Identification and characterisation of SMIM1 variants determining the Vel blood group

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Identification and characterisation of *SMIM1* variants determining the Vel blood group

Mikael Kronborg Christophersen

DOCTORAL DISSERTATION
by due permission of the Faculty of Medicine, Lund University, Sweden.
To be defended at 13:00 on Thursday 30th March 2017
at Biomedicinsk Centrum, Segerfalksalen

*Faculty opponent*
Associate Professor Morten Hanefeld Dziegiel
Identification and characterisation of SMIM1 variants determining the Vel blood group

Abstract
The Vel blood group antigen is present on red blood cells from all humans except rare Vel-negative individuals, who can form antibodies to Vel in response to transfusion or pregnancy. It was first described in 1952 as a high incidence antigen, while the molecular background was recently discovered to be a 17-bp deletion in Small Integral Membrane Protein 1, that causes a frame-shift mutation and abolishes SMIM1 expression, thus creating a Vel-negative phenotype.

This thesis contains one of the three original discovery studies reporting the Vel antigen-defining SMIM1-deletion. We analysed SNP microarray data from Vel-positive and -negative individuals and identified SMIM1 as a potential gene. We then found the 17-bp deletion to only be present in Vel-negative individuals and sought to finally prove SMIM1 as the genetic background for the Vel antigen by examining: 1) SMIM1 mRNA sequence and expression levels, 2) presence of SMIM1 and the Vel antigen in red blood cell membranes from Vel-positive and -negative individuals and 3) anti-Vel antibody reactivity in erythroleukaemia cells expressing wild type and mutant SMIM1 protein. Our discovery allowed Vel to be officially recognised by the International Society of Blood Transfusion as blood group system 034.

The SMIM1 deletion is the major determinant for Vel expression, yet even Vel-positive individuals (i.e. people carrying wild type SMIM1) show substantial variation in reactivity with anti-Vel antibody, creating a risk for Vel blood typing errors and transfusion reactions. We suspected the cause to be sequence variants in SMIM1 and found rs143702418 (insertion, C>CGCA) and rs1175550 (A>G) to independently influence expression of SMIM1, potentially mediated by the erythroid transcription factor TAL1 that binds preferentially to the high-expressing rs1175550G allele.

Lastly, we examined historical Vel-negative samples and sought to retroactively reconcile historic Vel designations to our current knowledge of the SMIM1 deletion variant. As such, we found the old Vel;−1,−2 designation, attributed to individuals phenotyped to have weak to no Vel antigen expression, to correspond to homozygosity for the SMIM1 deletion, while persons designated Vel;1,−2, low Vel antigen expression, matched with being heterozygous carrier of the deletion.

The Vel antigen was one of the few remaining clinically significant antigens with unknown genetic background, that causes severe haemolytic transfusion reactions. This thesis assisted in characterising the molecular background of the Vel antigen, which paved the way for molecular Vel typing in the clinic, as well as the prospect of manufacturing a synthetic monoclonal anti-Vel antibody to be used in diagnostics, blood group typing and research.

Key words Blood groups, Vel blood group system, genetic variation, transcriptional regulation

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Signature 

Date February 22nd 2017
Identification and characterisation of $SMIM1$ variants determining the Vel blood group

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Faculty of Medicine
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Division of Hematology and Transfusion Medicine

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Yet an experiment, were you to try it,
could free you from your cavil,
and the source of your arts' course
springs from experiment.

-Dante Alighieri: The Divine Comedy
Preface

The following PhD thesis details the results obtained through four years of research at the Division of Haematology and Transfusion Medicine, Lund University. It consists of three studies focusing on the Vel blood group system and one studying the disease multiple myeloma. Three studies have been published in *Nature Genetics*, *Scientific Reports* and *Blood* and one is an almost submission-ready manuscript.

The Vel blood group antigen is present on red blood cells from all humans except rare Vel-negative individuals who can form antibodies to Vel in response to transfusion or pregnancy. It was first described in 1952 as a high incidence antigen, while the molecular background was recently discovered to be a 17-bp deletion in Small Integral Membrane Protein 1, that causes a frame-shift mutation and abolishes *SMIM1* expression, thus creating a Vel-negative phenotype.

This thesis contains one of the three original discovery studies reporting the Vel antigen-defining *SMIM1*-deletion. We analysed SNP microarray data from Vel-positive and -negative individuals and identified *SMIM1* as a potential gene. We then found the 17-bp deletion to only be present in Vel-negative individuals and sought to finally prove *SMIM1* as the genetic background for the Vel antigen by examining: 1) *SMIM1* mRNA sequence and expression levels, 2) presence of SMIM1 and the Vel antigen in red blood cell membranes from Vel-positive and -negative individuals and 3) anti-Vel antibody reactivity in red blood cells from individuals homozygous or heterozygous for the 17-bp deletion and in erythroleukaemia cells expressing wild type and mutant SMIM1 protein. Our discovery allowed Vel to be officially recognised by the International Society of Blood Transfusion as blood group system 034.

The *SMIM1* deletion is the major determinant for Vel expression, yet even Vel-positive individuals show substantial variation in reactivity with anti-Vel antibody, creating a risk for Vel blood typing errors and transfusion reactions. We suspected the cause to be sequence variants in *SMIM1* and found rs143702418 (insertion, C>CAGA) and rs1175550 (A>G) to independently influence expression of *SMIM1*, potentially mediated by the erythroid transcription factor TAL1 that binds preferentially to the high-expressing rs1175550G allele.
Lastly, we examined historical Vel-negative samples and sought to retroactively reconcile historic Vel designations to our current knowledge of the SMIM1 deletion variant. As such, we found the old Vel;−1,−2 designation, attributed to individuals phenotyped to have weak to no Vel antigen expression, to correspond to homozygosity for the SMIM1 deletion, while persons designated Vel;1,−2, i.e. low Vel antigen expression, matched with being heterozygous carrier of the deletion.

This thesis assisted in characterising the molecular background of the Vel antigen, which paved the way for molecular Vel typing in the clinic as well as the prospect of manufacturing a synthetic monoclonal anti-Vel antibody to be used in diagnostics, blood group typing and research.

The final Study IV details the discovery of robust biomarkers for improved FACS sorting of multiple myeloma plasma cells in a bone marrow sample, a project I was involved in during the first year of my PhD.

The thesis is divided into four main sections: An introduction outlining haematopoiesis, erythroid transcription factors, blood group systems and genetic variation; the specific aims and a brief summary of the obtained results, and a discussion and conclusion section that puts the studies into a broader scientific context. The last section is the four articles included at the end of the thesis.
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Studies included in this thesis

I. **Homozygosity for a null allele of SMIM1 defines the Vel-negative blood group phenotype**
   * denotes shared authorship.

II. **SMIM1 variants rs1175550 and rs143702418 independently modulate Vel blood group antigen expression**
   Mikael Kronborg Christophersen, Magnus Jöud, Ram Ajore, Sunitha Vege, Klara W. Ljungdahl, Connie M. Westhoff, Martin L. Olsson, Jill R. Storry*, Björn Nilsson*.
   * denotes shared authorship.
   Scientific Reports, 2017. 7: 40451.

III. **Serologic And Molecular Studies Of The Vel– Phenotype In A Multi-ethnic Population**
    Manuscript in preparation

IV. **Robust isolation of malignant plasma cells in multiple myeloma**
    * denotes shared authorship.
## Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
</tr>
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<tbody>
<tr>
<td>ACKR1</td>
<td>Atypical chemokine receptor 1</td>
</tr>
<tr>
<td>BFU-E</td>
<td>burst forming unit-erythroid</td>
</tr>
<tr>
<td>C/EBPα</td>
<td>CCAAT-enhancer binding protein α</td>
</tr>
<tr>
<td>CFU-E</td>
<td>colony forming unit-erythroid</td>
</tr>
<tr>
<td>CLP</td>
<td>common lymphoid progenitor</td>
</tr>
<tr>
<td>CMP</td>
<td>common myeloid progenitor</td>
</tr>
<tr>
<td>CSF</td>
<td>colony stimulating factor</td>
</tr>
<tr>
<td>eTF</td>
<td>erythroid transcription factor</td>
</tr>
<tr>
<td>Epo</td>
<td>Erythropoietin</td>
</tr>
<tr>
<td>EpoR</td>
<td>Epo receptor</td>
</tr>
<tr>
<td>eQTL</td>
<td>expression quantitative trait loci</td>
</tr>
<tr>
<td>FACS</td>
<td>fluorescence-activated cell sorting</td>
</tr>
<tr>
<td>FOG-1</td>
<td>Friend of GATA-1</td>
</tr>
<tr>
<td>Gfi-1[B]</td>
<td>Growth factor independent 1[B]</td>
</tr>
<tr>
<td>GMP</td>
<td>granulocyte/macrophage progenitor</td>
</tr>
<tr>
<td>GWAS</td>
<td>genome-wide association study</td>
</tr>
<tr>
<td>GPA/B</td>
<td>glycophorin A/B</td>
</tr>
<tr>
<td>GYP A/B</td>
<td>glycophorin A/B genes</td>
</tr>
<tr>
<td>HBA</td>
<td>α-globin gene</td>
</tr>
<tr>
<td>HBB</td>
<td>β-globin gene</td>
</tr>
<tr>
<td>HBG1-2</td>
<td>γ-globin genes 1 and 2</td>
</tr>
<tr>
<td>HDAC</td>
<td>histone deacetylase</td>
</tr>
<tr>
<td>HSC</td>
<td>haematopoietic stem cell</td>
</tr>
<tr>
<td>HTR</td>
<td>haemolytic transfusion reaction</td>
</tr>
<tr>
<td>IL-3</td>
<td>interleukin-3</td>
</tr>
<tr>
<td>indel</td>
<td>insertion/deletion</td>
</tr>
<tr>
<td>Acronym</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>ISBT</td>
<td>International Society of Blood Transfusion</td>
</tr>
<tr>
<td>KLF1</td>
<td>Krüppel-like factor 1</td>
</tr>
<tr>
<td>LD</td>
<td>linkage disequilibrium</td>
</tr>
<tr>
<td>LDB1</td>
<td>LIM domain binding 1</td>
</tr>
<tr>
<td>LMO2</td>
<td>LIM-only Domain 2</td>
</tr>
<tr>
<td>LSD1</td>
<td>Lysine-specific histone demethylase 1A</td>
</tr>
<tr>
<td>MCHC</td>
<td>mean corpuscular haemoglobin content</td>
</tr>
<tr>
<td>MeCP1</td>
<td>Methyl-CpG-binding protein-1</td>
</tr>
<tr>
<td>MEP</td>
<td>megakaryocyte-erythocyte progenitor</td>
</tr>
<tr>
<td>MM</td>
<td>multiple myeloma</td>
</tr>
<tr>
<td>MPP</td>
<td>multipotent progenitor</td>
</tr>
<tr>
<td>NuRD</td>
<td>Nucleosome Remodelling Deacetylase</td>
</tr>
<tr>
<td>P-TEFb</td>
<td>Positive transcription elongation factor b</td>
</tr>
<tr>
<td>Pax5</td>
<td>Paired box 5</td>
</tr>
<tr>
<td>Pentameric</td>
<td>Pentameric eTF complex of GATA-1:TAL1:E2A:LMO2:LDB1</td>
</tr>
<tr>
<td>5'/3' RACE</td>
<td>(5' or 3') Rapid amplification of cDNA ends</td>
</tr>
<tr>
<td>RBC</td>
<td>Red blood cell</td>
</tr>
<tr>
<td>RUNX1</td>
<td>Runx-related transcription factor 1</td>
</tr>
<tr>
<td>SCF</td>
<td>stem cell factor</td>
</tr>
<tr>
<td>SMIM1</td>
<td>small integral membrane protein 1</td>
</tr>
<tr>
<td>SNP</td>
<td>Single nucleotide polymorphism</td>
</tr>
<tr>
<td>SLC4A1</td>
<td>Solute carrier family 4 member 1</td>
</tr>
<tr>
<td>Sp1</td>
<td>Specificity protein 1</td>
</tr>
<tr>
<td>STAT5</td>
<td>signal transducer and activator of transcription 5</td>
</tr>
<tr>
<td>TAL1</td>
<td>T-cell acute lymphocytic leukemia protein 1</td>
</tr>
<tr>
<td>TFRC</td>
<td>transferrin receptor</td>
</tr>
<tr>
<td>TMP</td>
<td>transmembrane protein</td>
</tr>
<tr>
<td>ZBTB7A</td>
<td>Zinc Finger And BTB Domain Containing 7A</td>
</tr>
</tbody>
</table>
Introduction

Blood is the life-giving fluid that flows through us all. Coursing around inside an intricate network of blood vessels, it supplies our body with vital nutrients and oxygen. It is often the first line of detection and defence against foreign objects and microorganisms, able to act swiftly and efficiently clear invaders before they make it past the circulatory system.

The formation of blood is a delicate balance of self-renewal and lineage choices. From the early haematopoietic stem cells to the strikingly different end products, from the enucleated doughnut-shaped erythrocytes carrying vital oxygen over the giant megakaryocytes shedding pieces of itself as part of its normal function, to the granulocytes with nuclei of creative shapes and forms, the haematopoietic system is a truly remarkable tissue.

But blood can also, inadvertently, become dangerous. With the delicate regulatory system comes an inherent risk of errors that can prove detrimental, causing loss of control and involuntary abandonment of normal function.

And in case the errors prove non-fatal, they will still make the blood unique. A seemingly harmless mutation may change the blood cells just enough to set them apart from the rest of the population. Meaning that if ever the need for administration of foreign blood arises, if it is not carefully selected, the first line of defence could become the enemy as, in its eagerness to defend, it will end up destroying not only the intruders, but also itself in the process.

Blood is life-giving. It is complex and it is crucial. And it can become treacherous and fragile. It makes us unique and sets us apart. As such, extensive knowledge of blood, the factors that govern its creation and differentiation, and the systems that control donor-recipient compatibility, is an imperative necessity.
Haematopoiesis

The haematopoietic system is a paradigmatic, stem cell-maintained organ with enormous cell turnover. Up to a staggering one trillion blood cells are produced every day\(^1\). The haematopoietic system has been extensively studied, documenting the process of going from haematopoietic stem cells (HSCs) through several checkpoints to a mature, differentiated blood cell.

![Diagram of Haematopoiesis]

**Figure 1 Haematopoiesis.** An overview of haematopoiesis with the progenitor and precursor cells and the factors that guide them as described in the main text. For simplicity, not all cell intermediates nor factors are included. Factors that drive lineage differentiation are in blue font, those that inhibit are in red. Abbreviations: HSC = haematopoietic stem cell; MPP = multipotent progenitor; CMP = common myeloid progenitor; GMP = granulocyte-macrophage progenitor; MEP = megakaryocyte-erythrocyte progenitor; CLP = common lymphoid progenitor; NK = natural killer; SL = small lymphocyte; T = T cell; B = B cell. Factor names are described in the main text.
Definitive haematopoiesis in the bone marrow

Every blood cell is derived from haematopoietic stem cells (HSCs) residing in the bone marrow. An intricate system of regulators, enhancers and suppressors guide the differentiating HSCs through the many cell fate decisions, checkpoints and branch points in the haematopoietic lineage tree in order to differentiate into a mature blood cell. The main endpoint cells in the blood are the platelet-producing megakaryocytes, the oxygen-carrying erythrocytes, the microorganism-fighting granulocytes (neutrophilic, basophilic and eosinophilic) and the immune system’s macrophages, T and B cells and plasma cells derived from the latter (Figure 1).

HSCs reside in the bone marrow, where they proliferate and are capable of self-renewal, maintained by T-cell acute lymphoblastic leukaemia protein (TAL1, also known as Stem cell leukaemia factor (SCL)), Runt-related transcription factor 1 (RUNX1) and stem cell factor (SCF) and others. From here a HSC differentiates into a multipotent progenitor (MPP) cell and starts down the haematopoietic lineage tree. The first major branch point is the lymphoid or myeloid lineage, with the former giving rise to the lymphocytes of the immune system and the latter the megakaryocytes and thrombocytes, granulocytes, erythrocytes and monocytes and macrophages. These cells are called common lymphoid/myeloid progenitors, respectively (CLP/CMP). The Ikaros transcription factor drives differentiation into the lymphoid lineage, and the lineage soon branches in two, one giving rise to natural killer cells and the other, the small lymphocyte, to the progenitors for the B and T cells. The T-cell lineage is promoted by Notch signalling, assisted by GATA-3 and finished T-cells are the active part of the adaptive immune system. Expression of transcription factor Paired box 5 (Pax5) is required for B cell formation and B cells and the plasma cells derived from them produce the antibodies that form the helper and memory part of the adaptive immune system.

Differentiation towards the CMP and the myeloid lineage is driven by PU.1 and CCAAT-enhancer binding protein α (C/EBPα) and has more branches than the lymphoid, leading to a more diverse set of cells. For simplicity, the lineage is perhaps best explained by describing the examples of tight competition going on in progenitor cells as two or more factors carry out dual roles of promoting their own lineage while antagonising another (Figure 1): GATA-1 drives CMPs to the megakaryocyte/erythroid progenitor (MEP) lineage, while PU.1 pushes cells toward the granulocyte/macrophage progenitor (GMP) lineage. Mirroring this, a study by Mancini et al. found Friend of GATA-1 (FOG-1) to drive CMPs towards the megakaryocyte/erythroid lineage while blocking further myeloid differentiation by antagonising C/EBPα. This study also implicated GATA-1 in the next step, driving MEPS toward the erythroid lineage, together with Krüppel-
like factor 1 (KLF1) which in turn also antagonises Fli-1’s push for megakaryocytic development\textsuperscript{17}. And, finally, PU.1 and Growth factor independent 1 (Gfi-1) antagonise each other while promoting the GMPs towards monocyte/macrophage or granulocyte differentiation, respectively\textsuperscript{18}.

Most of the haematopoietic differentiation is completely reversible (the enucleated mature red blood cells (RBCs) are an obvious exemption). Lineage commitment is strong and robust, but at the same time fragile due to the relatively few factors and the lack of redundancy in the haematopoietic system. Therefore the complete loss of a single factor can be catastrophic, and many of the haematopoietic malignancies (blood cancers like leukaemia and myeloma) arise from harmful genetic mutations disrupting a single or a few crucial factors’ normal function\textsuperscript{19-21}. But this dependency can also be exploited as a potential therapeutic strategy by reprogramming one terminally differentiated blood cell into another using only a few essential factors. This is possible due to a process called lineage priming; while cells differentiate into one lineage, they never lose the differentiation scheme of other lineages, it is only temporarily repressed\textsuperscript{22}. As such, researchers have been able to reprogram terminally differentiated blood cells into cells of another lineage by reactivating the differentiation scheme of that cell\textsuperscript{23,24}. And, taking it a step further, researchers have also successfully reprogrammed neural stem cells and differentiated fibroblasts into haematopoietic lineages\textsuperscript{25-27}.

**Plasma cell differentiation and Multiple Myeloma**

Plasma cells develop from mature, activated B cells and their main function is to produce large numbers of immunoglobulins (antibodies). Like all other aspects of haematopoiesis, plasma cell formation is a tightly regulated process of differentiation and lineage specification, characterised by an antagonistic relationship with the B cell lineage, not unlike those found in the myeloid lineage\textsuperscript{28,29}. Expression of Pax5 and a number of other factors and cytokines drive B cell lineage specification from the small lymphocyte to the mature, naive follicular B lymphocytes that populate the spleen and lymph nodes\textsuperscript{30,31}. Here they monitor the lymphatic system and when a B cell encounters a foreign antigen it is quickly activated and undergo proliferation, somatic hypermutation and immunoglobulin class switching. From here it differentiate into two populations; B memory cells capable of eliciting a quick antibody response, and long-lasting plasma cells that migrate back to the bone marrow and continually produce antibodies against the antigen that activated the B cell\textsuperscript{32}. Reactivation of Pax5-repressed genes is an early first step of plasma cell differentiation\textsuperscript{33}, followed by the upregulation of transcription factors X-box-binding protein 1 (XBP1), interferon-regulatory factor 4 (IRF4) and B lymphocyte-induced maturation protein 1 (BLIMP1), that are the main drivers of plasma cell maturation\textsuperscript{32}. Distinct
surface proteins are expressed on the mature plasma cell, a fact that is frequently exploited by researchers to sort and enrich plasma cell populations, which normally only constitute ~1% of the bone marrow cells. As such, CD27, CD38 and CD138 (Syndecan 1, SDC1) are highly expressed in mature plasma cells, while the common B cell markers CD19, CD20 and CD45 (Protein tyrosine phosphatase, receptor type C, PTPRC) are absent. Multiple myeloma (MM) is defined as an un-inhibited, clonal growth of plasma cells in the bone marrow, producing a monoclonal immunoglobulin (M protein) that can be detected in peripheral blood. It is preceded by a condition called monoclonal gammopathy of undetermined significance (MGUS), characterised by the presence of the M-protein, but absence of any MM clinical signs or symptoms. MGUS progresses slowly (~1% of cases per year) to MM, which is characterised by presence of the M component, clonal growth of plasma cells to comprise at least 10% of the cells in the bone marrow and evidence of organ damage, such as lytic bone lesions. The incidence of MM varies globally from about 1 in 100,000 in Asia, to 5-6 in 100,000 in Sweden and among Caucasians, to about 10 in 100,000 among Africans and in African Americans. In addition to geographic origin, the incidence is influenced by age (more common above 60 years old) and gender (more common among men). Several risk loci have been identified for MM and MGUS, and several studies have reported an aggregation of cases in families. There is currently no cure for MM, but therapies to extend the survival have been developed (recently reviewed by Bianchi et al.), including melphalan-prednisone treatment, autologous HSC transplantation, immunomodulatory drugs (e.g., thalidomide and lenalidomide), protease inhibitors (e.g., bortezomib) and monoclonal antibodies targeting plasma cell markers, such as CD38 (daratumumab).

MM differs from other haematological cancers in that the malignant cells comprise a small proportion of the bone marrow cell population at diagnosis. Plasma cell surface markers are therefore a major focus of MM research, as they can be used in fluorescence-activated cell sorting (FACS) to extract plasma cells from a bone marrow sample, to be used for classification, characterisation and research. Common markers used in MM cell sorting are CD19, CD45 and CD56, as multiple myeloma plasma cells (MMPCs) are normally negative for the two former, and positive for the latter, whereas the direct opposite is the case for normal plasma cells. Other markers include CD38 and CD138, with the latter preferred due to higher specificity towards plasma cells. However, CD138 is quickly degraded ex vivo, requiring immuno-phenotyping analyses to be done shortly after sampling, which is often not possible in a clinical setting. Additionally, the short half-life of CD138 limits its usability with biobanked samples. This prompted us to search for a more suitable plasma cell marker and we found CD319 and CD269. This discovery forms the basis of Study IV.
Erythropoiesis

Erythropoiesis is the branch of haematopoiesis that produces the RBCs crucial for transporting oxygen to all tissues in the body. Oxygen is transported in haemoglobin molecules within the RBCs, taking up as much as 98% of the entire cytoplasm. An estimated 2.5 billion RBCs are produced every day, with the system capable of production bursts when needed. The lifespan of an RBC is 120 days, in which it continually travels through the bloodstream and circulation, racing through the largest aortic blood vessels and squeezing through the smallest capillaries, made possible by the flexible cytoskeleton and the absence of a nucleus. However, before all this can be achieved, an intricate regulatory process has transformed the MEP cell into a mature RBC, where perhaps the most striking feature is the extrusion of the nucleus, thus irreversibly committing the resulting reticulocyte (as the cells are called at the penultimate stage) into the mature RBCs formed as the cells enter the blood stream.

The major drivers in differentiating HSCs into the erythroid lineage were described above, but several stages still remain before the RBC is fully matured (Figure 2). Apart from transcription factors, several cytokines and colony stimulating factors (CSFs) also influence the erythroid lineage, some of which will be highlighted in the following. Differentiation is initiated by GATA-2 and KLF1 together with interleukin-3 (IL-3), SCF and granulocyte-macrophage CSF (GM-CSF) that transform the MEP cell to a burst forming unit-erythroid (BFU-E). GATA-2 blocks further erythroid differentiation and instead facilitates proliferation and a surge in immature erythroblasts. Eventually, GATA-1 displaces GATA-2 in what is known as the GATA-switch, that facilitates the differentiation to colony forming unit-erythroid (CFU-E) cells, assisted by TAL1, SCF, IL-3 and GM-CSF, among others. At this point, expression of the prime erythroid cytokine erythropoietin (Epo) and its receptor (EpoR) increases and Epo replaces SCF as primary stimulatory cytokine. Epo:EpoR interaction triggers the signal transducer and activator of transcription 5 (STAT5). This initiates a signal transduction cascade that activates transcription of Bcl-x, transferrin receptor (TFRC, or CD71) and other transcription factors. Also in the nucleus the pentameric complex consisting of GATA-1, TAL1, E2A, LMO2 and LDB1 enhances transcription of erythroid genes (more on this later) to differentiate the CFU-E into a pro-erythroblast. Subsequently, the cell goes through a number of morphological changes known as basophilic erythroblasts, polychromatophilic erythroblasts and orthochromatophilic erythroblasts where Epo dependency decrease and upregulation of TFRC facilitates increased iron uptake and haem production in the cell. Haem in turn increases expression of globin genes which facilitate formation of the vital haemoglobin molecules. Simultaneously, production of several important cell membrane and cytoskeletal
INTRODUCTION

proteins, such as ankyrin-1, Band 3, Band 4.1, Band 4.2, glycoporphin A (GPA), Duffy, RhD and RhCE, is accelerated\(^{74}\). In the final stages of erythrocyte maturation histone deacetylation and DNA methylation increase as chromatin in the nucleus condenses; the cytoskeleton is rearranged and the nucleus is extruded from the erythroblast\(^{75,76}\). At this stage the cell is called a reticulocyte and enters the bloodstream, the remaining cell organelles and nucleus debris is cleared and differentiation into a mature biconcave RBC is complete\(^{58}\).

Figure 2 Erythropoiesis. Overview of erythropoiesis with the progenitor and precursor cells and the factors that guide them as described in the main text. For simplicity, not all cell intermediates nor factors are included. Abbreviations: MEP = megakaryocyte-erythrocyte progenitor; BFU-E = burst forming unit-erythroid; CFU-E = colony forming unit-erythroid; pro-EB = pro-erythroblast; baso-EB = basophilic erythroblast; poly-EB = polychromatic erythroblast; ortho-EB = orthochromatic erythroblast; 5meric complex = pentameric erythroid transcription factor complex consisting of GATA-1, TAL1, E2A, LMO2 and LDB1. Factor names are described in the main text.
Transcriptional regulation of erythropoiesis

Erythropoiesis is a complicated process governed by a multitude of factors and regulators. Some factors have broad regulatory potential and are found in many different areas of haematopoiesis, while others are more specialised and only exert their influence in a single lineage or transition. Prime erythroid transcription factors (henceforth abbreviated eTFs for simplicity) such as GATA-1, KLF1 and TAL1 drive the haematopoietic cells towards the erythroid lineage, but also act to suppress those that guide the other lineages, such as PU.1 and Fli-1. The regulation by these crucial factors takes many forms and can act via direct DNA:protein or protein:protein interaction or attraction of additional factors, that facilitate chromatin remodelling or histone modifications. eTFs may influence transcription near their target or from many kilobases away, evident from a recent study that showed that GATA-1, TAL1 and KLF1 prefer to occupy sites in gene regions that are intronic (~47% of all sites) or distal (~26%, defined as between 50kb and 1kb from nearest gene), rather than intergenic (~18%, more than 50kb from nearest gene)\textsuperscript{77}.

Presence or absence of a given factor is not sufficient to determine its influence on transcription. Concentration is also an important determinant, as illustrated in the stoichiometric relationship between TAL1 and ETO2 described below and that erythroid maturation is impaired by lowering the concentration of GATA-1\textsuperscript{78}.

Although tightly regulated, erythropoiesis is carried out by relatively few main factors. As noted previously, researchers were recently able to reprogram human fibroblasts towards induced erythroid progenitors and established the essential transcription factors to be GATA-1, TAL1, cMyc and LMO2\textsuperscript{27}.

The following is a brief presentation of the most important regulators of erythropoiesis, followed by a description of how these come together in large regulatory complexes that are crucial for erythroid differentiation and synthesis of RBCs, summarised in Table 1.
Table 1: Transcription factors and complexes involved in erythropoiesis.

Only those factors and target genes/effects described in the text are shown, each factor may have several other binding partners and related targets. To avoid overlap, complexes containing the same factors are only depicted for one of them, in no particular order.

<table>
<thead>
<tr>
<th>Factor</th>
<th>Binding partners</th>
<th>Target genes / effects</th>
<th>DNA-binding motif$^1$</th>
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<tbody>
<tr>
<td>GATA-1</td>
<td>STAT5</td>
<td>FOG1 $^\uparrow$</td>
<td>GATA1 $^\uparrow$</td>
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<td></td>
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<td>KLF1 $^\uparrow$</td>
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<td>EPOR $^\uparrow$</td>
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<td>GATA2 $^\uparrow$</td>
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<td>CR/HM$^3$</td>
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<td></td>
<td>SWI/SNF</td>
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<td>ZBTB7A $^\uparrow$</td>
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<td></td>
<td>CBP/p300</td>
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<td>FOG-1</td>
<td>GATA-1</td>
<td>GATA1 $^\uparrow$</td>
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<td>HBA $^\uparrow$</td>
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<td>GATA-1:MeCP1:NuRD</td>
<td>GATA2 $^\uparrow$</td>
<td>CR/HM</td>
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<tr>
<td>TAL1</td>
<td>Pentameric$^2$</td>
<td>P-TEFb recruitment</td>
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<td>LSD1:CoREST:HDAC1:HDAC2</td>
<td>EPB42 $^\uparrow$</td>
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<td>GATA2 $^\uparrow$</td>
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<td>Gfi-1B:E2A:ETO2</td>
<td>HBA $^\uparrow$</td>
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<td>GATA-1</td>
<td>Bcl-$\chi$,$\uparrow$</td>
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<td>GF11B $^\uparrow$</td>
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<tr>
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<td>CoREST:LSD1:HDAC1:HDAC2</td>
<td>GF11B $^\uparrow$</td>
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<td>c-Myc $^\uparrow$</td>
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<td>MEIS1 $^\uparrow$</td>
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<td>CR/HM$^3$</td>
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<td>GATA-1</td>
<td>Fi-1 $^\uparrow$</td>
<td>γ- to β-globin switch</td>
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<td></td>
<td>Pentameric (speculative)</td>
<td>Guidance/attraction to transcription factory</td>
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<tr>
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<td>Pentameric</td>
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<td>KLF1 $^\uparrow$</td>
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<tr>
<td>ZBTB7A</td>
<td>MeCP1:NuRD</td>
<td>Bin $^\downarrow$</td>
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<td></td>
<td></td>
<td>HBG1/2 $^\uparrow$</td>
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</table>

1: JASPAR database (http://jaspar.genereg.net/)
2: The pentameric complex consists of GATA-1, TAL1, E2A, LMO2 and LDB1.
3: CR = chromatin remodelling; HM = histone modification
**GATA-1**

GATA-1 is a member of a family of zinc-finger transcription factors and is involved in many stages of erythropoiesis. One of the first erythroid factors to be discovered and arguably the most investigated, GATA-1 is implicated in several erythroid processes, such as the STAT5 signalling cascade which facilitates upregulation of several erythroid and haemoglobin genes, including \textit{EPOR, KLF1} and \textit{FOG1}\textsuperscript{81,82}. GATA-1 also exhibits features that are common for the major regulators of haematopoiesis, namely the ability to activate erythroid-specific genes and repress or disrupt genes that drives other lineage. Examples of this include the antagonistic direct interaction between GATA-1 and the myeloid lineage-specific factor PU.1\textsuperscript{15,83} or, even more strikingly, that overexpression of GATA-1 results in complete reprogramming of lymphoid and myelomonocytic cells into early erythroblasts\textsuperscript{23,24}. GATA-1 is in the centre of the intricate erythropoietic network, by being involved in most of the complexes that govern it and loss of GATA-1 not surprisingly results in anaemia, thrombocytopenia and other malignancies\textsuperscript{84,85}.

The GATA family of transcription factors was named for their preferential binding to the DNA sequence motif (A/T)GATA(A/G), however it has more recently been discovered that GATA-1 does not always occupy these sites during erythropoiesis\textsuperscript{86,87}. These genome-wide occupancy studies also found that GATA-1 mainly occupies sites within the target gene or in intergenic regions, with only 10-15% binding to the proximal promoter, which corresponds fairly well with the study by Su \textit{et al.}\textsuperscript{77} referenced above, which found nearly half of all occupancy sites in intronic regions. The core GATA DNA binding motif is crucial, perhaps with the exception of the fourth and final nucleotide, as altering the motif to GATT had little effect on murine GATA-1 protein binding efficacy\textsuperscript{88}.

**FOG-1**

The direct protein interaction between GATA-1 and FOG-1 is indispensable for erythroid differentiation\textsuperscript{89}. FOG-1 is a co-regulator in the GATA-switch where GATA-1:FOG-1 blocks GATA-2 occupancy in its own \textit{GATA2} promoter abrogating its positive autoregulation\textsuperscript{62}. The switch also involves chromatin remodelling which shuts down \textit{GATA2} expression, thus driving progenitor cells further towards erythroid differentiation\textsuperscript{63}. Although no direct DNA-binding has been demonstrated for FOG-1, it is known to associate with GATA-1 and GATA-2 and the chromatin remodelling is not exclusive for the GATA-switch, leading researchers from Stuart Orkin’s group to propose FOG-1 as a new class of molecules functioning as chromatin occupancy facilitators (COFs)\textsuperscript{62}. 
**TAL1**

TAL1 is named after its discovery as a genetic translocation (t(1:14)(p32;q11)) causing T-cell acute lymphoblastic leukaemia\(^{90,92}\). TAL1 is expressed throughout haematopoiesis and is involved in many erythroid proliferation and maturation processes. TAL1 is important in haematopoiesis and erythropoiesis with specific DNA binding not necessarily required in the early stages of haematopoiesis\(^93\) yet critically required throughout most of erythropoiesis\(^94\). DNA binding is carried out through TAL1’s basic helix-loop-helix domain with the core DNA binding motif CANNTG, also called an E-box. Composite motifs with a GATA site 9 to 12 nucleotides from the E-box (designated CANNTG\(_{9,12}\)-GATA) are often found in genes directly regulated by TAL1, and it is the preferred binding motif for the pentameric complex\(^{95-97}\). Additionally, a shortened motif for TAL1, CTG, 9 nucleotides upstream of the GATA-1 motif was recently identified\(^98\). In general, GATA-1 and TAL1 have very similar expression patterns in haematopoiesis (i.e. mainly expressed in erythroid, megakaryocytic and mast cells\(^99\)), they are both crucial for correct erythropoiesis, and they often co-localise and affect gene transcription together. A TAL1 global occupancy network analysis determined its preference to intra- and intergenic over proximal promoter regions and that about half of the genes regulated by TAL1 are upregulated, with an added interesting notion that for those genes where TAL1 was found directly occupying the target DNA site, 75% were found to be upregulated, indicating that TAL1 is mainly a transcriptional activator\(^98\).

**Gfi-1B**

Gfi-1 and its parologue Gfi-1B are zinc-finger transcription factors with an N-terminal Snail/Gfi-1 (SNAG) repressor domain. They both bind a DNA sequence motif of TAAATCAC(A/T)GCA, core motif AATC\(^{100,101}\). Expression of both proteins occurs in early haematopoiesis, but diverges as the cells differentiate, with Gfi-1 mainly being expressed in the lymphoid and some myeloid lineages, and Gfi-1B mainly found in the erythroid and megakaryocytic lineages\(^{102,103}\), where it is an indispensable factor in early erythropoiesis promoting erythroblast proliferation, but not differentiation\(^{104}\). GATA-2 is known to activate Gfi-1B and repress lymphopoiesis, whereas Gfi-1 was recently shown to inhibit GATA-2 function, presenting an interesting three-way regulatory network governing cell fate of early haematopoietic development\(^{105}\). Gfi-1B acts as a repressor of transcription by direct DNA binding to target sites and recruitment of histone modification enzymes\(^{106}\), but also by direct interaction with GATA1 to repress Bcl-xl\(^{107,108}\) and GFI1B itself\(^{109}\).
**KLF1**

KLF1 was first described in 1993, originally named erythroid Krüppel-like factor (EKLF), based on similarity to a gene found in *D. melanogaster*\(^{110}\). It is exclusively found in erythroid and mast cells and is a crucial component of the erythrocyte/megakaryocyte lineage switch\(^{17,111}\). KLF1 preferentially binds to the DNA sequence motif of CCNCNCCCN, with the core motif being CACCC\(^{110}\).

Research in KLF1-deficient mice looking at various globin genes have established, that KLF1 is likely to participate in the switch from fetal γ-globin to adult β-globin\(^{112,113}\).

Together with GATA-1, KLF1 is probably the most well-studied eTF. In two gene expression profiling studies of KLF1-null cell lines, two-thirds of the differentially expressed genes were downregulated, and most of these genes showed direct binding of KLF1, indicating that KLF1 mainly acts as a transcriptional activator\(^{114,115}\), however examples of repressor function have also been demonstrated\(^{116,117}\).

An interesting discovery has coupled KLF1 to an emerging gene transcription paradigm: That chromosomal organisation inside the nucleus is highly plastic and chromosomes intertwine and flow freely in the nucleus, and that transcription is localised to focal spots, or transcription factories where the chromosomes are transported to, rather than the transcription machinery moving around inside the nucleus\(^{118-120}\). KLF1 was found to be an important factor, at least for haemoglobin genes, as it was shown to be capable of “pulling” and guiding target genes towards the factory\(^{121}\). This model may also explain KLF1’s long-range cis and trans regulation, and indeed, KLF1-enriched binding sites were shown to primarily be found more than 10kb away from the nearest known gene\(^{115}\).

**LMO2**

The LIM-only Domain 2 gene was discovered by investigating the 11p13 T-cell translocation cluster, which is a hotspot for translocations in T-ALL\(^{122}\). It contains two LIM-only class zinc-fingers and an N-terminal domain that is capable of transcriptional transactivation, *i.e.* to interact with other factors and activate them\(^{123}\). LMO2 is a critical eTF as absence of LMO2 leads to abrogated yolk sac erythropoiesis and embryonic lethality\(^{124}\).

To date, no direct DNA-binding has been described for LMO2, rather it is crucial as a bridging factor between TAL1 and GATA-1 in the pentameric complex also containing E2A (also known as transcription factor 3 (TCF3)) and LIM domain binding 1 (LDB1)\(^{125,126}\). This complex (or at least LMO2:GATA-1:TAL1) regulates expression of the *LMO2* gene itself\(^{127}\). Not surprisingly, the expression pattern of *LMO2* follows closely that of GATA-1 and TAL1 discussed above.
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**ZBTB7A**

Zinc Finger And BTB Domain Containing 7A (ZBTB7A) is also known as Leukaemia/Lymphoma Related Factor (LRF), as well as a number of more or less colourful names (e.g. “Pokemon” and “FBI”). As the names suggest, ZBTB7A contains four DNA-binding zinc finger domains and a Broad Complex, Tramtrack, and Bric-a-brac (BTB) domain that facilitates protein:protein interaction and functions as a transcriptional repressor. Upregulation of ZBTB7A is facilitated by GATA-1 binding directly to the ZBTB7A promoter and ZBT7A is essential in erythropoiesis to antagonise the proapoptotic factor Bim in late-stage erythroblasts. ZBTB7A has also been shown to repress fetal γ-globin, and is thus an important factor in early haematopoiesis. The DNA binding motif of ZBTB7A has been ambiguously reported as G(A/G)GGG(T/C)(C/T)(T/C)(C/T)\(^{128,129}\), \(\text{NGCGACCACCNN}^{133}\), \((G/A)(C/A)GACCCCCCCCC\)^\(^{129}\) and \(\text{GCGGGGGCGGGGC}^{130}\), but with the former remarking that the consensus motif seems to be fluid and preferential ZBTB7A binding to DNA quite flexible. The latter two motifs found by Maeda and colleagues do have similarities (if we assume that one of them is in reverse complement orientation), and since only these two studies investigated ZBTB7A’s role in erythropoiesis, this may indicate an erythroid-specific DNA binding motif.

**Erythropoietic transcription factor complexes**

Regulation of erythropoiesis by the above eTFs is never a one-factor task. Albeit a limited number of factors, they interplay and associate in various highly dynamic multimeric complexes (Figure 3) and may form, reform and dissolve throughout the differentiation from HSCs to mature erythrocytes. A given factor can thus either activate or repress transcription of its target genes, depending on which cofactors it associates with and where in the maturation process the cell is.

GATA-1 plays a central role in most of these complexes, and it was traditionally believed to primarily be a transcriptional activator. However, three genome-wide ChIP-seq studies found an estimated 41-60% of the genes, that GATA-1 directly influenced expression of, to be downregulated\(^{86,87,134}\). As expected, GATA-1’s role as a transcriptional activator or repressor depends on which cofactors it associates with.

Transcriptional activation of target erythroid genes is mediated mainly through the pentameric complex. As described earlier, GATA-1 and TAL1 bind directly to DNA sequences (the E-box/GATA composite motifs), with LMO2 acting as a bridge or spacer molecule\(^{126}\) and E2A functions as an obligate heterodimer of TAL1 to facilitate DNA-binding\(^{98,135}\). LDB1 is responsible for recruiting the kinase Positive transcription elongation factor b (P-TEFb) to the complex, which
in turn phosphorylates RNA Polymerase II, a crucial requirement for activation of the transcription machinery. Another proposed function of LDB1 is to facilitate long-range interaction between two pentameric complexes through its self-association domain. This is thought to be accomplished by the DNA bending or looping to bring the two distant complexes occupying promoter/intron and distal enhancer regions together and then tethered by direct LDB1:LDB1 interaction. This may explain why the pentameric complex is often found occupying sites several kilobases from the transcription start site of their target genes. The pentameric complex has been shown to activate important erythroid genes such as glycophorin A (GYP A), the α-globin locus (HBA) and KLF1.

GATA-1 and KLF1 also form an activation complex, illustrated by Tallack et al., who compared occupancy maps of GATA-1 and KLF1 and found that in nearly half of the sites they appeared within 1kb of each other with a peak distance of only 18 bp. Interestingly, they found a CTG sequence (TAL1 binding motif) often to be present between the two sites, however while GATA-1:TAL1 and GATA-1:KLF1 complexes were abundant, no trimeric GATA-1:KLF1:TAL1 complex was found. Not many genes have so far been found as direct targets of GATA-1:KLF1 complexes, but given the proximity of the two factors, it is speculated that KLF1 facilitates transcription of pentameric-bound genes by translocating the gene region into the transcription factories described earlier.

Not surprisingly, GATA-1 and FOG-1 associate readily, but the complex represents dual roles depending on additional co-factors in the complex. GATA-1:FOG-1-mediated activation has been reported in many genes, for example α- and β-globin (HBA and HBB), Solute carrier family 4 member 1 (SLC4A1/Band 3) and GATA1 itself, but these same studies also presented numerous examples of transcriptional repression. This is facilitated by association with the multimeric MeCP1:NuRD complex, which contains multiple proteins capable of both chromatin remodelling and histone deacetylation, leading to tighter chromatin packing and thus decreased expression. This complex was shown to bind directly to the distal promoter of GATA2, providing strong evidence that it is at least partly responsible for the decreased expression of GATA2 in erythropoiesis. While GATA-1 seems to be the one that tethers the complex to the DNA of the target gene, FOG-1 is responsible for attracting and binding the MeCP1:NuRD complex. Recently it was also shown that ZBTB7A interact directly with the NuRD-complex and represses foetal γ-globin genes (HBG1 and -2) keeping their expression low in adults.

Chromatin remodelling is the proposed function of the Gfi-1B:CoREST:LSD1:HDAC1:HDAC2 repressor complex as well. Gfi-1B is thought to bind target DNA and recruit the other proteins through its SNAG
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domain. Lysine-specific histone demethylase 1A (LSD1) then demethylates lysine 4 on histone H3 (H3K4), which leads to tighter chromatin packing and limits access to DNA. This repression affects the genes GFI1B itself (establishing a negative feedback loop), c-Myb and MEIS1.

Another Gfi-1B complex with cofactors TAL1, E2A and ETO2 associates in early erythroid progenitors (but not in megakaryocytes) and represses transcription of TAL1 target genes HBA, HBB and ALAS2. This complex is absent in late erythropoiesis, signifying a role as an important erythroid differentiation switch. The ETO-mediated repression is proposed as direct ETO2:E2A interaction which interferes with its association with transcriptional activator p300/CREB, while the de-repression and switch towards erythropoietic differentiation happens through an increase in the stoichiometric ratio of TAL1 to ETO2 present in the cells.

**Figure 3 Erythroid transcription factor complexes.** Selected complexes crucial for erythropoiesis. For simplicity, only the factors described in the main text are depicted, the complexes may contain several other co-factors. See also Table 1 for individual factors and binding motifs. a) GATA-1 binds to DNA and recruits FOG-1 to activate transcription of GATA1, HBA, HBB and SLC4A1. b) The GATA-switch. GATA-1:FOG-1 complex may also instigate transcriptional repression by attracting the MeCP1:NuRD complex that facilitate chromatin remodelling and histone modification, which limit transcription machinery access to GATA2 and other genes. c) The pentameric complex (GATA-1:TAL1:E2A:LMO2:LDB1) facilitates transcriptional activation by recruiting the kinase P-TEFb that phosphorylates polymerase II, a crucial requirement for transcription initiation. d) GATA-1 and Gfi-1B interact at AATC sites to facilitate downregulation of GFI1B and Bcl-XL. The precise mechanism of downregulation is not clear, but Gfi-1B has been shown to associate with the CoREST-complex (not shown) that facilitates chromatin remodelling and histone modification.

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Additional layers of erythroid transcriptional regulation

It goes without saying that a system as complex as erythropoiesis needs thorough and robust regulation beyond the level of transcription factor complexes. Indeed, epigenetics play a major role, as already described above, but other regulatory mechanisms are also present.

Apart from the epigenetic examples already presented, GATA-1 also interacts with the SWI/SNF chromatin remodelling complex and CBP/p300 histone acetyltransferases; Grass and colleagues showed broad histone deacetylation dependent on FOG-1 and Hu and coworkers presented a repressor complex consisting of TAL1, LSD1, CoREST and histone deacetylase 1 and 2 (HDAC1/2) playing a crucial role in the onset of erythroid differentiation through repression of GATA2 and EPB42.

Gene silencing, which is mediated by microRNAs and capable of degrading or sequestering newly transcribed mRNA, is a common posttranscriptional regulation process in cells. Several thousand microRNAs have been discovered so far, and some of them have been associated with erythropoiesis. As such, miR-223 was shown to target LMO2 and downregulation of this microRNA is crucial for terminal differentiation, while miR-191 downregulates Riok3 and Mxi1 and facilitates mouse erythroblast enucleation. It is beyond the scope of this thesis to go into more detail about this intriguing layer of regulation, additional knowledge can be gained from these excellent reviews.

Finally, the long-noncoding RNAs (lncRNAs) represent yet another layer of erythroid regulation. Knowledge of these abundant RNA elements, that play potentially crucial roles in gene regulation, is rapidly expanding and a few studies on erythroid-specific interaction have emerged. The LincRNA erythroid prosurvival (LincRNA-EPS) was found to promote erythropoiesis by halting apoptosis, while another study looked at genome-wide expression of lncRNAs in erythropoiesis and found major influences on RBC maturation and chromatin structure, and showed promoter occupancy by GATA-1, KLF1 and TAL1 at genes coding for lncRNAs, indicating erythroid-specific expression.
Blood group systems

Humans have been interested in blood transfusion since the Middle Ages, experimenting with more or less creative methods, including transfusing blood from dogs into live human beings, unsurprisingly with very limited success and very high mortality rate. Modern transfusion medicine owes much to the Austrian physician and scientist Karl Landsteiner, who at the turn of the 20th century conducted research into donor-recipient compatibility. By then it was known that mixing blood from two patients would cause the blood to clump, or agglutinate, but this was thought to be because of the sicknesses the patients had. Landsteiner was the first to mix blood from healthy individuals, and he quickly realised that the agglutination also happened here, yet not every time. Landsteiner also separated the RBCs from the plasma and was able to show, among other things, that RBCs from one individual could be mixed with plasma from another, but that same individual’s RBCs would clump when mixed with the first individual’s plasma. His work resulted in the classification of the first blood group system, which he dubbed ABC (later renamed to the ABO we know today) and established the antibody-antigen relationship that forms the basis of blood group systems and transfusion medicine to this day.

Certain RBC membrane proteins and molecules exhibit minor variations which are recognised as blood group antigens. They play an important role in blood transfusions, where donor-recipient compatibility is paramount. A potentially fatal haemolytic transfusion reaction (HTR) may occur if foreign RBC antigens of the donor blood trigger the recipient’s immune system to produce alloantibodies. In the most severe cases, these antibodies may bind complement and cause intravascular haemolysis. The alloantibodies reside in the plasma, and in the ABO blood group system, a type A individual will thus have the A antigen on the RBC surface and anti-B antibodies in the blood plasma, and vice versa for a type B individual. Type O has no A/B antigens on the RBC membrane, but has both anti-A and anti-B in the plasma, while type AB has both A/B antigens on the RBCs, and none of the antibodies in the serum. Apart from the obvious incompatibility between A and B, it follows that type O RBCs can be safely transfused to all other types, but can only receive RBCs from other type O individuals, while type AB can only be administered to other type AB individuals, yet a type AB individual can receive blood from any other type (since the recipient’s plasma contains neither anti-A or anti-B). It should be noted that even naïve individuals (never having received a blood transfusion) express alloantibodies against the non-self A/B antigen(s). It is commonly believed that we are exposed to microorganisms exhibiting structures very similar to A and B antigens early in life and thus have
alloantibodies against these readily available in the blood\textsuperscript{162,163}. This is in contrast to most other blood group antigens, where alloantibodies may be produced following transfusion with mismatched donor blood (or in some cases pregnancies of blood group negative mothers giving birth to positive babies), and as such will present a concern only on subsequent transfusions/pregnancies. The ABO system was the first blood group system to be described, and the antigen-antibody relationship resembles that of most other blood groups, yet the severity of mismatched transfusion reactions varies significantly.

**Blood group terminology and nomenclature**

Based on the breakthroughs made by Landsteiner and others, transfusion medicine became increasingly successful throughout the 20\textsuperscript{th} century. Yet with increased volumes of performed transfusions, reports of HTR incidents, where the recipient reacted strongly against the donor blood, also surged. This resulted in an ever growing number of new antigens and blood groups reported by transfusion specialists from all around the world and to bring order to the different reporting and naming schemes, the International Society of Blood Transfusion (ISBT) instigated the Working Party on Red Cell Immunogenetics and Blood Group Terminology in 1980 to come up with a uniform nomenclature for all antigens and blood groups. This resulted in a naming and numbering scheme of blood group systems and the antigens defining them, with the remaining antigens not belonging to any system divided into two groups according to their population frequency, the high and low incidence antigens\textsuperscript{164}. The working party has convened biannually ever since to characterise and catalogue the currently 36 blood group systems (Table 2), six collections (antigens with insufficient information to be assigned to a particular blood group), and two series (antigens waiting for their molecular background to be determined), all in all comprising 352 antigens.

The ISBT guidelines state that in order for antigens to be considered as a blood group system, the genetic background, \textit{i.e.} the sequence variant and/or mutation creating the antigen, which raises the alloantibody in the recipient, must be known. The molecular background of the ABO system was resolved in 1990, when Yamamoto \textit{et al.} found that the A and B antigens are determined by different sugar molecules attached to the RBC membrane by a specific glycosyltransferase enzyme, encoded by the \textit{ABO} gene\textsuperscript{165}. A mutation in this gene gives a slightly different version of the enzyme, which in turn affects which sugar molecule it attaches to the sugar backbone (also known as the H antigen) on the RBC membrane. People with the O blood type carry mutation(s) that completely abolishes the enzyme function, resulting in no sugar molecules being added to H, while type AB carry both variants of the enzymes and thus both A and B antigens can be found on the RBC membrane.
Table 2: Blood groups systems recognised by the ISBT, December 2016.

<table>
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<tr>
<th>#</th>
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<th>System symbol</th>
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<td>ACKR1</td>
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<td>SEM47A</td>
<td>6</td>
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</tr>
<tr>
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</table>

1) From Daniels 2011\(^{166}\). For CD59\(^{167,168}\) and for Augustine\(^{168}\).
Structure and function of antigens

Blood group antigens are found on well-defined structures on the surface of RBCs. These structures are often classified into six groups according to proposed or known function\(^{[170]}\): the glycocalyx, membrane transporters and channels, receptors and adhesion molecules, complement regulatory proteins, enzymes and cytoskeletal proteins (Table 2). The ABO antigens belong to the carbohydrates of the glycocalyx, sugar molecules that make up an outer protective layer of the RBC membrane. The structure of SMIM1, the protein responsible for the Vel blood group antigen, has been predicted to be a single-pass transmembrane protein (TMP)\(^{[171]}\), yet a function has not yet been described.

In contrast to their crucial role in transfusion medicine, most of the molecules that carry blood group antigens seem to be non-essential for the individual, as evidenced by the null individuals of various blood group systems (i.e. individuals homozygous for mutated and dysfunctional copies of the blood group gene, so that no antigen is expressed on RBCs), who show little to no signs of compromised health\(^{[172]}\). In most cases, other proteins take over and problems directly related to the lack of the antigen/protein only arise when the individual is exposed to stressful conditions\(^{[173]}\). This raises the question why we even have blood group systems, which is perhaps best answered by Geoff Daniels in his book Human Blood Groups, as he muses over a quote from Charles Darwin’s On The Origin of Species: That (antigen) polymorphisms may have risen through many centuries or possibly millennia of evolutionary selection pressure, which, in our modern world today, no longer exists, making blood group antigens largely vestigial as remnants of a long forgotten past (paraphrased from\(^{[166]}\)).

Pathological implications of blood group systems

There are several examples of blood antigens being remnants from selection pressures in early human history, and while modern medicine has definitely lessened or even removed some of this pressure, blood group antigens and certain pathological conditions remain closely linked to this day.

The majority of examples revolves around the malaria parasite, whose life cycle is intricately linked to human blood; the parasite infects RBCs by directly interacting with membrane receptors and multiply within before bursting the cells and finding new to infect\(^{[174]}\). Malaria is a crippling disease killing an estimated 429,000 people annually (World Health Organisation estimate for year 2015), and as such have been, and continues to be in some areas, the biggest evolutionary selection pressure in the world today\(^{[175]}\).

The ABO system seems intricately linked to the malaria species Plasmodium falciparum. Cserti and Dzik performed a thorough analysis of the numerous
published studies linking ABO to malaria (some of which deemed to be flawed, some well-designed) and showed a convincingly high prevalence of the O blood group accompanied by a lack of A individuals in many areas around the equator where *P. falciparum* is endemic\textsuperscript{176}. A functional explanation for this was reported by Carlson and colleagues\textsuperscript{177} showing that rosette-formation, *i.e.* clumping of infected RBCs to uninfected, is highest in blood group A and AB-individuals, intermediate in B and weakest in group O. The molecular biology behind rosette-formation is the parasite-derived *P. falciparum* erythrocyte membrane protein-1 (PfEMP-1) expressed on infected RBCs, and this associates strongly with the sugar moieties of blood group A and B\textsuperscript{178}. Subsequently, PfEMP-1 has also been shown to bind complement receptor 1 (CR1, also known as glycoprotein CD35), which is the gene associated with the Knops blood group system\textsuperscript{179}.

Several other blood group genes have been proven as receptors for *P. falciparum* infection: glycophorin A\textsuperscript{180} and B\textsuperscript{181} (blood group system MNS), glycophorin C\textsuperscript{182} (Gerbich) and basigin\textsuperscript{183} (Ok), all summarised in a recent review\textsuperscript{184}.

The Duffy blood group system, first described in 1950\textsuperscript{185} and later found to be determined by the Atypical chemokine receptor 1 (ACKR1)\textsuperscript{186,187}, has been linked to two other malaria species, *P. vivax* and *P. knowlesi*. A mutation in this gene results in the Duffy-null phenotype Fy(a\textsuperscript{–}b\textsuperscript{–})\textsuperscript{188}, which is highly prevalent in West Africa\textsuperscript{189}. As it happens, ACKR1 is the primary receptor used by *P. vivax* and *P. knowlesi*\textsuperscript{190,191}, which undoubtedly have represented a solid selection pressure and have been determined to be the cause for the extremely high prevalence of Fy(a\textsuperscript{–}b\textsuperscript{–}) in West Africa, and as a result, the very low occurrence of *P. vivax* and *P. knowlesi* in these regions\textsuperscript{190,192}.

Other malignancies with ties to blood groups include evidence that having a specific ABO allele increases the risk of developing certain cancers, where carrying any non-O allele makes an individual susceptible to develop gastric (60 years of research summarized in a meta-analysis by Wang *et al.*\textsuperscript{193}) and pancreatic cancer (multiple studies, summarized in\textsuperscript{194}), the latter presumably through increased glycosyltransferase activity of the A\textsuperscript{1} allele compared to A\textsuperscript{2} and B\textsuperscript{195,196}.

Furthermore, the ABO and Lewis blood group systems both contain histo-blood group antigens (A, B, H and Le\textsuperscript{a}) meaning that the antigens are present on other tissues than blood, and these systems must be considered with regards to organ transplantation and host-graft rejection. H and Le\textsuperscript{b} antigens have also been shown to be receptors for *Helicobacter pylori* bacterial infection\textsuperscript{197}.

Last but not least, and an exception to the rule that individuals with blood group null phenotypes are generally in good health, Kx-null individuals, carrying a mutation in the XK gene, suffer from McLeod syndrome presenting with acanthocytosis, neurological disorders and cardiomyopathy\textsuperscript{198}. 


The Vel blood group system

The Vel antigen was first described in 1952 in New York, when a female patient, Mrs. Vel, presented with an acute HTR after receiving a second transfusion of crossmatch-compatible group O RhD-negative blood[199]. Her serum was tested against 10,000 blood donors in New York and only four were compatible, establishing a Vel-negative (Vel−) frequency of 1 in 2,500 individuals and the antigen was thus placed in the high incidence antigen collection by the ISBT. Other screening studies in the UK, Australia, Norway, Finland and France found a total of 43 Vel− people out of 159,565, giving an estimated global incidence of 1 in 3,711200-205, summarised by Issitt and Anstee206. Interestingly though, a study performed by Cedergren et al.207 found a higher prevalence of 1 in 1,761 individuals in Northern Sweden, pointing towards a potential founder effect for this historically isolated and sparsely populated area.

The Vel antigen is a major concern in transfusion due to the severity of HTRs in some transfused individuals. Vel− individuals usually present without anti-Vel in their blood, but develop alloantibodies upon contact with the Vel antigen. The antibody present as both IgG and IgM, but rarely causes haemolytic disease of the foetus and newborn (HDFN)206. Antigen expression on the RBCs of Vel-positive (Vel+) individuals vary greatly, with some having such low values in serological tests that they are at risk of being typed as Vel− (henceforth referred to as Vel+ weak)171. Furthermore, due to the low incidence of Vel− individuals, a sufficient supply of anti-Vel for screening has not been available, which has made phenotyping, research and classification all the more demanding. In 1968, Issitt et al. reported a second Vel antibody, Vel 2, and presented cases characterised as Vel:1,−2 and Vel:−1,−2208, however the authors later disputed their own findings deeming the difference between the two antibodies to be purely quantitative206.

The Vel antigen remained a mystery until 2013 when we and two other groups independently, and more or less simultaneously, discovered that the Vel− phenotype is caused by homozygosity for a 17-bp deletion in the coding sequence of the erythroid gene small integral membrane protein 1 (SMIM1)171,209,210. This formally established Vel as a blood group system, recognised as system 034 by the ISBT in 2014211. Study I describes this discovery.
INTRODUCTION

The SMIM1 gene responsible for the Vel blood group system

SMIM1 is a small gene of 3,187 bp located on chromosome 1p36.32 (Figure 4). It is highly conserved with homologs in many different species and has four exons with the open reading frame spanning exon 3 and 4. As noted above, a 17-bp deletion in the open reading frame of exon 3 (c.64_80delAGCCTAGGGGCTGTGTC) results in a frame-shift mutation and stop codon skip, creating a dysfunctional protein, and thus a SMIM1-null phenotype. Other sequence variants are found throughout the gene, and in intron 2 in particular, in a region which is also rich in transcription factor binding sites, eTF occupancy (GATA-1, TAL1, ZBTB7A and others) and chromatin remodelling favouring active DNA transcription (as found by the Encyclopedia of DNA Elements (ENCODE)). Previous studies showed the sequence variant rs1175550 present in this region to directly influence the expression of the Vel antigen, making it a strong candidate to explain the large variation in Vel antigen expression routinely observed with Vel+ individuals. We confirmed this finding and also found an additional sequence variant, rs143702418 located only 96 bp upstream of rs1175550, which independently influence SMIM1 and Vel antigen expression (Study II of this thesis). An additional sequence variant in the coding region c.152T>G/A causes missense mutations p.Met51Arg and p.Met51Lys, respectively, and both mutations have been associated with lower Vel antigen expression.

The function of SMIM1 remains unknown; however, structure analysis predicted it to be a single-pass TMP with a GXXXG motif in the transmembrane domain, which is commonly associated with homodimer formation. An in silico gene network model showed SMIM1’s closest neighbours to be typical erythroid genes

Figure 4. SMIM1 gene, transcript and protein. Four exons make up the 78 amino acid protein. The genomic sequence contains at least four sequence variants influencing the expression of SMIM1. The red patch is the Vel-defining 17-bp deletion in exon 3. The SMIM1 protein is a single-pass transmembrane protein with several serines and threonines available for phosphorylation, three cysteine residues to facilitate protein:protein interaction and a GXXXG homodimer motif. The final three C-terminal amino acids Lys76-Cys77-Lys78 is hypothesised to have a role in Vel antigen epitope presentation. Adapted from Study I.
(RHD, XK, KEL, KLF1 etc.)\textsuperscript{171}. We originally proposed it to be a type I TMP, but a recent study reported findings to support a type II TMP with a very short C-terminal extracellular domain of only 3-12 amino acids\textsuperscript{216}. This study also pinpointed the final three amino acids (KCK, or Lys76-Cys77-Lys78) as crucial for the expression of the Vel epitope recognised by human anti-Vel. Finally, large GWA studies have linked rs1175550 with mean corpuscular haemoglobin content (MCHC)\textsuperscript{217} and blood copper concentration\textsuperscript{218}, suggesting that SMIM1 could play a role in copper transport, either as a main transporter or a cofactor of known copper transporters.
Large-scale analyses of genetic variation

Mutations in the genome occur naturally in the cell as the DNA is replicated, but these errors are quickly repaired or are often located in places that do not take part in active replication, transcription or regulation and thus are, apparently, silent. Only rarely do mutation have a detrimental effect on an individual’s health. Still, silent mutations contribute to changing the individual’s genome and since these mutations are passed on to the next generation, over time a certain population will contain a genomic profile that varies from other populations having been exposed to different mutation events and selection pressures. This gives rise to designations such as single nucleotide polymorphisms (SNPs), insertion/deletions (indels) and copy number variations (CNVs), collectively known as genetic variation, which has been a major research focus over the last decades. Starting with the multinational research project The Human Genome Project, that successfully sequenced the entire human genome several other studies have since followed; the UK10K aimed to map the genetic variation in 10,000 individuals from the United Kingdom with special focus on disease-causing variants; the African Genome Variation Project aimed at mapping the genetic diversity of 1,481 individuals in sub-Saharan Africa to gather information on this population group that has played an important role in ancient human history and continues to be under selection pressure; and the 1000 Genomes Project, a comprehensive study of global population variation in the full genomes of 2,504 individuals from five global population (African, Admixed American, European, East Asian and South Asian), divided into 26 subpopulations (such as African American, Finns, Japanese, Peruvians, Southern Han Chinese etc.). Additional noteworthy databases are the Encyclopedia for DNA Elements focused on transcription factors binding sites in the genome as well as various types of chromatin modelling affecting transcription; the Exome Aggregation Consortium focused on analysing whole exome data from 60,706 individuals, and dbSNP which collects and curates the continuous stream of new sequence variants being reported (currently contains more than 89 million validated human SNPs). The last few years have seen the scope of genomics projects significantly expanded with several planned projects aiming at sequencing individuals in the millions; the Qatar Genome, AstraZeneca’s multicentre collaboration (genomes of up to two million individuals to be analysed in the United Kingdom, Finland and the United States), the 100,000 Genomes Project in the United Kingdom, and the Precision Medicine Initiative (one million individuals) in the U.S.
Sequence variants such as SNPs and indels can only be described in relation to a given population. It is a locus (a single-base pair substitution for the former, and an insertion or deletion of multiple base pairs for the latter) in the genetic code, where a group of individuals varies from the rest of the population. These variants may confer a selective advantage in some cases, but are also one of the prime causes for diseases and disorders. With the completed Human Genome Project came the prospect of precisely pinpointing a disease to a gene region. However, the found sequence variants were in the millions, too great a number to test in large groups of patients. This was overcome by the discovery of haplotype blocks: During homologous recombination the DNA is broken at non-random sites leading to larger chunks of DNA (“haplotype blocks”) remaining intact through several generations. These haplotype blocks will thus contain several sequence variants that are said to be in linkage disequilibrium (LD) and inherited together. Mapping the entire genome’s haplotype blocks was done by the HapMap project. Combined with sequence variants collected in databases such as dbSNP, this facilitated the identification of index or tag SNPs representing only a single sequence variant but by proxy including the other variants in the haplotype block as well, significantly reducing the number of SNPs to investigate. From there the genome-wide association study (GWAS) was quickly developed, which combined index SNP analysis with case-control population cohorts, thus determining potential risk alleles, i.e. specific variants commonly associated with various diseases. The largest of the initial GWA studies investigated seven diseases in 14,000 cases and 3,000 controls and found several SNPs and risk loci; many have since followed, most relevant for this thesis is a study that found 75 SNPs (one of them rs1175550) that influence various red blood cell parameters. An approach similar to GWAS, but using gene expression profiles instead of case-control studies, yields expression quantitative trait loci (eQTLs) and combining the two to find overlapping reported SNPs greatly enhances the ability to characterise true association. This is achieved as GWAS looks at the phenotypic results of the variation on the organism as a whole, while eQTLs identifies the affected gene and thus gives a genetic background for the observed phenotype.

Trans-ancestry association studies

GWAS and eQTL studies were major breakthroughs and facilitated the discovery of disease-associated risk loci (i.e. causal variants) in numerous diseases including haematological disorders such as marginal zone lymphoma, acute lymphoblastic leukaemia and multiple myeloma. Yet, the techniques are inherently confined to discovering haplotype blocks, which makes it difficult to determine a specific causal variant locus if the LD in a region is strong. Likewise, due to the LD, a particular discovered index SNP may not be the disease-causing variant at all,
rather a proxy SNP for one of the other variants in the block that actually drives the trait. This is where trans-ancestry (or -ethnic) association studies come in, which can be seen as a way to mix GWAS/eQTLs with population genetics. Through generations of genetic code rearrangements and human migrations the unique genetic code of different populations spread across the globe and they continue to diverge (and occasionally converge) to this day. This means that haplotype blocks (and thus strength of LD) between sequence variants differ among population groups, and this can be exploited by combining GWAS or eQTL studies with population-based genetic variation, such as that reported by the 1000 Genomes Project. The last few years have seen an increase in high-throughput trans-ancestry association studies, and a very recent study identified seven loci for erythrocyte traits (including MCHC) using individuals from the African, European and South Asian superpopulations.

**Genetic variation in blood group systems**

A blood group antigen may arise when a sequence variation in the genetic code creates an immunogenic epitope that sets one individual apart from the general population. Such variation can be as small as a single or a few SNPs, as is the case for the *ABO* gene, or it can be indels of longer stretches of DNA in the genetic code, *e.g.* the 17-bp deletion in the coding region of *SMIM1*. The ISBT keeps track of and categorises all blood groups systems and antigens as they are discovered, and they are thoroughly described in the *Blood Group Antigen Factsbook*. Lastly, blood group systems and population genetics are obviously connected, and a recently published online database annotates sequence variants in blood group genes with population allele frequencies extracted from the 1000 Genomes catalogue.
The present investigation

Aims

This thesis investigated the gene SMIM1, which at the start of the thesis had only just been identified as having a potential role in the expression of the high incidence antigen Vel.

The overall aims were to:

- Confirm the putative erythroid gene SMIM1 as the gene underlying the Vel blood group system.

- Identify and characterise sequence variants in SMIM1 that modulate the expression of SMIM1 and the Vel blood group antigen.

- Investigate genotype-phenotype association in archived Vel-negative samples collected before the genetic background for Vel was discovered.
Summary of studies

Study I

Homozygosity for a null allele of SMIM1 defines the Vel-negative blood group phenotype

This study, together with two other studies published almost at the same time, determined the molecular background of the Vel antigen. The three studies used different approaches, but all reached the same conclusion: that a mutation in the gene SMIM1 was the prime cause of the lack of expression of the Vel antigen.

Our approach consisted of microarray-based SNP profiling of the genomes of 20 Vel– individuals from two Swedish families and comparing them to their Vel+ relatives. Comparing 5 Vel– individuals to their Vel+ siblings we narrowed the potentially causal SNPs from more than two million to 8,780 SNPs that segregated with the Vel– phenotype. After a second filtering combining the remaining 15 Vel– individuals with minor alleles frequencies in individuals of European descent in the 1000 Genomes catalogue\textsuperscript{223}, we found a region on chromosome 1p36, which was unique for all Vel– tested. This region contained five genes, where one in particular caught our attention, SMIM1, or, as it was known then, LOC388588. Protein structure and function algorithms predicted it to be a type I TM and an \textit{in silico} analysis of the gene network neighbourhood surrounding SMIM1 revealed the nearest genes to be largely erythroid, \textit{e.g.} KLF1, RHD and KEL. Furthermore, SMIM1 was found highly expressed in erythroleukaemia cell lines (Cancer Cell Line Encyclopedia\textsuperscript{240}) as well as upregulated in human CD34+ cells cultured towards erythroid differentiation\textsuperscript{241}. Sequencing of SMIM1 revealed that all 20 Vel– individuals were homozygous for a 17-bp deletion whereas all Vel+ had the consensus sequence. This deletion is located within the coding sequence and introduces a frame-shift that skips the stop codon. We took the following steps in our efforts to determine if SMIM1 was indeed responsible for the long sought after Vel antigen:

- 5’ and 3’ Rapid Amplification of cDNA Ends (RACE) experiments found the consensus SMIM1 mRNA sequence when using total RNA from Vel+ individuals, and no detectable sequence with RNA from Vel– individuals.
- Western blot with human anti-Vel and rabbit polyclonal antibodies raised against the presumed extracellular domain of SMIM1 detected bands of ~20kDa on Vel+ RBC membranes, but nothing on Vel– RBC membranes.
• Flow cytometry with a human anti-Vel gave a strong reaction with samples homozygous for wild type SMIM1, less so for heterozygous samples and virtually no reaction for samples homozygous for the SMIM1 17-bp deletion.

• Overexpression of SMIM1 wild type in the erythroleukaemia cell line K562 greatly increased human anti-Vel reactivity assayed by flow cytometry, whereas cells expressing the deletion mutant did not. This was also seen in Western blots of the cell membranes using the rabbit anti-SMIM1 antibody.

Conclusions
On top of the conclusions already made, the above steps determined that:

• SMIM1 encodes an erythroid single-pass transmembrane protein with a GXXXG homodimer domain and a 17-bp deletion causing a frameshift mutation exclusively present in Vel− individuals.

• No translatable SMIM1 mRNA is created in Vel− individuals.

• Human anti-Vel and rabbit anti-SMIM1 both recognise the same protein on Vel+ RBC membranes; they do not detect anything on Vel− RBCs.

• SMIM1 deletion zygosity affects the expression of the Vel antigen on the RBC surface (i.e. evidence of a gene dosage effect).

• The SMIM1 open reading frame encodes a protein that reacts strongly with human anti-Vel antibody and the deletion abolishes this reactivity.

Based on this, SMIM1 was established to be the gene responsible for the elusive Vel blood group antigen, and that the Vel− phenotype is caused by a 17-bp deletion resulting in abolished SMIM1 expression. This determined the molecular background for the Vel antigen, prompting it to be recognised as the Vel blood group system, and be given number 034 by the ISBT.
Study II

SMIM1 variants rs1175550 and rs143702418 independently modulate Vel blood group antigen expression

Having found SMIM1 to be the gene responsible for the Vel blood group system, we then set out to examine it in detail. A genome-wide search for sequence variants affecting binding of eTFs located the SNP rs1175550 (A>G) in intron 2 of SMIM1. This SNP changes a non-canonical binding site of GATA-1, GATT, to GGTT, and could potentially influence binding of this prime erythroid transcription factor to SMIM1. rs1175550 was reported to influence mean corpuscular haemoglobin content\textsuperscript{217} and was later shown to have a role in Vel expression\textsuperscript{210,213}, however no functional explanation for this was given.

Examining 150 samples from Swedish Vel+ donors we:

- Confirmed that rs1175550G is associated with high expression as determined by mRNA expression levels, human anti-Vel reactivity and SMIM1 protein expression on RBC membranes.
- Discovered a new sequence variant, rs143702418 – a trinucleotide insertion (C>CGCA), located 97 bp upstream of rs1175550.

The two variants were in near-perfect linkage disequilibrium (LD) making it impossible to pinpoint the causal variant. By consulting the 1000 Genomes catalogue, we realised that the two SNPs were frequently unlinked in individuals of African descent. Thus, repeating the above experiments in a collection of 202 African American samples, we found:

- rs1175550G still correlated with high expression.
- rs143702418 was more mixed and no direct correlation was found.
- Regression analysis conditioned on rs1175550 showed an independent effect with rs143702418C being the high expressing allele.

We expanded our analysis with functional studies of the two variants. We created four luciferase constructs corresponding to the four possible combination alleles existing between the two bi-allelic variants and confirmed that the rs1175550G and rs143702418C are the most transcriptionally active variant alleles. In gel shift assays with antibody supershifts and with probes spanning either allele of the variants we found that GATA-1 and Gfi-1B bind near the rs1175550 variant in both alleles, whereas a complex containing TAL1 preferentially binds only to the rs1175550G allele. Assays for rs143702418 showed differential binding patterns, but no antibody supershifts.
Conclusions:
Analysing 150 Swedish and 202 African American blood samples, we found that:

- rs1175550 and rs143702418 independently influence SMIM1 and Vel antigen expression, partly explaining the large variation in anti-Vel reactivity routinely observed for Vel+ individuals.
- TAL1 is likely to mediate the increased expression as it binds exclusively to the high-expressing rs1175550G allele.
- GATA-1 and Gfi-1B are mainly associated with the low-expressing rs1175550A allele.
- Using population genetics to deconvolve tightly linked variants we demonstrated a small-scale trans-ancestry association analysis, which is a promising new tool for molecular geneticists.
Study III

SeroLogic And Molecular Studies Of The Vel– Phenotype In A Multi-ethnic Population

With the molecular background determined, we examined 40 historic Vel– blood samples characterised as Vel:1,–2 and Vel:–1,–2\textsuperscript{208} for the 17-bp deletion. We found:

- 22 individuals originally classified as Vel:–1,–2 were homozygous for the deletion.
- 14 individuals originally classified as Vel:1,–2 were heterozygous for the deletion.
- One sample was heterozygous for the c.152T>G mutation in the coding sequence.
- The remaining three samples did not carry the deletion or the point mutation.

We sequenced the entire SMIM1 gene region in the three outlying samples but found no apparent sequence variants to explain the Vel– phenotype. We then performed whole exome sequencing and preliminary results indicate potentially interesting variants in KLF1, where mutations have previously been linked to the rare In(Lu) phenotype of the Lutheran blood group system\textsuperscript{242}.

Conclusions

Investigating the archived Vel– samples with the current knowledge of the Vel blood group system, we found:

- The historic classification Vel:–1,–2 corresponded 100% to samples homozygous for the 17-bp deletion (i.e. true Vel– as we know now).
- The historic classification Vel:1,–2 is indicative of heterozygous carriers of the mutation, which corresponds to very weak Vel antigen expression.
- Likewise, the one sample carrying the point mutation causing an amino acid change (p.Met51Arg) also had very weakly expressed Vel antigen.
- No apparent reason for the three outlying samples has been found; whole exome sequencing data is currently being analysed.
- The 17-bp deletion is present in individuals of African, Hispanic and European descent indicating the mutation is globally distributed and happened early in human history.
Study IV

**Robust isolation of malignant plasma cells in multiple myeloma**

This study focused on biomarker shedding in multiple myeloma plasma cells. Membrane protein CD138 is a well-established marker\(^{51,52}\), yet it has one major disadvantage: It quickly disappears from plasma cells *ex vivo*\(^{52,53}\). This makes it unsuitable for delayed analysis or when working with biobank or frozen samples. We therefore set out to discover a more stable MMPC marker.

We retrieved gene expression profiles of 1,285 MM samples from microarray data in the NCBI Gene Expression Omnibus and compared expression of individual genes to that observed in 3,164 non-MM haematological malignancies to find differentially expressed genes specific for MM. This *in silico* search was further narrowed by gene ontology terms, literature and database crossmatches, potential cell surface expression, and availability of FACS-suitable antibodies. This yielded seven potential targets, CD269 (NFRSF17/BCMA), CD319 (SLAMF7/CS1), GPRC5D, FKBP11, CD208 (LAMP3), ITGA8, and CD307e (FCRL5).

We then did FACS analyses of bone marrow cells either freshly sampled, stored at 8°C or frozen, using antibodies against these seven targets plus CD138 and found:

- CD319, CD269 and CD307e consistently sorted distinct cell populations in all three experimental conditions. The remaining four did not.
- These populations were confirmed to be MMPCs by morphology, clonal excess, and DNA copy number microarray (chromosomal abnormalities).
- CD138 was confirmed to be suitable for fresh samples, yet ineffective in samples that had been stored or frozen.

**Conclusions**

Narrowing a large list of potential MMPC-specific proteins to seven putative MM-specific surface markers, we found that:

- CD269 and CD319 are robust markers suitable for isolating MMPCs from bone marrow cells, regardless of storage conditions.
- These are solid replacements for CD138 as MMPC markers in FACS analysis.
- Robust isolation of MMPCs with handling methods suitable in a clinical setting enables thorough analysis and characterisation of MMPCs.
Methodology

A list of methods utilised in this thesis is presented in Table 3. No new methodology was developed, but a gene-specific PCR capable of quickly detecting the SMIM1 17-bp deletion (and thus Vel status) in a single PCR run was designed, as well as a new marker (CD319) for immunophenotyping MMPCs in FACS analysis was characterised.

Table 3. Methods employed in this thesis.

<table>
<thead>
<tr>
<th>Name</th>
<th>Study</th>
<th>Brief description of usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>5‘ and 3‘ RACE</td>
<td>I</td>
<td>Determine the start and end point of SMIM1 mRNA from Vel+ and Vel− individuals.</td>
</tr>
<tr>
<td>Cell culture</td>
<td>I, II</td>
<td>Maintain and passage K562 and HEL cells.</td>
</tr>
<tr>
<td>Cloning and transfection</td>
<td>I, II</td>
<td>Insert the SMIM1 coding sequence or a DNA fragment into GFP/luciferase vectors, then transflect into K562 or HEL cells before flow cytometry analysis/luciferase assays.</td>
</tr>
<tr>
<td>Flow cytometry</td>
<td>I, II</td>
<td>Analysis of transfected K562 cells or human RBCs coated with a human anti-Vel antibody.</td>
</tr>
<tr>
<td>PCR</td>
<td>I, II, III</td>
<td>Wide usage to amplify various SMIM1 regions.</td>
</tr>
<tr>
<td>Gene-specific PCR</td>
<td>I</td>
<td>Primer design and parameter optimisation of a PCR able to determine SMIM1 17-bp deletion zygosity.</td>
</tr>
<tr>
<td>Sanger sequencing</td>
<td>I, II, III</td>
<td>Identify genetic variants in SMIM1 intron 2 or full gene; in silico electropherogram analysis.</td>
</tr>
<tr>
<td>Total RNA extraction</td>
<td>II</td>
<td>Phenol-chloroform extraction from whole blood samples of Swedish and African-American origin.</td>
</tr>
<tr>
<td>RT-qPCR</td>
<td>II</td>
<td>cDNA synthesis from total RNA samples, then quantitative PCR to examine SMIM1 expression.</td>
</tr>
<tr>
<td>Gel shift assay with antibody supershift</td>
<td>II</td>
<td>Nuclear extracts from K562 and HEL cells, mixed with biotin-labeled oligos corresponding to the found SMIM1 genetic variants and antibodies recognising select eTFs.</td>
</tr>
<tr>
<td>Luciferase assays</td>
<td>II</td>
<td>Luciferase constructs transfected into K562 or HEL cells, then harvested, lysed and analysed by chemiluminescence.</td>
</tr>
<tr>
<td>Whole genome amplification</td>
<td>III</td>
<td>Unspecific, unbiased amplification of the full genome of the outlying Vel− samples in preparation for whole exome sequencing.</td>
</tr>
<tr>
<td>Bioinformatic analyses</td>
<td>I, II, III</td>
<td>UCSC genome browser, 1000 Genomes database and dbSNP search and filtering, sequencing data, JASPAR TF motif scan, whole exome sequencing data.</td>
</tr>
<tr>
<td>FACS</td>
<td>IV</td>
<td>Sort bone marrow cells coated with fluorophor-conjugated antibodies.</td>
</tr>
</tbody>
</table>
Discussion

The Vel blood group system has now been solidly linked to the SMIM1 gene. With the 17-bp deletion known, we designed a gene-specific PCR that amplifies a region of 178 bp spanning the deletion, allowing for quick identification of deletion carriers, and while several studies have reported on various aspects of SMIM1, such as Study II of this thesis and the study of the Vel antigen epitope on SMIM1 by Arnaud and co-workers\(^\text{216}\), a specific function remains to be found.

Population genetics of Vel/SMIM1

As noted in the introduction, several studies were performed in the decades after the Vel antigen was reported in 1952, most of them on Caucasian population groups, but also two small studies suggesting a much higher incidence of Vel– individuals (1 in ~80) in both Thais\(^\text{243}\) and Chilcotin Indians in Canada\(^\text{244}\). The discovery of the 17-bp deletion in SMIM1 opened up for genetic screens for the Vel antigen and we used our gene-specific PCR on a collection of 520 random blood donors to establish a Vel– incidence in Southern Sweden of 1 in ~1,200\(^\text{171}\), which is significantly higher than the global incidence, but this might be explained by the close relation to the hypothesised potential founder population in Northern Sweden described by Cedergren et al. in 1976. Similar gene-specific PCRs were employed to establish a Vel– incidence of 1 in ~2,100 individuals in southern Germany\(^\text{245}\) and 1 in ~4,700 in a Brazil\(^\text{246}\).

The apparent high incidence of Vel– individuals in Thais and Chilcotin Indians could be the result of a founder effect but it should also be noted that Vel status in these early studies were determined by serology with human anti-Vel. There is considerable variation in anti-Vel reactivity even among Vel+ individuals and it has been shown that individuals heterozygous for the SMIM1 17-bp deletion always show very low reactivity, even if the Vel antigen is present on their RBCs. Thus, the unusually high frequency of Vel– in Thais and Indians can possibly be explained by a high number of Vel+\(^\text{weak}\) individuals wrongfully typed as true Vel–.

In support of this hypothesis, at least in a Southeast Asian population, no 17-bp deletion carriers were found in a study of 325 individuals of Malaysian and Indian ancestry (personal communication; Veera Sekaran Nadarajan). Allele frequency for the low SMIM1-expressing rs1175550A allele was 97.5% in these samples, which is in line with the allele frequencies of 99.8% in the East Asian
superpopulation in the 1000 Genomes catalogue. This could mean that there is a high frequency of Vel$^{+}\text{weak}$ individuals in the Asian population, leading to a potentially higher amount of individuals falsely identified as Vel$^-$ in serotyping screens. There are unfortunately no Native American populations in the 1000 Genomes catalogue, nor have any SMIM1 genetic screens been performed, so we cannot know if the high incidence here is due to true Vel$^-$ or Vel$^{+}\text{weak}$ individuals.

Unfortunately, the 17-bp deletion is not called in the 1000 Genomes catalogue presumably due low sequencing coverage in the region and that indels are inherently hard to impute from SNP microarray data. This precludes a thorough analysis of the global distribution of the Vel-defining sequence variant, yet we were able to find the deletion in a few samples of Hispanic, Caucasian and African descent in Study III, and combined with the studies of Asian and Brazilian population groups, there is ample evidence to assume that the mutation is distributed worldwide, and that it thus presumably arose early in human history. Evidence for an ancestral allele was also presented by Ballif and coworkers who found the SNP rs71634364, located 240 bp upstream of the 17-bp deletion, to be in LD with the deletion in 67 out of 68 Vel$^-$ cases$^{209}$. And we and the third group that discovered SMIM1 have also observed that rs1175550A and the 17-bp deletion almost always exist on the same allele, pointing at potential LD between the two (unpublished data and personal communication with Ellen van der Schoot).

*Vel antigen presentation and SMIM1 function*

As described earlier, SMIM1 was recently characterised as a type II TMP with a short extracellular C-terminal and the final three amino acid residues crucial for proper Vel antigen presentation$^{216}$, yet questions remain regarding how the antigen is presented on the RBC surface and what the specific function of SMIM1 is. The find directly opposes our conclusion in Study I that SMIM1 is a type I TMP and in the following, I will try to reconcile these two seemingly contradictory characterisations, and argue that they are not necessarily mutually exclusive. But first, a bit about antibody specificity and antigen presentation.

A recent study reported the production of a monoclonal anti-Vel antibody (SpG213Dc) raised from plasma anti-Vel found in a Vel$^-$ individual and propagated in a human myeloma cell line$^{247}$. This detects a 32kDa protein in Western blots under non-reducing conditions and a 20kDa when reduced. This is in line with results presented by Ballif et al.$^{209}$ and in our own Study I with anti-Vel from immunised Vel$^-$ individuals. Since the predicted molecular weight of SMIM1 is 8.75kDa it can be speculated that the 20kDa band found under reducing conditions might be a homodimer probably held together by the reported transmembrane GXXXXG motif$^{71}$, and protected from reducing agents by the RBC
membrane, as is the case for GPA, which forms detergent-resistant transmembrane dimers through its GXXXG motif\textsuperscript{15,248}. The size difference may also be due to posttranslational modifications, yet we found SMIM1 to not be O-glycosylated or sialylated. The SMIM1 32kDa band observed in native conditions can be thought to be a multimeric complex consisting of homodimeric SMIM1 and one or more unknown SMIM1 partner(s), interacting through domains located outside the transmembrane region. Indeed, SMIM1 contains two cysteine residues N-terminal of this region (located in the cytoplasm if SMIM1 is a type II TMP) capable of forming disulphide bonds, however these cysteines were shown to be expendable for proper human anti-Vel reactivity by Arnaud et al.\textsuperscript{216}. Another cysteine residue (Cys77) is present as the penultimate residue in the C-terminal, and this might very well be crucial for direct interaction with one or more proteins to form the Vel antigen epitope, as absence of this and the two lysine amino acids flanking it (referred to as the KCK motif in the following) abolished anti-Vel reactivity entirely\textsuperscript{216}. In Study 1, we raised an anti-SMIM1 antibody by immunising a rabbit with a polypeptide corresponding to a region in the N-terminal of SMIM1\textsuperscript{171}. In contrast to the bands detected above, we observed two bands of ~9-10 and ~20kDa when examining solubilised RBC membranes under reducing conditions. And when we treated whole RBCs with the protease chymotrypsin, the bands disappeared, indicating that the anti-SMIM1 epitope presumably sits on the RBC surface. This finding seemingly contradicts SMIM1 being a type II TMP as anti-SMIM1 would then have been raised against the intracellular part.

So the questions remain: What do the different antibodies recognise, and what is the correct orientation of SMIM1 in the RBC membrane. Regarding the first question; our anti-SMIM1 recognises a linear sequence of amino acids in SMIM1, but not necessarily the Vel antigen epitope (indeed, despite multiple attempts, we were unable to haemagglutinate RBCs using anti-SMIM1\textsuperscript{171}), while the human anti-Vel and the SpG213Dc antibodies obviously recognise the Vel antigen epitope (since it is pulled directly from plasma from immunised Vel– individuals). But since antigen epitopes can be dependent on conformation and protein fold as much as amino acid sequence, it follows that anti-Vel does not necessarily bind specifically to a linear stretch of amino acids in the SMIM1 protein, but rather to whatever native conformation and/or complex SMIM1 exists in the RBC membrane. The anti-Vel can even be thought to specifically recognise another protein entirely, but a protein (complex) that is critically dependent on a SMIM1 anchor on the RBC membrane for proper antigen presentation. Such membrane multicomplexes are not a new concept for blood group systems, where the proteins of several systems come together in one massive macrocomplex. For example, GPA and GBP (MNS blood group system), RhAG (RHAG), RHD and RHCE (Rh), LW (Landsteiner-Wiener) and CD47 all form complexes that are anchored to the cell membrane through Band 3 (Duffy), Protein 4.2 and Ankyrin. Absence
of Band 3 results in significant reduction of MNS, Rh and RhAG antigens on the RBC surface. If SMIM1 is indeed a scaffold/anchor protein, it follows that absence of SMIM1 would mean no antigen epitope formation and display, leading to a Vel– phenotype.

The question of SMIM1 membrane orientation in the RBC membrane is not straightforward to answer, based on the few studies published so far. Either antibody (anti-SMIM1 and anti-Vel) supposedly recognise epitopes on either side of the transmembrane region and both antibodies presumably recognise something on the extracellular surface of RBCs. And both detect proteins that can be deduced to be SMIM1 or complexes that SMIM1 is a part of. Interestingly, papain treatment resulted in a reduction of ~4kDa with SpG213Dc anti-Vel, and decreased band intensity with anti-SMIM1, meaning that either conformation complex is sensitive to protease degradation of some of its components.

In summary, there is evidence that SMIM1 exists as both a type I and a type II TMP in the RBC membrane. As a type I monomer and (homo-)dimer, but not necessarily presenting the Vel antigen and as a type II with a short extracellular C-terminal and the final three amino acids important for antigen presentation. Future research should focus on finding SMIM1 binding partners, which would ascertain if either of these orientations can be excluded and shed light on the actual nature of the Vel antigen presentation.

On a final note, even various online protein sequence analysis tools predict the membrane topology differently. All tools predicted a single-pass TMP without signal peptide, but eight out of ten (e.g. CCTOP and Philius) predicted SMIM1 to have the N-terminal on the cytoplasmic side (i.e. Type II TMP), while Phobius and TMHMM predicted the opposite orientation. The lack of a predicted signal peptide conflicts with classifying SMIM1 as a type I TMP, but there are examples of extracellular N-terminal proteins with reverse signal anchors that function as a signal peptide, but are not cleaved when the proteins are inserted in the ER membrane (these are also referred to as type III TMPs).

**SMIM1 as an invasion receptor for malaria?**

In Study I we hypothesise that SMIM1 may be a receptor for *P. falciparum* and there is some evidence to support this. First, SMIM1’s structure as a single-pass TMP with a GXXXG motif is also characteristic for GPA, which, as noted in the introduction, has been shown to be a receptor for *P. falciparum* infection. Second, we found SMIM1 to not be sialylated, yet resistant to trypsin and very sensitive to chymotrypsin treatment, three characteristics of an unknown erythroid receptor (receptor Z) for the *P. falciparum* ligand *P. falciparum* reticulocyte binding homologue 2b (PfRH2b) which has been described, but not yet linked to a certain protein in the RBC membrane. And third, SMIM1 was found to be
DISCUSSION

phosphorylated on Ser22 and Ser28 (see Figure 4 on page 37) in a study of the global phospho-proteome in RBCs infected with *P. falciparum*, along with 25 other human proteins, several of which are well-known erythroid proteins, such as Band 3, Glycophorin C, Protein 4.1, Kell, CD44 and Ankyrin-1. It seems likely that SMIM1 plays some form of role in malaria pathogenesis, yet at this point, more research is needed.

**Sequence variation in SMIM1**

In Study II, we found that both rs1175550 and rs143702418 in SMIM1 intron 2 influence SMIM1 and Vel expression, partially explaining the large variation in anti-Vel reactivity observed among Vel+ individuals. We initially performed a scan of sequence variants disrupting eTF binding sites, thus potentially altering expression of the target gene and found rs1175550 due to its position in a non-canonical binding site for GATA-1; GATT, with the minor allele changing this to motif GGT. rs1175550G is the effector allele responsible for high Vel expression. But how is this regulation achieved? Our luciferase assays in Study II showed that the region surrounding the variant is certainly capable of attracting factors that affect transcription, and reports have shown that GATA-1 bound to the GATT motif can lead to transcriptional activation in testicular cells. However when GATA-1 is associated with Gfi-1B (whose binding motif is AATC, the reverse-complement of GATT), it leads to repression in erythroid cells of the genes Gfi-1B and Bcl-xL. So maybe GATA-1/Gfi-1B are responsible for repressing SMIM1 transcription when bound to rs1175550A? Our gel shift assays were inconclusive, with GATA-1 not binding directly to GATT, however Gfi-1B was shown to not bind a complex specific for rs1175550G. However, and more intriguingly, we found TAL1 to specifically bind to this complex, consistent with a very recent report showing that TAL1 binds near rs1175550 and preferentially to the G allele. We thus assumed that TAL1 is a major factor governing the high expression of rs1175550G. Again, how is this achieved? Obviously the A>G shift disrupts Gfi-1B binding and potentially inhibits its repressor function. But what about TAL1 binding sites? Shortly upstream of rs1175550 is a variant of the TAL1/GATA-1 composite motif described earlier – (CTG(n)GATT) – yet this would suggest that the rs1175550A should be the high-expressing allele, so presumably this motif is not used. Another TAL1-like motif is also present, CAGCCT, (degenerate from the canonical CAGNTG motif) and might be what TAL1 binds to in our gel shift assays. However the most compelling motif in the region is actually found on the antisense strand where an exact match for the composite motif can be found, starting six basepairs upstream from rs1175550 (motif on the sense strand is TTATCA(n7)CAG, Figure 5). Although GATA-1 binding to GATA/GATT motifs seems fairly redundant for transcriptional regulation, TAL1 and the
composite motif is a good predictor for transcriptional activation, so it is very plausible this motif is used in vivo. This theory can easily be tested by running gel shift assays with probes spanning the composite motif and antibody supershifts with anti-GATA-1, -Gfi-1B and -TAL1. Binding specificity can then be examined by using probes with the specific binding sites mutated.

The question remains why rs1175550 is an eQTL in this regard, if TAL1 binding is not directly affected by it. I would argue that since the CAG and GATT motifs are only separated by five basepairs, the occupancy could very well be governed by steric hindrance; for rs1175550A the GATT (AATC) motif is occupied by GATA-1 and/or Gfi-1B while the substitution to rs1175550G disrupts this binding and allows TAL1 to bind immediately upstream, displacing or blocking Gfi-1B (Figure 5).

**Figure 5** rs1175550 alleles and proposed bound erythroid transcription factor complexes. Expression variation is hypothesised to be governed by physical blocking of one eTF complex by another. The rs1175550A allele allows binding of the transcriptional repressor complex consisting of Gfi-1B and GATA-1 (and possibly the CoREST multicomplex, not shown). rs1175550G disrupts this site, allowing TAL1 and the transcriptional activation pentameric complex to bind to the composite TAL1-GATA motif only six basepairs upstream. Transcriptional regulation resulting in high or low expression of SMIM1 and the Vel antigen thus becomes a matter of steric hindrance by two opposite-acting complexes.

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We suspected the other variant in Study II, rs143702418 (C>GCA) to also be a potential eQTL, since it is located within a binding motif for one of the other major cTFs, KLF1. The variant changes the core CACCC sequence to CACGCACC and even though this resembles the extended motif CCNCCNC (see Table 1, page 23), we found that the tri-nucleotide insertion correlates with lower SMIM1 expression and transcriptional activity, so presumably the motif is altered sufficiently to disrupt KLF1 binding. Unfortunately, we were unable to show a specific binding of KLF1 to the probes spanning the variant in our gel shift assays, but this may not necessarily rule out KLF1-binding entirely. Supershifts in gel shift assays are technically challenging and often require an antibody specifically tested for this method, which was not the case for the antibody we used. We did see an altered binding profile for the two probes (one for each variant allele) without antibody supershifts, indicating that different proteins bind to either probe. We did not find solid evidence of other erythroid proteins (i.e. GATA-1, TAL1, Gfi-1B or ZBTB7A, these antibodies worked well in the gel shift assays for rs1175550) binding at or near the rs143702418. This is not surprising as the probes do not contain any binding motifs for the antibodies being tested (Table 1), perhaps with the exception of ZBTB7A’s alternative GC-rich motifs reported by Maeda and coworkers. If KLF1 does not bind to rs143702418, which protein could then be responsible for the transcriptional regulation? The KLF-related ubiquitous transcription factor Specificity protein 1 (Sp1) binds to GC-rich regions and particularly CCCCCCCCC (JASPAR motif) and this may very well also play a role in SMIM1 expression, as Sp1 has been shown to interact and recruit GATA-1 to DNA in the absence of GATA sites. In summary, while we were unable to detect a supershift with KLF1, this does not necessarily rule out binding in this region. This could be investigated by optimising antibody binding conditions in the gel shift assay or by performing chromatin immunoprecipitation (ChiP) of erythroleukaemia cells.

KLF1 and TAL1/GATA-1 binding sites are often found within 1kb of each other and the 96 bp between rs143702418 and rs1175550 fits well with this prediction. This distance could be sufficient for rs143702418-bound KLF1 to loop the DNA and interact with the pentameric complex bound to rs1175550 to bring the SMIM1 gene into the transcription factory, as described in the introduction. This could be tested by mixing the gel shift assay probes with nuclear extracts and then co-immunoprecipitate with KLF1 and one of the components of the pentameric complex.
We were surprised to find the two SMIM1 variants tightly linked in our Swedish samples and quickly realised that with the strong LD, separating their effects would be statistically impossible. We originally planned to separate the two variants in a cell line using CRISPR-Cas9 genome editing with homology-directed repair, but soon realised that the two variants had significantly different allele frequencies in the African superpopulation in the 1000 Genomes catalogue. In other words, the LD was most likely not as strong as in Swedes. Therefore, we analysed a set of African American samples, and were able to separate the effects of the two variants using nothing but the naturally occurring population variation.

Genetic variation profile for SMIM1 full gene

In Study III, we investigated three outlying Vel samples not carrying the 17-bp deletion, but originally typed to be Vel−. Knowledge of the three samples is somewhat limited (one of them was archived in 1976), but we know that they were all Caucasian and that at least one of them presented with Vel antibodies in the plasma (i.e. probably a true Vel− individual).

We sequenced the entire SMIM1 gene plus up- and downstream regions, 6,000 bp, in all three outlying samples and a Vel+ control, and even with a small sample size and DNA region we got an interesting glimpse of the substantial interindividural variation that exist in a population. The region contains 37 sequence variants with a minor allele frequency above 1% in dbSNP version 147 and in 13 (~35%) of these, at least one sample had a different genotype than the others. Unfortunately, no variant was found to be shared for all three Vel− samples and not the control, yet we can not necessarily assume that their Vel status is caused by the same sequence variant.

There are a few known eQTLs that we already know causes low to no Vel antigen expression. The first is obviously the 17-bp deletion, but all three samples were homozygous wild type. Then there is the rs1175550 and rs143702418 described in Study II. All three samples did not contain the trinucleotide insertion (rs143702418) and were homozygous rs1175550A. The latter would confer low antigen expression, which could explain a potential mistyping, however at least one of the individuals is supposed to be true Vel−, and not Vel+weak. And finally, all three samples were also homozygous wild type for the triallelic variant c.152T>G/A, where the two minor alleles results in lowered SMIM1 expression.

We did find a 97-bp deletion compared to the curated RefSeq sequence, only 200 bp upstream of the transcription start site, which was present in all four samples. Interestingly, the sequence variant rs367810010 (minor allele is a deletion of 16 bp GCCCCCGCCCCCCTCTC, minor allele frequency in Europeans: 23%) is located
at the very start of this deletion. The region is quite GC-rich and contains several repeated sequences, so we cannot rule out either sequencing or annotation errors in the RefSeq or our sequencing reaction, nor can we say if the rs367810010 is actually longer than the 16 bp reported by dbSNP. Unfortunately, low sequence coverage in the 1000 Genomes catalogue prevented us from examining if this is a variant calling artefact and if there are samples in this database that also contains the 97-bp deletion instead of only 16 bp. Some form of correlation to the variant can be assumed though, it seems too coincidental to find two deletion variants starting at the same basepair, but of varying length. Interestingly, the deleted stretch contains several binding motifs for Sp1 (and possibly KLF1), and given it’s proximity to the transcription start site, this could potentially have functional relevance for SMIM1 expression (as discussed above). It would not explain the lack of Vel antigen expression in the three outlying samples, since the Vel+ control was also homozygous for the deletion, but it is an intriguing observation nevertheless.

The question remains of what caused these samples to type as Vel−. We found no cis-eQTLs, so the answer must lie in the remaining genome. The samples come from healthy individuals, so we do not expect any detrimental mutation(s) in common TFs or genes, rather a (heterozygous) mutation in any of the eTFs or other erythroid genes. We are currently analysing data from whole exome sequencing performed on the three outlying samples and a Vel− control. We have found potential variants in erythroid genes such as GFI1B, RB1, YPEL4, ZBTB7A and KLF1. The KLF1 mutations are particularly interesting as mutations in this gene are known to be implicated in the rare blood group phenotype In(Lu)\textsuperscript{242} and cause reduced expression of CD44 (Indian blood group)\textsuperscript{261} and Kell, Duffy, RhD and other blood group antigens\textsuperscript{114,262,263}. 

DISCUSSION
Conclusions and Future Perspectives

In the work comprising this PhD thesis, we:

- established SMIM1 as the gene behind the Vel blood group antigen and devised a gene-specific PCR to detect the Vel– defining 17-bp deletion.
- characterised two sequence variants as determinants of SMIM1 and Vel antigen expression and found indications that the transcriptional activator TAL1 is involved in transcriptional regulation.
- determined a Vel incidence in southern Sweden and found that the 17-bp deletion is globally distributed and probably arose early in human history.

But several questions still remain: What is SMIM1’s functional role in the RBC membrane? Is SMIM1 the Vel antigen epitope, a carrier molecule or a scaffold protein for a membrane complex? Which other factors bind to the two sequence variants and what is the mechanism of regulation? Is SMIM1 the missing chymotrypsin-sensitive P. falciparum receptor? And why did the outlying samples type as Vel–?

Future research should focus on characterising the function of SMIM1. It is likely that SMIM1 interacts with proteins in the RBC membrane and determining these binding partners, perhaps by mass spectrometry, will give valuable insight and likely a major hint of SMIM1’s function.

Determining the factors binding to the two sequence variants rs143702418 and rs1175550 is also an important task. Their eQTL effects are substantial and it is very possible that the factors would not only shed light on SMIM1 expression, but on erythroid transcriptional regulation in general.

This prospect is also present for the study of the outlying Vel– samples. Since we found nothing in the SMIM1 gene to explain the Vel– phenotype, it will be interesting see if our whole exome sequencing data will uncover potential trans-acting eTFs.

This thesis assisted in characterising the molecular background of the Vel antigen, which is a clinically significant antigen, that can cause severe haemolytic transfusion reactions. This enabled the development of molecular Vel blood group
typing in the clinic as well as the prospect of creating a synthetic monoclonal anti-Vel antibody to be used in diagnostics, blood group typing and research. Furthermore this PhD found TAL1 to be potentially involved in the expression of SMIM1 and the Vel antigen. Lastly, this thesis demonstrated how pre-existing population genetics can be exploited to deconvolve potentially causal sequence variants locked in tight LD, a powerful approach to solve the inherent limitations of GWAS and eQTL studies.

As always in scientific endeavours, solving one long-standing mystery only lead us to dozens more intriguing new questions, puzzles and conundrums, which, in essence, is what science is all about – the most exciting thing is not the destination, but the road you take to get there.
Populærvideneskabelig sammenfatning

Denne Ph.d. afhandling beskriver resultater opnået gennem fire års laboratoriearbejde på Afdelingen for Hæmatologi og Transfusionsmedicin, Medicinsk Fakultet på Lunds Universitet. Den består af tre studier med fokus på den sjældne blodtype Vel. To af studierne er publiceret i de videnskabelige tidsskrifter *Nature Genetics* og *Scientific Reports*.

Mens de fleste kender til ABO og Rh blodtypesystemerne, er det nok de færreste, der ved, at der faktisk findes yderligere 34 blodtypesystemer. ABO-systemet blev beskrevet af den østrigske biolog og læge Karl Landsteiner i starten af 1900-tallet, og med det blev grundstenene til den moderne transfusionsmedicin lagt. Paradigmet, som er gældende for de fleste blodtypesystemer, er kort fortalt, at alle mennesker har visse strukturer, antigener, på overfladen af de røde blodlegemer, som vores immunsystem genkender. Men for nogen individer kan disse strukturer have et lidt andet udseende, eller helt være fraværende. Hvis blodlegemer fra én person bliver blandet med blod fra én, som har et andet antigen, så kan immunsystemet ikke genkende strukturen på de indkommende blodlegemer og vil reagere som om, det var en fremmed mikroorganisme og danne antistoffer imod det fremmede antigen. Dette kan i værste tilfælde forårsage kraftige og dødelige hæmolytiske transfusionsreaktioner.

Blodtypen Vel er ét af de mere sjældne systemer. Hvis man er Vel-positiv, sidder Vel antigenet på overfladen af de røde blodlegemer, hvis ikke, er man Vel-negativ, og kroppens immunforsvar danner antistoffer imod Vel antigenet – disse kaldes anti-Vel. På verdensplan siger man, at 1 ud af 4.000 er Vel-negative, alle andre er Vel-positive. Der er evidens for en højere frekvens af Vel-negative personer i Norden, hvor vi i denne afhandling bl.a. fandt 1 Vel-negativ person ud af 1.200 i Region Skåne.

Vel antigenet blev første gang beskrevet hos en 66-årig kvinde, Mrs. Vel, i 1952 i New York, der fik en hæmolytisk transfusionsreaktion efter en blodtransfusion. I de efterfølgende årtier blev Vel antigenet fundet hos patienter over hele verden; i Sverige, Australien, Thailand, blandt indianere i Canada og afroamerikanere i USA, men den molekylære årsag, det vil sige det gen eller den mutation, der er skyld i at nogen er Vel-negative, forblev i det uvisse.
Da jeg startede på min Ph.d., var forskningsgruppen allerede godt på vej til at have løst det 60 år gamle mysterium om Vels molekylære baggrund. Man havde bl.a. sammenlignet mutationer i hele genomet hos 20 Vel-negative personer med deres Vel-positive familiemedlemmer som kontrolpersoner og derved fundet frem til et område i genomet, hvor der var en ophobning af mutationer kun fundet hos de Vel-negative. I dette område var der fem potentielle gener, som kunne danne baggrund for Vel. Vi undersøgte nu disse gener for diverse typiske karakteristika for gener udtrykt i blodceller og fandt frem til, at genet Small Integral Membrane Protein 1 (SMIM1) var den mest oplagte kandidat af de fem. Ved computeraffald blev SMIM1’s nærmeste naboer påvist at være andre gener, der koder for proteine i røde blodlegemer, og ved at sekventerne alle SMIM1’s DNA sekvens fandt vi en mutation, som var tilstede hos alle de Vel-negative og ingen af de Vel-positive. Denne mutation er en deletion af 17 basepar (byggesten) i DNA’et, som bevirket, at SMIM1 proteinet ikke kommer til udtryk i røde blodlegemer hos Vel-negative personer.

Vi gik derfra videre for at forsøge at bevise sammenhængen mellem SMIM1 og Vel antigenet. Vi undersøgte tilstedevarsel af mRNA (et mellemstadiimellem DNA og protein) for SMIM1 og fandt det kun hos Vel-positive individer. Vi fandt ligeledes kun Vel antigenet samt SMIM1 proteinet udtrykt på overfladen af røde blodceller fra Vel-positive personer. Derudover kiggede vi på sammenhængen mellem personer, som havde nul, én eller to kopier af SMIM1 mutationen (på de såkaldte alleller, som vi har to af for alle gener i genomet), og hvor kraftigt anti-Vel antistoffet reagerede med deres blodceller. Her fandt vi, at dem som ikke havde SMIM1 deletionen på nogen allel reagerede kraftigt med anti-Vel; dem, der havde deletionen på den ene allel, reagerede middel, og dem, der havde deletionen på begge alleller, reagerede svagt – altså et tegn på at SMIM1 og Vel antigenet har noget med hinanden at gøre. Sidst, men ikke mindst reagerede anti-Vel antistoffet kraftigt mod et kunstigt fremstillet SMIM1 protein, mens det nærmest ikke reagerede med det samme protein, som man indeholdt deletionen. Med andre ord, når SMIM1 genet blev udtrykt, opførte anti-Vel sig, som det ville, hvis det blev blandet med blod fra Vel-positive, mens når SMIM1-genet blev muteret, opførte anti-Vel sig, som det ville gøre med blod fra Vel-negative.

Disse forsøg fremførte kraftige argumenter for, at SMIM1 er baggrund for Vel blodtype antigenet, og at mutationen i SMIM1 bevirker, at man ikke udtrykker Vel antigenet på sine røde blodlegemer, og dermed at man siges at være Vel-negative.

Samtidig med os havde to andre forskningsgrupper også fundet frem til mutationen i SMIM1 og påvist sammenhængen mellem SMIM1 og Vel, og dermed kunne Vel officielt godkendes som et blodtypesystem af International Society of Blood Transfusion i 2013. Projekt I i denne afhandling omhandler de omtalte forsøg.
Med Vel blodtypesystemet på plads kiggede vi herefter på et velkendt fænomen i blodlaboratorier: At selv om man er Vel-positiv, så er der meget stor variation i, hvor kraftigt ens røde blodlegemer reagerer med anti-Vel antistoffet. Det kan komme så vidt, at en svagt-reagerende Vel-positiv person fejlagtet bliver bestemt til at være Vel-negativ, hvilket kan få store konsekvenser ved en blodtransfusion. Vi ville derfor undersøge, hvad der kunne være årsagen til denne store variation og mistænkte en region i SMIM1 genet, hvor der foregår meget såkaldt genregulering. I denne region fandt vi en anden mutation, en såkaldt enkeltnukleotidpolymorfi, forkortet SNP fra det engelske single nucleotide polymorphism, hvor DNA-nukleotidet A (et nukleotid er et af DNA’s byggesten – to nukleotider danner sammen et basepar) er skiftet ud med et G i en vis procentdel af befolkningen. Denne SNP, kaldet rs11755550, var tidligere vist at have indflydelse på Vel udtryk, og vi satte os for at undersøge den nærmere. I blodprøver fra 150 svensere fandt vi, at de individer, der havde G-nukleotidet i deres DNA-sekvens, viste klart højest reaktivitet med anti-Vel, dvs. de udtrykker meget Vel antigen på celleoverfladen. Og vice versa, personer med A-nukleotidet havde lav forekomst af Vel antigen og lav anti-Vel reaktivitet. Vi undersøgte nu DNA regionen omkring rs1175550 og fandt flere bindingssteder, hvor såkaldte transkriptionsfaktorer kan binde. Det er proteiner, som regulerer hvor meget af et givent protein, der bliver udtrykt i cellen. Gennem flere forsøg med binding af transkriptionsfaktorer til sekvensen omkring rs1175550 fandt vi, at særligt én faktor, kaldet TAL1, så ud til at binde specifikt til DNA-regionen, kun når G-nukleotidet og ikke A var til stede. Eftersom TAL1 oftest aktiverer de gener den binder til og derved fremmer produktion af det resulterende protein, så er det vores hypotese, at TAL1 er en af grundene til, at personer med G-nukleotidet i deres DNA-sekvens har højere ekspression af SMIM1 og dermed Vel antigen end personer med A-nukleotidet. Disse forsøg udgør Projekt II.

Det sidste studie om Vel blodtypen i denne afhandling omhandlede bl.a. en tidligere metode at klassificere Vel på: I 1968 blev det påvist, at der fandtes to Vel antigener, Vel 1 og Vel 2, og man klassificerede personer, der var svagt reaktive med anti-Vel antistof som Vel;1,–2 (alså antog man at personen havde Vel 1, men manglede Vel 2), mens personer med meget svag til ingen reaktion med anti-Vel var Vel;1,–2. Med den molekylære baggrund for Vel antigenet kendt, var vores mål for projekt III at klassificere disse gamle prøver ved hjælp af den nye viden. Til det formål fandt vi 40 gamle prøver fra Vel-negative personer i New York Blood Center’s blodbank, som alle var blevet Vel blodtypebestemt med det gamle to-antigen system. Vi undersøgte DNA fra prøverne for 17-bp deletionen i SMIM1, hvor tilstedeværelse af deletionen som bekendt er ensbetydende med lavere eller helt fraværende Vel antigen udtryk. Vi fandt frem til, at samtlige prøver, som oprindeligt var blevet bestemt til at være Vel;1,–2, havde én kopi af deletionen i deres DNA. Og for Vel;–1,–2 prøverne, som var vist at have lav til
ingen Vel antigen ekspression, fandtes deletionen på begge alleller, og altså kunne SMIM1 proteinet ikke udtrykkes hos disse personer. Vi fandt med andre ord komplet overensstemmelse imellem mængden af udtrykt Vel antigen og tilstedeværelsen af 17-bp deletionen i SMIM1 på én eller begge alleller. Herudover fandt vi tre prøver, som oprindeligt blev klassificeret som Vel negative, men som ikke indeholdt 17-bp deletionen på nogen af allellerne. Dette er højest usædvanligt, og noget der kun er rapporteret få gange tidligere, og vi er derfor nu i gang med at analysere disse prøver nærmere for at finde en eller flere yderligere mutationer, som kan forklare det lave Vel antigen udtryk.

Vel antigenet var én af de få resterende klinisk vigtige antigener på grund af dens potentielt kraftige transfusionsreaktioner, og opdagelsen af den molekylære baggrund har åbnet op for hurtigt at kunne blodtypebestemme ved hjælp af standard laboratorieteknikker. Anti-Vel antistof egnet til klinisk forskning har gennem historien været svært at få adgang til, men med den molekylære baggrund kendt åbnes nu op for at kunne fremstille kunstigt anti-Vel antistof, som kan bruges til diagnosticering, blodtypebestemmelse og forskning.
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