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New gene functions in megakaryopoiesis and platelet formation

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Abstract

Platelets are the second most abundant cell type in blood and are essential for maintaining haemostasis. Their count and volume are tightly controlled within narrow physiological ranges, but there is only limited understanding of the molecular processes controlling both traits. Here we carried out a high-powered meta-analysis of genome-wide association studies (GWAS) in up to 66,867 individuals of European ancestry, followed by extensive biological and functional assessment. We identified 68 genomic loci reliably associated with platelet count and volume mapping to established and putative novel regulators of megakaryopoiesis and platelet formation. These genes show megakaryocyte-specific gene expression patterns and extensive network connectivity. Using gene silencing in *Danio rerio* and *Drosophila melanogaster*, we identified 11 of the genes as novel regulators of blood cell formation. Taken together, our findings advance understanding of novel gene functions controlling fate-determining events during megakaryopoiesis and platelet formation, providing a new example of successful translation of GWAS to function.

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To discover novel genetic determinants of megakaryopoiesis and platelet formation, we performed meta-analyses of GWAS for mean platelet volume (MPV) and platelet count (PLT). Our analyses included 18,600 (13 studies, MPV) and 48,666 (23 studies, PLT) individuals of European descent, respectively, and up to \sim 2.5 million genotyped or imputed single nucleotide polymorphisms $(SNPs)^{1}$. Briefly, we tested within each study (Supplementary Table 1) the associations of MPV and PLT with each SNP using an additive model; we then combined these study-specific test statistics in a fixed-effects meta-analysis. To reduce the risk of spurious associations, we applied common stringent quality control filters and the genomic control method² to the meta-analysis, which shows no evidence for residual inflation of summary statistics (Supplementary Fig. 1).

A total of 52 genomic loci reaching statistical significance at the genome-wide adjusted threshold of $P \quad 5 \times 10^{-8}$ were discovered in this stage 1 analysis; 55 additional loci reached suggestive association (5×10^{-8} < P 5×10^{-6}). We tested one SNP per locus in a stage 2 analysis that included *in silico* and *de novo* replication data in up to 18,838 individuals from 12 additional studies, confirming 15 additional loci (Supplementary Table 2). One further independent locus (TRIM58) associated with PLT was identified through detection of secondary association signals. Overall, 68 independent genomic regions were associated with PLT and MPV with P 5 × 10⁻⁸, of which 52 are new and 16 were described previously in Europeans³⁻⁶ (Table 1). Of the 68 loci, 43 and 25 loci were associated significantly with PLT and MPV, respectively; 16 of them reached genome-wide significance with both traits (Supplementary Fig. 2). This partial overlap reflects the negative correlation of both traits (gender-adjusted $r = -0.49$, Fig. 1a) that results from the tight control of platelet mass ($PLT \times MPV$)⁷. The association of some loci with both PLT and MPV may reflect this negative correlation between the two traits or independent pleiotropic effects of a locus on megakaryopoiesis and platelet formation. The different statistical power at the two traits and small effect sizes at many loci reduce our power to discriminate among loci controlling MPV and PLT through analysis of platelet mass. Their testing will require the collection and analysis of PLT and MPV in large independent homogeneous cohorts. Some loci, however, have a clear-cut effect. For instance, BAK1 affects PLT specifically, compatible with its role in apoptosis and platelet lifespan.

We further tested the association of the 68 loci in 7,949 (MPV) and 8,295 (PLT) samples of south Asian and 14,697 (PLT) samples of Japanese⁸ origin. We detected substantial overlap of association signals, with effect size and direction highly concordant with findings in Europeans (Supplementary Fig. 3 and Supplementary Table 3). In the south Asian sample, 15 of the 68 (22.1%) loci were significant after adjustment for multiple testing ($P \sim 7 \times$ 10−4). In the Japanese sample, 13 of 55 (23.6%) PLT loci showed significance. Moreover, 73 of 84 (87%, South Asians) and 45 of 55 (82%, Japanese) SNPs showed associations with effect estimates directionally consistent with Europeans. Such concordance is highly unlikely to be due to chance $(P = 2.3 \times 10^{-12}$ and $P = 2.1 \times 10^{-6}$), and provides independent validation of the locus discovery in Europeans.

The 68 loci cumulatively explain 4.8% of the phenotypic variance in PLT and 9.9% in MPV, accounting respectively for average increases of 2.57×10^9 l⁻¹ PLT and 0.10 fl MPV per copy of allele. These levels of explained variance are in accordance with other GWAS of complex quantitative traits⁹. Our results indicate that many other common variants of similar or lower effect size, rare variants as well as structural variants may also contribute to the variation of both platelet traits. We used the method of ref. 10 to estimate the number of additional PLT- and MPV-associated loci having effect sizes comparable to those observed in our analysis. The method (with caveats discussed in the Supplementary Information) predicted that 137 and 81 such loci exist for PLT and MPV respectively, accounting for 9.7% and 18.3% of the total phenotypic variance.

Gene-prioritization strategies

Evidence from recent, highly powered meta-analyses suggests that the association peaks are enriched for genes controlling key underlying biological pathways^{11,12}. In our case, a large proportion of the association signals (46 out of 68) had the most significant SNP in stage 1 ('sentinel SNP') mapping to within a gene-coding region, including several key regulators of haemostasis (ITGA2B, F2R, GP1BA), megakaryopoiesis (THPO, MEF2C) and platelet lifespan (BAK1). Through an unbiased analysis of our GWAS results, we estimated that PLT-associated SNPs are significantly more likely to map to gene regions than expected by chance ($P < 0.05$, Supplementary Fig. 4), suggesting that we may prioritize the search of additional yet unknown genes controlling these processes in the associated regions. To define a univocal rule to study the enrichment of functional relationships in associated genes, we made the choice to focus on a set of 54 'core' genes selected as either containing the sentinel SNP or mapping to within 10 kb from an intergenic sentinel SNP (Table 2). This selection strategy is designed to obtain unbiased hypotheses producing interpretable biological inference for genes near the association signals, but has reduced sensitivity for genes that map further from the sentinel SNP. For instance VWF, a key regulator of haemostasis, maps to 55 kb from the sentinel SNP (Supplementary Fig. 3 and Supplementary Table 4) and is therefore not considered as a core gene. We further note that this selection strategy does not imply knowledge of the location of causative variants, which is currently incomplete. A detailed SNP survey showed that at 15 loci the sentinel SNPs either encoded, or were in high linkage disequilibrium (LD, r^2 0.8) with, a nonsynonymous variant (Supplementary Table 5); another 11 either matched or were in high linkage disequilibrium with SNPs associated with expression levels of core genes (or ciseQTLs, Supplementary Table 6), indicating that other loci may exert their effect through regulation of gene expression 13 . The validation of suggestive causative effects, as well as the identification of more complex interactions involving other genomic loci (trans eQTLs), will require a more comprehensive discovery in appropriately powered genomic data sets.

As a first effort to characterize biological connectivity among the core genes, we applied canonical pathway analyses (see [http://www.ingenuity.com\)](http://www.ingenuity.com), detecting a highly significant over-representation of core genes in relevant biological functions such as haematological disease, cancer and cell cycle (Supplementary Table 7). Encouraged by these results, we extended this effort to construct a comprehensive network of protein-protein interactions incorporating the core genes. This effort integrated information from public databases (principally Reactome and IntAct) with careful manual revision of published evidence and high-throughput gene expression data. The resulting network, which includes 633 nodes and 827 edges, showed extensive connectivity between the proteins encoded by the core genes with an established functional role in megakaryopoiesis and platelet formation and those encoded by genes hitherto unknown to be implicated in these processes (Fig. 1b).

Transcriptional patterns of core genes

We next considered whether this connectivity was also reflected in the regulation of core gene transcription, and whether expression patterns were unique to megakaryocytes. Despite high levels of correlation in gene expression between different blood cell types (median 5 0.8; median absolute deviation = 0.1)¹⁴, we found that core genes tend to have significantly greater expression in megakaryocytes than in the other blood cells ($P = 7.5 \times 10^{-5}$, Supplementary Fig. 5a). This observation is compatible with the notion that ultimate steps in blood cell lineage specification are accompanied, or driven, by the emergence of increasing numbers of lineage-specific transcripts. To explore this assumption, we used genome-wide expression arrays to determine changes in global transcript levels during in vitro differentiation of umbilical-cord blood-derived haematopoietic stem cells to precursors of

blood cells. We considered five different time points and two cell types, erythroblasts (the precursors of red blood cells) and megakaryocytes. Notwithstanding high levels of correlation of gene expression between erythroblasts and megakaryocytes¹⁴, core gene transcripts showed a significant increase over time in megakaryocytes ($P = 1.5 \times 10^{-6}$) but not in erythroblasts ($P = 0.77$, Fig. 1c, d; see also Supplementary Fig. 5b). Taken together, these patterns of core gene expression are consistent with a different regulation of their transcription in megakaryocytes versus erythroblasts, and with their centrality in megakaryopoiesis and platelet formation. This hypothesis is also consistent with the observation that only 5 of the 68 sentinel SNPs exert a significant effect on erythrocyte parameters (HBS1L-MYB, RCL1, SH2B3, TRIM58 and TMCC2, Supplementary Table 8).

Gene silencing in model organisms

To assess whether core genes are indeed implicated in haematopoiesis, we interrogated the function of 15 genes using gene silencing in *D. rerio* and *D. melanogaster*, and supported empirical data with published evidence on knockout models in M. musculus (Table 2 and Supplementary Table 4). In D. rerio, we applied morpholino constructs to silence the expression of six genes (Fig. 2 and Supplementary Fig. 6) selected to have >50% homology with the human counterpart and no previous evidence of involvement in haematopoiesis. Silencing of four genes in *D. rerio (arhgef3, ak3, rnf145, jmjd1c)* resulted in the ablation of both primitive erythropoiesis and thrombocyte formation. Silencing of tpma, the orthologue of TPM1 that is transcribed in megakaryocytes but not in other blood cells, abolished the formation of thrombocytes but not of erythrocytes. Silencing of ehd3 did not yield a haematopoietic phenotype. We also screened *D. melanogaster* RNA interference (RNAi) knockdown lines for quantitative alterations in the two most prevalent classes of blood elements: plasmatocytes and crystal cells. The repertoire of blood cells in D. melanogaster, consisting of about 95% plasmatocytes and 5% crystal cells, is less varied than in vertebrates. Transcription factors and signalling pathways regulating haematopoiesis have, however, been conserved throughout evolution¹⁵, making the RNAi knockdown studies a relevant first step towards a better understanding of the putative role of these GWAS genes in haematopoiesis. Four core-gene *D. melanogaster* lines (shibire (DNM), ush (ZFPM2), rpn9 (PSMD13), Brf (BRF1)), as well as five others (sun (ATP5E), CG3704 (XAB1), Su(var)205 (CBX5), dve (SATB1) and RpL6 (RPL6)), displayed highly reproducible differences in the numbers of these two cell types (Table 2 and Supplementary Table 4). Despite widespread differences between mammalian and insect haematopoietic lineages¹⁶, our findings from D. melanogaster provide new and supporting examples of functional conservation in the control of blood cell formation in invertebrates and vertebrates¹⁷⁻¹⁹.

New gene and functional discoveries

The data from studies in *D. rerio* by us and in *M. musculus* by others (see Supplementary Table 4) provided proof-of-concept evidence that our prioritization strategy is appropriate for selecting novel genes controlling thrombopoiesis and megakaryopoiesis, respectively. More detailed insights and additional implicated genes will be revealed through the systematic silencing of all genes in the associated regions. For instance, RNAi knockdown of dve in D. melanogaster reduces plasmatocyte numbers and increases the number of crystal cells, thus providing supporting evidence that its non-core genehuman homologue SATB1 should be prioritized in functional studies. However, the results of the knockdown study in D. rerio do not clarify at which hierarchical positions in thrombopoiesis and erythropoiesis the genes exert their effect, requiring further assessment in conditional knockout models in M. musculus with lineage-specific regulation of gene transcription. Nevertheless, our results have already allowed novel insights into the genetic control of these processes. Signalling cascades initiated by thrombopoietin (THPO) and its receptor

cMPL via the JAK2/STAT3/5A signalling pathway are key regulatory steps initiating changes in gene expression responsible for driving forward megakaryocyte differentiation²⁰. Our study highlights several additional signalling proteins implicating potentially important novel regulatory routes. For instance, two genes encoding guanine nucleotide exchange factors (DOCK8 and ARHGEF3) were identified. Mendelian mutations of the former are causative of the hyper-IgE syndrome, but its effect on platelets had not yet been identified. The silencing of the latter gene in *D. rerio* resulted in a profound haematopoietic phenotype characterized by a complete ablation of both primitive erythropoiesis and thrombocyte formation, demonstrating its novel regulatory role in myeloid differentiation. In a parallel and in-depth study we demonstrated its novel role in the regulation of iron uptake and erythroid cell maturation²¹. A second class of genes also known to critically control early and late events of megakaryopoiesis are transcription factors. For instance, MYB silencing by microRNA 150 determines the definitive commitment of the megakaryocyte–erythroblast precursor to the megakaryocytic lineage¹⁵. A further 10 core genes identified in this study are implicated in the regulation of transcription. Among these, we have demonstrated here that silencing of $rnf145$ and $jmjdlc$ in *D. rerio* severely affects both lineages.

In conclusion, this highly powered study describes a catalogue of known and novel genes associated with key haematopoietic processes in humans, providing an additional example of GWAS leading to biological discoveries. We further showed that for a large proportion of these known and new genes, functional support is achieved from model organisms and by overlap with genes implicated in inherited Mendelian disorders and in human cancers because of acquired mutations. In-depth functional studies and comparative analyses will be necessary to characterize the precise mechanisms by which these new genes and variants affect haematopoiesis, megakaryopoiesis and platelet formation. Furthermore, we provide extensive new resources, most notably a freely accessible knowledge base embedded in the novel protein-protein interaction network, with information about the identified platelet genes being implicated in Mendelian disorders and results from gene-silencing studies in model organisms. We anticipate that these resources will help to advance megakaryopoiesis research, to address key questions in blood stem-cell biology and to propose new targets for the treatment of haematological disorders. Finally, MPV has been associated with the risk of myocardial infarction^{22,23}. The contribution of the new loci to the aetiology of acute myocardial infarction events will require assessment in a prospective setting.

METHODS SUMMARY

A summary of the methods can be found in Supplementary Information and includes detailed information on: study populations; blood biochemistry measurements; genotyping methods and quality control filters; genome-wide association and meta-analysis methods; gene prioritization strategies for functional assessment and network construction; proteinprotein interaction network; in vitro differentiation of blood cells; experimental data sets and analytical methods for gene expression analysis; zebrafish morpholino knockdown generation; assessment of other model organism resources.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Protein-protein interaction network and gene transcription patterns

a, Negative correlation between PLT and MPV in a UK sample. The gender-adjusted correlation coefficient r and trend line are shown. **b**, Protein–protein interaction network of platelet loci. For the nodes, genes are represented by round symbols, where node colour reflects gene transcript level in megakaryocytes on a continuous scale from low (dark green) to high (white). Grey-coloured round symbols identify first-order interactors identified in Reactome and IntAct. Core genes not connected to the main network are omitted. The 34 core genes are identified by a blue perimeter. Yellow perimeters identify five additional genes (VWF, PTPN11, PIK3CG, NFE2 and MYB) with known roles in haemostasis and megakaryopoiesis and mapping to within the association signals at distances greater than 10 kb from the sentinel SNPs. These genes, which do not conform to the rule for inclusion into the core gene list, are not considered in further analyses presented in Fig. 2c, d and Supplementary Fig. 5 and are shown here for illustration purposes only. Network edges were obtained from the Reactome (blue) and IntAct-like (red) databases and through manual literature curation (black). The network including the 34 core genes alone contains 633 nodes and 827 edges; after inclusion of the 5 additional genes, the network (shown here) includes 785 nodes and 1,085 edges. The full network, containing gene expression levels and other annotation features, is available in Cytoscape²⁵ format for download (Supplementary Data 1). **c**, **d**, Time course experiments of gene expression in megakaryocytes and erythroblasts. Expression of core genes in $log₂$ transformed signal intensities ($log₂$ SI) during differentiation of the haematopoietic stem cells into megakaryocytes (**c**) or erythroblasts (**d**), segregated by their trends of statistically significant increasing (red), decreasing (blue) or unchanged (grey) gene expression. The corresponding gene list for the three classes is given in Supplementary Data 1.

Figure 2. Functional assessment of novel loci in *D. rerio*

Gene-specific morpholinos were injected into wild-type and $Tg(cd41:EGFP)$ embryos at the one cell stage (Supplementary Fig. 6) to assess alterations in erythropoiesis and thrombopoiesis. a, Control D. rerio embryo at 72 h post fertilization (h.p.f.); the boxed region corresponds to images in the middle panels of **b**–**h**. **b**–**h**, Left: o-dianisidine staining was used to assess the number of mature erythrocytes at 48 h.p.f.: *ehd3* (**h**) morpholinoinjected embryos showed normal haemoglobin staining, whereas embryos injected with $ak3$ (**c**), rnf145 (**d**), arhgef3 (**e**) or jmjd1c (**g**) morpholinos showed a decrease in the number of haemoglobin-positive cells compared to control embryos (**b**). Embryos injected with tpma morpholinos (**f**) showed normal numbers of erythrocytes but unusual accumulation dorsally in the blood vessels (compatible with cardiomyopathy). Middle: haematopoietic stem-cell and thrombocyte development was assessed using the transgenic $Tg(cdd1:EGFP)$ line at 72 h.p.f. Embryos injected with the $ehd3$ (h) morpholino had a normal number of $GFP¹$ cells in the caudal haematopoietic tissue and circulation, when compared to control embryos (**b**). However, GFP¹ cells were absent in ak3 (**c**), rnf145 (**d**), arhgef3 (**e**), tpma (**f**) and jmjd1c (**g**) morpholino-injected embryos. Right: One-cell-stage embryos were injected with the standard control morpholino (**b**) or gene-specific morpholino (**c**–**h**) and monitored during development. No gross lethality or developmental abnormalities were observed at 72 h.p.f. in gene-specific morpholino-injected embryos (**c**–**h**) compared with the control (**b**). **a** and middle and right panels of **b**–**h**, lateral view, anterior left; left panels of **b**–**e, g, h**, ventral view, anterior up; left panel of **f**, dorsal view, anterior up. The genes appear to be nonspecifically expressed during embryogenesis as shown by patterns deposited in the ZFIN resource ([http://zfin.org\)](http://zfin.org).

Summary of loci associated with platelet count and mean platelet volume in Europeans Summary of loci associated with platelet count and mean platelet volume in Europeans

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Results are provided for the 68 loci and 84 sentinel SNPs reaching genome-wide significant ($P 5 \times 10^{-8}$) association with PLT or MPV. Results for stages 1 and 2 of the analysis in Europeans are provided P 5 × 10^{−8}) association with PLT or MPV. Results for stages 1 and 2 of the analysis in Europeans are provided Results are provided for the 68 loci and 84 sentinel SNPs reaching genome-wide significant (in Supplementary Table 2. MPV, mean platelet volume; PLT, platelet count. in Supplementary Table 2. MPV, mean platelet volume; PLT, platelet count.

Alleles are indexed to the forward strand of NCBI build 36. Alleles are indexed to the forward strand of NCBI build 36.

† Effect sizes in ln(fl) for MPV and 10 9 l^{−1} for PLT.

 t^* All P values are based on the inverse-variance weighted meta-analysis model (fixed effects). P values are based on the inverse-variance weighted meta-analysis model (fixed effects).

[§] TRIM58 identifies the only secondary signal identified in this study, derived from a genome-wide secondary signal discovery effort carried out by conditioning the discovery GWAS on all SNPs reaching significance in th TRIM58 identifies the only secondary signal identified in this study, derived from a genome-wide secondary signal discovery effort carried out by conditioning the discovery GWAS on all SNPs reaching P values reported are obtained in the secondary analysis. The corresponding values in the stage 1 analysis are effect (s.e.)=2.721 (0.542) and $P=4.06 \times 10^{-7}$. Further details of this analysis are given in the Supplementary Information. significance in the stage 1 meta-analysis. The effects (s.e.) and

P=4.06 × 10⁻⁷. Further details of this analysis are given in the Supplementary Information.

 $^{\prime\prime}$ THPO narrowly misses the level required for nominal significance (P<5 × 10⁻⁸) in Europeans, but shows genome-wide significance in Japanese. THPO narrowly misses the level required for nominal significance (P<5 × 10−8) in Europeans, but shows genome-wide significance in Japanese.

Rep. indicates replication of European stage 1+2 results in non-Europeans (Supplementary Table 3): yes, if association P value is at least in one non-European population <0.0007 (to account for multiple P value is at least in one non-European population <0.0007 (to account for multiple ¶ Rep. indicates replication of European stage 1+2 results in non-Europeans (Supplementary Table 3): yes, if association testing of 68 loci). testing of 68 loci).

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Relevant references are indicated. Relevant references are indicated.

Information is given only for genes with a haematopoietic phenotype. A more extensive annotation of genes within associated intervals is presented in Supplementary Table 4. Information on variants associated with gene expression is presented in Supplementary Table 6. No evidence for a haematopoietic effect was associated with the following core genes: rs3811444 (PLT) (TRIM58 tripartite motif-containing 58 (0)); rs4305276 (MPV) (ANKMY1 ankyrin repeat and MYND domain containing 1 (0)); rs3792366 (PLT) (PDIA5 protein disulphide isomerase family A, member 5 (0)); rs10512627 (MPV) (KALRN kalirin, RhoGEF kinase (0)); rs11734132 (MPV) (KIAA0232 (5,628)); rs441460 (PLT) (LRRC16A leucinerich-repeat containing 16A (0)); rs13300663 (PLT) (RCLI RNA terminal phosphate cyclase-like 1 (0)); rs3731211 (PLT) (CDKN2A cyclindependent kinase inhibitor 2A (0)); rs2950390, rs941207 (MPV, PLT) (PTGES3 prostaglandin E synthase 3 (cytosolic) (1,871); BAZ2A bromodomain adjacent to zinc finger domain, 2A (0)); rs7961894 (MPV, PLT) (WDR66WD repeat domain 66 (0)); rs8022206 (PLT) (RAD51L1 RAD51-like 1 (S. cerevisiae) (0)); rs8006385 (PLT) (ITPKI inositol-tetrakisphosphate 1-kinase (0)); rs2297067, rs944002 (PLT, MPV) (C14orf73 exocyst complex component 3-like 4 (0)); rs8076739, rs559972 (MPV, PLT) (TAOKI TAO kinase 1 (3,357, 0)); rs1697127 (MPV) (AP2B1 adaptor-related protein complex 2, beta 1 subunit (0)); rs11082304 (PLT) (CABLES1 Cdk5 and Abl enzyme substrate 1 (0)); rs8109288 (MPV, PLT) (TPM4 tropomyosin 4 (0)), rs17356664 (PLT) (EXOC3L2 exocyst complex component 3-like 2 (3,301)); rs2015599 (MPV) (FAR2 fatty acyl CoA reductase 2 (0)); rs397969 (PLT) (AKAP10 A kinase (PRKA) anchor protein 10 (4,506)); rs11789898 (PLT) (BRD3 bromodomain containing 3 (0)).

* Core genes are defined as either containing a sentinel SNP or as mapping at less than 10 kb from an intergenic SNP. Distance from nearest gene is calculated as the absolute distance between SNP and transcription start site of the gene or 3′ end of last exon.

 ϕ^{\dagger} Phenotypes are defined from exhaustive search of the OMIM (Online Mendelian Inheritance in Man) database, published *in vitro* studies for humans and knockout and knockdown experiments for model organisms for both core and non-core genes. ¹² values are calculated from the HapMap phase 2 CEU panel. Drosophila indicates Drosophila melanogaster.