

MYELOID NEOPLASIA

Germ line *ERG* haploinsufficiency defines a new syndrome with cytopenia and hematological malignancy predisposition

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KEY POINTS

- Germ line *ERG* LOF variants predispose to cytopenias and HMs.
- Somatic genetic rescue of *ERG* pathogenic variants in hematopoietic tissues impacts diagnosis, disease severity, and potential for correction.

The genomics era has facilitated the discovery of new genes that predispose individuals to bone marrow failure (BMF) and hematological malignancy (HM). We report the discovery of ETS-related gene (*ERG*), a novel, autosomal dominant BMF/HM predisposition gene. *ERG* is a highly constrained transcription factor that is critical for definitive hematopoiesis, stem cell function, and platelet maintenance. *ERG* colocalizes with other transcription factors, including RUNX family transcription factor 1 (*RUNX1*) and GATA binding protein 2 (*GATA2*), on promoters or enhancers of genes that orchestrate hematopoiesis. We identified a rare heterozygous *ERG* missense variant in 3 individuals with thrombocytopenia from 1 family and 14 additional *ERG* variants in unrelated individuals with BMF/HM, including 2 de novo cases and 3 truncating variants. Phenotypes associated with pathogenic germ line *ERG* variants included cytopenias (thrombocytopenia, neutropenia, and pancytopenia) and HMs (acute myeloid leukemia, myelodysplastic syndrome, and acute

lymphoblastic leukemia) with onset before 40 years. Twenty *ERG* variants (19 missense and 1 truncating), including 3 missense population variants, were functionally characterized. Thirteen potentially pathogenic erythroblast transformation specific (ETS) domain missense variants displayed loss-of-function (LOF) characteristics, thereby disrupting transcriptional transactivation, DNA binding, and/or nuclear localization. Selected variants overexpressed in mouse fetal liver cells failed to drive myeloid differentiation and cytokine-independent growth in culture and to promote acute erythroleukemia when transplanted into mice, concordant with these being LOF variants. Four individuals

displayed somatic genetic rescue by copy neutral loss of heterozygosity. Identification of predisposing germ line *ERG* variants has clinical implications for patient and family diagnoses, counseling, surveillance, and treatment strategies, including selection of bone marrow donors and cell or gene therapy.

Introduction

Tightly controlled regulation of hematopoiesis is essential to ensure adequate supply of healthy blood cells and the ability to respond to increased demand. Transcription factors (TFs), including ETS-related gene (*ERG*), TAL bHLH transcription factor 1, erythroid differentiation factor (*TAL1*), LYL1 basic helix-loop-helix family member (*LYL1*), LIM domain only 2 (*LMO2*), GATA binding protein 2 (*GATA2*), RUNX family transcription factor 1 (*RUNX1*), Meis homeobox 1 (*MEIS1*), Spi-1 proto-oncogene (*SPI1/PU.1*), Fli-1 proto-oncogene, ETS transcription factor (*FLI1*), and growth factor independent 1B transcriptional repressor (*GFI1B*), form part of a network that, in varying combinations, control the regulation of hematopoietic stem or progenitor cells.^{1,2} This TF network is important for regulating cellular self-renewal, lineage specification, differentiation, and migration. Dysregulation can lead to cellular malfunction, impaired differentiation programs, and aberrant stem cell self-renewal, all of which have implications for human disease. Highlighting this, recurrent somatic variants within several of these TFs, including chromosomal rearrangements, point mutations, and insertions and deletions, are detected in hematological malignancies (HM). Furthermore, germ line pathogenic variants in *RUNX1* (familial platelet disorder with predisposition to myeloid malignancies, *RUNX1-FPD*; Monarch Disease Ontology [MonDO]: 0011071), *GATA2* (*GATA2* deficiency with susceptibility to myelodysplastic syndrome [MDS]/acute myeloid leukemia [AML]; MonDO: 0042982), and *FLI1* (bleeding disorder, platelet-type, 21; Online Mendelian Inheritance in Man: 617443) are associated with the development of hematological disorders typified by an increased risk for cytopenias and/or HMs. The roles of other members of this key hematopoietic TF network in predisposing to HMs remain to be identified.³

ERG, a member of the ETS TF family, was first reported to be critical for normal hematopoiesis in 2008.⁴ Like most ETS TFs, *ERG* demonstrates various homeostatic functions by binding to specific GGA(A/T) motifs to regulate genes in hematopoietic and nonhematopoietic contexts. This includes binding at gene regulatory sites with other key hematopoietic transcriptional regulators, including with products of the HM predisposition genes *GATA2* and *RUNX1*.⁵ In normal hematopoiesis, *ERG* is essential for maintaining quiescence and preventing differentiation of hematopoietic stem cells (HSCs),⁶ thereby promoting HSC self-renewal after hematopoietic stress (eg, bone marrow [BM] transplantation)⁷ and supporting definitive hematopoiesis, adult HSC function, and the maintenance of peripheral blood (PB) platelet numbers.⁴

The perturbation of hematopoiesis by aberrant *ERG* expression and the contribution of *ERG* overexpression to HMs have been well documented,⁸⁻¹⁸ particularly in AML.¹⁹⁻²² The consequences of dysregulated *ERG* are also evident in other diseases, including cardiovascular disease,²³⁻²⁶ prostate cancers,^{27,28} Ewing's sarcoma,²⁹ and B-cell acute lymphoid leukemia (B-ALL),³⁰ the latter 3 via chromosomal translocations and genomic rearrangements.

The first report of disease owing to a germ line *ERG* variant was heterozygous *Erg* (S322P) that causes thrombocytopenia in a mouse model.⁴ More recently, germ line loss-of-function (LOF) *ERG* variants have been associated with a predisposition to primary lymphedema.³¹

To our knowledge, we report for the first time germ line *ERG* variants in patients with a range of malignant and nonmalignant hematological phenotypes. This discovery marks the identification of the third ETS TF with autosomal dominant pathogenic germ line variants and HM and/or bone marrow failure (BMF) (in addition to ETS variant transcription factor 6 [*ETV6*]³² and *FLI1*^{33,34}) and adds to the growing list of master hematopoietic TFs already included in germ line-targeted sequencing panels for HM predisposition (*RUNX1*, *CEBPA*, *GATA2*, *ETV6*, *MECOM*, *PAX5*, and *IKZF1*).³⁵ Our findings suggest a pathogenic role for *ERG* haploinsufficiency or hypomorphic actions in contrast with the prototypical oncogenic nature of *ERG*, thereby defining *ERG* deficiency syndrome as a new disease entity.⁹ This paradox implies that the consequence of dysregulated *ERG* may not be consistent across all spatial and temporal cellular contexts, a phenomenon that is not uncommon in hereditary HM predisposition (eg, *GATA2*).³⁶ It also implies a strict expression threshold at which point dysregulation of *ERG* may upset the stoichiometry within a TF complex, leading to predisposition for and/or initiation of HM-related disease. We systematically examined the functional implications of *ERG* variants on DNA binding, subcellular localization, and transactivation of gene expression and focused on specific variants to demonstrate their effect on *ERG*-mediated myeloid differentiation and cytokine independence *ex vivo* and their impact on *ERG*-driven leukemia in an *in vivo* murine model.

Methods

Human and animal ethics

Samples were obtained from the Australian Familial Haematological Conditions Study, which was approved by the Women's and Children's Health Network Human Research Ethics Committee, Adelaide, Australia (approval 2020/HRE00981). All other patient samples and data were covered by local institutional human research ethics committees. Mouse experiments were done with approval from the Hudson Animal Ethics Committee in conjunction with the Monash University Animal Research Platform and the Monash Health Translation Precinct Animal Facility.

Gene discovery

Genomic DNA was extracted (QIAamp DNA Mini Kit, Qiagen), exonic sequences captured using xGen (Integrated DNA Technologies), and libraries sequenced using NextSeq 550 (Illumina) to an average depth of 50× (hair) and 100× (blood). In addition, polymerase chain reaction-free, short-read whole genome sequencing was performed at the Australian Genome Research

Facility (Melbourne, Australia). Variant calling was performed using GATK (v3) (details in the supplemental Methods).

Identification of additional *ERG* variants in hematological cohorts

To identify additional kindreds with blood phenotypes and rare *ERG* variants, we contacted collaborators with existing genomic data obtained through routine clinical testing and research studies. GeneMatcher³⁷ was used to further expand our patient cohort. All patient samples and data were covered by the local institutional human research ethics committees in accordance with the Declaration of Helsinki.

Luciferase assay

An integrin, alpha-2b (*ITGA2B*), in a pGL4.10-Luc vector was kindly donated by Marie-Christine Kopp (University of Sydney).³³ K562 cells were seeded, transfected (Lipofectamine 2000), and then lysed after 20 hours (Dual-Luciferase Reporter kit, Promega). pUC18 was used to normalize the amount of transfected DNA. Luciferase levels were measured using the Explorer Multimode Microplate Reader (Promega).

Immunofluorescence staining

COS-7 cells were transiently transfected with pcDNA3.1-*ERG* (Myc-tag) wild-type (WT) or variant expression vector, fixed after 20 hours (4% paraformaldehyde), and probed with anti-Myc antibody (9B11; New England Biolabs) and Alexa Fluor 488 Rabbit anti-mouse (A27023) secondary antibody. Fluorescent cells (100) were classified as nuclear (protein only in nucleus) or cytoplasmic (protein in nucleus and cytoplasm).

EMSA

Human embryonic kidney 293 (HEK293) cells were seeded, transiently transfected (Lipofectamine 3000), and lysates were prepared (RIPA 9806S, Cell Signaling Technology). Both biotin-labeled and unlabeled double-stranded DNA oligonucleotides containing an *ERG* binding site were synthesized (5' biotin-GGCACTCACTTCGGGCTTGCCGTCGA-3'). Electrophoretic mobility shift assays (EMSAs) were performed using the LightShift Chemiluminescent EMSA kit (ThermoFisher) with 20 pmol probe.

ERG overexpression in FLCs

Fetal liver cells (FLCs) harvested from embryonic day 14.5 WT C57BL/6 mouse embryos were transduced (supplemental Methods, available on the *Blood* website) with MSCV-IRES-mCherry-based retrovirus containing either the *ERG* WT or *ERG* variants (P116R, M219I, D345N R370P, Y372*, and Y373C).⁸ Transduced cells were cultured in StemSpan (STEMCELL Technologies) supplemented with interleukin-3 (IL-3) (10 ng/mL), IL-6 (10 ng/mL), Flt3L (50 ng/mL), SCF (50 ng/mL), and TPO (50 ng/mL). Fluorescence-activated cell sorter analysis was performed weekly for 4 to 6 weeks. For cytokine independence assays, all cytokines were removed when cell populations reached 80% to 90% mCherry⁺. Cell viability and survival were measured every 3 days for 12 days.

In vivo leukemia model driven by *ERG* overexpression

MSCV-*ERG*-IRES-mCherry retrovirus-transduced FLCs (method as per “*ERG* overexpression in FLCs”) were cultured for 3 days

before IV injection into sublethally irradiated 8-week-old C57BL/6 mice. The mice received neomycin water for 3 weeks after irradiation, and their blood was monitored every 2 weeks for mCherry expression.

Results

Identification of BMF and/or HM families and individuals with rare *ERG* variants

Family 1 presented with a range of hematological abnormalities that included thrombocytopenia, neutropenia, and AML (Figure 1A-B; supplemental Figure 1). Patient I-2 developed AML at 27 years of age. The patient entered morphologic remission, subsequently developed therapy-related MDS, and died at the age of 36 years (supplemental Figure 1). A gene panel analysis (supplemental Table 1) was conducted for all 3 affected family members and showed no germ line pathogenic variants in known BMF/HM predisposition genes. Whole exome sequencing analysis on the unaffected father confirmed the absence of any germ line variants that may explain the phenotypes seen in both children. Cytogenetic analysis identified a constitutional mosaic trisomy 8 in patient I-2 (supplemental Table 2), which was not present in either of her children and may have contributed to the myeloid malignancy progression. A pathogenic somatic *RUNX1* (G165V) variant was identified in individual II-1 at a low variant allele frequency (VAF) (2%) that may be an early indicator or marker of clonal progression to malignancy. Platelet morphology studies for individual II-1 identified minor platelet abnormalities, including a slight increase in alpha granule numbers, a slight dilation of the open canalicular system, and mildly enlarged platelets, despite no noticeable mean platelet volume abnormalities (supplemental Figure 2A-B).

Whole exome sequencing analysis of all 4 individuals revealed heterozygosity of a novel *ERG* variant (chr21q.38383725T>C (hg38); c.1118A>G; p.Y373C) in the 3 affected individuals in that a highly conserved amino acid was altered, and it segregated with thrombocytopenia (Table 1; supplemental Figure 3A-B). This variant was absent in the general population (Genome Aggregation Database [gnomAD])³⁸ and was predicted to disrupt DNA binding (supplemental Figure 3C). Interestingly, single-nucleotide polymorphism arrays identified copy neutral loss of heterozygosity (cnLOH) events on chromosome 21q that favored the *ERG* WT copy in all 3 carriers, strongly suggesting somatic genetic rescue (SGR) of the germ line deleterious *ERG* variant (c.1118A>G; VAF 30% [I-2], 17% [II-1], and 44% [II-2]; Figure 1C; supplemental Figure 3D). Intriguingly, cnLOH encompassed the entire *RUNX1* gene in 2 patients (I-2 and II-2), and the recombination breakpoint in the most telomeric event in individual II-1 was within the *RUNX1* gene (Figure 1C). With the addition of whole genome sequencing, we confirmed the presence of the cnLOH events, and the absence of deleterious variants in *RUNX1*, including noncoding and structural variations, that might explain the phenotype (supplemental Figure 4). Note, cnLOH favoring the WT is not a commonly described mechanism of somatic reversion in *RUNX1*-FPD.³⁹

Through international collaborations and GeneMatcher³⁷ (genotype matching resource), we identified an additional 13 rare *ERG* heterozygous variants associated with BMF- or HM-related disease and 2 associated with lymphedema that were

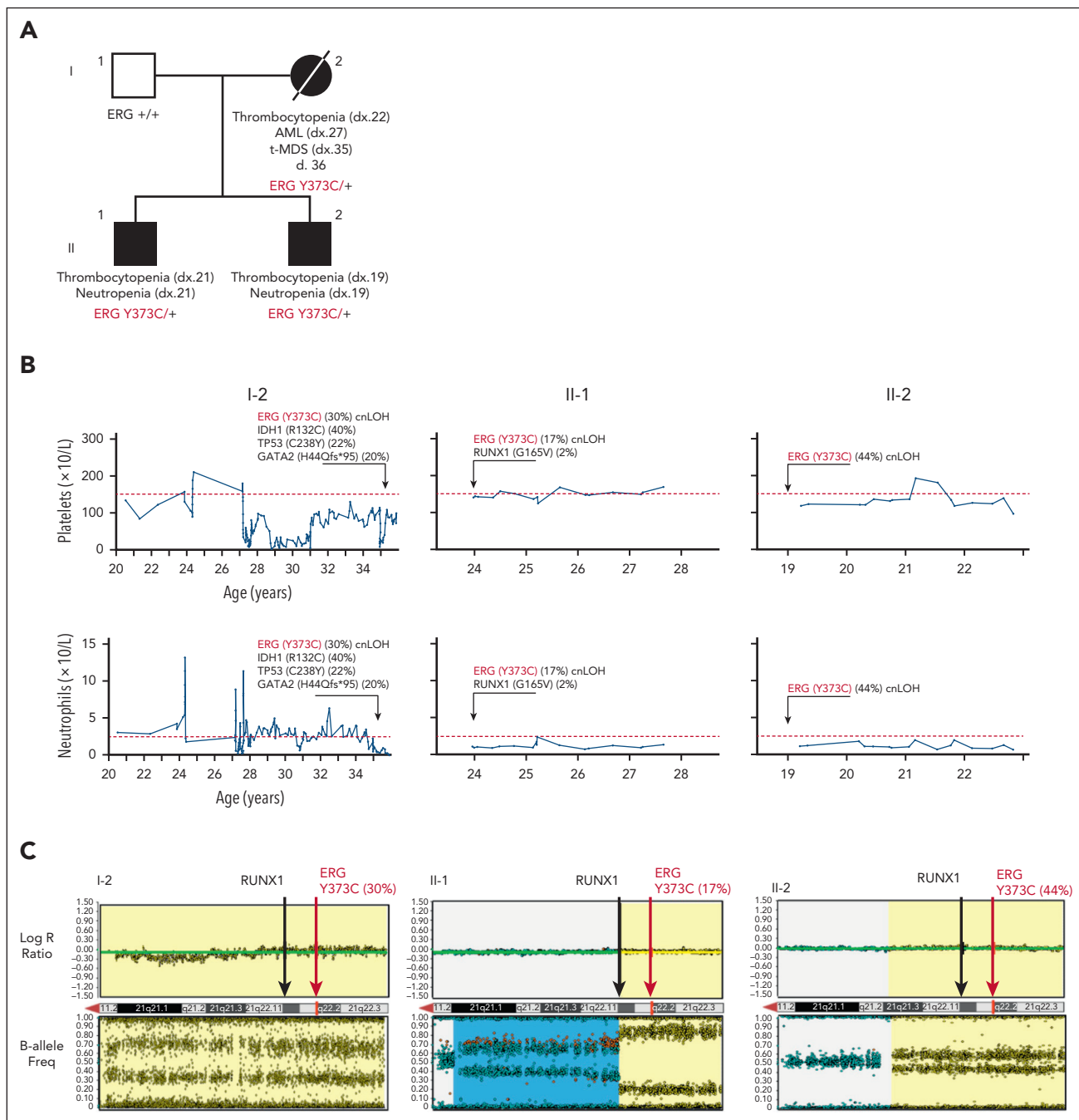


Figure 1. Germ line ERG variant identified in a family with hematological conditions. (A) Pedigree of family 1 containing the ERG (Y373C) variant that segregates with thrombocytopenia and AML. Age of onset (years). (B) History of platelet and neutrophil counts of affected family members (patient 15, 16, and 17) (I-1, II-1, and II-2). Absolute neutrophil or platelet counts per microliter of blood from complete blood examinations plotted with age (years). Normal lower limit for neutrophils and platelets is marked with a dotted red line. (C) Log R ratio and B-allele plots of single-nucleotide polymorphism–array analysis. All individuals show a cnLOH event that encompasses the entire ERG gene (yellow highlight). Individual II-1 (patient 16) had a second cnLOH event (blue highlight). Log R ratios (top panels) show no loss of copy number, and B-allele frequencies (bottom panels) show regions of LOH, together demonstrating cnLOH. Samples used: patient I-2, BM cytogenetic pellet; II-1 and II-2, PB mononuclear cells. d, died; dx, diagnosed.

identified in a primary lymphoedema cohort in the 100 000 Genomes Project.⁴⁰ This included 15 probands; 12 were confirmed germ line variants, including 2 de novo (Table 1; supplemental Table 2; supplemental Figure 5), and germ line samples were not available for the remaining 3 variants. Three variants were predicted to cause premature protein termination and 10 were missense variants that clustered within the ETS domain (310-395 aa). For 4 missense variants, the Rare Exome

Variant Ensemble Learner (REVEL) scores indicated pathogenicity (score >0.85; Table 1; supplemental Figure 6). To determine the prevalence of rare ERG variants in different study cohorts, we tabulated the number of patients screened, phenotypes encompassed, and number of ERG variants found, including multiple cohorts without rare ERG variants (supplemental Table 3). Because of the wide range of phenotypes covered by different research cohorts, the associated

Table 1. Rare heterozygous ERG variants and population variants

Patient no.	Sex	ERG variant (479 aa)	VAF (%)	gnomAD v4.0.0	REVEL	Hematological-related phenotype	Nonhematological phenotype	Age onset of first phenotype (y)	Germ line ERG (inherited/de novo) (Sample)	Somatic variants	Chr 21q cnLOH
1	M	E20Vfs*13	60	0	NA	MDS		38	Yes (MSC)		No
2	F	P116R	56	69 0.004%	0.36	Thrombocytopenia, thrombocytopathy, and platelet aggregation disorder	Hypertension, diabetes, cataract	73			No
3		V127Efs*82	50	0	NA	ALL		<18	Yes		
4		I126T	59	1 0.00016%	0.512	Chronic thrombocytopenia AML	Not reported in clinical records	<14 40	Yes (SF)	FLT3-ITD and IDH1	
5	M	R302C	40	10 0.00068% (1 × R302L) (17 × R302H)	0.365	CLL	Not reported in clinical records	Unknown	Yes (hair)		
6	F	P306L	44	0 (6 × P306W)	0.394	None	Lymphedema	<1	Yes (inherited) (PB)		No
7	M	P306L	57	0 (6 × P306W)	0.394	None	Lymphedema		Yes		
8	F	M341V	45	0	0.45	Severe congenital aplasia and abnormal B cells	Prematurity for acute fetal distress (33 wk)	0	Yes (inherited) (SF)	None	
9	F	M341V	45	0	0.45	MDS		56			No
10		D345N	24	1 0.00012%	0.3149	MDS					No
11		D363A	34	0	0.6209	MDS					No
12	M	R370H	45	0	0.881	Neutropenia Pancytopenia		0 18	Yes (SF)	None (blood at 18 y)	
13	M	R370P	44	0	0.888	MDS (asymptomatic thrombocytopenia and leukopenia)	Deformation (avascular necrosis) of femoral head, severe aortic valve insufficiency with secondary heart failure	29	Yes (hair)	None	No
14	M	Y372*	48	0	NA	Congenital pancytopenia, and BMF		0	Yes (de novo) (PB)		
15	F	Y373C	30	0	0.852	AML, thrombocytopenia, and t-MDS (RAEB2)	Not reported in clinical records	27	Yes (hair)	IDH1 (R132C) (40%) TP53 (C238Y) (22%) GATA2 (H442Qfs*95) (20%)	Yes

Rare ERG variants (NP_891548.1) from similar phenotypic groups, including BMF and/or HM and lymphedema, among population variants (gnomAD >200),³⁸ a germ line thrombocytopenic mouse mutation,¹ a paralogous *ETV6* pathogenic variant (thrombocytopenia),⁴⁹ and COSMIC mutation (somatic).⁴¹ Information unavailable (blank); gnomAD v4.0.0, alternative variants at the same amino acid position (brackets).

Chr, chromosome; CLL, chronic lymphocytic leukemia; COSMIC, Catalogue Of Somatic Mutations In Cancer; DLBCL, diffuse large B-cell lymphoma; F, female; hom, homozygous; M, male; NA, not applicable; MSC, mesenchymal stromal cells; PB, peripheral blood; REVEL, Rare Exome Variant Ensemble Learner; SF, skin fibroblasts; t-MDS, therapy-related myelodysplastic syndrome.

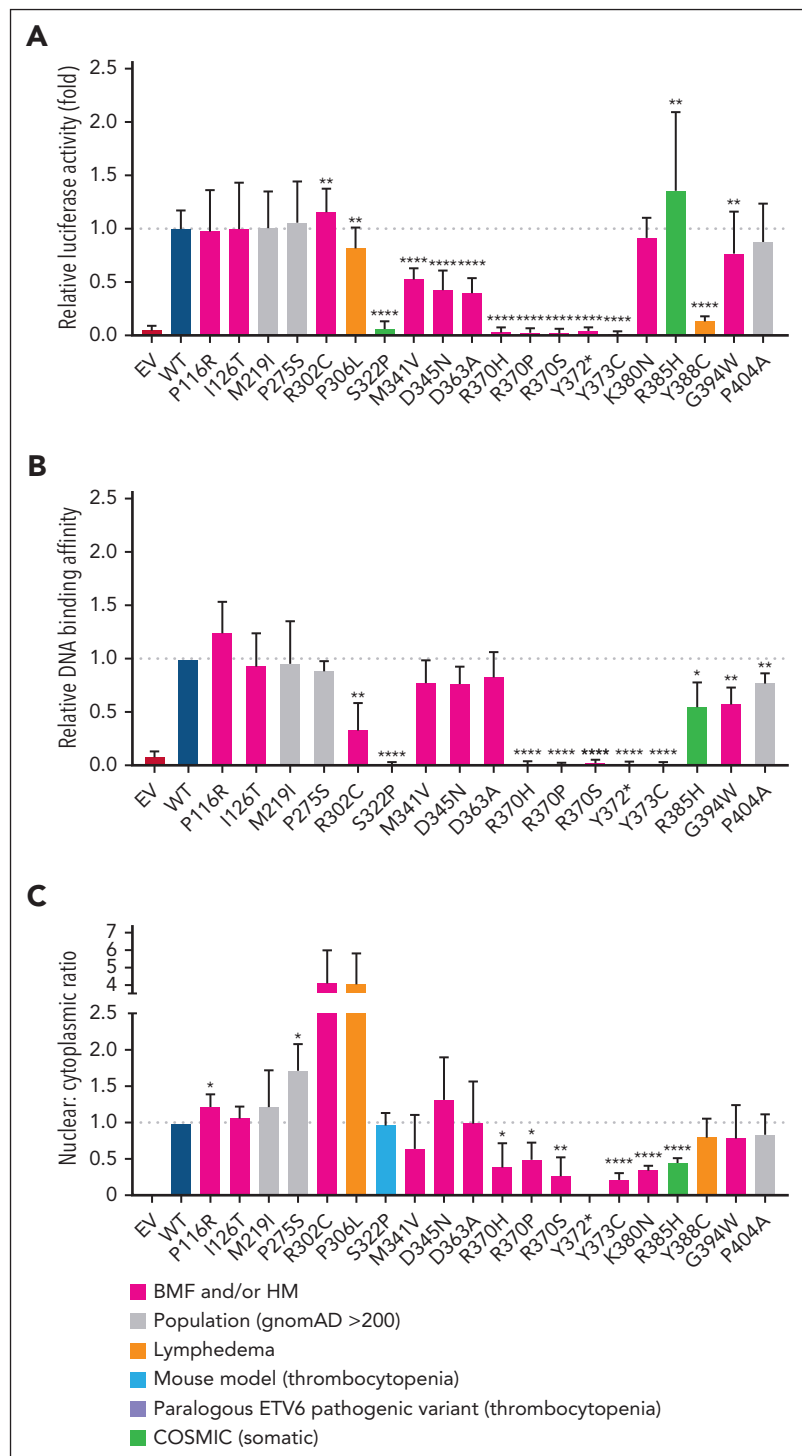
Table 1 (continued)

Patient no.	Sex	ERG variant (479 aa)	VAF (%)	gnomAD v4.0.0	REVEL	Hematological-related phenotype	Nonhematological phenotype	Age onset of first phenotype (y)	Germ line ERG (inherited/de novo) (Sample)	Somatic variants	Chr 21q cnLOH
16	M	Y373C	17	0	0.852	Thrombocytopenia and neutropenia	Not reported in clinical records	21	Yes (inherited) (hair)	RUNX1 (G165V) (2%)	Yes (2 events)
17	M	Y373C	44	0	0.852	Thrombocytopenia and neutropenia	Not reported in clinical records	19	Yes (inherited) (hair)	None	Yes
18		K380N	46	0	0.642	Anemia, thrombocytopenia, pancytopenia, macrocytic anemia, abnormality of the spleen, and eosinophilic infiltration of the esophagus	Hemangioma, hepatosplenomegaly, gastrointestinal inflammation, erythema, and capillary malformation	>18	Yes (de novo) (PB)		No
19	F	Y388C	50	0	0.90	None	Lymphedema	8	Yes (inherited) (PB)		No
20	F	Y388C	33	0	0.90	None	Lymphedema (bilateral intermittent lower limb swelling)	50	Yes (PB)		Yes
21		G394W	15	0 (3× G394R)	0.661	t-AML after treatment for DLBCL and prostate cancer	DLBCL and prostate cancer		Yes (PB)		No
Population		M219I		372 0.023% (1 hom)	0.058	NA			Yes		
Population		P275S		254 0.016% (3 hom)	0.223	NA			Yes		
Mouse		S322P		0	0.574	ALL and thrombocytopenia			Yes		
ETV6		R370S		0	0.803	ALL			No		
COSMIC		R385H		0 (2× R385C)	0.674	NA	2× BrCa, 1× biliary, 1× upper aerodigestive tract		No		
Population		P404A		236 0.015%	0.086	NA			Yes		

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Chr, chromosome; CLL, chronic lymphocytic leukemia; COSMIC, Catalogue Of Somatic Mutations In Cancer; DLBCL, diffuse large B-cell lymphoma; F, female; hom, homozygous; M, male; NA, not applicable; MSC, mesenchymal stromal cells; PB, peripheral blood; REVEL, Rare Exome Variant Ensemble Learner; SF, skin fibroblasts; t-MDS, therapy-related myelodysplastic syndrome.

Figure 2. Functional characterization of ERG variants. (A) ERG variants residing in the ETS domain reduce transactivation. K562 cells were transfected with pcDNA3 empty vector (EV) or pcDNA3-ERG (WT or variants). All constructs were cotransfected with a luciferase reporter plasmid driven by an *ITGA2B* promoter (quadruplicate replicates, repeated 3 times). Fold change (mean \pm standard error of the mean [SEM]) in comparison with the WT is plotted. Pairwise comparisons are shown ($*P < .05$ in comparison with WT). (B) ETS domain variants reduce ERG DNA binding affinity. EMSA of WT ERG and variants. Transfected HEK293 whole cell lysates were prepared and bound to an oligonucleotide containing an ETS DNA consensus sequence. Probes were visualized using chemiluminescence. Pairwise comparisons are shown ($*P < .05$ compared with WT). (C) ERG ETS domain variants alter subcellular localization. A Myc-tag was added to WT ERG and variants. COS-7 cells were transfected with pcDNA3 EV, pcDNA3-ERG-Myc (WT), and pcDNA3-ERG-Myc variants. Cells were stained for a Myc-tag and DAPI (4',6-diamidino-2-phenylindole). The nuclear to cytoplasmic ratio of each variant was quantified and the fold change (mean \pm SEM) in comparison with the WT was plotted. Pairwise comparisons are shown ($*P < .05$ compared to WT). Nuclear to cytoplasmic ratio was obtained from 3 independent experiments. In all comparisons, a Student *t* test was used ($*P < .05$; $**P < .01$; $***P < .001$ in comparison with WT).



ascertainment biases, and the inconsistent variant filtering strategies, determining the prevalence or penetrance of germ line pathogenic ERG variants to different phenotypes remains to be determined and requires the incorporation of additional well-defined cohort studies and systematic analyzes.

In vitro functional characterization of ERG variants

The impact of ERG variants on transactivation, DNA binding, and nuclear localization was examined (Figure 2). Western blot analysis showed all ERG variants produced protein, except for Y372*,

which was unstable (supplemental Figure 7). To assess transactivation ability, assays were performed using a platelet-specific *ITGA2B* promoter-luciferase reporter in K562 myeloid cells. Most ETS domain variants showed either complete LOF (S322P, R370H/P/S, Y372*, Y373C, and Y388C) or were hypomorphic (M341V, D345N, and D363A) when compared with the WT (Figure 2A). In contrast, a somatic variant (R385H), reported in multiple cancers in the COSMIC (Catalogue Of Somatic Mutations In Cancer),⁴¹ and R302C increased transactivation. ERG variants observed more than once in the population (ie,

>0.00012% gnomAD²⁷; supplemental Figure 8) showed no impact on the transactivation ability in this assay (Figure 2A; Table 1).

EMSA were performed to measure the effect of ERG variants on DNA binding (Figure 2B; supplemental Figure 9). Several ETS domain variants (S322P, R370H/P/S, Y372*, and Y373C) entirely ablated DNA binding, consistent with R370 and Y373 being critical contacts for DNA binding (supplemental Figures 3C and 10).⁴²

Immunofluorescence was used to quantify the effect of ERG variants on subcellular localization (Figure 2C). R302C and P306L displayed an increased nuclear to cytoplasmic localization ratio, but this effect was not statistically significant. Conversely, ETS domain variants R370H/P/S, Y372*, Y373C, K380, and R385H reduced nuclear protein localization (**P* < .05; Figure 2C; supplemental Figure 11A). Equivalent amino acid residues to ERG (R370, Y373) in this highly conserved ETS domain in ETV6 (R399)⁴³ and FLI (Y343),⁴⁴ respectively, are similarly critical for proper nuclear localization of these TFs (supplemental Figure 11B).

ERG LOF variants failed to drive myeloid differentiation and expansion of cytokine-independent stem or progenitor cells in FLC cultures

Retroviral-driven expression of WT ERG in FLCs drives the expansion of an immature stem or progenitor cell population with megakaryocytic features as demonstrated by intermediate cKIT⁺ and high CD41⁺ cell surface expression⁸ (Figure 3A-B). Several representative variants, including 3 complete LOF variants, 1 hypomorphic variant, and 2 WT-like variants from *in vitro* assays, were chosen to study their effect on *ex vivo* ERG-driven megakaryocytic expansion and cytokine-independent cell growth. Like the WT, P116R, a population (ie, benign) variant (M219I), and a hypomorphic variant (D345N) also drove this expansion. Strikingly, the ETS domain complete LOF variants (R370P, Y372*, and Y373C) did not show expansion of this immature stem or progenitor cell population after 4 to 5 weeks in culture, thereby replicating nontransduced FLCs or FLCs transduced with an empty MSCV-IRES-mCherry control retrovirus empty vector.

Furthermore, we observed that ERG WT and the P116R, M219I, and D345N variants were each able to induce cytokine independence in FLCs after 6 weeks in culture, thereby driving cell survival in the absence of essential cytokines. Conversely, ETS domain complete LOF variants (R370P, Y372*, and Y373C) were not able to drive cytokine-independent growth with all cells dying within 9 days of cytokine removal (Figure 3C), thereby replicating nontransduced FLCs.

Complete LOF ERG variants failed to drive leukemia in an ERG overexpression mouse model

Overexpression of ERG in murine FLCs, followed by transplantation into sublethally irradiated C57BL/6 mice, led to the development of a well-described erythro-megakaryocytic leukemia that is characterized by an accumulation of immature erythroblasts that infiltrate the BM, spleen, liver, and lung.^{8,10,45} Enforced expression of ERG WT, a WT-like variant (P116R), and a population variant (M219I) in murine FLCs similarly drove the

development of erythro-megakaryocytic leukemia (Figure 4A) by mCherry engraftment (Figure 4B) within 220 days after transplantation in 10 of 15 animals. Moribund mice displayed a large liver and spleen (supplemental Figure 12A), and flow cytometric analysis of the BM and spleen revealed a cKit⁺CD71⁺mCherry⁺ leukemic cell population consistent with ERG-driven erythro-megakaryocytic leukemia (Figure 4C-D), as previously described.⁸ Histology further confirmed leukemic cell infiltration in the BM, spleen, and liver (supplemental Figure 12B). In contrast, all mice (20/20) that received the ERG ETS domain complete (R370P, Y372*, and Y373C) or hypomorphic (D345N) variants showed no signs of disease or mCherry⁺ PB cells in the first 220 days, thereby demonstrating LOF in this *in vivo* assay. Notably, 1 variant (D345N) is hypomorphic in transactivation assays (Figure 2A), acts WT-like in cytokine independence assays (Figure 3A,C), and has complete LOF in this murine erythro-leukemia assay (Figure 4A). Hence, the choice of assay(s) is critical in ERG functional studies, especially when applying to variant classification for diagnostic and clinical applications.

Classification of germ line ERG variants and their phenotypes

Based on clinical, *in silico* predictive and functional data, all ERG variants were classified using the American College of Medical Genetics and Genomics (ACMG) and the Association for Molecular Pathology criteria and using heterozygous LOF variants (including missense) and the consequent haploinsufficiency as the disease mechanism⁴⁶ (supplemental Table 2). The ERG ETS domain is highly conserved (average constraint score, 0.26; supplemental Figure 13; supplemental Table 4),⁴⁷ and LOF variants across the protein are rare in the normal population (LOEUF = 0.23 and pLI = 1, gnomADv4.0.0; supplemental Figure 8),³⁸ suggesting intolerance to haploinsufficiency.

Five variants associated with BMF/HM (R370H, R370P, Y372*, Y373C, and K380N) demonstrated complete LOF in one or more functional assays and were classified as pathogenic, whereas Y388C (lymphedema) was classified as likely pathogenic (Figure 5; supplemental Table 5). Consistent with other BMF/HM predisposition genes, we observed variability in the clinical presentation associated with these variants. Thrombocytopenia, neutropenia, or pancytopenia was diagnosed in all patients with a likely pathogenic or pathogenic ERG variant (Table 1; supplemental Figure 14A; supplemental Table 2). Among this cohort, 28% developed an HM (AML and/or MDS) with a median age of onset of 29 years (supplemental Figure 14B). Longitudinal monitoring of individuals 10, 11, 15 (II-1), and 16 (II-2) consistently showed thrombocytopenia (platelets <150 × 10⁹/L) and neutropenia (<2.0 × 10⁹/L; supplemental Figure 14C-D).

Ten variants were classified as variants of uncertain significance (VUS), including 4 variants (M341V, D345N, D363A, and G394W) that demonstrated complete LOF or were hypomorphic in transactivation, DNA binding, and/or erythro-leukemia assays and 1 variant (V127Efs*82) that is likely to be pathogenic based on complete LOF of other premature termination variants (ie, Y372* and lymphedema variants³¹; Figure 5). Seven VUSs were associated with HM (MDS, AML, ALL, chronic lymphocytic leukemia, diffuse large B-cell lymphoma), 3 with BMF, and 1 with lymphedema. Note, 1 patient with thrombocytopenia who carried a germ line I126T variant also harbored a

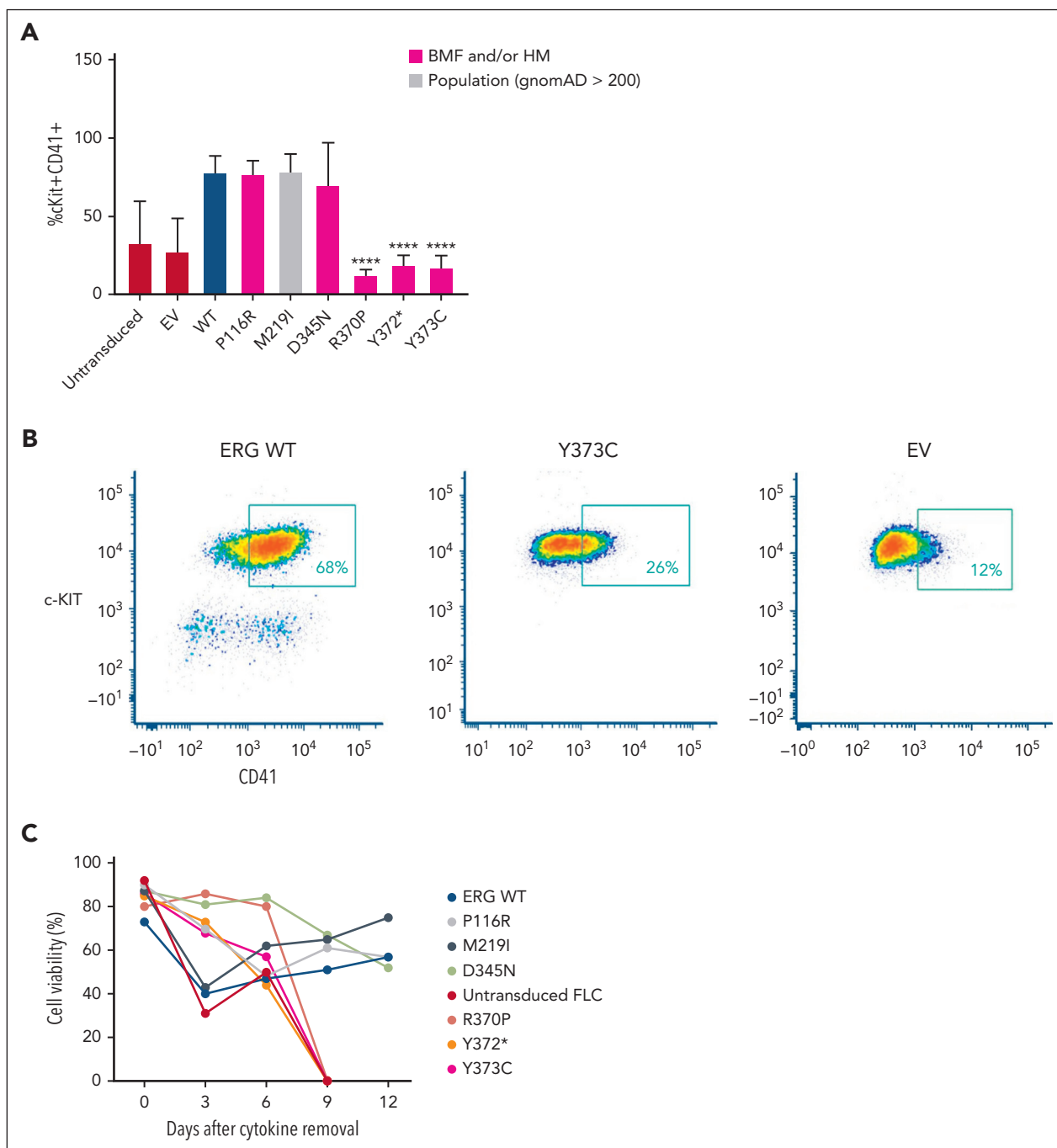


Figure 3. ERG LOF variants fail to drive megakaryocytic differentiation and cytokine independence. (A) In vitro differentiation of FLCs that overexpress WT ERG and variants. Immature stem or progenitor cell population with megakaryocytic features were stained using antibodies against cKIT⁺ and CD41⁺ with the percentage of double positive cells measured by flow cytometry (6 independent experiments). Pairwise comparisons are shown (**** $P < .001$ in comparison with WT). In all comparisons, a Student *t* test was used. (B) Example of gating strategy for cKIT⁺ and CD41⁺ expression on ERG WT- or variant-transduced FLCs. Cells were first gated for viability and mCherry expression. (C) Cell viability of FLC culture after cytokine removal measured over time.

germ line pathogenic *RUNX1* whole gene deletion (supplemental Table 2), likely explaining the phenotype; whether the *ERG* variant also contributed to the phenotype is unclear. Two variants (D345N and D363A) could not be verified as germ line because of unavailability of samples and have been reported multiple times as somatic with D363 being a hot spot for somatic variants in multiple cancers (supplemental Figure 15; supplemental Table 6). Somatic analysis of BM or blood identified variants in patients 2, 3, 7, 14, 15, and 16, 2 of

whom (3 and 16) acquired *IDH1* variants; no variants were found in patients 11 and 12 (Table 1).

Somatic *ERG* variants in HM and other cancers

The COSMIC was filtered for *ERG* variants that occurred ≤ 3 times (ie, very rare) in gnomAD v4.0.0. There were 64 unique ETS domain missense variants. Screening for somatic missense variants at the same amino acids affected by germ line ETS domain variants in this study revealed that all except 1 (Y373C) were considered to be

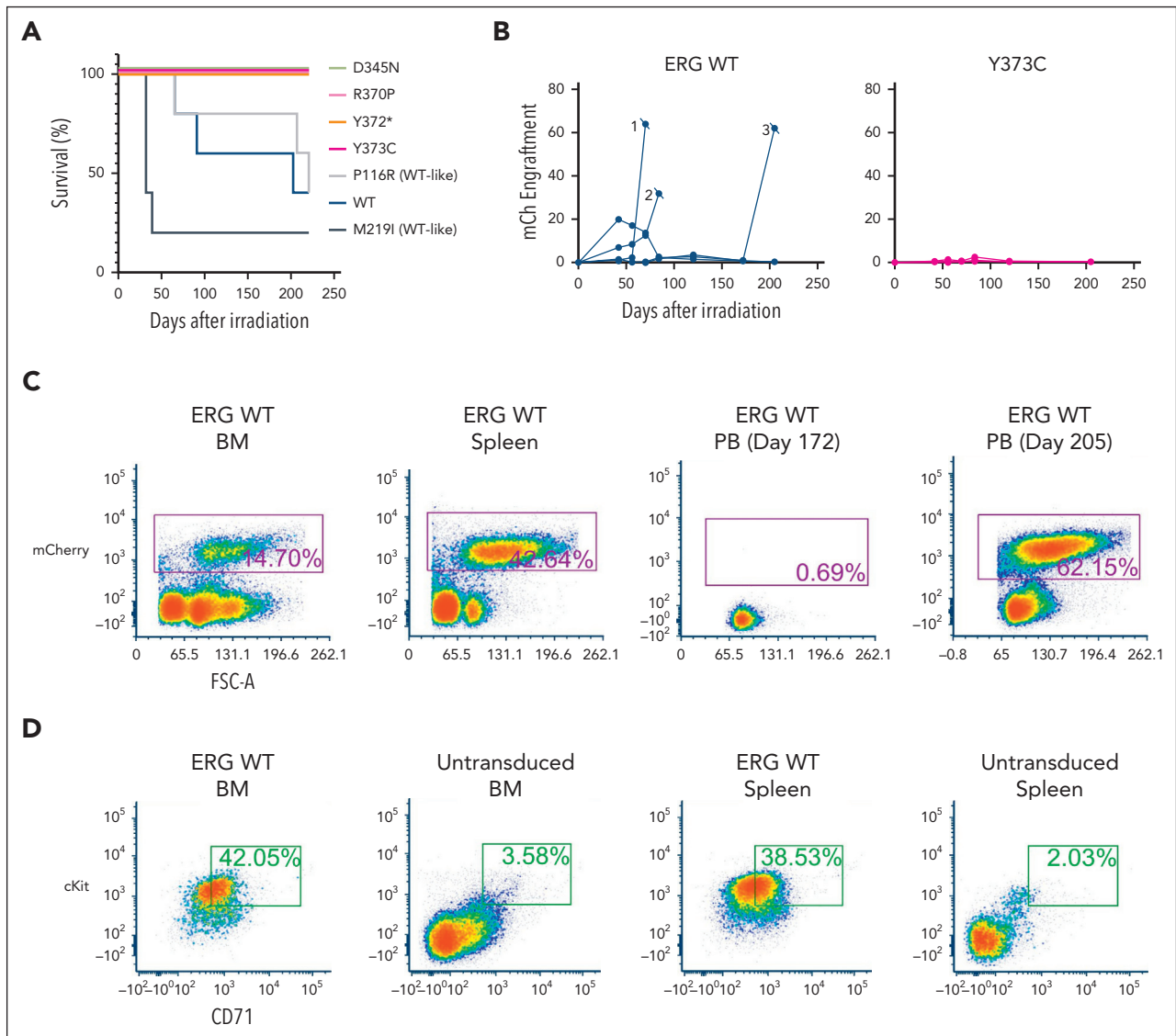


Figure 4. ERG variants are LOF in an in vivo leukemia model driven by ERG overexpression. (A) Enforced expression of ERG WT and WT-like variants (P116R, M219I) in mice led to the development of erythro-megakaryocytic leukemia within 220 days. (B) mCherry engraftment over time in the PB of mice. Numbering refers to 3 ERG WT mice that succumbed to disease (Figure 5A). (C) Infiltration of ERG WT/mCherry⁺ transplanted FLCs into recipient mouse BM, spleen, and PB as measured by fluorescence-activated cell sorter (FACS) analysis. Recipient mouse number 3 shown is representative of all ERG WT and WT-like (P116R, M219I) mice. (D) ERG WT and WT-like variant mice developed mCherry⁺ erythro-megakaryocytic leukemia with cKit⁺ and CD71⁺ expression. FACS plots of a representative ERG WT leukemia are shown. Nonleukemic (ie, mCherry⁻) BM and spleen cells are shown as comparison.

somatic (supplemental Figure 15), adding weight to these amino acid substitutions being drivers of malignancy.

In a separate pediatric familial cancer cohort (St Jude Children's Research Hospital), a single germ line *ERG* (V127Efs*82) variant was identified (Table 1), in addition to several predicted pathogenic somatic *ERG* variants, predominantly in children with B-ALL, but also AML, non-Hodgkin lymphoma, and some solid cancers (supplemental Table 6). Interestingly, *ERG* (V127Efs*82) was identified both as a germ line and a somatic variant in different patients with B-ALL.

Discussion

We reported the discovery of *ERG* as a new autosomal dominant BMF and HM predisposition gene, expanding its disease

phenotypes beyond recently reported primary lymphedema.³¹ *ERG*-based clinical disease resembles that of *RUNX1*,⁴⁸ *ETV6*⁴⁹ and *ANKRD26*⁵⁰ (thrombocytopenia, myeloid, and lymphoid HM), *GATA2*^{36,51} (BMF, myeloid HM, lymphedema), and *ERG*'s most closely related ETS family member *FLI1* (thrombocytopenia, but to date without reports of HM).^{33,34,52,53} These overlapping, but nonidentical, phenotypes are consistent with many of these TFs being part of a complex homeostatic network that is crucial for normal hematopoiesis.⁵ Furthermore, consistent with a role for *ERG* and *GATA2* in the lymphatics and predisposing to lymphedema, in single cell studies, there is high expression of *ERG* and *GATA2* in BM endothelial cells in contrast with *RUNX1* for which no primary lymphatic phenotype has been described.⁵⁴

Intriguingly, cnLOH favoring the WT allele was identified in all 3 *ERG* (Y373C) carriers in index family 1. SGR events such as these

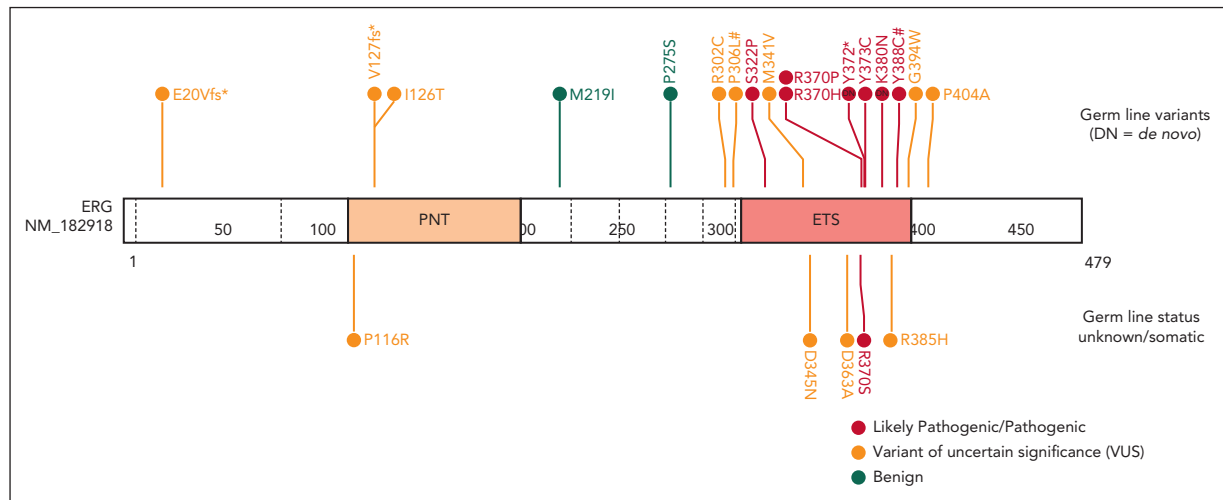


Figure 5. ACMG classification of germ line *ERG* variants. *ERG* variants were classified using the ACMG and the Association for Molecular Pathology criteria for cytopenias, HMs, and/or lymphedema (#). Based on functional assay criteria, variants with complete LOF in at least 1 functional assay were classified as PS3_Strong, variants with hypomorphic activity (<50%) in 1 or more functional assays were classified as PS3_Moderate, and variants with hypomorphic activity (>50%) in 1 or more assays were classified as PS3_Supporting. The PS3 and BS3 criteria were not applied when variants showed no change in functional assays or were not tested. PNT, pointed domain.

can complicate the detection of germ line variants from hematopoietic tissues by mimicking the lower VAFs of somatic variants. Consistent with this, individuals with either Y388C or S182Afs*22 (recently published familial germ line *ERG* variant in a patient with lymphedema³¹) had cnLOH but no hematological disease (supplemental Figure 16). Notably, the patient with S182Afs*22 presented with 4% VAF in the PB,³¹ which, upon further investigation, was caused by a cnLOH event that corrected 92% of cells. SGR may lead to asymptomatic carriers, milder symptoms, a later age of onset, or a missed diagnosis (hidden predisposition) and reveals the potential for disease prevention via cell and/or gene therapy.^{55,56} Hence, identified *ERG* variants should be tested in germ line, nonhematopoietic tissues, such as fibroblasts, mesenchymal stromal cells or hair, to confirm germ line or somatic status.

Functional studies demonstrated that all rare *ERG* variants within the ETS domain led to complete or hypomorphic LOF in transactivation, DNA binding, and/or subcellular localization, and selected variants similarly exhibited LOF in fetal liver hematopoietic cell growth assays ex vivo and leukemogenic assays in vivo. This is in contrast with the gain-of-function nature of common *ERG* fusions^{19-21,27,28} and hence shows that *ERG* can act as both a tumor suppressor gene and an oncogene. Notably, disease manifestations stemming from germ line LOF mechanisms extend to other members of the ETS family (*ETV6* and *FLI1*).^{33,34,52,53} *ETV6* and *FLI1* variants often cluster in the ETS domain,³² and several are analogous to pathogenic *ERG* variants in our study (supplemental Figure 11B). For example, a single consanguineous family with 2 siblings affected with moderate thrombocytopenia and a lifelong bleeding history was described to have a rare *FLI1* ETS domain missense variant in homozygosity (R324W, NM_002017.4).³³ This variant (analogous to *ERG* R354) was hypomorphic in in vitro analyzes,³³ similar to 3 variants in our study in the same region (M341V, D345N, and D363A) (Figure 2A). To date, we have not seen patients with homozygous *ERG* germ line variants or evidence of autosomal recessive disease, although this might be possible for hypomorphic variants.

Structural modeling shows that several complete LOF variants (in DNA binding and transactivation assays) affect residues that contact DNA (supplemental Figure 9).⁴² In addition, the nuclear localization 2 region of the *FLI1* protein is identical to this ETS domain region in *ERG*.^{44,57} Unsurprisingly, 6 variants within this paralogous region were unable to appropriately localize to the nucleus (Figure 2C; supplemental Figure 10). Therefore, because of the high conservation of the ETS domain in paralogous TFs, reported function-disrupting variants in 1 ETS TF are likely to affect protein function at the corresponding amino acid position in others. Indeed, paralogous amino acids cause similar loss of DNA binding and nuclear localization and several have been classified in ClinVar as pathogenic for their relevant disease phenotypes. Pathogenic variants and their functional analyzes in highly conserved paralogous genes and domains, such as ETS TFs and ETS domains, is topical for variant curation expert panel discussion in generating gene-specific guidelines.

In addition to germ line variants, *ERG* somatic variants are found in a proportion of sporadic cancers, including HM (B-ALL, MDS, diffuse large B-cell lymphoma, AML, and T-cell acute lymphoblastic leukemia) and solid cancers (predominantly skin, gastrointestinal, breast, lung). Interestingly, the majority of the missense variants in HMs are located within the ETS domain with several variant hotspots, including ones we tested (ie, D345N, D363A), that were hypomorphic in our system; this raises the likelihood of a strict threshold for *ERG* activity in certain cellular contexts, including concurrent with other somatic variants. For germ line *ERG* HM cases, as for *GATA2*, we have not identified somatic variants on the other allele (ie, biallelic) as occurs for *RUNX1*,⁵⁸ *CEBPA*,^{58,59} and *ETV6*.⁴³

Our data establishes a gene-disease association for *ERG* in the pathogenicity of BMF and HM, in addition to lymphedema. Six ETS domain variants in this study were classified as pathogenic or likely pathogenic, whereas 10 variants were classified as VUS (Figure 5). Addition of functional data criteria changed the ACMG classification of 5 *ERG* variants (3 VUS to likely pathogenic, 2 VUS to benign; supplemental Table 5), highlighting

Table 2. Summary of functional assays for phenotypic ERG variants in this study

Variant	Phenotypes	Transactivation	DNA binding	Subcellular localization	FLC myeloid differentiation	FLC cytokine independence	Leukemogenesis assay	ACMG classification
WT	NA	✓✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	NA
I126T	AML	✓✓✓✓	✓✓✓✓	✓✓✓✓	ND	ND	ND	VUS
P116R	Cytopenia	✓✓✓✓	✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	VUS
R302C	CLL	✓✓✓✓	ND	✓✓✓✓	ND	ND	ND	VUS
K380N	Cytopenia	✓✓✓✓	✓✓✓	✓	ND	ND	ND	P
P306L	Lymphedema	✓✓✓	ND	✓✓✓✓	ND	ND	ND	VUS
G394W	AML	✓✓✓	✓✓✓✓	✓✓✓✓	ND	ND	ND	VUS
M341V	Cytopenia	✓✓	✓✓✓✓	✓✓✓✓	ND	ND	ND	VUS
D345N	MDS	✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	✓✓✓✓	✗	VUS
D363A	MDS	✓✓	✗	✓✓✓✓	ND	ND	ND	VUS
Y388C	Lymphedema	✗	ND	✓✓✓✓	ND	ND	ND	LP
R370H	Cytopenia	✗	✗	✓✓	ND	ND	ND	P
R370P	Cytopenia + MDS	✗	✗	✓✓	✗	✗	✗	P
Y373C	Cytopenia + AML	✗	✗	✓	✗	✗	✗	LP
Y372*	Cytopenia	✗	✗	✓	✗	✗	✗	P
E20Vfs*13	MDS	ND	ND	ND	ND	ND	ND	VUS
V127Efs*82	ALL	ND	ND	ND	ND	ND	ND	VUS

ERG variants are ordered in descending order of transactivation ability. Ticks (✓) indicate degree of WT-like activity in transactivation, DNA binding, subcellular localization, FLC myeloid differentiation, FLC cytokine independence, and leukemogenesis assays. ✗ indicates complete LOF in these assays.

CLL, chronic lymphocytic leukemia; LP, likely pathogenic; NA, not applicable; ND, not determined; P, pathogenic.

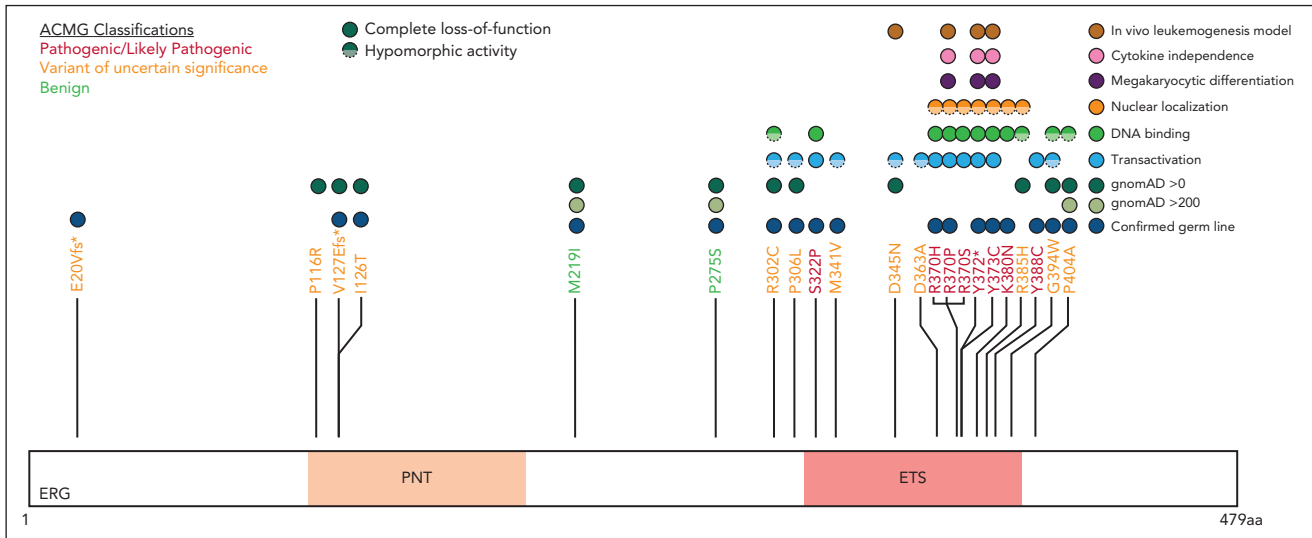


Figure 6. Functional consequences of rare ERG variants. Rare ERG variants from similar phenotypic groups, including BMF and/or HM and lymphedema, among population variants (gnomAD >200),³⁸ a germ line thrombocytopenic mouse variant,¹ a paralogous ETV6 pathogenic variant (thrombocytopenia),⁴⁹ and Catalogue Of Somatic Mutations In Cancer mutation (somatic)⁴¹ are mapped onto the ERG protein (isoform, NP_891548.1; transcript, NM_182918.4). Functional characterization of each variant via transactivation, DNA binding, subcellular localization, FLC myeloid differentiation, FLC cytokine independence, and leukemogenesis assays are displayed.

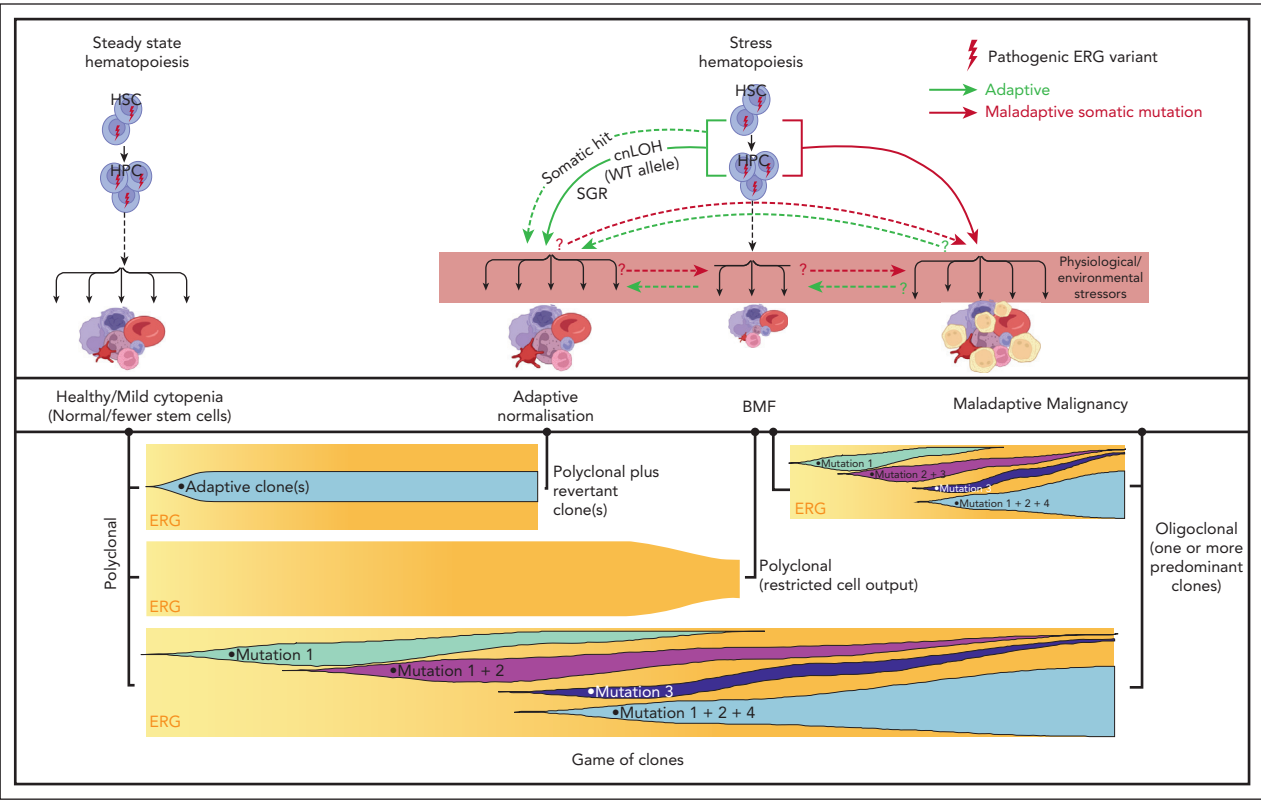


Figure 7. Proposed model for ERG deficiency syndrome-associated phenotypes showing adaptive/maladaptive events in ERG heterozygous carriers. It is proposed that ERG carriers harbor a threshold of activity that is insufficient under certain physiological and environmental stressors, thus leading to a range of phenotypic outcomes (top panel). Malignant cells (yellow cells). Maladaptive and adaptive events observed in our study (arrow). Other potential (maladaptive/adaptive) somatic events seen in other BMF and/or HM predisposition genes⁵³⁻⁶⁵ (dotted arrow) or hypothetical events (question mark preceding dotted arrow). Game of clones (bottom panel). The development of different physiological outcomes in patients is the consequence of a game of clones in which context-dependent competitive fitness of the clone(s) determines progression and the observed outcome. Notably, adaptive and maladaptive events and subsequent clonal selection may occur simultaneously. The clonal output shown is based on mature blood cells, reflecting collectively the impact on mature and stem and progenitor cells.

the importance of generating faithful functional assays. For the VUSs, despite their seemingly WT-like behavior in functional assays, it remains plausible that some may have critical functional consequences that were not detected in our over-expression systems. The functional consequences of these variants may impact via temporal and spatial mechanisms, and therefore caution is required when interpreting WT-like VUS because protein-protein interactions or target binding may vary in different cellular contexts. It is probable, however, that some variants classified as VUS (ie, I126T and P116R) are not monogenic, fully penetrant variants given their lack of functional consequence in the overexpression assays used in this study.

Our data (Table 2; Figure 6) defines *ERG* as a new predisposition gene to be added to germ line screening panels for BMF and HM syndromes. Demonstration of de novo germ line variants emphasizes testing for inheritance for family planning and counseling. We stress the importance of screening true germ line samples of patients and family members because we posit that SGR and the use of hematopoietic samples may mask asymptomatic or mildly symptomatic carriers, potentially missing more severe clinical presentation in other family members, including perinatal and neonatal lethality as has been observed in *MECOM*.^{60,61} Enticingly, SGR highlights the potential for preemptive cell therapy and/or gene editing strategies to prevent or alleviate *ERG*-related disease in the blood stem cell compartment. We only observed several inherited cases (4 small families), which may indicate low penetrance, mild phenotypes, and/or incomplete family data. However, of the 3 families in the Genomics England Rareservoir study with primary lymphedema carrying *ERG* premature termination variants³¹ and 2 missense *ERG* variants reported here, 4 showed familial inheritance and 1 was de novo. Whether lymphedema is a highly penetrant phenotype and hence seen in family units remains unanswered. Although *ERG* variants in lymphedema and HM phenotypes have not yet coincided, we anticipate that with a larger, more defined cohort study they will overlap, mirroring the history of *GATA2* variants and Emberger syndrome (lymphedema and MDS).⁵¹

Our current disease model (Figure 7) proposes that, in *ERG* carriers, the interplay between physiological and environmental stressors impacts critical threshold-sensitive gene expression or biologic pathways that are required for normal hematopoiesis, which leads to BMF and adaptive (SGR) and/or maladaptive (hematological malignancy) selective processes. It is possible for both adaptive and maladaptive clones to be present in the same individual for whom context-dependent competitive fitness of the clone determines the physiological outcome, a game of clones. Notably, *ERG* expression is high in primary human HSCs and it drops during transition to hematopoietic progenitor cells (HPCs)⁵² and during differentiation. Analysis of the *ERG* downstream effectors and pathways impacted in primary BM and blood cells in *ERG* carriers and patients with BMF/HM is warranted to better understand disease initiation and progression. Clearly, describing these disease mechanisms, the phenotypic and mutational spectrum, and the natural history of diseases caused by *ERG* germ line variants has only just commenced for what may well be another pleiotropic and protean transcriptopathy. As population-scale genomic studies,

such as the UK Biobank and All of Us, become popular, our study demonstrates that they should not become de rigueur. Careful clinical and laboratory observations with professional networking will remain important in describing new *ERG*-associated disease and other disease entities.

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Authorship

Contribution: J.R.Z. wrote the manuscript and was involved in all aspects of the project including research design, manuscript preparation, and collecting and analyzing experimental and clinical data; A.L.B., C.L.C., H.S.S., and C.N.H. were involved in the research design, data analysis, manuscript writing and preparation, American College of Medical Genetics and Genomics-variant classification, and providing scientific insight; C.C.H., P.A., and P.V. contributed to different aspects of design and analysis of clinical and experimental data and manuscript writing and preparation; X.L., S.J.S., and L.T. helped to collect experimental data; M.B. and P.J.B. helped with sample preparation; L.A.-M. and K.S.K. performed bioinformatic analyses; S. Moore, R.H., W.T.P., and H.N. performed cytogenetic, somatic next-generation sequencing, and/or diagnostic arrays for family 1; S.B. and M.W.W. were involved in figure generation; S.F., L.L., F.S.d.F., S. Demirdas, S.d.M., H.A.-P., B.B., S. Mansour, K.G., A.K., S. Dobbins, P.G.N.J.M., K.E.N., N.O., D.D., R.M., A.C., J.M., D.B., J.F.-D., M.W.D., K.P., N.K.P., G.M.B., D.P., P.O., A.S., L.A.G., D.M.R., D.K.H., and J.S. provided clinical and variant information of patients; M.W.W. and the *ERG* Variant Research Network confirmed the absence of rare *ERG* variants in their cohorts; and all authors critically reviewed and approved the manuscript.

Conflict-of-interest disclosure: A.C. is an employee of and stockholder in Illumina, Inc. The remaining authors declare no competing financial interests.

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Footnotes

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Original data are available on request from the corresponding authors, Christopher N. Hahn (chris.hahn@sa.gov.au) and Hamish S. Scott (hamish.scott@sa.gov.au).

The online version of this article contains a data supplement.

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