

Cytokine pathway variants modulate platelet production: *IFNA16* is a thrombocytosis susceptibility locus in humans

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Key Points

- A cytokine pathway array has identified variants and functional interactive networks regulating platelet formation in stress thrombocytosis.
- *IFNA16* genetic variants behave as susceptibility loci in essential (but not reactive) thrombocytosis, limited to *JAK2*^{V617F}-negative cohorts.

Inflammatory stimuli have divergent effects on peripheral platelet counts, although the mechanisms of thrombocytopenic and thrombocytotic responses remain poorly understood. A candidate gene approach targeting 326 polymorphic genes enriched in thrombopoietic and cytokine signaling pathways was applied to identify single nucleotide variants (SNVs) implicated in enhanced platelet responses in cohorts with reactive thrombocytosis (RT) or essential (myeloproliferative neoplasm [MPN]) thrombocytosis (ET). Cytokine profiles incorporating a 15-member subset, pathway topology, and functional interactive networks were distinct between ET and RT, consistent with distinct regulatory pathways of exaggerated thrombopoiesis. Genetic studies using aggregate (ET + RT) or ET-restricted cohorts identified associations with 2 *IFNA16* (interferon- α 16) SNVs, and the ET associations were validated in a second independent cohort ($P = .0002$). Odds ratio of the combined ET cohort ($n = 105$) was 4.92, restricted to the *JAK2*^{V617F}-negative subset (odds ratio, 5.01). ET substratification analysis by variant *IFNA16* exhibited a statistically significant increase in IFN- α 16 levels ($P = .002$) among 16 quantifiable cytokines. Recombinantly expressed variant IFN- α 16 encompassing 3 linked non-synonymous SNVs (E⁶⁵H⁹⁵P¹³³) retained comparable antiviral and pSTAT signaling profiles as native IFN- α 16 (V⁶⁵D⁹⁵A¹³³) or IFN- α 2, although both native and variant IFN- α 16 showed stage-restricted differences (compared with IFN- α 2) of IFN-regulated genes in CD34⁺-stimulated megakaryocytes. These data implicate *IFNA16* (IFN- α 16 gene product) as a putative susceptibility locus (driver) within the broader disrupted cytokine network evident in MPNs, and they provide a framework for dissecting functional interactive networks regulating stress or MPN thrombopoiesis.

Introduction

Megakaryocytopoiesis and proplatelet formation represent progressively linked stages of hematopoietic stem cell (HSC) development that maintain the normal circulating pool of platelets, critical components of normal hemostasis, pathologic thrombosis, and host adaptive immunologic responses.¹⁻³ Mechanisms regulating normal thrombopoiesis are distinct from those occurring during stress, with evidence that a genetically distinct subset of megakaryocyte (MK)-committed progenitors are mobilized during

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Requests for original data may be submitted to the corresponding author (Wadie F. Bahou; e-mail: wadie.bahou@stonybrook.edu).

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

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inflammatory stimuli.⁴ Indeed, pathologically exaggerated platelet production (reactive thrombocytosis [RT]) is generally classified as a stimulus-dependent phenomenon occurring in the setting of cytokine release.^{5,6} Similarly, inflammatory dysregulation is evident in thrombocytosis-associated myeloproliferative neoplasms (MPNs) occurring in conjunction with constitutive activation of the Jak/Stat pathway.⁷⁻⁹ Whether cytokines in MPNs function as disease drivers in the setting of predisposition alleles¹⁰⁻¹² or as sequelae of clonal hematopoietic expansion remains unestablished. Inflammatory triggers on the background of *JAK2* susceptibility loci have been suggested as initiating stimuli for sustained expansion of *JAK2*^{V617F} mutant clones, either secondary to hypermutable somatic mutagenesis or expansion of cells with preexisting mutant alleles. Interestingly, the *JAK2* susceptibility locus has also been associated with Crohn's disease,¹³ further implicating dysregulated cytokine stimulation as pathogenetic drivers for inflammatory bowel disease.

Thrombopoiesis is primarily regulated by the thrombopoietin (TPO)/*c-mpl* receptor/ligand system, although there is little explanation for the genetic variability of platelet counts in normal and diseased subjects. The normal range of platelet counts in humans is broad ($1.5 \times 10^8/\text{mL}$ to $3.5 \times 10^8/\text{mL}$) and remarkably constant over time, with coefficients of variation of 6.7% in male subjects and 8.6% in female subjects.¹⁴ Twin studies indicate that genetic factors account for normal variation of platelet counts in the healthy population,^{15,16} although the associated genes remain unidentified. Limited studies have focused on polymorphisms within *c-mpl* as platelet modifier genes,¹⁷ and genome-wide association studies (GWAS) have identified candidate single nucleotide variants (SNVs) that may influence the number and size (mean platelet volume) of platelets¹⁸ and/or red blood cells.¹⁹ More recent studies have expanded the subset of SNV loci that account for 18% and 30%, respectively, of platelet count and mean platelet volume variability.²⁰ Despite this progress, a preponderance of genetic modifiers regulating physiological platelet production remains uncharacterized, and identification of genetic loci that modulate stress-associated platelet production is specifically unknown.

Inflammation induces widely variable patterns of cytokine release, although downstream cellular effects are mediated by shared cytokine and hematopoietic signaling pathways.^{4,21} The current study applies a candidate gene approach to dissect the role of cytokine pathway SNVs as genetic modifiers of enhanced thrombopoiesis. A custom SNV array encompassing nonredundant cytokine signaling pathway genes identified variants and distinct functional networks differentially associated with RT or essential thrombocytosis (ET) cohorts, and a predominant *IFNA16* (interferon- α 16) predisposition locus in ET validated in a second, unrelated cohort. These data provide the first comprehensive study of cytokine pathway variants implicated in exaggerated thrombopoietic responses in humans, and they provide a framework for further dissecting the complex interplay of regulatory networks linked to inflammatory stimuli.

Materials and methods

Human subjects

All subjects (ET, $n = 106$; RT, $n = 56$; healthy control subjects, $n = 224$) were enrolled in a protocol approved by the Stony Brook University Institutional Review Board that was conducted in accordance with the Declaration of Helsinki,²² applying well-established

clinical criteria for phenotypic classification.^{23,24} ET recruitment was completed in 2 phases, Cohort 1 ($n = 52$ was used for initial genotyping and association studies), followed by a second validation phase (Cohort 2, $n = 54$), under the auspices of Institutional Review Board #88960 (Platelet Systems Biology in Health and Disease).

SNV selection, genotyping, and statistical analyses

Peripheral blood high-molecular weight genomic DNA was used for genotyping, *CALR* insertion/deletion detection,²⁵ and *JAK2*^{V617F} allelic quantification. Genotyping was completed by using a custom SNV array encompassing discrete pathways implicated in hematopoietic cytokine signaling²¹ and thrombopoiesis: (1) the TPO/*c-mpl* (TPO/TPO receptor) pathway; (2) the JAK/STAT pathway; (3) a steroidogenic pathway previously implicated in thrombocytosis²⁶; and (4) platelet-modifier SNVs implicated in platelet function²⁷ or thrombopoiesis (mean platelet volume or number) based on human GWAS.¹⁸ Details of the 326 gene/1056 SNV list are provided in supplemental Table 1. Samples were blinded to case-control, and call rates ranged between 97.0% and 99.5%; platform reliability and quality control were established by using 10 randomly duplicated samples with concordance >98%. Pyrosequencing was used for *JAK2*^{V617F} allelic determination (defining homozygosity by >95% mutant A alleles) and as a secondary validation step for a subset of eight SNVs (genes and oligonucleotide primers provided in supplemental Table 2). Assay design and allelic discrimination were completed by using PyroMark software version 2.0.1.15 (Qiagen, Germantown, MD), with sensitivity of mutant allele detection ~3%; concordance between genotyping and pyrosequencing was >98%. Focused nucleotide confirmation of *IFNA16* was established by dideoxy sequence analysis.

For each SNV, genotype frequencies were delineated and excluded from association studies for sample call rates <70% or for failure to meet Hardy-Weinberg equilibrium. SNV case-control data were analyzed by using the χ^2 test, Fisher's exact test, or the Cochran-Armitage trend test, using distinct genetic models (allelic, dominant, and recessive), substratified according to phenotype (ET alone, RT alone, or ET + RT) and a common control subset; Cohort 1 and Cohort 2 recruitment proceeded independently with no subject overlap. Association testing of ET, odds ratios (ORs), and confidence intervals (CIs) were computed in R package (version 3.6.2). Sample size validation of association testing was established by including five *JAK2* SNVs as internal controls for ET genetic susceptibility, with a post hoc power analysis of 0.77 at the significance levels of 0.05 (one-sided).¹⁰ Statistical comparisons were completed by using analysis of variance or *t* tests (or their nonparametric counterparts if the normality assumptions were not met) at the significance level of $P < .05$ adjusted for the false discovery rate.²⁸

Cytokine measurements

Cytokines analyzed in this study were selected from a broad group of inflammation-related cytokines based on 3 criteria: (1) demonstrated role in thrombosis-related inflammation^{29,30}; (2) dysregulated expression patterns in ET and/or RT^{29,31,32}; and (3) availability of sensitive and robust detection methods. Plasma cytokines were measured by using multiplex, bead-based immunoassays (ProcartaPlex; Thermo Fisher Scientific, Waltham, MA), or IFN- α 16 was quantified by using capture enzyme-linked immunosorbent assay

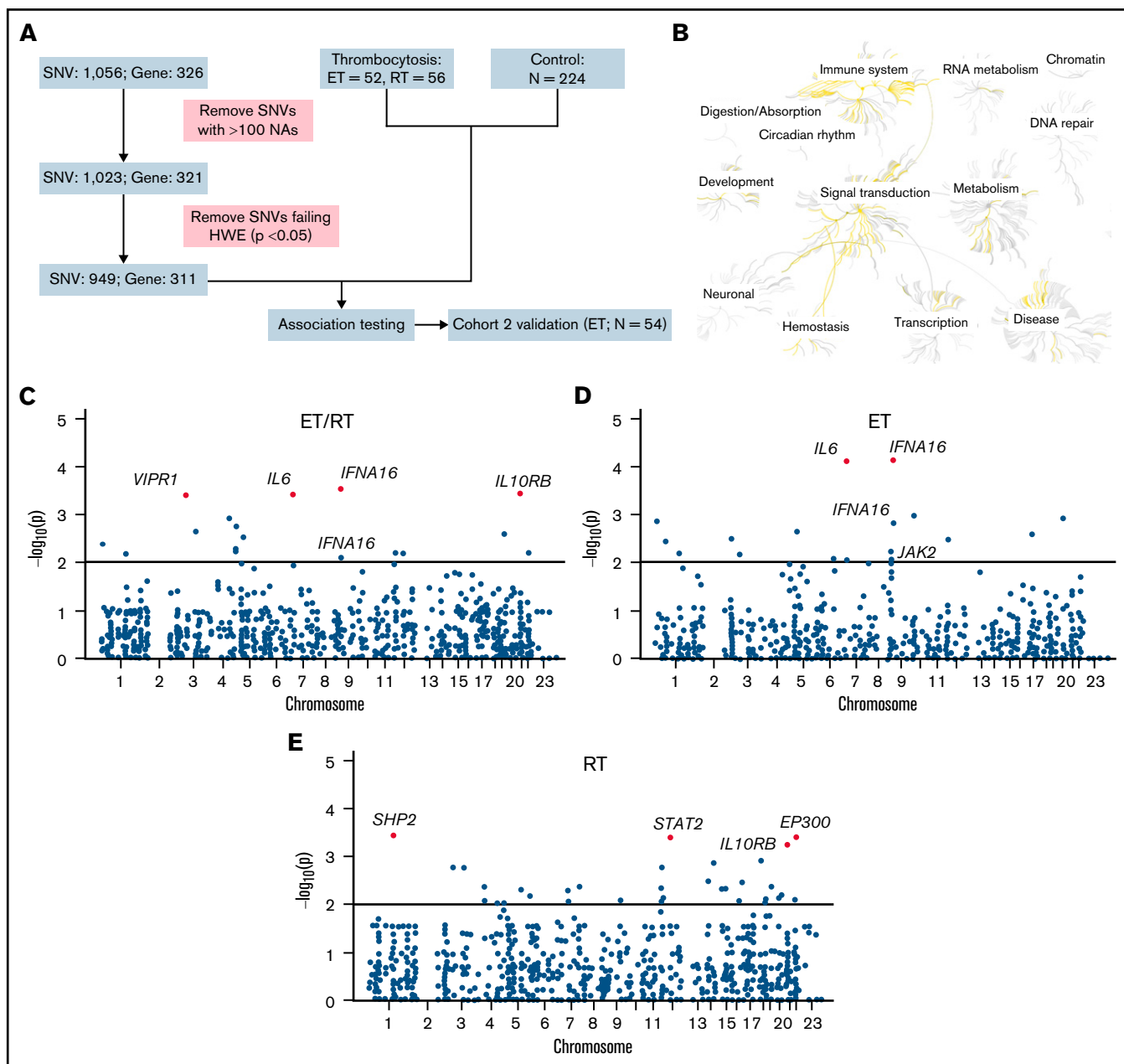


Figure 1. Genotyping overview. (A) Work-flow schema showing clinical characteristics and genotyping strategy. (B) Genetic overrepresentative analysis (N = 311 genes) was completed by using Reactome,³⁶ and enriched pathways are highlighted in yellow. (C-E) Genome-wide association plots (aggregate ET and RT [C], ET [D], and RT [E]) detailing significant gene/SNVs (at $P < .01$) identified by dominant, allelic, or recessive models; the most significant gene/SNVs ($P < .001$) are in red (Table 1). HWE, Hardy-Weinberg equilibrium; NA, not available.

(MyBioSource, Inc., San Diego, CA). Sample concentrations were determined in duplicate by using standard curves run in parallel.

Expression and functional characterization of IFN- α 16

Recombinant human IFN- α 16 wild-type (IFN- α 16^{WT}) or mutant (IFN- α 16^{MUT}) were expressed and purified as C-terminal His₆-Strep chimeric proteins in *Escherichia coli* Origami2 DE3 cells. Recombinant proteins were serially purified by tandem affinity purification, and

after cleavage from carriers, bacterial endotoxin was removed by using a high-capacity endotoxin removal spin column (Thermo Fisher Scientific). IFN- α protein concentrations were quantified, stabilized with purified bovine serum albumin (final concentration 0.1%), and stored at -80°C . IFN- α antiviral activity was determined by using adenovirus type 5–infected human diploid fibroblasts (HDF/*Tert1*), whereas cellular pSTAT1 activation was established by using A549 (lung epithelial), human diploid fibroblast (HDF/*tert1*), or primary human CD34⁺ cells incubated for 1 hour with 500 U/mL human universal type I IFN- α 2 (PBL Assay Science, Piscataway, NJ), or

Table 1. Gene/SNV association studies according to cohort

ET and RT				ET				RT			
SNV	Gene	P	Model*	SNV	Gene	P	Model	SNV	Gene	P	Model
rs28368163	<i>IFNA16</i>	.000283	A	rs28368163	<i>IFNA16</i>	7.80×10^{-5}	A	rs8191984	<i>SHP2</i>	.000373	A
rs17860266	<i>IL10RB</i>	.000368	A	rs2069824	<i>IL6</i>	8.34×10^{-5}	A	rs12628803	<i>EP300</i>	.000381	A
rs2069824	<i>IL6</i>	.000377	A	rs2280583	<i>STAM</i>	.001066	D	rs2066809	<i>STAT2</i>	.000406	A
rs72862555	<i>VIPR1</i>	.000392	A	rs6072296	<i>PLCG1</i>	.001215	D	rs17860266	<i>IL10RB</i>	.000559	A
rs3087209	<i>IL2</i>	.001225	D	rs2336384	<i>MFN2</i>	.001395	A	rs581950	<i>PI3K</i>	.001202	A
rs3822430	<i>SRD5A1</i>	.001728	R	rs28368160	<i>IFNA16</i>	.001505	D	rs28691160	<i>CALM1</i>	.001378	A
rs35835913	<i>CBLB</i>	.002254	A	rs34417936	<i>IL6ST</i>	.002258	D	rs3217921	<i>CCND2</i>	.001664	A
rs149698066	<i>BLVRB</i>	.002581	D	rs1045000	<i>AP-1</i>	.002473	A	rs35835913	<i>CBLB</i>	.001664	A
rs34417936	<i>IL6ST</i>	.002914	A	rs6723506	<i>UGT1A1</i>	.003189	D	rs72862555	<i>VIPR1</i>	.001671	A
rs2336384	<i>MFN2</i>	.004046	A	rs10849023	<i>CCND2</i>	.003374	R	rs35504537	<i>SOCS4</i>	.003172	A
rs10076470	<i>SRD5A1</i>	.005248	A	rs8179183	<i>LEPR</i>	.003642	A	rs75887164	<i>P2RX1</i>	.003301	A
rs166049	<i>SRD5A1</i>	.005722	A	rs12340895	<i>JAK2</i>	.005952	A	rs140463378	<i>TBXAS1</i>	.004211	A
rs12628803	<i>EP300</i>	.006147	A	rs3737224	<i>PEAR1</i>	.006491	R	rs149698066	<i>BLVRB</i>	.00428	D
rs3217921	<i>CCND2</i>	.006147	A	rs72862555	<i>VIPR1</i>	.006491	D	rs7689953	<i>TEC</i>	.004382	A
rs8191984	<i>SHP2</i>	.00647	A	rs1008084	<i>CCDC162P</i>	.008285	R	rs4252249	<i>IL10RA</i>	.004565	D
rs2066809	<i>STAT2</i>	.006477	A	rs1800797	<i>IL6</i>	.008654	R	rs1051442	<i>THBS1</i>	.004586	R
rs28368160	<i>IFNA16</i>	.007803	D	rs12343867	<i>JAK2</i>	.008962	A	rs2572207	<i>DENND4A</i>	.004676	A
rs7732059	<i>GHR</i>	.010701	D	rs3780367	<i>JAK2</i>	.009638	A	rs35695978	<i>HSD17B4</i>	.004871	A
rs4252249	<i>IL10RA</i>	.011014	A	rs904011	<i>FDFT1</i>	.010497	D	rs13244259	<i>WBSCR22</i>	.005213	A
rs1800797	<i>IL6</i>	.011145	R	rs166049	<i>SRD5A1</i>	.011011	A	rs74474807	<i>GNAS</i>	.006456	A

The top SNVs (N = 20 per cohort) rank-ordered according to P value.

*Genetic model applied for association testing: allelic (A), dominant (D), and recessive (R).

equimolar concentrations (50 ng/mL) of IFN- α 16^{WT} or IFN- α 16^{MUT}. Primary antibodies were directed against phosphorylated STAT1 (Y701) and total STAT1 (Cell Signaling Technology, Danvers, MA), or α -tubulin (as loading control; MilliporeSigma, Burlington, MA), followed by immunodetection using appropriately conjugated secondary antibodies.

Hematopoietic and cellular studies

Human CD34⁺ HSCs obtained from umbilical cord blood and CD34⁺ immunoselection contained >95% CD34⁺ cells at the start of individual experiments. CD34⁺ cells were cultured in serum-free expansion medium II expansion medium for 24 to 48 hours followed by terminal MK differentiation using serum-free expansion medium media containing MK expansion supplement CC220 (Stemcell Technologies, Vancouver, BC, Canada), supplemented (or not) every 48 hours with individual IFNs (500 U/mL human universal type I IFN- α 2), or equimolar concentrations (50 ng/mL) of IFN- α 16^{WT} or IFN- α 16^{MUT}. At distinct time points, cells were stimulated (or not) with the same concentration of IFNs for 60 minutes, followed by RNA isolation and quantitation using fluorescence-based real-time polymerase chain reaction technology (TaqMan Real-Time PCR; Applied Biosystems, Foster City, CA).³³ Oligonucleotide primers are provided in supplemental Table 2. Cell differentiation was monitored by flow cytometry, gating on live 7-actinomycin D–negative cells for immunophenotypic quantification and lineage specification, using fluorescein isothiocyanate–conjugated anti-CD41a (integrin α _{IIb}, MK) or Annexin V as a marker of phosphatidylserine exposure and apoptosis.^{34,35}

Computation and bioinformatics

Pathway enrichment³⁶ and functional interactive networks were generated by using the set of phenotype-restricted genetic variants associated with ET or RT. Because multiple SNVs were variably represented within the same gene, interactive networks were built by using the global adaptive rank truncated product method, which allows for different modes of inheritance and multiple SNVs associated with individual genes.³⁷ Sources for construction of the interactive networks included well-curated databases (Reactome, Kyoto Encyclopedia of Genes and Genomes, and Human PPI), which were trained and validated by using a naive Bayes classifier.

All statistical analyses, including gene expression, cytokine quantification, and principal component analysis (PCA), were conducted in the R program environment as previously described,^{9,34} using normalized (log₂-transformed) data and clustering (on Euclidean space) for cross-group comparisons, defining a significance level of $P < .05$ adjusted for the false discovery rate.²⁸

Additional detailed Methods are provided in the supplemental Materials and methods.

Results

Targeted gene/SNV identification

We dissected the role of cytokine pathway SNVs as genetic modifiers of enhanced thrombopoiesis by generating a custom SNV array incorporating 1056 SNVs within 326 genes collectively

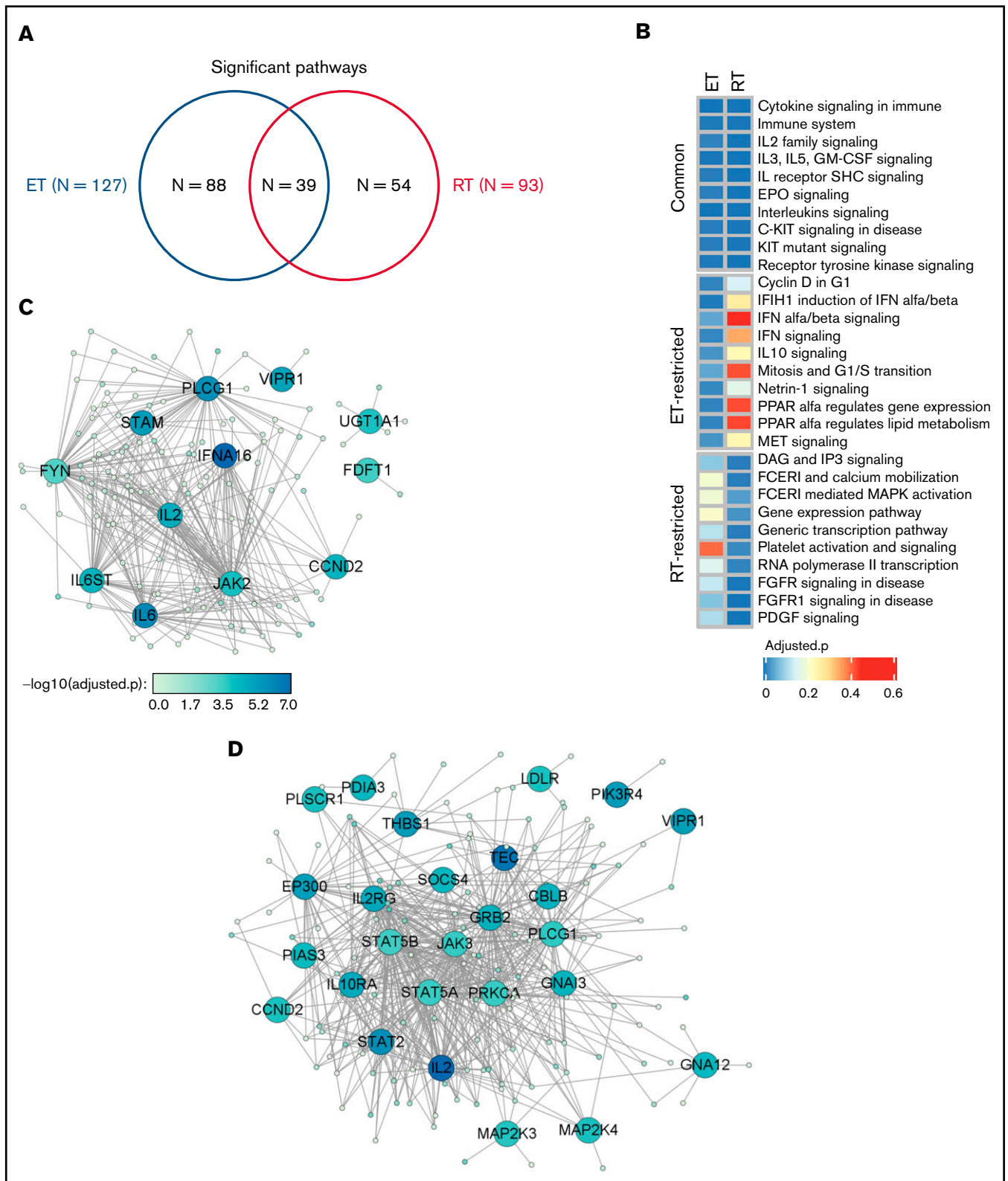
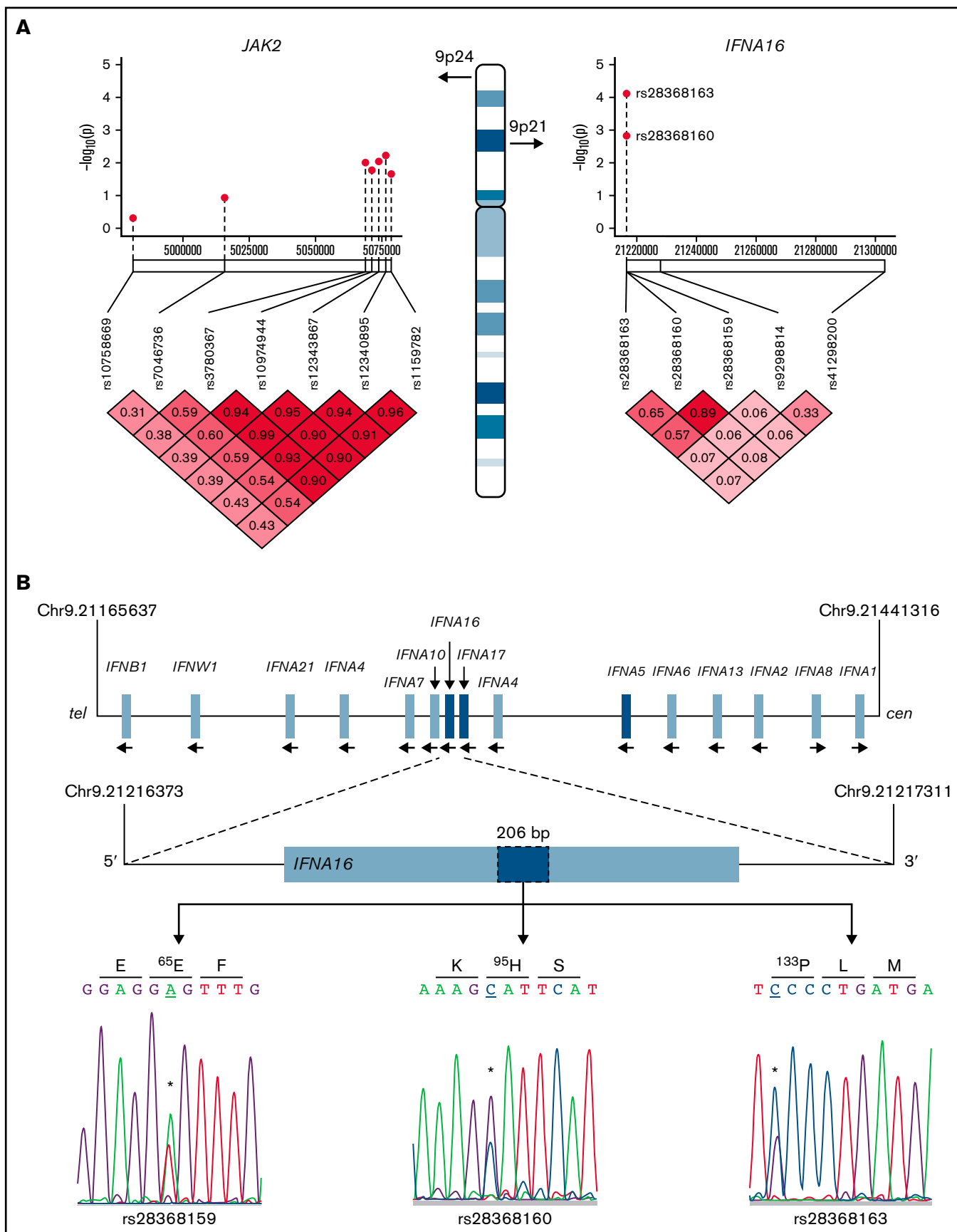


Figure 2. Pathway topology and functional interaction networks in ET and RT. (A-B) Venn diagram delineating pathway overrepresentation analysis using significantly associated ($P < .05$) ET and RT gene/SNVs (A) and top 10 pathways each for common, ET-, or RT-restricted phenotypes (B). Scale bar (adjusted. P) is shown. (C-D) Functional interaction networks in ET and RT were generated by applying global adaptive rank truncated product method to the SNV association data.³⁷ For the nodes, genes are represented by round symbols, and node color reflects the adjusted P value²⁸ on a continuous scale from low (light green) to high (dark blue). Gray-colored round symbols identify first-order interactors, and core genes not connected to the main network are omitted. The ET network includes 12 core genes (encompassing 127 nodes and 362 edges), and the RT network includes 26 core genes (encompassing 150 nodes and 633 edges). Scale bar [$-\log_{10}(\text{adjusted}.P)$] is shown. FGFR, fibroblast growth factor receptor; PDGF, platelet-derived growth factor.



implicated in hematopoietic cytokine receptor signaling. The non-redundant gene/SNV list was modified to incorporate quantitative trait loci associated with parameters of platelet function²⁷ or megakaryocytopoiesis,^{18,26} and included SNVs within the *JAK2* susceptibility locus,¹⁰ which served as known association controls for ET predisposition. Of the 1056 SNVs, 949 encompassed within 311 genes passed technically required screening characteristics using Illumina chips (San Diego, CA), and they conformed to Hardy-Weinberg equilibrium predictions based on SNV distributions formulated from the 1000 Genomes database (Figure 1A). Overrepresentation analysis using the curated gene list confirmed enrichment of genetic pathways involved in immune regulation, signal transduction, development, and hemostasis (Figure 1B).

Genotype association studies

Genotyping was initially completed on 224 healthy control subjects and 108 subjects with thrombocytosis. Thrombocytosis cohorts were phenotypically categorized according to etiology as a means of further establishing risk stratified by reactive (RT, $n = 56$) or MPN-associated (ET, $n = 52$) subtypes. Case-control association studies using dominant, allelic, or recessive models applied to the aggregate (ie, ET and RT) data set showed the strongest associations with immune-regulated (*IFNA16* rs28368163) or cytokine (*IL6* rs2069824, *IL10RB* rs17860266) pathway SNVs (Figure 1C-E; Table 1; supplemental Table 3). Associations with *IL6* rs2069824 and *IFNA16* rs28368163 reached greater statistical significance in the ET cohorts than in the aggregated cohort; technical validation of the most significant *IFNA16* rs28368163 SNV was confirmed by pyrosequencing. Not surprisingly, the RT cohort displayed gene/SNV associations that were distinct from ET, most significant for *SHP2* rs8191984, *EP300* rs12628803 (validated by pyrosequencing), *STAT2* rs2066809, and *IL10RB* rs17860266; *IL10RB* rs17860266 identified in the aggregate cohort retained its strongest association in the RT (and not ET) cohort. The protein tyrosine phosphatase SHP2 (PTPN11), adenovirus E1A-associated cellular transcriptional coactivator EP300, and STAT2 collectively regulated cellular proliferation, with evidence that EP300 functionally regulates megakaryocytopoiesis in *mpl*^{-/-} (myeloproliferative leukemia protein) mice.³⁸

Pathway topology identifies functionally diverse thrombocytosis interaction networks

The initial association data imputed the presence of distinct pathways regulating enhanced platelet formation in pathogenetically divergent thrombocytosis phenotypes; these results were confirmed by pathway enrichment analyses³⁶ applied to the ET and RT gene sets. The phenotypes converged on a limited subset of overlapping pathways ($n = 39$) encompassing immune and cytokine signaling (Figure 2A-B), although the most significant RT- or ET-restricted pathways were distinct. Pathways enriched in RT (but not ET)

included platelet-derived growth factor, fibroblast growth factor, and calcium signaling, whereas the 10 most significant ET-restricted pathways included IFN signaling ($n = 3$) or regulation by peroxisome proliferator-activated receptor α ($n = 2$).

Because genes rarely act alone, we applied the gene/SNV data to reconstruct synergistic effects of variants on functional interaction networks defining thrombopoiesis in RT and ET (Figure 2C-D). The ET network incorporates a smaller subset of core genes ($n = 12$) compared with the RT network ($n = 26$), implying that molecular pathways regulating thrombopoietic stimuli in RT display greater redundancy and are less centralized. Predicated on the network plot, ET-regulated thrombopoiesis appears to be largely controlled by integrated cytokine interactions involving *IL2*, *IL6* (and its intracellular effector *IL6ST*), and immune mediators (ie, IFNs) that presumably synergize to amplify the thrombopoietic signaling repertoire linked to *JAK2*.

An *IFNA16* susceptibility genotype in ET

Type 1 IFNs include 17 subtypes that encompass 13 IFN- α isoforms,³⁹ whose genes are located within the IFN gene cluster on chromosome 9 (Figure 3). Two non-synonymous *IFNA16* SNVs rs28368163 (397G>C, Ala¹³³Pro) and rs28368160 (283G>C, Asp⁹⁵His) were significantly associated with both aggregate and ET phenotypes but not with the RT phenotype; a contiguous *IFNA16* SNV rs28368159 (194T>A, Val⁶⁵Glu) included in the genotyping data were excluded from the association testing because of failed Hardy-Weinberg equilibrium. The SNV chip included 2 additional markers within the IFN gene cluster (*IFNA17* [rs9298814] and *IFNA5* [rs41298200]), allowing for association testing across these 5 markers; comparative testing across the *JAK2* susceptibility locus served as a genetic standard for MPN risk (Figure 3A). The three *IFNA16* markers (but neither centromeric *IFNA5* nor *IFNA17* SNVs) were in linkage disequilibrium, results confirmed by dideoxy sequencing of the ET cohort (Figure 3A-B); unavailability of parental genotyping data did not allow us to assign haplotypes. *IFNA16* SNVs represented strong susceptibility alleles (Table 2): the *IFNA16* rs28368163 OR was 5.40 (CI, 2.32-12.53), and the *IFNA16* rs28368160 OR was 4.71 (CI, 1.77-11.81). ORs for both SNVs were greater than those of the *JAK2* susceptibility SNVs (rs12340895 [OR, 2.13; CI, 1.14-3.97], rs12343867 [OR, 2.07; CI, 1.11-3.91], rs3780367 [OR, 1.95; CI, 1.04-3.65], rs10974944 [OR, 1.89; CI, 1.01-3.56], and rs1159782 [OR, 1.84; CI, 0.99-3.46]) and were in excellent agreement to those previously described (supplemental Table 4).¹⁰⁻¹² RT was not associated with *JAK2* predisposition alleles (or with *JAK2*^{V617F}). *IFNA16* genotyping substratified according to *JAK2*^{V617F} status exhibited predisposition risk restricted to *JAK2*^{V617F}-negative ET cohorts, evident for both rs28368163 (OR, 8.60; CI, 2.75-26.94; $P = 2.01 \times 10^{-5}$) and rs28368160 (OR, 7.84; CI, 2.35-26.21; $P = .0001$).

Figure 3 (continued) IFNA16 genetic characterization. (A) Chromosome 9 structure shows *JAK2* and *IFNA16* genomic regions and SNVs used for association analyses in the P value ($-\log_{10}$) scatter plots ($n = 51$). Note the stronger association of both *IFNA16* SNVs compared with those of *JAK2* SNVs. The accompanying linkage disequilibrium heatmap was generated by using pairwise χ^2 test for 7 evaluable *JAK2* SNVs or 5 evaluable *IFNA* SNVs (3, *IFNA16*; 1, *IFNA17*; 1, *IFNA5*) using the MPN genotyping data. Boxes indicate D values for SNV pairs. (B) Genomic organization of the IFN gene cluster and *IFNA16* dideoxy sequence analysis delineating the 3 SNVs and their predicted translation products (all displayed 5'-3' orientation using RefSeq NM_002173.3 as reference). For all sequences, the polymorphic transition is underscored and delineated by the asterisk. Chromosomal positions are based on National Center for Biotechnology Information Build 37 (GRCh37).

Table 2. IFNA16 SNV ORs substratified according to JAK2^{V617F}

Gene	SNV	Alleles	P	OR*	95% CI		
Cohort 1 (n = 52)							
<i>IFNA16</i>	rs28368163	G/G†	GC	CC			
Aggregate		38 (74.5%)	12 (23.5%)	1 (2.0%)	2.23 × 10 ⁻⁵	5.40	2.32-12.53
JAK2 ^{V617F+}		24 (47.0%)	5 (9.8%)	1 (2.0%)	.49‡	0.26‡	0.00-14.69
JAK2 ^{V617F-}		11 (21.6%)	6 (11.8%)	0 (0%)	2.01 × 10 ⁻⁵	8.60	2.75-26.94
<i>IFNA16</i>	rs28368160	G/G	GC	CC			
Aggregate (N = 46)		37 (80.4%)	9 (17.7%)	0 (0%)	.0004	4.71	1.87-11.81
JAK2 ^{V617F+}		25 (54.4%)	4 (7.8%)	0 (0%)	.36‡	0.18‡	0.00-10.09‡
JAK2 ^{V617F-}		12 (26.1%)	5 (9.8%)	0 (0%)	.0001	7.84	2.35-26.21
Cohort 2 (n = 54)							
<i>IFNA16</i>	rs28368163	G/G	GC	CC			
Aggregate		42 (77.8%)	12 (22.2%)	0 (0%)	.0002	4.51	1.92-10.56
JAK2 ^{V617F+}		25 (46.3%)	9 (16.7%)	0 (0%)	.62‡	0.37‡	0.00-20.1‡
JAK2 ^{V617F-}		17 (31.5%)	3 (5.6%)	0 (0%)	.12	2.78	0.72-10.73
Cohort 1+cohort 2 (n = 105)							
<i>IFNA16</i>	rs28368163	G/G	GC	CC			
Aggregate		80 (76.2%)	24 (22.9%)	1 (0.96%)	3.23 × 10 ⁻⁶	4.92	2.40-10.10
JAK2 ^{V617F+}		49 (46.7%)	14 (13.3%)	1 (0.96%)	.31‡	0.55‡	0.00-16.45‡
JAK2 ^{V617F-}		28 (26.7%)	9 (8.6%)	0 (0%)	.0002	5.01	1.98-12.94

*The χ^2 test using genotyped controls (N = 218).

†C, minor (variant) allele; G, major allele.

‡Haldane-Anscombe correction.

For confirmation, we applied *IFNA16* rs28368163 as a tag SNV in a second ET validation cohort (n = 54) and reconfirmed a highly significant association ($P = .0002$) and OR (4.51; CI, 1.92-10.56), nearly identical to that found in Cohort 1 (Table 2). Consistent with results in Cohort 1, substratification revealed no association in *JAK2^{V617F+}* cohorts; in contrast, association within the *JAK2^{V617F-}* negative cohort was less robust, approaching statistical significance ($P = .12$) with a less robust OR (2.78; CI, 0.72-10.73). Combined *IFNA16* rs28368163 data incorporating both cohorts (n = 105 subjects) confirmed the overall association (OR, 4.92; CI, 1.92-10.56) and the specificity with *JAK2^{V617F-}* negative ET (OR, 5.01; CI, 1.98-12.94). These collective data showed that: (1) *IFNA16* SNVs represent significant ET predisposition alleles¹⁰; and (2) ET susceptibility was largely restricted to *JAK2^{V617F-}* negative ET. These results were distinct from those previously described for the *JAK2* predisposition haplotype, which selectively confers hypermutability to *JAK2^{V617F}* expansion.¹⁰⁻¹²

Comparative cytokine profiles across the thrombocytosis spectrum

We characterized RT and MPN-associated “inflammasomes” by measuring 15 cytokines in a randomly selected subset of ET (n = 20), RT (n = 18), and healthy cohorts (n = 14), focusing on IFNs (α -, β -, γ -), various interleukins (ILs), and components of the tumor necrosis factor (TNF) receptor pathway previously identified in thrombosis-related inflammation^{29,30} or thrombocytosis.^{29,31,32} Unsupervised hierarchical clustering using the aggregate cytokine data confirmed general segregation according to phenotype (Figure 4A), with statistically significant cross-group

differences for the majority (13 of 15) of cytokines measured. PCA designed to reduce the data dimension revealed overlapping clusters of healthy control and RT cohorts, and relative segregation of ET cohorts that were broadly distributed along the PCA spectrum (Figure 4B). ET cytokine determinations generally displayed a broader intersample dynamic range than those from RT or healthy control subjects. Compared with both RT and healthy control subjects, cytokines were generally elevated in ET subjects, and pairwise comparisons established that 11 of 15 cytokines were statistically different between the ET and RT cohorts (Figure 4C). Binary logistic Least Absolute Shrinkage and Selection Operator regression reduced the cytokine profile to a 4-member subset (CD30, IL-10, TNF receptor 1, and TNF- α) that optimally distinguished ET from healthy cohorts, and a distinct 4-member subset that distinguished ET from RT (Tweak, IL1RA, TSLP, and TNF receptor 2 cytokine). Within the ET cohort, repeat PCA substratified according to *IFNA16* rs28368163 or *JAK2^{V617F}* genotypes failed to generate meaningful separation (Figure 4D-E), imputing the lack of *IFNA16*- or *JAK2^{V617F-}* restricted mean differences for the cytokines studied. Thus, although the cytokine profiles are clearly different between ET and RT, these data further established that neither *JAK2^{V617F-}* nor *IFNA16* rs28368163 independently partitioned the cytokine abnormalities evident in ET.

Evolutionary duplication of 13 highly homologous IFN- α subtypes results in a largely redundant mechanism for viral neutralization and cellular effects. We saw no differences in aggregate serum IFN- α levels in ET cohorts using a multiplex bead assay (discussed earlier)

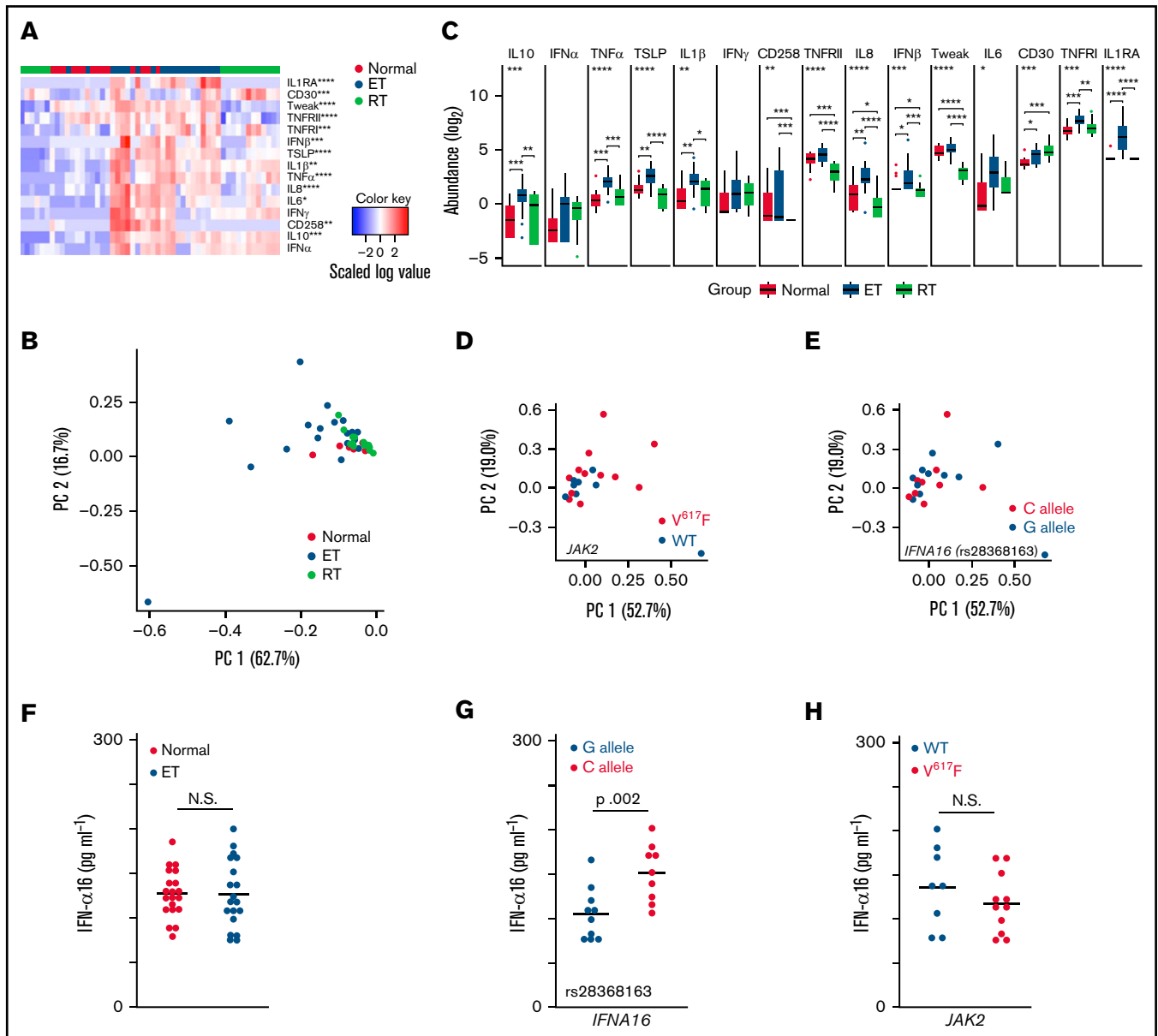


Figure 4. Cytokine studies in thrombocytosis. (A) Cytokines ($n = 15$) were quantified from ET ($n = 20$), RT ($n = 18$), or healthy controls ($n = 14$), and the normalized data (centered and scaled) were used for unsupervised hierarchical clustering using aggregated data; P values corresponding to group differences were determined by using Kruskal-Wallis tests; scale bar is shown. (B) PC analysis was applied to cytokine levels for data reduction, and iteratively applied to ET, RT, or healthy cohorts; note the general segregation of ET from RT and healthy controls. (C) Box plots show normalized cytokine levels by cohort; group-wise (upper panels) and statistically different pairwise P values are shown. (D-E) PC analysis applied to cytokine levels was iteratively applied to ET cohorts delineated by *JAK2*^{V617F} (D) or *IFNA16* rs28368163 genotype (E). In both panels D and E, mutant alleles are in red. (F-H) IFN- α 16 was quantified by enzyme-linked immunosorbent assay from ET ($n = 19$ [1 of 20 samples was censored for a technical limitation]) and healthy controls ($n = 20$), substratified according to *IFNA16* or *JAK2*^{V617F} genotype. Mutant alleles in panels G and H are in red. P values using the t test are shown. * $P < .05$, ** $P < .01$, *** $P < .001$, **** $P < .0001$. *CD258*, tumor necrosis factor superfamily member 14; *CD30*, TNF receptor superfamily member 8; *IFN*, IFNs α , β , γ ; *IL*, interleukins 6, 8, 10, or 1 β ; *IL1RA*, IL-1 receptor antagonist; N.S., not significant; *TNF α* , tumor necrosis factor α ; *TNFR1*, TNF receptor type 1 cytokine; *TNFR2*, TNF receptor 2 cytokine; *TSLP*, thymic stromal lymphopoietin; *Tweak*, TNF-like weak inducer of apoptosis.

but expanded these studies using an enzyme-linked immunosorbent assay designed to specifically quantify IFN- α 16 (Figure 4F-H). Although IFN- α 16 levels between ET and normal were nearly identical, substratification according to *IFNA16* rs28368163 revealed statistically greater concentrations in ET cohorts expressing the

mutant C allele ($P = .002$); in contrast, IFN- α 16 differences substratified according to *JAK2*^{V617F} were not evident. These results established that among the cytokines quantified, only ET IFN- α 16 concentrations were associated with the presence of the mutant *IFNA16* SNV.

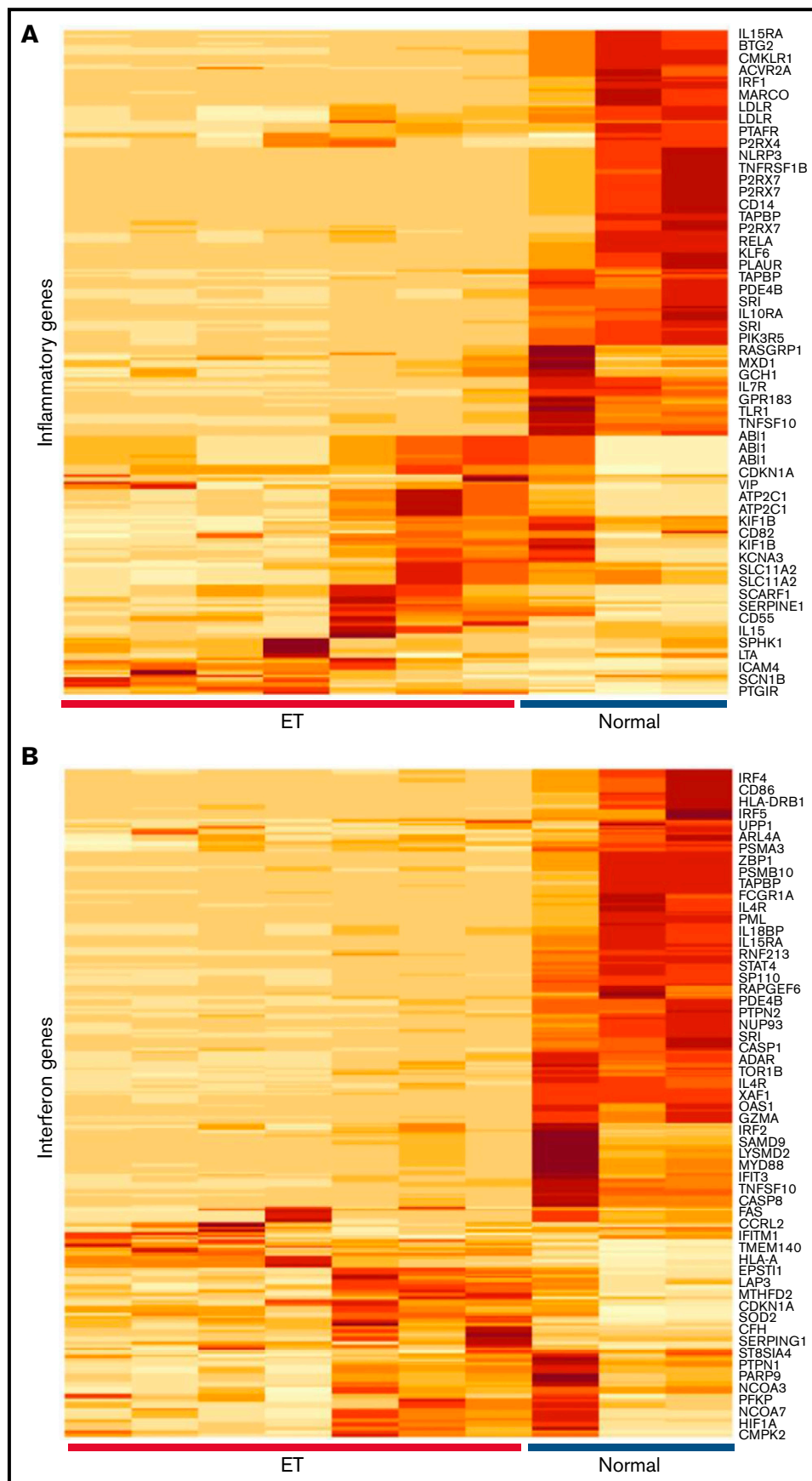


Figure 5. Characterization of inflammatory gene profiles in platelets. (A-B) RNA-Seq data from ET (N = 7) or healthy control (N = 3) platelets³⁵ were filtered by using Hallmark⁴⁰ inflammatory response (A, n = 200) or IFN response (B, n = 224) genes, followed by unsupervised hierarchical clustering by Euclidean distance

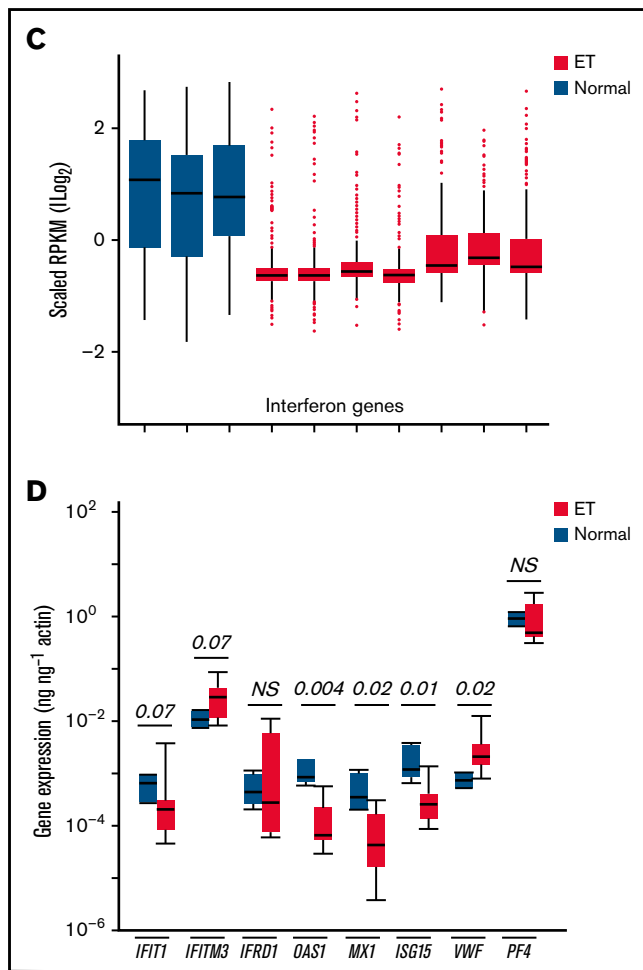


Figure 5 (continued) (differentially expressed genes are shown). Note clear segregation of phenotypes in both panels. (C) In silico expression studies of aggregated IFN response genes ($n = 224$) by phenotype (normal, $n = 3$; ET, $n = 7$); data are expressed as mean \pm standard error of the mean (SEM) of reads per kilobase per million (RPKM) normalized and scaled to mean 0. (D) Platelet quantitative polymerase chain reaction of select IRGs or platelet-specific genes (*VWF* and *PF4*) from a secondary cohort of healthy controls ($n = 4$) or ET ($n = 8$), expressed as mean \pm SEM normalized to actin; box plots (panels C-D) represent the within-group interquartile range encompassing 50% of the values, whereas the 95% CIs and outliers are depicted. The horizontal bar within each box represents the group median. P values (t test) are shown. NS, not significant.

Expression studies of IFN-response genes in ET platelets

Given the evidence for immune and cytokine disruption in ET, we applied inflammatory and IFN-restricted gene sets⁴⁰ to initially characterize downstream cellular effects on ET platelet signatures using previously generated RNA-Sequencing (RNA-Seq) data³⁵ (Figure 5A-B); these initial analyses were designed to address putative cellular responses in targets (MKs/platelets) exposed to altered cytokine signaling. Unsupervised hierarchical clustering showed clear segregation of ET and healthy control platelets using both gene sets, with further evidence for downregulation of IFN genes in ET platelets (Figure 5C). For a subset of

IFN-regulated genes (IRGs) (Figure 5D),^{41,42} we validated these patterns in a second unrelated cohort (ET, $n = 8$; normal, $n = 4$) using a gene subset that is induced during stress hematopoietic models (*IFIT1*, *IFITM3*, *MX1*, *ISG15*, *IFRD1*, and *OAS1*).⁴ Consistent with the RNA-Seq data, the majority (four of six IRGs [*IFIT1*, *OAS1*, *MX1*, and *ISG15*]) were downregulated in ET compared with normal platelets ($P = .07$ to $P = .004$); only *IFITM3* was upregulated in ET (2.8-fold) and approached statistical significance ($P = .07$). We saw no evidence for differential platelet factor 4 (*PF4*) expression, although *VWF* (von Willebrand factor) as a marker of MK-biased stress hematopoiesis identified in inflammatory disorders⁴ was upregulated in ET (4.4-fold; $P = .02$). These genetic expression studies established that the dysfunctional ET cytokine profiles were associated with concomitant target cellular effects in MKs/platelets, with further evidence for defective IFN IRG signatures, possibly mediated by IFN- α 16.

Comparative studies of IFN- α 2 and IFN- α 16

IFNs are functionally characterized by their antiviral and antiproliferative properties, and IFN- α cellular responses are mediated by binding to a common IFNAR1/IFNAR2 heterodimeric receptor complex.^{39,43} To study putative cellular effects mediated by IFN- α 16, we used a tandem affinity tag to express and purify wild-type ($V^{65}D^{95}A^{133}$) and mutant IFN- α 16 as recombinant proteins in *E. coli*. Because *IFNA16* mutant SNVs were found in linkage disequilibrium ($T^{194}A$, $G^{283}C$, $G^{397}C$) (Figure 3), mutant IFN- α 16 was specifically designed to incorporate the triple mutation ($E^{65}H^{95}P^{133}$) in the IFN- α 16 backbone (supplemental Figure 1A). Both wild-type (IFN- α 16^{WT}) and mutant (IFN- α 16^{MUT}) exhibited comparable solubility and purification characteristics as *E. coli* recombinant proteins, excluding the likelihood that $E^{65}H^{95}P^{133}$ mutant affected protein stability. An adenoviral replication system designed to quantify antiviral properties exhibited comparable inhibitory effects between wild-type and mutant IFN- α 16, and antiviral properties were functionally equivalent to that of clinically used recombinant human IFN- α 2a (Hu-IFN- α 2a) (supplemental Figure 1B). Functional integrity of IFNAR1/IFNAR2 signaling was established by using both A549 lung epithelial cells (not shown) and HDF/*Tert1* skin fibroblasts (supplemental Figure 1C), with no differences in STAT1 phosphorylation between recombinant IFN- α 16^{WT} and IFN- α 16^{MUT}, and nearly identical to IFN- α 2. These initial data excluded the possibility that mutant IFN- α 16 represented a gain-of-function or loss-of-function mutation of key antiviral or cell-signaling mechanisms.

IFN- α 16 effects on MKs

Given the sustained cytokine disruption in ET, we applied an in vitro model system to characterize effect(s) of chronic IFN exposure on gene expression and patterns of IRG expression during megakaryocytopoiesis. CD34⁺ HSCs were differentiated in the absence or presence (supplemented every 48 hours) of IFN- α 2, IFN- α 16^{WT}, or IFN- α 16^{MUT} over a 14-day MK differentiation protocol (Figure 6A). At the concentration(s) of IFNs studied, we saw no evidence for disparate effects on CD41⁺ (integrin α _{IIb}) or Annexin V expression, the latter used as a quantitative marker of phosphatidylserine exposure that accompanies terminal megakaryocytopoiesis (Figure 6B).³⁵ To determine if the IFN-regulated gene profiles were differentially modulated by IFN isoforms, we quantified gene expression over time, applying the same IRG subset used in the ET/normal platelet expression studies (discussed earlier). In the absence of interferon,

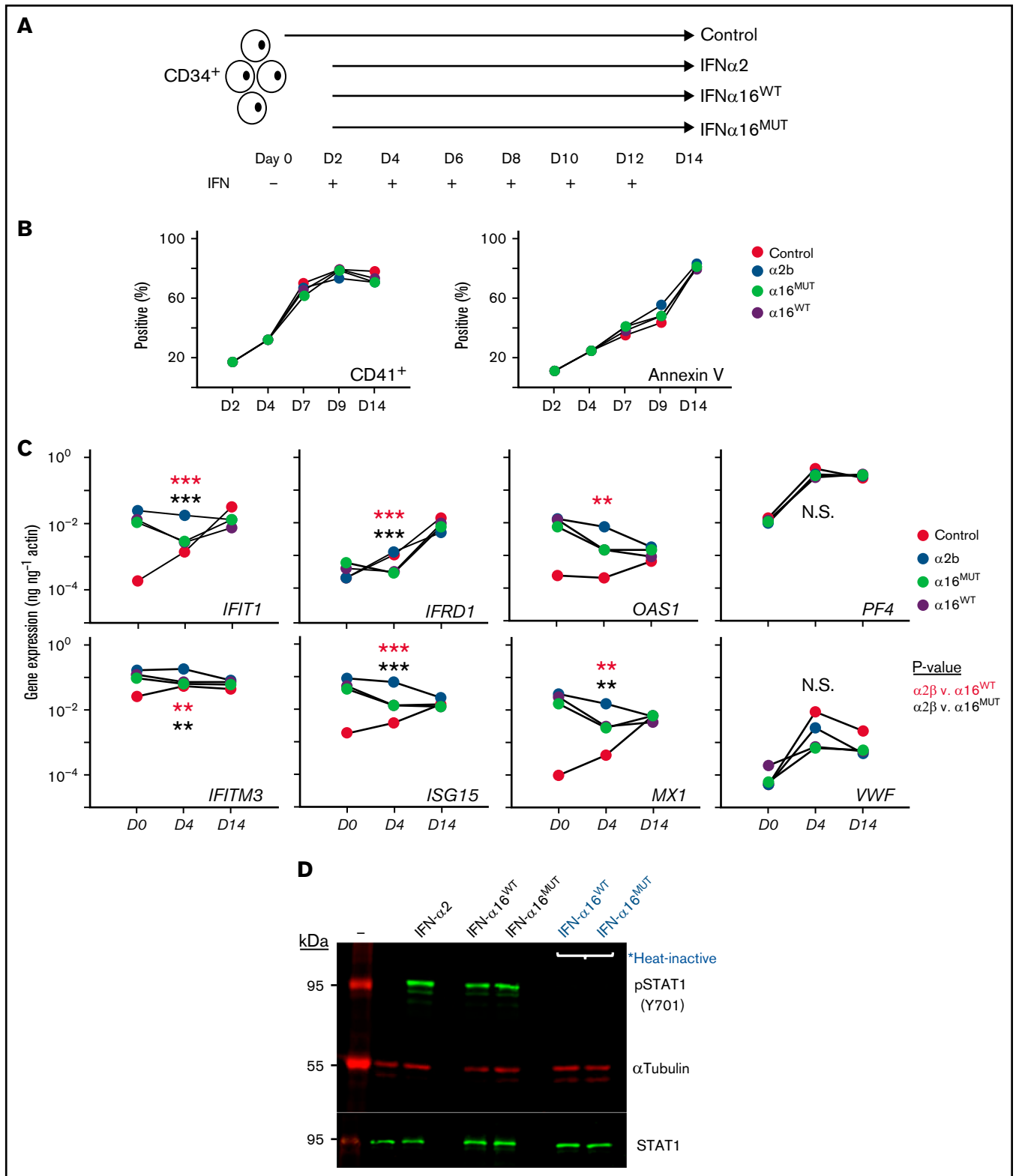


Figure 6. Functional and cellular characterization of *IFN-α2* and *IFN-α16* in hematopoietic stem cells. (A-B) CD34⁺ HSCs were expanded in TPO-conditioned media supplemented (or not) with IFNs α2b (500 U/mL) or equimolar concentrations (50 ng/mL) of IFN-α16^{WT} or IFN-α16^{MUT} at 2-day intervals (A), followed by flow cytometric quantification of CD41⁺ (integrin α_{IIb}) and Annexin V (phosphatidylserine) at designated time points (B). Data represent the mean ± standard error of the mean (SEM) from triplicate determinations of a single representative experiment, repeated once. (C) Differentiating CD34⁺ HSCs at specific time points (day 0 [D0], D4, and D14) were stimulated with IFN-α2a (500 U/mL) or equimolar (50 ng/mL) concentrations of IFN-α16^{WT} or IFN-α16^{Mut} for 60 minutes, followed by RNA isolation and

5 of 6 genes (*IFIT1*, *MX1*, *ISG15*, *IFRD1*, and *OAS1*) exhibited time-dependent induction corresponding to maximal CD41⁺ positivity at day 14 (Figure 6C). IRG induction was most striking for *IFIT1* (175-fold), *IFRD1* (65-fold), and *MX1* (65-fold); the most abundant gene (*IFITM3*) exhibited minimal (1.7-fold) change.

To determine if there were stage- or isoform-restricted differences after IFN stimulation, cells were stimulated with corresponding IFNs for 60 minutes, and IRG responses were quantified at baseline (day 0), at an early stage of MK differentiation (day 4), or at terminal MK differentiation (day 14). Across the IFN isoforms ($\alpha 2$, $\alpha 16^{WT}$, and $\alpha 16^{MUT}$), maximal and comparable gene induction was evident at day 0, with less robust responses at day 14; indeed, day 14 gene induction was minimal for all IRGs and essentially comparable to maximal expression evident during terminal differentiation in the absence of IFN. At a time corresponding to modest CD41⁺ acquisition (~30% at day 4), HSCs retained their capacity for IFN induction, although we noted clear differences in IRG responses between the $\alpha 2$ and $\alpha 16$ isoforms. Although gene induction by $\alpha 16^{WT}$ and $\alpha 16^{MUT}$ was nearly identical, both responses were consistently attenuated compared with that of the $\alpha 2$ isoform. These differences were not due to accompanying divergence in pSTAT phosphorylation (Figure 6D), confirming the presence of a functional IFNAR axis, and excluding the possibility of cell desensitization with continuous IFN exposure; heat-inactivated IFN- $\alpha 16$ (WT and MUT) failed to induce pSTAT1 phosphorylation, establishing specificity of the IFN- α /IFNAR signaling axis, and suggesting progressive (or transient) loss of pSTAT1 signaling during cell differentiation in the absence of an acute IFN stimulus. These collective data provide evidence for divergent IFN- α isoform- and stage- restricted IRG responses during MK lineage speciation, and they are consistent with prior evidence for a STAT-independent mechanism regulating IRG responses.^{4,3}

Discussion

We have identified a subset of gene/SNVs associated with thrombocytosis that provides a framework for dissecting molecular mechanisms of enhanced platelet formation during inflammatory triggers. These interactions are distinct from those during steady-state thrombopoiesis, and they provide insight into genetic variants that regulate disparate platelet responses to stress in the general population. We applied the SNV data to better define epistatic interactions in thrombocytosis, with evidence for both overlapping and functionally diverse pathways that comprise the ET and RT interactive networks. These distinct genetic networks are consistent with the pathophysiologically divergent mechanisms of enhanced thrombopoiesis in clonal and stress thrombocytosis. In contrast to rare single-gene defects resulting in predominant phenotypes (ie, hemophilia and sickle cell disease), complex phenotypes such as thrombopoietic responses likely result from epistatic gene–gene interactions exerting synergistically additive effects within overlapping pathways⁴⁴;

these data provide a framework for further dissecting the complex interplay of regulatory genetic networks linked to inflammatory and clonal thrombopoiesis.

The majority of cytokines were statistically different in cross-group comparisons (13 of 15), and non-overlapping 4-member subsets effectively distinguished ET from normal (CD30, IL10, TNF receptor 1, and TNF- α) or ET from RT (Tweak, IL1RA, TSLP, and TNF receptor 2 cytokine). These observations highlight the important pathophysiological differences in inflammatory responses in phenotypically divergent thrombocytosis models. There is considerable interest in identifying polymorphic changes that influence cytokine levels in various disease states,⁴⁵ although literature review using the 48-member ET-associated SNV list (supplemental Table 3) failed to implicate these gene/SNVs as modulators of cytokine levels in ET; similarly, we saw no differences in cytokine profiles according to age or sex. Interestingly, gene/SNVs involving both *IL6* and *IL6ST* (signal transducer encoding the IL-6 β subunit) were prominently identified in the ET network but not in the RT network where IL-6 has been identified as a driver of stress thrombopoiesis.⁵ IL-6 is nonselectively elevated in a number of inflammatory (and malignant) conditions,⁴⁵ although *IL6* polymorphisms (including those in the promoter) inconsistently influence serum levels or confer disease risk in various disease states⁴⁶⁻⁴⁹; thus, the lack of *IL6* SNV associations in RT is not unexpected. Consistent with prior data,⁶ serum IL-6 was statistically different among the 3 phenotypes (and highest in ET), and the *IL6* SNV *rs1800797* associated with ET has been associated with higher serum IL-6 levels, putatively linking cytokine levels with genetic susceptibility.⁵⁰ Interestingly, RT was associated with both *IL10RA* (*rs4252249*) and *IL10RB* (*rs17860266*) SNVs, which are encompassed within both polypeptide chains (α and β) of the IL-10 receptor, and known to be required for IL-10 signaling. Although IL-10 is largely considered an anti-inflammatory cytokine (and statistically different in cross-group comparisons), data using recombinant human IL-10 show that it retains pro-inflammatory effects in infection⁵¹ and Crohn's disease,⁵² with putative adverse effects on platelet production.⁵³ Neither IL-10 nor its cognate receptor complex is studied in the context of RT, although associations with both *IL10RA* and *IL10RB* SNVs impute previously uncharacterized functions relevant to inflammation, receptor signaling, and thrombopoiesis.

Our initial genetic studies identified polymorphic *IFNA16* as a novel susceptibility allele in aggregate (ET and RT) thrombocytosis. The *IFNA16* association was most exaggerated in the ET cohort (to the exclusion of RT), which also revealed overrepresentation of IFN-regulated pathways. These data extend previous GWAS in MPN cohorts that implicate various SNVs in risk susceptibility,⁵⁴⁻⁵⁷ pathogenesis,⁵⁸ or prognosis,^{59,60} although the majority of loci conferring familial MPN risk remain unidentified.⁵⁷ Limitations of our study are the relatively small sample size compared with aggregated and

Figure 6 (continued) quantification of selected IRGs (*IFIT1*, *IFITM3*, *IFRD1*, *OAS1*, *MX1*, and *ISG15*) or MK-specific genes (*VWF* and *PF4*). Data are from a single representative experiment (mean \pm SEM from triplicate determinations normalized to actin), repeated once; statistically significant differences ($\alpha 2$ to either $\alpha 16^{WT}$ [red asterisks] or $\alpha 16^{MUT}$ [black asterisks]) only seen at D4 are shown (*t*-test; ***P* < .01, ****P* < .001). (D) D4 HSCs were stimulated for 60 minutes with IFN- $\alpha 2a$ (500 U/mL) or equimolar (50 ng/mL) concentrations of IFN- $\alpha 16^{WT}$ or IFN- $\alpha 16^{Mut}$, followed by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and pSTAT1 detection by immunoblot analysis (20 μ g/lane); α -tubulin and total cellular STAT1 are shown as loading controls. Heat-inactivated IFN- $\alpha 16^{WT}$ and IFN- $\alpha 16^{MUT}$ exhibit no pSTAT1 activation. N.S., not significant.

long-term data reported for the *JAK2* susceptibility locus.¹⁰⁻¹² Nonetheless, our conclusions are supported by a sequential study design that incorporated an initial and secondary (validation) cohort for independent confirmation. Unlike the *JAK2*-susceptibility haplotype, the risk was greater in the initial subset of *JAK2*^{V617F}-negative cohorts, although substratification data from the validation cohort were not as robust as those from the initial cohort. Aggregated *IFNA16* rs28368163 data incorporating both cohorts (n = 105 subjects) confirmed the overall association (OR, 4.92) and the specificity with *JAK2*^{V617F}-negative ET (OR, 5.01). Predicated on these initial studies, variant *IFNA16* represents a stronger susceptibility allele than that previously described for SNVs within the *JAK2* locus.^{9,10,35} Because of our limited sample size, we were unable to further substratify according to *c-MPL*, *CALR*, or triple-negative MPN.

We were unable to show that variant IFN- α 16 displayed cellular or antiviral properties that diverged from those of native IFN- α 16, although plasma IFN- α 16 levels were higher in subjects expressing the mutant *IFNA16* allele. Unexpectedly, both IFN- α 16^{WT} and IFN- α 16^{MUT} displayed stage-restricted differences in IRG responses compared with therapeutically used IFN- α 2 at a confined developmental time point of MK differentiation. These results are paradoxical given the redundancy and evolutionary conservation of 13 type 1 IFN- α isoforms that collectively use the identical IFNAR1/IFNAR2 heterodimer for downstream signaling.^{39,43} Although type I IFNs share a common docking mode for receptor binding, recent data suggest that ligand discrimination occurs through distinct energetics of shared receptor contacts. Indeed, the extent of pSTAT activation as measured by tyrosine phosphorylation does not fully explain the level of gene expression,⁴³ results that are consistent with our CD34⁺-differentiated IRG data, which exhibit dichotomous IFN- α 16/IFN- α 2 genetic changes (at day 4) in the setting of comparable pSTAT patterns. Although IFN- α 2 induces complete hematologic responses in subsets of MPN patients,⁶¹ the mechanism(s) remain incompletely understood, and our data suggest that its effect may be at a discrete stage of lineage development. Whether alternative IFN isoforms display differential efficacy in IFN-responsive disorders remains speculative⁶² but supported by these collective observations.

Our data suggest that chronic IFN stimulation may be sufficient as an initiator for clonal expansion, although the complex cytokine profiles identified in ET confound linear causal relationships. Elevated IFN- α 16 levels were restricted to the subgroup expressing the *IFNA16* rs28368163 mutant (C) allele, although the mechanism(s) whereby this variant affects circulating levels remains unestablished. We did not complete expression studies in eukaryotic systems, although variant IFN- α 16 displayed comparable solubility and yield characteristics as the naive recombinant protein in *E coli*, minimizing a mechanism related to intrinsic protein stability or cellular expression as identified with polymorphic IL-6.⁴⁶⁻⁴⁹ The low minor allele frequency (~0.03) in healthy control subjects was insufficient to establish comparable effects on serum IFN- α 16 levels in otherwise healthy individuals. Inflammatory stimuli induce variable effects on peripheral platelet counts, observations that are compounded by the complex patterns of cytokine release evident in various autoimmune, inflammatory, or malignant processes. In mice, IFN (IFN- α 2) results in expansion of a genetically biased pool of MKs designed to rapidly regenerate the platelet pool,⁴ although such inflammatory models fail to

incorporate MK genetic responses in diseases of chronicity or those displaying complex patterns of cytokine release (eg, MPNs).

Variability in platelet inflammatory and IFN genes was evident in our ET cohort, consistent with transcriptomic changes in the setting of chronic cytokine dysfunction. Interestingly, evidence for an "interferogenic" platelet profile has been established in patients with systemic lupus erythematosus,⁶³ presumably due to immune complex-mediated stimulation of plasmacytoid dendritic cells resulting in type I IFN release and downstream effects on MKs. Three of the genes (*PRKRA*, *IFITM1*, and *CD69*) had concomitant increased changes in protein expression and were associated with vascular disease in systemic lupus erythematosus. Paradoxically, IFN genes were downregulated by using aggregate RNA-Seq data from an initial ET cohort, observations that were validated in a second ET cohort for an IRG subset (*MX1*, *ISG15*, *OAS1*, and *IFIT1*; $P = .004$ to $P = .07$). CD34⁺ experiments focusing on the differentially expressed ET genes confirmed that IRG subsets are generally induced during megakaryocytopoiesis. Furthermore, MKs retain the capacity for IRG induction during in vitro models of IFN supplementation, most striking in undifferentiated (day 0) CD34⁺ HSCs and progressively attenuated during terminal MK differentiation. The limited changes in IRGs with IFN stimulation evident in terminally differentiated MKs may provide an explanation for the decreased IRG expression levels evident in ET, although lack of information on differential mRNA decay rates limits firm conclusion.⁶⁴ Similarly, whether these altered expression patterns have functional effects on the platelet thrombohemorrhagic phenotype remains unestablished.²⁶ Interestingly, MKs retain the capacity for IFN- α secretion and paracrine MK stimulation of *IFITM3* (IFN-induced transmembrane 3 restriction factor), defining novel function(s) in antiviral neutralization, innate immunity, and IFN-dependent hematopoietic responses.⁶⁵ Among the IRGs characterized, *IFITM3* displays abundant and relatively stable expression in differentiating MKs, and is upregulated in ET platelets (2.8-fold; $P = .07$), imputing a comparable function of ET platelets that may regulate innate immunity and thromboinflammatory responses.

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Authorship

Contribution: D.V.G., S.W., W.Z., and W.F.B. conceptualized the study and formulated the hypothesis; S.W., D.V.G., W.Z., and W.F.B. designed the research; S.W., L.E.M., A.F., P.H., S.-Y.S., Z.L., D.V.G., W.Z., Y.H., and W.F.B. performed the research, analyzed the data, and generated images; and W.F.B. directed the overall research and finalized the manuscript preparation.

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