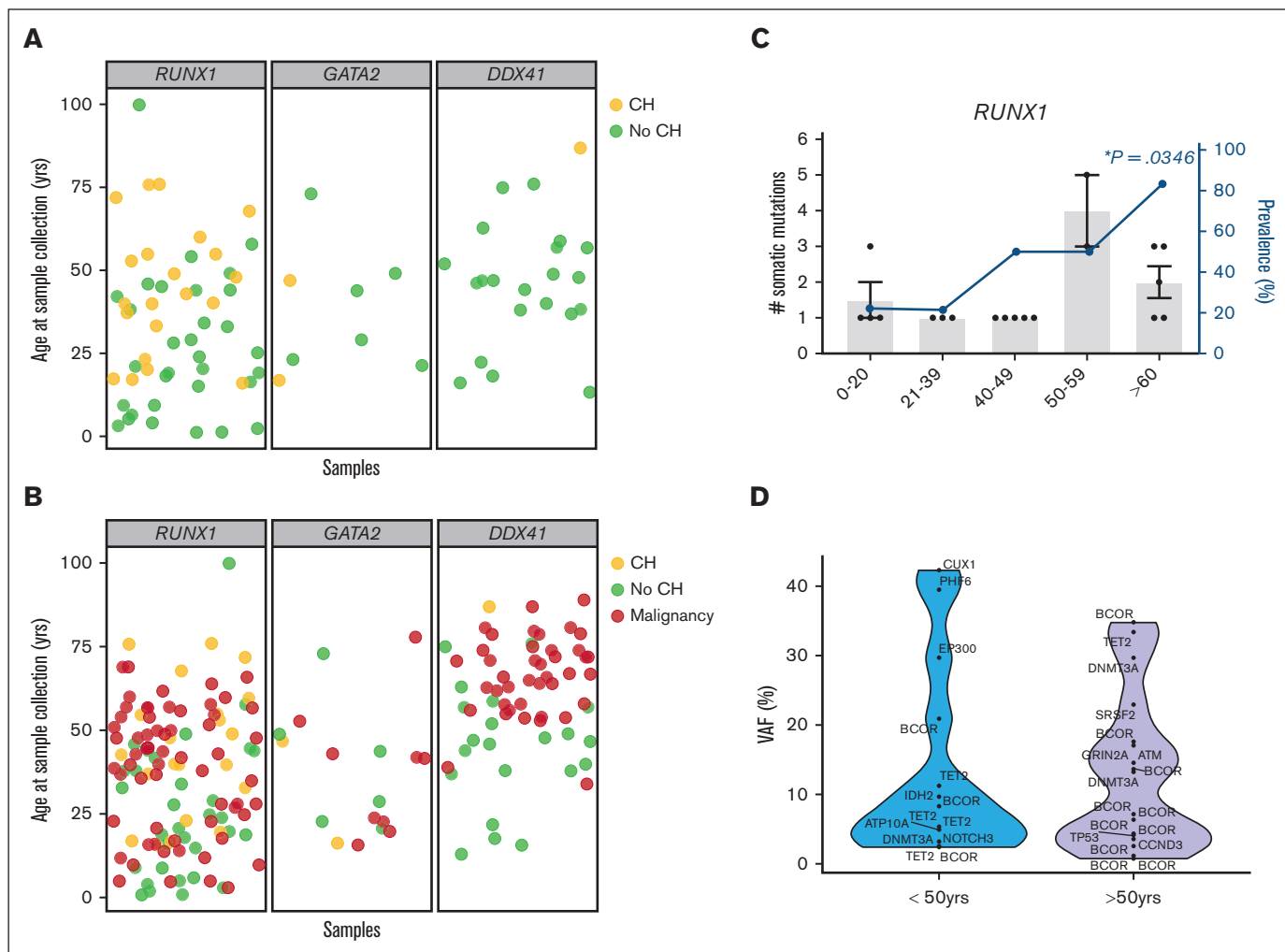


Figure 2. Defining the spectrum of CH in germline *GATA2*, *RUNX1*, and *DDX41* carriers-without HM. (A) Each individual in the carriers-without HM cohort was defined as having CH (yellow) or no identifiable CH (no-CH, green), based on the identification of somatic clinically relevant variants, driver somatic variants. The age of the individual at the time of sample collection is indicated. (B) Correlation of the ages of malignancy development (HM, red) observed in the germ line malignancy cohorts with the carriers-without HM cohort, with (yellow) and without CH (green). (C) Demographics of individuals with CH variants in the *RUNX1* germ line carriers-without HM cohorts. Column graph shows the number of somatic variants identified in individuals with CH. Error bars show the standard error of the mean. Line graphs show the prevalence of CH in the carriers-without HM germ line cohort in different age groups. * $P < .05$, logistic regression model. (D) Violin plots showing the distribution of VAFs of driver somatic variants (shown in panel A) in the germ line *RUNX1* carriers-without HM cohort in individuals under the age of 50 years or >50 years old. VAFs for X-chromosome genes were normalized in male individuals to compensate for ploidy, enabling comparison with autosomal genes. HM, hematologic malignancy.

DDX41 cohort was independent of age, as the age distribution of samples was overlapping between HHM cohorts (supplemental Figure 1A). For germ line *RUNX1*, CH was identified in all age groups, and the prevalence of CH significantly increased with age (Figure 2C, $P = .0267$, logistic regression). In the *RUNX1* cohort, 5 of 6 (83%) individuals 60 years of age and older had at least 1 CH variant (Figure 2C). The number of variants increased with age, as 92% of individuals under the age of 50 years with CH had only 1 CH variant, whereas 71% of patients over the age of 50 years had 2 or more CH variants ($P = .001$, Figure 2C,E). The median VAF of CH variants did not change significantly with age (Figure 2D). For all cohorts, no CH was identified in any individual younger than 16 years ($n = 9$).

CH is increased in *RUNX1* carriers-without HM relative to population controls

We then compared the prevalence of CH in our cohort of *RUNX1* carriers-without HM to population controls from Jaiswal et al and Genovese et al ($n = 27\,783$).^{32,33} The prevalence of CH was higher in *RUNX1* carriers-without HM in every age group (Figure 3A, Z test of proportions, $P < .0001$). The prevalence of CH was 0.2% in controls between the ages of 19 and 29 years but was 22.2% in *RUNX1* carriers-without HM in the same age group. In individuals aged 60 years or older, CH was detectable in 7% of controls and 83% of *RUNX1* carriers-without HM, demonstrating that *RUNX1* carriers-without HM have an increased prevalence of CH at all ages

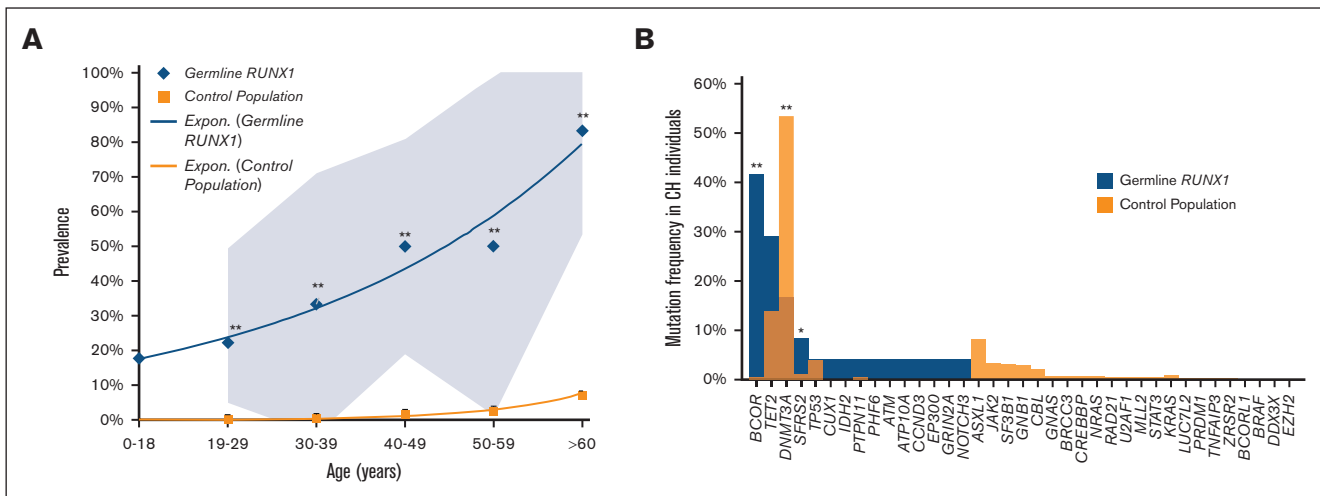


Figure 3. Spectrum of CH in age-related CH compared with the germ line *RUNX1* carriers-without HM cohort. (A) Prevalence of CH in the control population compared with the germ line *RUNX1* carriers-without HM cohort. The control population includes cohorts from Jaiswal et al.³² and Genovese et al.³³ (n = 27 783). The 0 to 18 age group is only available for the *RUNX1* cohort. (B) Mutational spectrum of CH in the control population compared with the germ line *RUNX1* carriers-without HM cohort. Graph shows the frequency of variants in individuals with CH in each cohort. The control population includes cohorts from Desai et al.³⁶ Abelson et al.³⁵ and Jaiswal et al.³² **P* < .05, ***P* < .001, 2-proportions Z test, approximate to normal distribution, ±95% confidence interval.

compared with population controls. We investigated the frequency of variants in prototypical CH genes. Variants in the epigenetic regulators *DNMT3A* (54%), *TET2* (29%), and *ASXL1* (8%) are the most frequent CH-related genes in the general population.^{32,35,36} Surprisingly, the most frequently mutated CH-related gene in *RUNX1* carriers-without HM was *BCOR* (42%), which is mutated in only 0.6% of population controls with CH (Figure 3B, *P* < .0001).^{32,35,36} *DNMT3A* was mutated in 17% (*P* < .0001), *TET2* in 14% (*P* = .2182), and *ASXL1* was not mutated in *RUNX1* carriers-without HM with CH (Figure 3B). These findings demonstrate that the mechanism of CH in *RUNX1* carriers-without HM is distinct for this syndrome as compared with population controls.

Clonal structure and evolution in *RUNX1* carriers-without HM

Further examining the clonal composition of somatic variants in carriers, we postulated the order of mutation acquisition in samples with multiple variants by using relative VAFs (Figure 4A). We observed that *BCOR*, as well as being the most frequently mutated gene, was also present across the entire age spectrum, from carriers as young as 16 years to 76 years, found as a first hit (Figure 4A; Table 1). Consistent with the overall data, there was a general increase in *BCOR* VAF with age and additional mutations, which could be both additional *BCOR* mutations as well as mutations in other genes including *TP53* and *ATM*, with 1 case where *DNMT3A* was antecedent to *BCOR* variants (Figure 4A; Table 1). Three *RUNX1* carriers-without HM had longitudinal peripheral blood samples available, which allowed us to track the temporal evolution of CH (Figure 4B). Case 1: a male with thrombocytopenia and a germ line *RUNX1* p.R169I variant had a somatic *DNMT3A* p.F751fs variant detected at a VAF of 3.1% at 43 years of age. The clone increased to a VAF of 5.0% over 5 years without any clinical-level changes, including leukemogenesis, or the development of additional clones. Case 2: a female with thrombocytopenia and a germ line *RUNX1* p.R320* variant who

developed a *TET2* p.Y1598* somatic variant that persisted for more than 7 years, increasing from a VAF of <1% to 5%, without clinical changes. Case 3: a female with a germ line *RUNX1* c.351+1G>A splicing variant and thrombocytopenia who developed AML 3 years later. We identified 3 somatic variants (*DNMT3A*, *SRSF2*, and *TET2*) in the patient's initial sample, collected at age 53 years. These variants persisted for 2 years with persistent thrombocytopenia but no leukemogenesis. The patient then developed AML with additional somatic *RUNX1* and *STAG2* variants at 56 years of age. The initial *DNMT3A* and *SRSF2* CH-related variants remained stable, whereas the *TET2* variant was outcompeted during leukemogenesis.

Somatic variants in germ line *RUNX1*, *GATA2*, and *DDX41* malignancy samples

We next sought to define the landscape of driver somatic variants in our carriers-with HM cohorts who had developed malignancies. In the *RUNX1* carriers-with HM cohort, at least 1 driver somatic variant was detected in 46 of 52 (88%) individuals diagnosed with an HM. No association between the number of driver somatic variants and the histologic subtype of malignancy was observed (supplemental Figure 2C). Driver somatic variants were identified in 64 unique genes, and 22 genes were mutated in more than 1 individual (Figure 5A; supplemental Figure 3A). Second hits in *RUNX1* were the most frequent somatic mutations, with variants detected in 18 individuals (41% of patients with complete sequencing coverage of *RUNX1* [supplemental Methods]). Three types of somatic *RUNX1* variants were identified: small indels and SNVs (unique from the germ line variant, 72%), copy neutral loss of heterozygosity variants (17%), and trisomy 21 (somatic amplification of the germ line *RUNX1* variant, 11%). Somatic second hits in *RUNX1* included 12 missense variants in the exons coding for the RUNT domain as well as a splice-site variant (c.507_508+1dupAGG, Figure 6A-B). Cytogenetic analyses identified 2 individuals with +21 (VAF > 60%) and 3 individuals

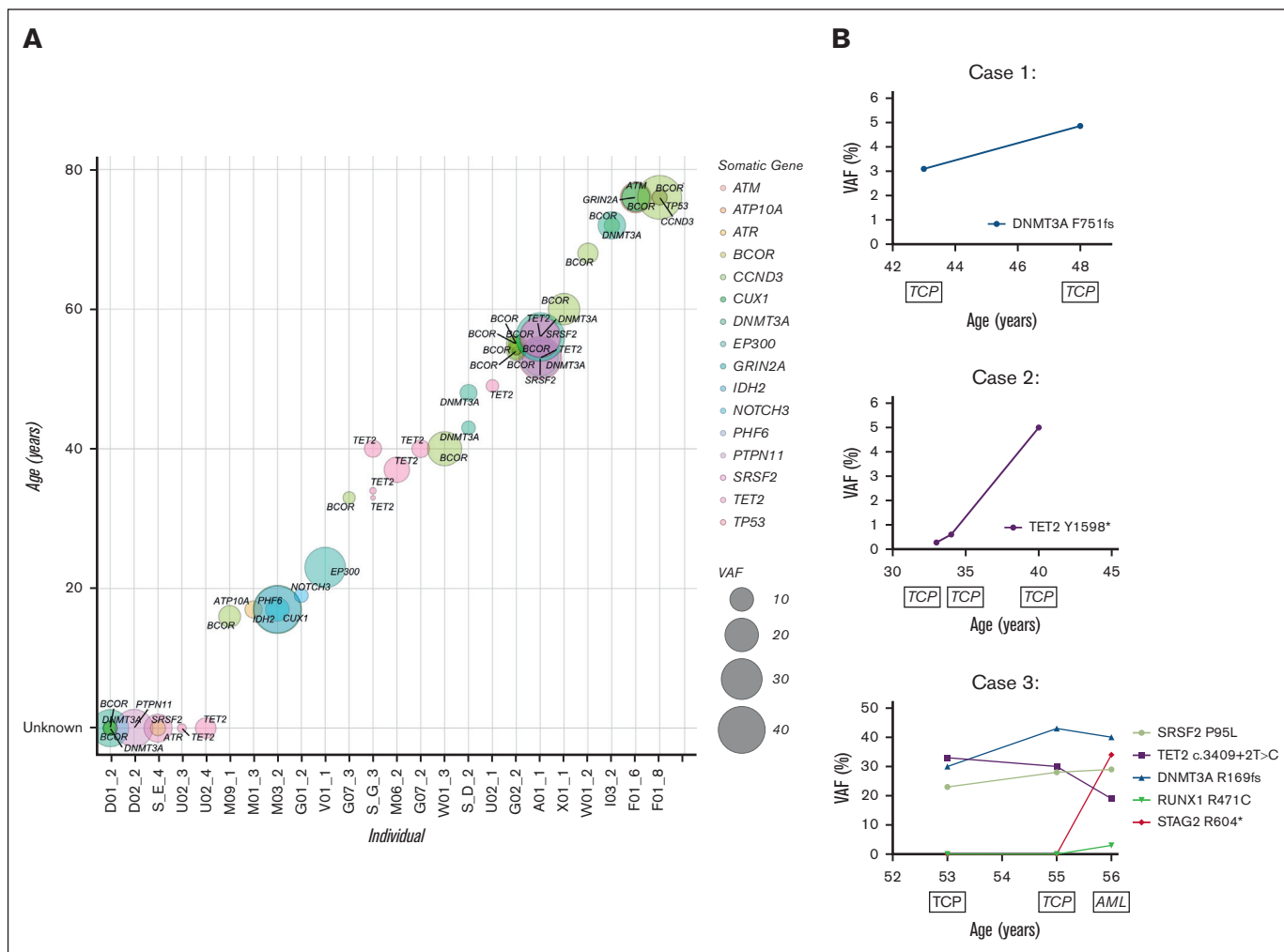


Figure 4. Molecular monitoring of germ line *RUNX1* carriers with CH. (A) Driver somatic variants identified in *RUNX1* carriers-without HM individuals across age. Circle size = increasing VAF, color = gene. Individuals with unknown age were given the value 0 (D01_2, D02_2, S_E_4, U02_3, U02_4). (B) Longitudinal case studies with the VAF of detected driver somatic variants plotted across years. Clinical diagnosis at the time of monitoring is indicated below the age of the individual at which the sample was collected. AML, acute myeloid leukemia; TCP, thrombocytopenia.

with a mutant VAF >80% (copy neutral loss of heterozygosity variants) (supplemental Figure 3A). We did not identify associations between individual germ line and driver somatic variant pairs. Most individuals (78%) with a somatic *RUNX1* variant were female ($P = .02$, Figure 6C). Somatic *RUNX1* variants were identified in all age groups, and no association was established between individual somatic *RUNX1* variants and the age of HM diagnosis (Figure 6D). A female sex bias for HM was observed in all age groups (Figure 6D). Besides second hits in *RUNX1*, a series of established cancer genes were mutated in the HM cohort: *PHF6* (21%), *BCOR* (20%), *TET2* (13%), *SH2B3* (11%), and *SRSF2* (11%) (Figure 5A; supplemental Figure 3A). AML was the predominant malignancy in germ line *RUNX1* variant carriers, with a sex bias for female AML diagnosis (23 of 29 females, 9 of 23 males, $P = .004$, supplemental Figure 4). Among individuals with somatic *RUNX1* variants, 15 (83%) had AML, and 12 of 15 (80%) were females. These data from germ line *RUNX1* variant carriers support a female sex bias for AML leukemogenesis driven by somatic *RUNX1* variants.

No somatic second hits in *GATA2* were detected in our cohort of 13 germ line *GATA2* variant carriers (Figure 5B; supplemental Figure 3B). We detected at least 1 driver somatic variant in 69% (9 of 13) of germ line *GATA2* variant carriers who had developed malignancies. Analysis of the *GATA2* cohort was limited by low sample numbers (supplemental Figure 4A), but the lack of second hits in *GATA2* suggests biallelic variants are not a common leukemogenic mechanism in germ line *GATA2* variant carriers.³⁷

In the *DDX41* carriers-with HM cohort, we identified at least 1 driver somatic variant in 10 unique genes in 18 individuals (62%, Figure 5C; supplemental Figure 3C). Only 3 genes were mutated in more than 1 individual (*DDX41*, *ASXL1*, and *JAK2* p.Val617Phe). The most frequent somatic event was a second hit in *DDX41*, which was observed in 62% ($n = 18$) of individuals with HM. Apart from a single splice-site variant (c.1621+1G>A), all somatic *DDX41* variants were missense variants in the DEAD-box domain (3 of 18) or the recurrent p.R525H variant in the helicase C domain (14 of 18 *DDX41* somatic variants, 78%, Figure 7A-B). We

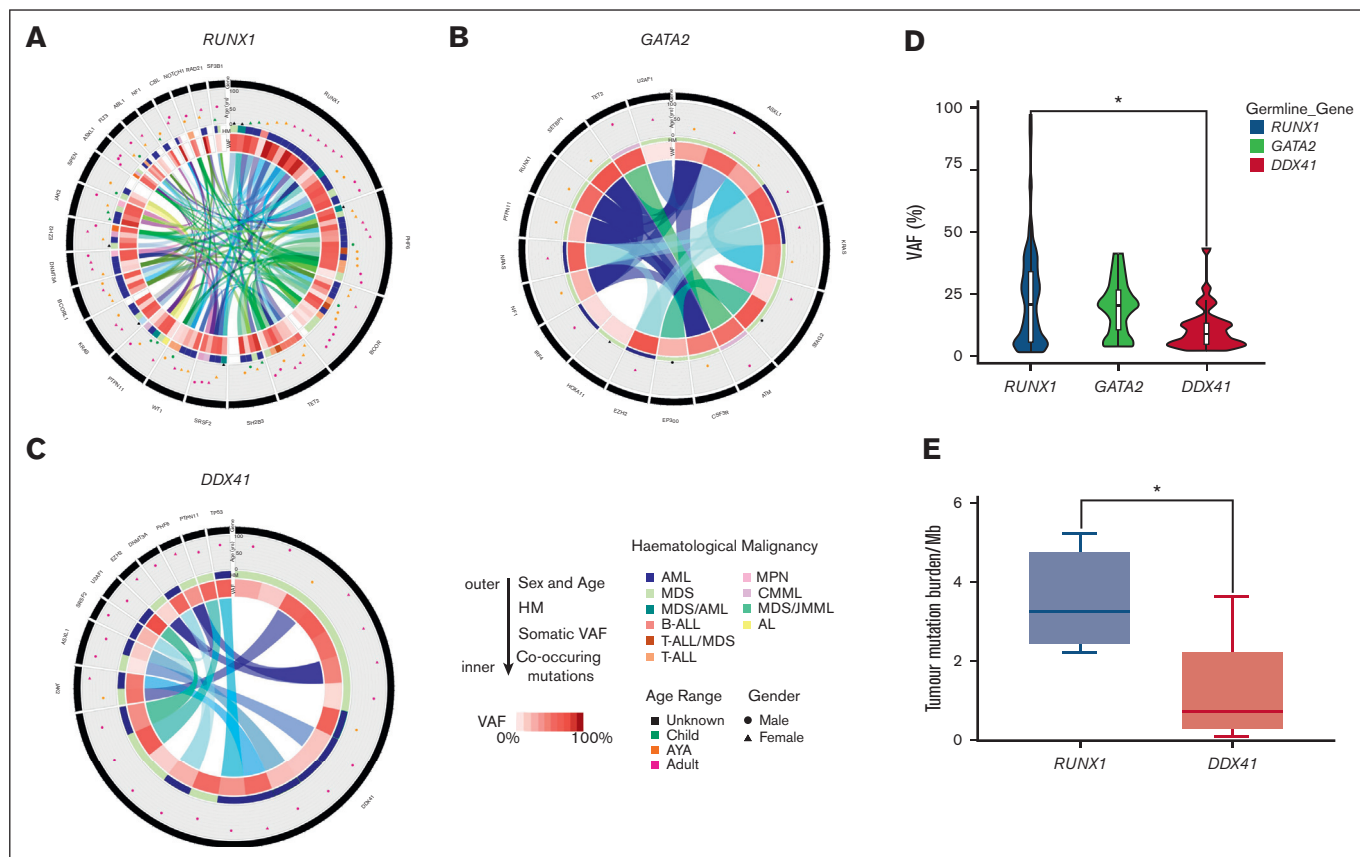


Figure 5. Clinically relevant somatic variants identified in the germ line carriers-with HM cohorts. Distribution of the clinically relevant somatic variants, driver somatic variants, identified in the carriers-with HM cohorts. From outside to inside: 1. Gene with the somatic variant. The length of the black bar indicates the frequency of variant within the germ line carriers-with HM cohort. 2. The age and sex of the individual with the somatic variant. Triangle = female, circle = male. Age groups are indicated by colors (green = child [≤ 14 years], orange = AYA [15-39 years], adult = pink [≥ 40 years], black = the age of the individual is unknown). 3. The type of HM is indicated by the color of the bar. 4. VAF of the somatic variant in the sample as represented on a sliding scale (darker = high VAF, lighter = low VAF). 5. The inner ring depicts the association of different somatic variants in the sample. The colored ribbon depicts a unique sample and the associated somatic variants observed in the sample. (A) Germ line *RUNX1* carriers-with HM cohort. Only shown are the genes that are somatically mutated in 2 or more individuals. (B) Germ line *GATA2* carriers-with HM cohort, showing all driver somatic variants and (C) Germ line *DDX41* carriers-with HM cohort showing all driver somatic variants. (D) Violin plot displaying the distribution of driver somatic variant VAFs observed in the germ line carriers-with HM cohorts. Boxes represent the 25th and 75th percentiles, with the horizontal line in the middle indicating the median, and the vertical lines representing the 95th percentile cohorts. $*P < .05$, 1-way analysis of variance of log-transformed values, with Tukey multiple comparison test. (E) TMB in germ line *RUNX1* and *DDX41* carriers-with HM cohorts. TMB is the number of SNV and INDELS divided by 38Mb coding region. Only malignancy samples where we had available a matched germ line control tissue were used for analysis. Boxes represent the 25th and 75th percentiles, with the horizontal line in the middle indicating the median, and the vertical lines representing the max and min values. $*P < .05$ nonparametric Mann-Whitney *U* test. AYA, adolescents and young adults; AML, acute myeloid leukemia; AL, acute leukemia; B-ALL, B-cell acute lymphoblastic leukemia; CML, chronic myeloid leukemia; CMML, chronic myelomonocytic leukemia; JMML, juvenile myelomonocytic leukemia; MPN, myeloproliferative neoplasms; T-ALL, T-cell acute lymphoblastic leukemia.

observed a significant sex bias for *DDX41* malignancies (3:1 male:female, $P = .0002$), which correlated with males presenting with a somatic *DDX41* variant (14 of 18 males, Figure 7C-D). No association between specific somatic variants and germ line *DDX41* variants, age of malignancy diagnosis, or histologic subtype of malignancy was observed.

Mutational burden in germ line *RUNX1*, *GATA2*, or *DDX41* malignancy samples

To better understand the somatic mutational burden in each syndrome, we evaluated the VAF of all driver somatic variants. A large distribution of VAFs was observed in *RUNX1* carriers-with HM

(median VAF = 22.4%, mean = 27.0%, mode = 5.7, 34%) and *GATA2* carriers-with HM (median = 21.0%, mean = 20.6%, mode = 8, 27.2%). VAFs among *RUNX1* carriers-with HM showed the largest distribution (Figure 5D). The *DDX41* cohort harbored low VAF driver somatic variants (median VAF = 8.9%, mean 13.4%, Figure 5D). No association between age at malignancy diagnosis and the VAF of driver somatic variants was observed in any cohort (supplemental Figure 5). TMB was calculated for *DDX41* ($n = 14$) and *RUNX1* ($n = 4$) carriers-with HM with matched germ line/tumor samples. *DDX41* had a lower TMB (0.75 mutations/Mb) than *RUNX1* malignancies (3.3 mutations/Mb; $P = .01$, Figure 5E).

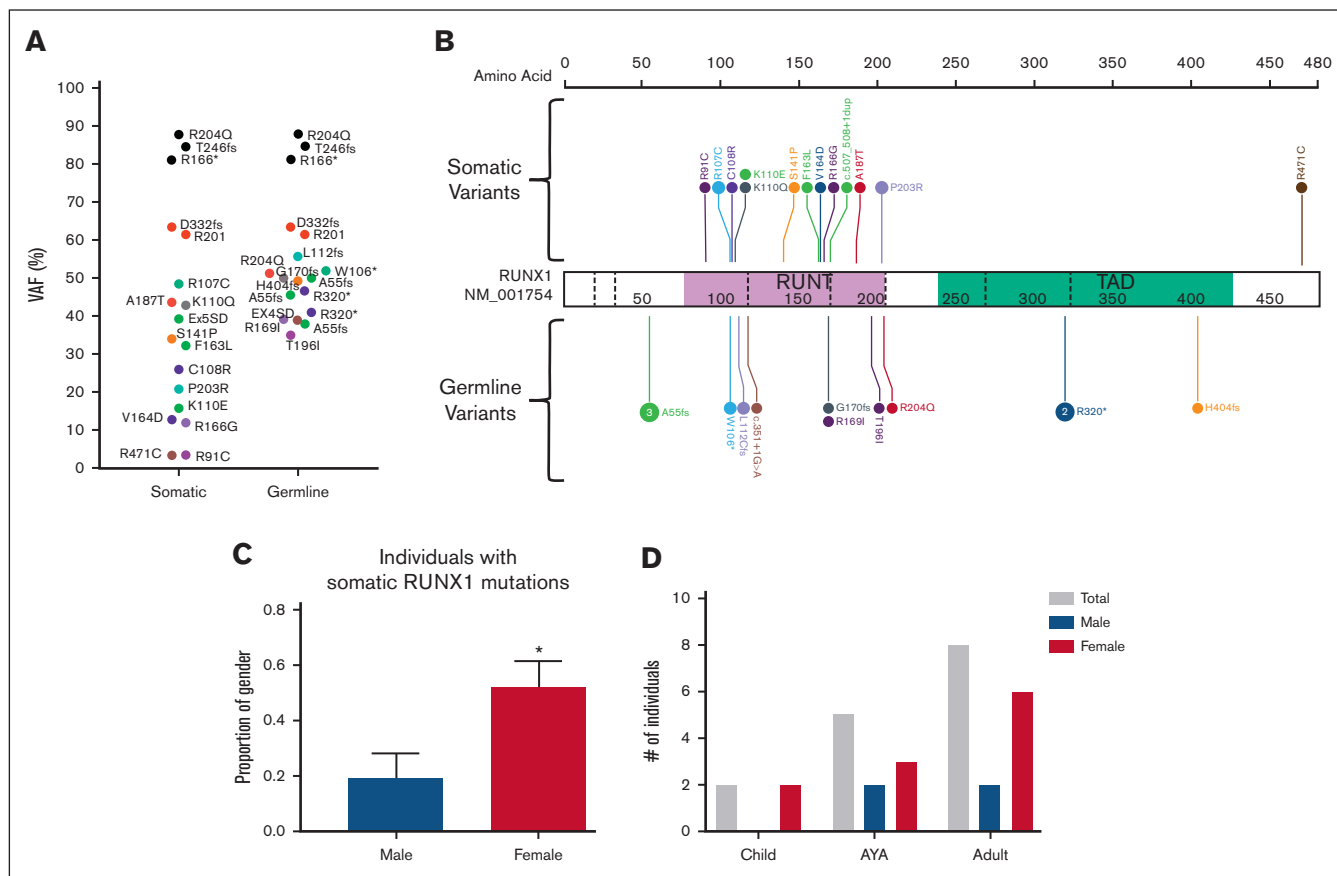


Figure 6. Somatic variants in *RUNX1* are the most common event in the germ line *RUNX1* carriers-with HM cohort. (A) Plot of acquired somatic *RUNX1* variants and associated germ line *RUNX1* variants. Data points are colored according to the somatic and associated germ line variant observed in the patient. VAF of more than 60% indicates a copy neutral loss of heterozygosity (CNLOH) or Trisomy 21. (B) Somatic and germ line *RUNX1* variants are visualized using the ProteinPaint web application.³⁰ Variants are colored according to the somatic and associated germ line variant observed in the patient. The number of probands for each variant is indicated within the circle where the number is greater than 1. All variants are annotated to *RUNX1*; NM_001754.4; LRG_482. (C) The proportion of male and females harboring a somatic *RUNX1* variant is significantly different. * $P < .05$. (D) Sex and age distribution of individuals with a somatic *RUNX1* variant; adult ≥ 40 years, AYA = 15 to 39 years, children ≤ 14 years. Data points are colored according to the somatic and associated germ line variant observed in the patient. AYA, adolescents and young adults.

Discussion

The prevalence of HHMs is estimated to range from 7% to 14% in cohorts of patients with myeloid malignancies.^{38,39} Although the clinical recognition of these syndromes has improved since *RUNX1*-driven HHMs were first described,⁸ questions remain regarding the optimal approach to monitoring carriers-without HM and how malignancy-directed treatments may be individualized for affected patients. Currently it is challenging for clinicians to provide tailored risk-assessment to patients as the natural history of carriers-without HM is not well understood and there has been no approach to identify HHM individuals at highest risk for leukemogenesis. To address this gap, we have leveraged our HHM international collaborative network and assembled and characterized the most extensive cross-sectional comparative cohort of carriers-without HM and carriers-with HM germ line *RUNX1*, *GATA2*, or *DDX41* variants ($n = 191$, 102 probands, Figure 1). We demonstrate *RUNX1*, *GATA2*, and *DDX41* germ line variant carriers experience highly variable risk for CH and unique somatic drivers during CH relative to population controls. Each HHM is remarkable

for mutational profiles during frank leukemogenesis that are also unique to each HHM syndrome.

The most significant risk factor for CH in the general population is aging, with $\sim 10\%$ of individuals over the age of 70 years having detectable CH.^{32,33} Several studies investigating CH in the background of inherited bone marrow failure have shown an increased risk for CH.^{34,40,41} Interestingly, individuals without HM with HHM germ line variants have been shown to have variable risk for CH in a series of small studies (*ANKRD26*, *ETV6*, *RUNX1*).^{18,19,42,43} We have now performed the largest collective analysis of CH in *RUNX1*, *GATA2*, and *DDX41* carriers without HM to date. This analysis extends studies of CH in the HHMs to novel phenotypes (*DDX41*) and suggests that HHM predisposition in *GATA2*, and *RUNX1* carriers-without HM, may be driven by early-onset CH (22.2% in *RUNX1* and 25% in *GATA2*). Recently, larger cohorts of patients with germ line *GATA2* without HM,^{20,21} have also shown CH is common in patients without HM, with CH associated with a hypocellular marrow. Further investigation is required to determine if CH also correlates with cytopenias in germ line *RUNX1* cohorts. In

longitudinal, lineage tracing, and single-cell sequencing studies are required to determine if these are initiating events in malignancy development and the timeline to disease progression.

Interestingly, both germ line *RUNX1* and *DDX41* cohorts presented with a sex bias for HM development, but this did not correlate with differences in X-linked somatic variants. Sex bias was not observed in our *GATA2* cohort, as we have also observed previously.¹⁴ *RUNX1* genomic alterations have a high correlation with hormone-related cancers, especially cancers common in female patients, and with estrogen known to play a role in hematopoiesis,^{48,49} we hypothesize that disruption of specific estrogen signaling pathways in germ line *RUNX1* carriers could predispose females to AML.⁵⁰⁻⁵³ In the germ line *RUNX1* malignancy cohort, recurrent somatic gene variants are involved in epigenetic regulation and epigenetic dysregulation and can occur in leukemogenesis, with sex-specific differences in methylation observed in hematopoietic tissue.^{50,54,55} The innate immune response is also known to be increased in females relative to males.⁵⁶ For *DDX41*, given its role as an intracellular pattern recognition receptor that triggers the innate immune response,⁵⁷ a dysregulated immune response could exaggerate existing differences in innate immunity between males and females, contributing to the observed sex bias in malignancy penetrance. Further investigation is warranted to understand the interplay of these mechanisms on tumorigenesis, which may ultimately inform the development of sex-specific therapies that optimize outcomes for patients with HHM.

Despite a lack of definitive guidelines, limitations, and ongoing debate, molecular monitoring in clinical practice is becoming more widespread.⁴³ Findings from this study have implications for clinical surveillance and counseling for different patients with HHM. For example, in *RUNX1* and *GATA2* HHMs, regular targeted sequencing of CH genes, even in younger carriers-without HM, will provide a tool to monitor the evolution of the clonal burden associated with these variants. In contrast, given the low VAF and high frequency of somatic *DDX41* variants in *DDX41* HHMs, serial high-depth sequencing of *DDX41* for the common R525H mutation may be a preferred approach in *DDX41* carriers-without HM. Although in the aging population, CH is a risk factor for leukemic transformation, the presence of CH in inherited bone marrow failure is in some situations associated with somatic rescue or normalization of HSPC fitness. Therefore, it is important to discriminate CH events that are associated with risk for leukemic transformation from CH, which results in normalization of function.⁵⁸ Given that CH variants feasibly confer a step toward HM,³⁶ the high frequency of *BCOR* and *TET2* variants in our cohort of *RUNX1* HHM malignancies and their presence in *RUNX1* carriers-without HM, at least in the research study setting, warrant monitoring of these genes as potential molecular biomarkers of leukemogenesis. Changes in CH trajectory may eventually inform clinical decision-making, such as the timing of repeat bone marrow biopsies. These decisions will be made in conjunction with more classic clinical tools, such as the monitoring of peripheral blood cell counts.⁴³

This study highlights the immense benefit of international collaboration and data sharing within the HHM and rare disease communities. We have established the framework for the continued accumulation of patient data, including longitudinal molecular

monitoring, which is required to define the different risk states associated with leukemogenesis across these disorders. With continued progress, this work may lead to the establishment of a defined molecular risk stratification for leukemia progression in carriers and, with it, the ability to design and test in trials interventions to halt progression to full-blown HM in vulnerable HHM-variant carriers. For instance, with regular clinical surveillance, it may be possible to detect individuals who develop second hits in *DDX41* or *RUNX1* before a clinical diagnosis of HM. These individuals may benefit from intensive clinical surveillance or low-toxicity prophylactic therapies. In contrast, defining *TET2*, *BCOR*, or other epigenetic regulators as emerging vulnerabilities opens an avenue for the development of prophylactic treatments for HHM carriers via TET inhibitors, histone deacetylases (HDAC) inhibitors, hypomethylating agents, and combinatorial therapies that do not carry the morbidity and mortality of stem cell transplant. This study provides the most comprehensive investigation of leukemogenic molecular mechanisms in HHMs to date, informing the next generation of studies into the clinical management and surveillance of these disorders as well as potential insights into personalized and preemptive therapies for carriers.

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Authorship

Contribution: C.C.H., M.W.D., and K.Y. were involved in all aspects of the project including designing the research, manuscript preparation, collecting next-generation sequencing (NGS) and clinical data, American College of Medical Genetics and Genomics (ACMG)-variant classification, somatic-variant analysis, and curation and analysis of the data; A.L.B., L.A.G., P.L., and H.S.S. designed the research, contributed NGS and clinical data, manuscript preparation, and ACMG-variant classification; J.F., L.A.-M., M.J.P., K.E.M., T.H., M.A., P.W., A.W.S., E.K., and R. Sood designed bioinformatic pipelines and analysis; D.M.L. designed VariantGrid software used for somatic and germ line variant curation (VariantGrid); P.V., P.A., S.L.K.-S., J.C., and C.N.H. curated somatic and germ line variant data; B.P. advised on statistical analysis; C.B., A.B.C., M.C., E.D., C.D.D., N.D., R.F., S.F., A.R.-M., B.P., J.M.K., A.K., M.K., J.L., N.V.M., G.N., C.O., K.P.P., C.P., H. Raslova, H. Rienhoff, T.R., R. Susman, K.T., E.V., E.K., R. Schulte, A.P.H., S.M.H., K.P., N.K.P., M.B., A.H.W., C.F., H.M.F., I.D.L., J.C., R. Sood, L.C.F., P.B., D.S., D.H., B.Y., L.M., A.L.B., and C.N.H. contributed NGS data, clinical patient information, and scientific insight; and all authors critically reviewed and approved the manuscript.

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