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BOSTON UNIVERSITY

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Dissertation

CHARACTERIZATION OF A TRANSCRIPT FOUND WITHIN THE *HBS1L-MYB* INTERGENIC REGION AND ITS ROLE IN HEMOGLOBIN REGULATION IN ERYTHROID CELLS

by

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Submitted in partial fulfillment of the

requirements for the degree of

Doctor of Philosophy

2017

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First Reader

David H. K. Chui, M.D. Professor of Medicine

Second Reader

Martin H. Steinberg, M.D. Professor of Medicine "I want to make sure we use all our talent, not just 25 percent. Don't let anyone rob you of your imagination, your creativity, or your curiosity. It's your place in the world; it's your life. Go on and do all you can with it, and make it the life you want to live." —Mae C. Jemison (first African American woman in space)

DEDICATION

This doctoral work is dedicated to my nieces and nephews. From oldest to youngest: Eddie, Pearl, Shafiq, Marcel, Johnny III, Terrance Jr, Malachi, Makaia, Troy, Camille, Christian, Jahseem, Zoe, Myah, Malia and Tyler. Because of you, I strive to excel in order to pave the way for your future accomplishments. Know that there is nothing in this world you can't do. So dream big, stay focused and disciplined, don't hold on to past mistakes, love and be loved, don't be scared of change, don't be scared to be different, and NEVER EVER give up. Life can be interesting sometimes, and you may find it difficult to climb up the ladder of success. But keep climbing...keep moving...stay positive...follow your dreams, not your fears...all things will work together for the good. And know that Aunt Tasha (aka Titi) has your back and loves you so very much.

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CHARACTERIZATION OF A TRANSCRIPT FOUND WITHIN THE *HBS1L-MYB* INTERGENIC REGION AND ITS ROLE IN HEMOGLOBIN REGULATION IN ERYTHROID CELLS

TASHA ALEASE MORRISON

Boston University School of Medicine, 2017

Major Professor: David H. K. Chui, M.D., Professor of Medicine

ABSTRACT

Sickle cell disease (SCD) is one of the most common hemoglobinopathies worldwide. It is caused by a homozygous mutation in codon 6 of the beta globin gene (*HBB*), which leads to polymerization of the variant hemoglobin and sickled red blood cells that obstruct blood vessels and reduce oxygen delivery to tissues. Patients with SCD have multiple clinical problems, including pain crises, anemia and organ damage. However, not all patients with SCD display all these clinical manifestations. One major factor for reduced occurrences of symptoms is fetal hemoglobin (HbF). HbF is the main hemoglobin in the fetus, and declines one year after birth to less than one percent of total hemoglobin. Nevertheless, there are individuals who continue to have high levels of HbF into adulthood, which is beneficial for an individual with SCD because HbF reduces the amount of sickle polymer in red blood cells. There are three major quantitative trait loci (QTL) associated with high HbF. However, these QTL account for 20-45% of HbF variance. Therefore, further investigation is required to fully understand how HbF is regulated.

The *HBS1L-MYB* intergenic polymorphism (HMIP) on chromosome 6q23 is one of the major QTL associated with high HbF. This region is also known to regulate other erythroid-specific traits due to an enhancer element that promotes the expression of the downstream gene, *MYB*, which controls hemoglobin expression and erythroid proliferation and maturation. The presence of RNA polymerase II binding and a 50-bp transcript suggested that a long noncoding RNA (lncRNA) is transcribed from this region. LncRNAs are non-protein-coding transcripts greater than 200 nucleotides and are involved in gene regulation. Therefore, it was hypothesized that a lncRNA is transcribed from the enhancer of *MYB* and regulates hemoglobin expression.

I characterized a novel lncRNA, 1283 bp in length that was differentially expressed among various tissue types, among erythroid progenitor cells with different hemoglobin makeup, and also during erythroid differentiation. Furthermore, knockdown of this lncRNA, named the *HBS1L-MYB* intergenic long noncoding RNA (*HMI-LNCRNA*), significantly increased HbF. Taken together, these observations suggest that *HMI-LNCRNA* can be a possible therapeutic target to increase HbF expression in patients with SCD and β -thalassemia.

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LIST OF ABBREVIATIONS

3'	3-1	prime
		L
5'	5-]	prime

adenine	A
alpha	α
3 ₂ adult hemoglobin A	$\alpha_2\beta_2$
$S_2^{S_2}$ sickle hemoglobin	$\alpha_2\beta^S{}_2$
³ γsickle-gamma hemoglobin	$\alpha_2\beta^S\gamma$
<i>j</i> ₂ adult hemoglobin A ₂	$\alpha_2\delta_2$
2Hemoglobin Gower II	$\alpha_2 \epsilon_2 \dots$
¹ 2 fetal hemoglobin	$\alpha_2 \gamma_2$
AAfrican American	AA
3Maorta-gonad-mesonephros	AGM
cRNAantisense lncRNA	alncR
Oantisense oligonucleotide	ASO

β	beta
β^{S}	sickle beta globin
Baso	basophilic erythroblasts
Bcl11a	B-cell lymphoma/leukemia
bp	base pair

BFU-E	burst-forming unit erythroid
BU	Boston University

C	cytosine
cDNA	complementary DNA
CD	cluster of differentiation
CD235a	glycophorin A
CD71	transferrin receptor
CFU-E	colony-forming unit erythroid
ChIP	chromatin immunoprecipitation
ChIRP-seq	chromatin isolation by RNA purification sequencing
chr	chromosome
CLP	common lymphoid progenitor
СМР	common myeloid progenitor
cPPT	central polypurine tract
CRISPR	clustered regularly interspaced short palindromic repeats
CSSCD	
Ct	

δ	delta
Δ	delta
$(\delta\beta)^0$ -thalassemia	delta-beta zero thalassemia

DMEM	Dulbecco's Modified Eagle's Medium
DNA	deoxyribonucleic acid
dsRNA	double-stranded RNA

epsilon
enhanced chemiluminescence
ethylenediaminetetraacetic acid
erythro-myeloid progenitor
Encyclopedia of DNA Elements
erythropoietin
erythropoietin receptor
enhancer RNA
primitive erythroid progenitor cell

FACS	fluorescence-activated cell sorting	
FDA	Food and Drug Administration	

γ	gamma
g	gravitational
G	guanine
GAG	group-specific antigen
GFP	green fluorescent protein

GTP	guanosine triphosphate
GWAS	genome-wide association study

H ₂ O	water
H3K4Me1	histone 3 lysine 4 monomethylation
HBA	alpha globin protein
HBA1	alpha globin gene
HbA ₁	adult hemoglobin A
HbA ₂	adult hemoglobin A ₂
HBB	beta globin protein
HBB	beta globin gene
HBD	delta globin gene
HBE	epsilon globin gene
HbF	fetal hemoglobin
HBG	gamma globin protein
HBG	gamma globin gene
HbS	sickle hemoglobin
HBZ	zeta globin gene
HIV	human immunodeficiency virus
HMG	high-mobility group
HMIP	
HMI-LNCRNA	HBS1L-MYB intergenic long noncoding RNA

	HMIT
hereditary persistence of fetal hemoglobir	HPFH
hypersensitive site	HS
hematopoietic stem cel	HSC
	HUDEP

IL-6	interleukin 6
ilncRNA	intronic overlapping lncRNA
IMDM	Iscove's modified Dulbecco's medium
iPSC	induced pluripotent stem cell

kb	kilobase

LB	Luria broth
LCR	locus control region
LD	linkage disequilibrium
LDS	lithium dodecyl sulfate
lincRNA	intergenic lncRNA
LNA	locked nucleic acid
IncRNA	long noncoding RNA
LTR	long terminal repeat

molar	M
murine embryonic stem cells	mESC
minute	min
Mistral	MIRA
milliliter	ml
mixed lineage leukemia 1	MLL1
millimolar	mM
multiplicity of infection	MOI
messenger RNA	mRNA

n	sample size
NaCl	sodium chloride
ncRNA	noncoding RNA
ng	nanogram
NGS	next generation sequencing
NHLBI	National Heart, Lung, and Blood Institute
NIH	National Institutes of Health

OrthoE	orthochromatic	erythroblasts
--------	----------------	---------------

PBMC	
	1 1
PBS	phosphate-buffered saline

PCR	polymerase chain reaction
plncRNA	pseudogene lncRNA
Pol II	
poly(A)	polyadenylation
PolyE	polychromatophilic erythroblasts
ProE	proerythroblasts
PVDF	polyvinylidene difluoride
qPCR	quantitative polymerase chain reaction
QTL	quantitative trait loci
RACE	rapid amplification of cDNA ends
RBC	red blood cell
RISC	RNA-induced silencing complex
RLM	RNA ligase mediated
RNA	ribonucleic acid
RNA ChIP	RNA-chromatin immunoprecipitation
RNA-Seq	RNA sequencing
RPM	revolutions per minute
RPMI	Roswell Park Memorial Institute
rRNA	ribosomal RNA
rs	reference SNP ID

RSV	
RT-PCR	reverse transcription polymerase chain reaction

S	sample
SCD	sickle cell disease
SCF	stem cell factor
sec	second
SFEM	serum-free expansion medium
shlncRNA	small RNA host lncRNA
shRNA	short hairpin RNA
siRNA	small interference RNA
SNP	single nucleotide polymorphism
sRNA	small RNA
SV40	simian virus 40

Τ	thymine
T87Q threonine to g	lutamine substitution at amino acid 87
TALENtransc	ription-activator like effector nuclease
TAP	tobacco acid pyrophosphatase
TOI	transcript-of-interest
Tris-HCL	tris hydrochloride
tRNA	transfer RNA

TSStra	anscription	start site
--------	-------------	------------

U	units
μg	microgram
μl	microliter
μΜ	micromolar
US	United States
USA	United States of America

ζ	zeta
ζ ₂ ε ₂	Hemoglobin Gower I
ZFN	zinc finger nuclease

<	less than
>	greater than
=	equal
%	percent
+	positive
°C	degree Celsius

CHAPTER 1. INTRODUCTION

1.1 Red Blood Cells

Cellular respiration—the set of metabolic processes that convert the nutrients we consume into energy—is required to sustain all the biological functions of the human body. However, energy can only be provided if oxygen is available. The presence of oxygen is controlled by the circulatory system, in which the heart pumps blood throughout the body. Three types of cells are found in the blood—leukocytes, thrombocytes and erythrocytes. Erythrocytes, also known as red blood cells (RBCs), are red, round, biconcave cells, are the most common cell type in the blood, and carry oxygen to tissues throughout the body. The process by which this occurs begins in the lungs, where the oxygen that is inhaled through the lungs is absorbed by RBCs. Once in the RBC, oxygen binds to the heme group that is attached to globin.¹ Next, RBCs circulate through the vascular system, releasing oxygen to all vital organs. Since RBCs are so important to human life, it is essential to know and understand how they are developed—a process called erythropoiesis—and the main protein content responsible for their function—hemoglobin.

1.1.1 Erythropoiesis

The ontogeny of erythropoiesis changes while the human embryo is developing in the womb. Collectively, there are two types of erythropoiesis—primitive and definitive—and three sites of origin for RBCs—the yolk sac, liver and bone marrow.^{2–4} For both primitive and definitive erythropoiesis, the basic order of these processes is to expand or proliferate erythroid precursors from progenitor cells, promote expression of hemoglobin, decrease cell size, enucleate and function as a red blood cell in circulation. However, the type of progenitors these cells are derived from and the location of erythroid differentiation differs between these two processes (**Figure 1.1A**).

Primitive erythropoiesis begins 15 days after fertilization within the yolk sac.^{2,3} Hemangioblast progenitor cells derived from the mesoderm differentiate in the yolk sac into primitive erythroid progenitor cells (EryP-CFC), which form into clusters of cells called "blood islands."⁵ Next, these erythroid progenitor cells are released into blood vessels and differentiate further as four different erythroblast precursors that are named based on the morphology of the cells. In the order of maturation, they are proerythroblasts basophilic erythroblasts (BasoE), (ProE), polychromatophilic erythroblasts (PolyE) and orthochromatic erythroblasts (OrthoE), during which the erythroblasts' nuclei condense, the cell size decreases and hemoglobin production increases (Figure 1.1B).^{2,6} As OrthoE, the erythroblasts enucleate and become reticulocytes, and then achieve terminal maturation into RBCs indicated by the biconcave disc shape of the cell membrane.^{7,8}

Definitive erythropoiesis occurs in both the liver and bone marrow.² However, the source of progenitor cells is different in both locations. In the liver, erythroblast precursors called burst-forming units erythroid (BFU-E) are originally from the yolk sac.^{9,10} In the yolk sac, at around 4 to 5 weeks of gestation, erythro-myeloid progenitors (EMPs), which are multipotent hematopoietic progenitor cells, differentiate into BFU-E. Once expansion of BFU-E occurs, they are released into circulation and migrate to the

liver. Concurrently, hematopoietic stem cells (HSCs) begin to emerge from the aortagonad-mesonephros (AGM) region, and also travel to and expand in the fetal liver. At 10 to 11 weeks of gestation, HSCs begin to travel from the liver to the bone marrow, where they will permanently stay. This transition continues until after birth, when the bone marrow becomes the only source of red blood cells for the duration of life. In both the fetal liver and bone marrow, HSCs give rise to BFU-E, which will give rise to another erythroid precursor called colony-forming unit erythroid (CFU-E). Both are named based on their morphology when they are cultured *in vitro* in semisolid medium. Next. erythroid maturation occurs in the same order as primitive erythropoiesis. However, maturation does not occur in the blood vessel. It occurs in the fetal liver and bone marrow within erythroblastic islands, which are made up of erythroblasts that are physically attached to macrophages.¹¹ Once the OrthoE enucleate, reticulocytes are formed and released into the bloodstream where they terminally maturate into RBCs. The lifespan of a RBC is 120 days, and thereafter senesce and are removed by macrophages located in the spleen.²



Figure 1.1. Erythropoiesis.

(A) Illustration of the direction of erythropoiesis from progenitor cells in the fetal liver and bone marrow (definitive), and progenitor cells in the yolk sac (primitive). During definitive erythropoiesis, erythroid differentiation and maturation occurs in the fetal liver or bone marrow until enucleation occurs, and the reticulocyte is released into the blood to undergo terminal maturation into a red blood cell. During primitive erythropoiesis, progenitor cells are released from the yolk sac and into the blood where they undergo erythroid differentiation from erythroblasts to red blood cells. Image adapted from Palis, Frontiers in Physiology 2014. (B) Morphology of human erythroblasts as proeythroblasts (ProE), basophilic erythroblasts (BasoE), polychromatophilic erythroblasts (PolyE) and orthochromatic erythroblasts (OrthoE). Image adapted from Palis, Blood 2014.

The most important driver of erythroid maturation is erythropoietin (EPO).^{2,9,12} EPO is a cytokine that is released from the kidneys when RBC count is low, and binds to the erythropoietin receptors (EPOR) of CFU-E to promote differentiation. During erythroid maturation, two markers of maturation are expressed—transferrin receptor (CD71) and glycophorin A (CD235a).¹⁰ CD71 is a marker for early erythroid maturation, and it is found in the plasma membrane of erythroblasts to promote uptake of iron within the cell.¹³ This iron will help create the heme group that will be attached to globin to form hemoglobin that can reversibly bind oxygen. As the erythroblasts mature they also begin to express CD235a, which is another protein found in the plasma membrane of erythroblasts, as well as mature RBCs. CD235a contains sialic acid, which makes the RBCs hydrophobic. This allows the RBCs to flow through the blood without adhering to blood vessels and other cells. Once erythroid cells become fully mature RBCs, they lack expression of CD71 and only express CD235a.

1.1.2 Hemoglobin switching

Hemoglobin is a protein expressed by RBCs to bind to the oxygen that will be delivered to tissues in the entire body. It is made up of four subunits of globin chains, with each subunit bound to a heme group, which is necessary for oxygen transportation. From the development of the first RBCs derived from the yolk sac to the RBCs that are ultimately made from the bone marrow, the type of hemoglobin expressed changes, and this is known as "hemoglobin switching."^{14–16} Hemoglobin switching can also occur as the erythroblasts mature into RBCs, and is called "maturational switching."¹⁷ This is due to the regulation of gene expression on two loci—the alpha (α) globin gene cluster on

chromosome (chr) 16p13.3, and the beta (β) globin gene cluster on chr11p15. The functional genes of the α -globin gene cluster are made up of zeta (ζ) globin (*HBZ*) and two α -globin genes (*HBA2*, *HBA1*)¹⁸; and the β -globin gene cluster contains epsilon (ϵ) globin (*HBE*), two gamma (γ) globin (*HBG2*, *HBG1*), delta (δ) globin (*HBD*), and β globin (HBB) genes.¹⁹ On both loci, the direction of globin gene expression moves from the 5' end to the 3' end. Hemoglobin is composed of two subunits from the α -globin gene cluster and two subunits from the β -globin gene cluster. During primitive erythropoiesis, erythroblasts derived from the yolk sac undergo a maturational switch that occurs between five to seven weeks of gestation.¹⁷ The switch is only made within the α globin gene cluster, from hemoglobin Gower I ($\zeta_{2}\varepsilon_{2}$) to hemoglobin Gower II ($\alpha_{2}\varepsilon_{2}$). At around three months of gestation, when the fetal liver begins to take over RBC production, the switch is now within the β -globin gene cluster, and the cells make fetal hemoglobin (HbF) ($\alpha_2\gamma_2$). HbF expression continues when the location of definitive erythropoiesis changes to the bone marrow. However, HbF does not remain the most abundant hemoglobin, and another switch occurs within the β -globin locus. After birth, HbF declines while adult hemoglobin (HbA₁) ($\alpha_2\beta_2$) increases as the main type of hemoglobin expressed in RBCs. Another type of adult hemoglobin (HbA₂) ($\alpha_2\delta_2$) is also made, but in small quantities. Eventually, the hemoglobin makeup in an adult becomes less than 1% HbF, 2.5% HbA₂ and 95.5% HbA₁.



Figure 1.2. Hemoglobin Switching.

(A) Globin chains are transcribed from both the alpha globin gene cluster on chromosome 16 and the beta globin gene cluster on chromosome 11. (B) Genes on both loci are differentially expressed during gestation, and translates into individual globin subunits that form into a tetramer to make embryonic, fetal or adult hemoglobin (C). Chart in (B) is adapted from Forget, *N Engl J Med 2011*.
1.1.3 Transcription factors that regulate erythropoiesis and hemoglobin switching

Primitive and definitive erythropoiesis, and hemoglobin switching are regulated by the expression pattern of key transcription factors. *GATA1*, *KLF1*, *TAL1*, *LMO2* and *LDB1* are all expressed during primitive and definitive erythropoiesis.^{2,4,20,21} They not only regulate hemoglobin expression but also erythroid maturation. GATA1 promotes erythropoiesis by regulating the expression of erythroid-specific genes. It does so by working in complex with TAL1, LMO2 and LDB1. Loss of GATA1 inhibits the maturation of erythroblasts.²² KLF1 is another regulator of erythropoiesis as well as hemoglobin production.²³ It controls the expression of other erythroid-specific genes to promote expression of both embryonic and adult hemoglobin, stabilizes α -globin, ensures the biosynthesis of heme, and regulates other transcription factors.

The differences between primitive and definitive erythropoiesis are the expression of additional transcriptions factors only during the definitive stage (**Table 1.1**). These transcription factors work mostly to regulate expression of genes on the β -globin gene locus via the locus control region (LCR) that is located upstream this region. The LCR is a super enhancer with five DNase I-hypersensitive sites (HSs), which interacts with promoters of the downstream globin genes to promote their expression.²⁴ However, which promoter it will interact with depends on the expression of key transcription factors. c-MYB is one of these transcription factors that have been shown to promote erythropoiesis and work in complex with other transcription factors on the β -globin locus to reduce expression of γ -globin genes *HBG1* and *HBG2*.^{25–29} The next transcription factor, SOX6, is a Sry-related high-mobility group (HMG) box transcription factor, and has been shown to directly bind to the promoter of *HBE* to repress its expression.³⁰ The last transcription factor, BCL11a, contains zinc fingers that bind directly to the LCR and multiple sites on the β -globin gene cluster to specifically reduce expression of *HBG* and increase expression of the *HBB*.^{31–34} It has been shown to do so by working with other transcription factors, including SOX6.³⁵

Table 1.1. Globin genes and transcriptions factors that are expressed duringprimitive and definitive erythropoiesisTable adapted from Palis, *Blood 2014*.

	Primitive Erythropoiesis	Definitive Erythropoiesis
Globin Genes	<i>HBZ</i> and <i>HBA1</i> <i>HBE</i> and <i>HBG</i>	HBA1 HBB
Transcription Factors	GATA1, KLF1, TAL1, LMO2 and LDB1	GATA1, KLF1, TAL1, LMO2, LDB1, c-MYB, SOX6, and BCL11a

Taken together, RBCs are vital to human life, and require regulated processes at both the biological and molecular levels to ensure that healthy RBCs are developed and have the ability to effectively transport oxygen throughout the body.

1.2 Sickle Cell Disease

Genetic variants or mutations within both the α -globin and the β -globin loci, and transcription factors that regulate erythroid-specific traits can threaten erythropoiesis, hemoglobin expression and/or the viability of RBCs, and cause various blood disorders

or hemoglobinopathies.^{18,36,37} One of these blood disorders is called sickle cell disease (SCD).

SCD was first described in 1910 by a cardiologist named James B. Herrick who saw that the appearance of RBCs from his patient was abnormal. However, Herrick could not decipher whether the patient had a primary blood disorder or whether the peculiar shaped cells occurred because of another clinical disorder.^{38–40} It was not until 1945 that Linus Pauling hypothesized that sickled RBCs were due to defects within the hemoglobin, and in 1949 Pauling and others showed by gel electrophoresis that hemoglobin from sickle cells migrate differently than hemoglobin from normal cells, thus proving that his hypothesis was indeed correct.^{40–42} Later that year, it was established that SCD can be inherited and as an autosomal recessive trait. However, the exact mutation of SCD was still not known. Ten years later, Vernon Ingram established that SCD is due to a mutation of codon 6 of *HBB*, which causes a change in an amino acid of adult hemoglobin (HbA₁) (glutamic acid to valine).⁴³

Since then, with more understanding of erythropoiesis and hemoglobin switching, scientists were able to establish the pathophysiology of SCD. One aspect of SCD that was not quite understood during the earlier years of study is the sickled shape. The glutamic acid-to-valine change turns HbA₁ to sickle hemoglobin (HbS), where the two β -chains found in HbA₁ ($\alpha_2\beta_2$) becomes sickle β -globin (β^S).^{44,45} This modifies the properties of hemoglobin produced in RBCs because valine is more hydrophobic than glutamic acid. When an RBC releases its oxygen, it is deoxygenated. In a RBC with normal hemoglobin, deoxygenation does not modify the structure of the hemoglobin

molecule. However, in a RBC with HbS, deoxygenation causes the more hydrophobic hemoglobin to polymerize. This leads to disruption of the cytoskeleton and protrusion of HbS polymer through the cell membrane, and is the reason sickle RBCs display many abnormal morphological characteristics. Damage to the cell membrane alternates the transmembrane proteins and the lipid bilayer, consequently changing many properties of the RBC. Normal RBCs can flow freely in circulation, without attaching to blood vessels and other cells. However, sickled RBCs undergo hemolysis, and attach to both the endothelium of blood vessels and other cells, and this promotes vascular occlusion (Figure 1.3).^{46–48} Hemolysis and vaso-occlusion of sickle RBCs leads to acute pain crises, anemia and organ damage due to tissue oxygen deprivation. Other symptoms of SCD are swelling of the hands and feet, bacterial infection and acute chest syndrome. SCD is associated with premature death. Another aspect of SCD is the timeline when symptoms occur. Babies born with SCD do not show clinical evidence of the disease until sometime after the first three to six months of life. This is due to hemoglobin switching. After birth, the main type of hemoglobin a baby makes is HbF. However, during that first year of life, HbF significantly reduces while HbA₁ becomes the main type of hemoglobin expressed in RBCs. In an infant homozygous for the mutation of codon 6 in HBB, it is HbS, not HbA₁ that is mainly expressed. Therefore, clinical symptoms related to SCD are delayed while HbF is still high, and do not start until HbF is replaced by HbS.



Figure 1.3. Sickle Cell Disease.

(A) Peripheral blood smear from a patient with sickle cell disease, stained with May-Grunwald-Giemsa. Red blood cells displaying a sickled shape (S) are distinct from normal round red blood cells. Image adapted from Rees et al, *The Lancet 2010*. (B) Illustration of how red blood cells with normal hemoglobin can flow freely through blood vessels, compared to red blood cells with sickle hemoglobin, which cause vaso-occlusion. Image adapted from NIH/NHLBI website (https://www.nhlbi.nih.gov/health/health-topics/topics/sca).

1.2.1 Epidemiology of SCD

SCD has become one of the most common inherited hemoglobinopathies worldwide.^{44,45,49} Originally, SCD was mostly found in populations of people located in regions where malaria was endemic.⁵⁰ When individuals heterozygous for the sickle mutation—also known as having sickle cell trait—are infected with malaria, they have increased survival and are able to reproduce. Public health initiatives to decrease the incidence rate of malaria and other infectious diseases, and to promote better clinical practices within these regions have also increased the lifespan of people with SCD. Furthermore, the rise in human migration from these regions has now made SCD a global health problem. SCD is a prime example of a genetic balanced polymorphism; increased fitness (survival, reproduction) of the heterozygote is accompanied by decreased fitness of the homozygote.

SCD is mostly found in people of African, Arabian and Indian descent. Globally, about 300,000 newborns are born with SCD each year, and it has been projected that it will rise to about 400,000 newborns by 2050.⁴⁹ The highest incidence rate of SCD is found in West Africa, with up to 90,000 babies born with the disease per year in Nigeria alone (the highest rate worldwide).⁵¹ In the United States of America (USA), approximately 100,000 people have SCD; one out of 365 African American (AA) babies are born with SCD; and one out of 13 AA babies are born with the sickle cell trait.⁵²

1.2.2 HbF and SCD

The delayed appearance of sickle RBCs and the clinical manifestations of disease that hematologists observed throughout the first year after birth of infants with SCD led them to consider HbF as a key factor for this delay. Subsequently, laboratory studies discovered that HbF inhibits the polymerization of HbS by obstructing the formation of HbS polymer in RBCs.⁵² The tetramer that makes up HbS is $\alpha_2\beta_2^{S}$. However, when γ -globin is also expressed, the HbF molecule $\alpha_2\gamma_2$ is excluded from the HbS polymer, and the mixed hybrid tetramer $\alpha_2\beta_{\gamma}^{S}\gamma$, has reduced incorporation into the polymer. Together, this reduces the polymerization potential of HbS leading to more normal RBCs.

SCD patients display a wide range of disease severity. Also different among SCD patients is the percent of HbF—some with nearly normal HbF, and others with higher levels of HbF. Epidemiological studies, which looked at the natural history of SCD, found that patients with SCD and high HbF have lower disease burden than SCD patients with lower HbF. One such study is the Cooperative Study of Sickle Cell Disease (CSSCD), which prospectively studied the natural history of about 4,000 patients in 23 health facilities around the US.^{53–55} Funded by the National Heart, Lung, and Blood Institute (NHLBI) at the National Institutes of Health (NIH), the CSSCD found that various complications of SCD, including death, were reduced in patients with HbF > 8.6%. Taken together, SCD patients benefit from having high expression of HbF, and this is due to the inhibitory effects it has on the polymerization of HbS.

1.2.3 Prevention and treatment of SCD

There are preventive measures to reduce the incidence rate of SCD. Programs for premarital, antenatal and neonatal screenings determine if both individuals who are engaged to get married have the sickle cell trait, and if an unborn child or a newborn child has SCD.⁴⁴ This will allow couples to make an informed decision about whether to

continue with their nuptials or to have children if they are both carriers of the sickle cell trait; allow for antenatal diagnosis that will provide a choice to continue with the pregnancy if the fetus is found to have SCD; and will alert clinical professionals to newborns born with SCD in order to monitor and start treatment at an earlier time.

As for therapeutic treatments for SCD, there are two United States (US) Food and Drug Administration (FDA)-approved drugs to treat SCD. First approved is hydroxyurea. Hydroxyurea was initially used as a chemotherapeutic agent to treat myeloproliferative disorders. Scientists found that hydroxyurea treatment increased HbF levels in baboons⁵⁶, and therefore started clinical trials to determine if it is helpful for patients with SCD, leading to it being the standard of treatment.^{57–61} However, the mechanism of how hydroxyurea upregulates HbF is not clearly understood. Furthermore, not all SCD patients respond to hydroxyurea treatment. Recently, L-glutamine, given as daily supplement, was associated with reduced vaso-occlusive crises and was FDA approved.⁶² Whether or not this will change the standard of care is unclear.

Blood transfusions are another form of treatment, which reduces the percent of sickle cells in circulation by giving the patient healthy RBCs. Even though blood transfusions are effective, it is a burden because patients have to frequently have this procedure.^{63–65} Also, patients who undergo frequent blood transfusions can develop antibodies against red cell antigens from transfused blood. Furthermore, only individuals living in countries with a good health care structure will be able to access this treatment.

A curative form of treatment is bone marrow transplantation, which replaces the bone marrow of SCD patients with bone marrow from healthy individuals, thus substituting abnormal HSCs with normal HSCs.⁶⁶ The limitations with this procedure are that complications can be fatal, it is expensive and there are few suitable donors. An approach taken to address the lack of donors is gene therapy. Gene therapy takes autologous HSCs, infects them with a self-inactivating lentiviral vector that encodes a variant for β -globin chain (T87Q substitution), which inhibits the polymerization of HbS, and engraft the transduced HSCs into the bone marrow of the patient.⁶⁷ Recently, a case study was published about the 15-month follow-up of an SCD patient who underwent gene therapy.⁶⁸ Not only did high expression of the variant β -globin persist, the patient had no sickle crises. Gene therapy is a promising treatment, but the remaining two limitations of bone transplantation still is a concern.

1.3 HbF Variability

The observation that HbF is differentially expressed in patients with SCD, and that it is beneficial for patients to have higher levels of HbF, have steered researchers to determine the causes for this occurrence. What has been elucidated as the main causes for high HbF are two types of genetic variants—deletions along the β -globin gene locus, and single nucleotide polymorphisms (SNPs) found in various regions of the genome.⁶⁹ These deletions have been named based on sequence of discovery, or the geographical location or ethnic group the variant is mostly prevalent.

1.3.1 $(\delta\beta)^{0}$ -thalassemia and hereditary persistence of fetal hemoglobin

Numerous deletions have been discovered within the β -globin gene cluster. Two types of deletions— $(\delta\beta)^0$ -thalassemia and hereditary persistence of fetal hemoglobin

(HPFH)—are associated with high HbF.^{70,71} Multiple deletions have been characterized for each type; they vary in size, and new deletions continue to be discovered. The deciding factors for what makes a deletion either a $(\delta\beta)^0$ -thalassemia or an HPFH are the percentage of HbF that is expressed and the distribution of HbF among RBCs. Adults with $(\delta\beta)^0$ -thalassemia have HbF of 5-15% and it is distributed heterocellularly among RBCs, which means that not all cells have the same amount of HbF. Conversely, adults with HPFH have higher levels of HbF (15-30%) and is distributed evenly or pancellularly among RBCs.

The location and size variability has led to discoveries of cis-acting elements essential for globin switching. The first element is a 3.5 (kilobase) kb intergenic region 5-prime (5') to the δ -globin gene (*HBD*), as has been found to be essential for the silencing of *HBG* expression via binding of Bcl11a and other transcription factors within this region.⁷² Consequently, a deletion within this region will increase *HBG* expression. Next, is the 3-prime (3') HS1, which is a DNase1 hypersensitive site (HS) located downstream of *HBB* and is thought to be an insulator against enhancers for *HBG*.⁷³ Deletion of this region is also thought to increase *HBG* expression. Last, are the three 3'enhancers, which are also downstream of *HBB* and have been found to enhance *HBG* expression.⁷³ A deletion through *HBB* to a position 5' to anyone of these enhancer regions is thought to bring the enhancer closer to *HBG* and increase its expression.

1.3.2 The 3 major quantitative trait loci

SNPs are single nucleotide variants and are found everywhere in the genome. Some SNPs do not change gene expression or epigenetic regulatory elements, while others do. Genome-wide association study (GWAS) is a tool used to determine which SNPs are significantly associated with a specific trait. Since HbF is a trait that is beneficial for individuals with SCD, investigators have used GWAS to determine which SNPs are associated with high HbF, with the intent to determine if these SNPs could lead them to understand how HbF is regulated.⁷⁴ Using GWAS, three regions of the genome have been revealed to be associated with high HbF. These are the *HBG2* promoter region, *BCL11A*, and the *HBS1L-MYB* intergenic region, and together are the three major quantitative trait loci (QTL) for HbF.

First of these QTL to be discovered was the Xmn1-HBG2 C-T polymorphism located within the promoter of HBG2 158 base pairs (bp) upstream from the HBG2 transcription start site (TSS). This SNP creates an Xmn1 restriction site and is associated with increased expression of HBG2.^{19,75} The mechanism of action or the functional basis of this association is currently unknown. The next region is *BCL11A* found on chr2p16, which encodes the transcription factor B-cell lymphoma/leukemia 11A (BCL11a). As previously discussed (section 1.1.3.) BCL11a is a transcription factor that is important for SNPs found in intron 2 of BCL11A inhibit the γ -globin to β -globin switch.^{76–79} transcription of the gene, and therefore the hemoglobin switch does not occur, causing constitutive expression of *HBG*. The last QTL is the HBS1L-MYB intergenic polymorphisms (HMIP) found on chr6q23, which consists of three HMIP trait-associated blocks with SNPs that are in linkage disequilibrium (LD).^{80,81} LD implies that SNPs are likely to be inherited with each other. This QTL will be discussed further in the next section.

1.4 The HBS1L-MYB Intergenic Region

The HBS1L-MYB intergenic region is 126 kb in length, and is in between the genes HBS1L and MYB. HBS1L is a member of the GTP-binding elongation factor family with no known association with erythropoiesis or hemoglobin expression. As discussed before (section 1.1.3), MYB encodes for the transcription factor c-MYB, and is known to regulate erythroid proliferation and maturation, and hemoglobin expression.^{25–27} There are many SNPs within this intergenic region, and for a while it was not understood why these SNPs were associated with high HbF. Our lab was one of many laboratories to confirm the association of these SNPs with high HbF.⁸² Among a group of non-related Chinese individuals heterozygous for β-thalassemia, HbF variability was associated with known SNPs such as rs9399137. What was novel was the discovery of a 3-bp deletion polymorphism (rs66650371), which is in LD with rs9399137 and therefore also associated with high HbF. Additionally, it was discovered by chromatin immunoprecipitation (ChIP) around the 3-bp deletion that in K562 cells, an immortalized erythroid cell line, are binding sites for four transcription factors that are involved in erythroid cell differentiation, GATA, TAL1, E47 and RUNX1. Also, Encyclopedia of DNA Elements (ENCODE) datasets revealed binding of RNA polymerase II and other factors such as Brg1 signaling, which signifies transcriptional activity, histone H3 lysine 4 monomethylation (H3K4Me1) activity (marker for enhancer regions), and expression of a 50-bp transcript. To confirm that this region was an enhancer, the fragment was cloned into a vector with an HBG2 promoter and a luciferase reporter gene. Cells transfected with the fragment containing this region showed higher levels of luciferase

activity than cells cloned without this region. We concluded that the 3-bp deletion within HMIP was associated with increased HbF production, had signatures of transcription activity very similar to erythroid-specific transcription activity, and contained an enhancer element.

This work was later confirmed and extended by others who found that within the HMIP region was an enhancer region, which included the 3-bp polymorphism and binding sites for erythroid-specific transcription factors, and that the enhancer element actually regulated the expression of *MYB*.^{83,84} They determined the mechanism of how polymorphisms in this region were associated with high HbF, and showed that when this enhancer region is intact, it has the ability to bind to the promoter of *MYB* and upregulate its expression. However, if the 3-bp polymorphism was present, the enhancer cannot bind to the promoter, resulting in a reduction in the expression of *MYB* and increase in HbF expression.

1.5 Long Noncoding RNA

The genome contains genes that are transcribed and processed into messenger RNAs (mRNAs), and these mRNAs are then translated into proteins to regulate the multiple processes in the cell. However, it was discovered that while 50-70% of the genome is transcribed, only 2% is translated into protein.⁸⁵ Except for transfer RNAs (tRNAs) and ribosomal RNAs (rRNAs), the other remaining untranslated RNAs were considered to have no biological function and were labeled as "junk RNAs." This was puzzling because many less complex species translate more of their genome, while more complex species, such as humans, translate less. As next generation sequencing (NGS)

became a popular scientific tool to look at total RNA expression, scientists discovered that noncoding RNAs (ncRNAs) are differentially expressed during development and within various disease models.⁸⁶ The discovery of changes in ncRNA expression patterns led to functional analyses to determine if modulation of ncRNA expression affect any downstream molecular or cellular activity. Their findings showed that ncRNAs actually do regulate molecular and cellular activities, and therefore changed the dogma of ncRNAs—they do have biological functions. There are various types of ncRNAs and among them are the long noncoding RNAs or lncRNAs.

LncRNAs are transcripts of 200 nucleotides or greater that do not code for protein.^{87,88} They are transcribed from any region in the genome; low in abundance; mostly tissue-specific (more so than mRNA); located mostly in the nucleus of the cell; and can have many signatures of mRNA, such as the 5'-cap, 3'-polyadenylation and the ability to be spliced.⁸⁹ Furthermore, they have been shown to regulate cellular processes at various levels.^{90,91} They can recruit epigenetic factors, control pluripotency, regulate transcription (in *cis* or *trans*) and translation, and be involved in post-transcriptional processing of mRNA. There are various types of lncRNAs that are named based on location relative to neighboring protein-coding genes.^{92,93} Intergenic lncRNAs (lincRNAs) are located between two protein coding genes; antisense lncRNAs (alncRNAs) are transcribed complementary to a mRNA; intronic overlapping lncRNAs (ilncRNAs) are transcribed from the intronic region of mRNA; small RNA (sRNA) host lncRNAs (shlncRNAs) have sRNAs transcribed from their intronic regions; pseudogene

lncRNAs (plncRNAs) are transcribed from pseudogenes; and enhancer RNAs (eRNAs) are transcribed from the enhancer of an mRNA.

1.5.1 Enhancer RNAs

Enhancers are regions in the genome that when bound by transcription factors, are able to change the conformation of the chromatin in order to activate transcription of a gene. Active enhancers are known to have transcriptional activity. There are three reasons for enhancer transcription—noise, transcription-dependent effects, or RNA-dependent effects.⁹⁴ Noise is due to collision of Pol II with accessible chromatin, and does not have any biological function. A transcription-dependent effect is when movement of Pol II causes the conformation of chromatin to change, and epigenetic processes such as histone acetylation and methylation occur. This is eRNA-independent. eRNA-dependent effects are when transcripts derived from enhancer regions function to regulate transcription either in *cis* or *trans*.^{95–99} This effect is eRNA-dependent, where eRNAs recruit or bind coactivators or transcription factors, and at act as a scaffold to ensure binding of these proteins at the promoter of neighboring protein-coding genes to promote gene expression (**Figure 1.4**).⁹⁵



Figure 1.4. The function of an enhancer RNA (eRNA).

In this illustration, long noncoding RNA, ncRNA-a7, is transcribed from the enhancer element for *SNAI1* and acts as a scaffold to transcription factors bound to the gene's promoter to promote transcription of *SNAI1*. Image adapted from Ong and Corces, *Nature Reviews, Genetics, April 2011*.

1.5.2 Hematopoietic/Erythroid-associated long noncoding RNAs

There are multiple lncRNAs found to regulate either hematopoiesis or erythropoiesis.^{93,100–111} Most of what is known has been studied mostly in murinederived cells. Hu et al found that more than 400 putative mouse lncRNAs fluctuate during erythropoiesis.¹¹² Functional studies on one lncRNA, named long intergenic noncoding RNA (lincRNA) erythroid prosurvival (LincRNA-EPS), discovered that it inhibits apoptosis to promote erythroid differentiation. Other work characterized a variety of lncRNAs by using microarray analyses from the ENCODE datasets.¹¹¹ Since the HOXA gene cluster is differentially expressed during granulocytic differentiation and is myeloid-specific, they looked for expression of intergenic transcripts within this cluster in human cell lines that are able to undergo granulocytic differentiation. This laboratory was able to find a lncRNA, which they named HOTAIRM1, located between HOXA1 and HOXA2 genes. During granulocytic differentiation, HOTAIRM1 is upregulated, and was dependent on the expression pattern of both HOXA1 and HOXA2. Bertani et al looked for lncRNAs that are involved in epigenetic activation via Mixed lineage leukemia 1 (MLL1), which is known to be vital for embryonic development and hematopoiesis. RNA-chromatin immunoprecipitation (RNA ChIP) along with DNA microarray (RNA ChIP-on-chip) assay showed that the lncRNA Mistral (MIRA) binds and recruits Mll1 to activate transcription of homeotic genes HOXA6 and HOXA7.¹¹³ When Hoxa6 and Hoxa7 are expressed, they are able to activate expression of germ layer marker genes to induced differentiation of murine embryonic stem cells (mESC). Finally, work done by Paralker et al, used RNA sequencing in erythroblasts from both murine fetal liver and

human CD34⁺ cells derived from cord blood, identified 1109 and 594 polyadenylated lncRNAs, respectively.¹⁰⁸ For both species, more than 50% of the lncRNAs were unannotated. Data from mouse erythroblasts showed that the expression of many of these lncRNAs are regulated by Gata1 and Tal1, which are key erythroid-specific transcription factors; 15% of lncRNAs are expressed in human erythroblasts; and knockdown of 7 out of 21 of the most abundant lncRNAs inhibited terminal erythroid differentiation (6 of the 7 were not expressed in human erythroblasts). Only 15% of the lncRNAs in human erythroblasts were expressed in mouse erythroblasts, suggesting that lncRNAs expressed in erythroid cells are highly species-specific.

1.5.3 Disease-associated lncRNAs

LncRNAs have also been implicated as key players in disease pathophysiology, including Alzheimer's disease, cancer and cardiovascular disease.^{114–119} β -secretase-1 (*BACE1*)-*AS* is an antisense lncRNA transcribed complementary to the gene *BACE1*, which is known to be necessary for the formation of beta-amyloid and amyloid plagues found in patients with Alzheimer's disease.¹²⁰ Further investigation into the function of *BACE1-AS* discovered that this lncRNA actually interacts with and stabilizes the mRNA of *BACE1*, causing the upregulation of BACE1 protein levels. This implies that *BACE1-AS* is the key target to stop the production of amyloid plagues for treatment of Alzheimer's disease. In cancer, many lncRNAs have been found to be differentially expressed compared to their noncancerous counterparts. *MALAT1* is expressed in normal tissue; however, it is increased in multiple types of cancer.^{121–123} Functional analyses for *MALAT1* show that when it is downregulated in cancer cells, they no longer have the

ability to proliferate, migrate and invade distal tissues. In cardiovascular disease, SNPs found in lncRNAs such as *ANRIL*, have been found to be associated with myocardial infarction.^{124,125} Certain SNPs upregulated the expression of *ANRIL*, and promoted the production of peripheral blood mononuclear cells within the blood and buildup of atherosclerotic plagues within blood vessels, which can lead to a heart attack and stroke.

LncRNAs are now considered important regulators of gene expression and processes of the cell, including erythroid cells. Moreover, lncRNAs are implicated in disease pathophysiology that suggests the possibility that they might have some role in hemoglobinopathies such as SCD.

1.6 Rationale and Hypothesis

SCD is a genetic disease that affects the functions of both hemoglobin and RBCs. Not all individuals with SCD display the same disease phenotypes. One factor known to affect the phenotype of disease is HbF. The three known QTL only account for 20-45% of HbF variability.^{77,126,127} Therefore, more is to be discovered about the regulation of HbF.

The *HBS1L-MYB* intergenic region is a key HbF-associated QTL. Within this region is an enhancer element for *MYB*, which is known to regulate erythroid-specific traits. Querying the ENCODE datasets in erythroid cells revealed the presence of a 50-bp RNA transcript from this enhancer element. Knowing that enhancers can express functional lncRNAs, we hypothesized that this 50-bp RNA might be part of an eRNA transcribed from within the *HBS1L-MYB* intergenic region, specifically from the enhancer for *MYB*, and represents a putative lncRNA that regulates *MYB* and its

downstream targets. If true, this might contribute to a better understanding of the regulation of erythropoiesis and hemoglobin expression, and this lncRNA could possibly be considered a therapeutic target for the treatment of SCD.

1.7 Specific Aims

There are two specific aims to address this hypothesis. The first aim is to characterize the transcript by determining the following: the full length, cellular location, relative expression compared to protein-coding genes, tissue-specificity, and the expression pattern among erythroid cells with different hemoglobin makeup and during erythroid differentiation. The second aim is to address the functional aspect of the transcript, and determine if the transcript regulates *MYB* and hemoglobin gene expression.

CHAPTER 2. CHARACTERIZATION OF THE TRANSCRIPT LOCATED WITHIN THE *HBS1L-MYB* INTERGENIC REGION

2.1 Introduction

Since the year 2000, when the International Human Genome Project made the assembly of the human genome available to the public, it became easier for investigators to know the location, length and sequence of genes of interest.^{128,129} However, the caveat to knowing these characteristics is that these genes would have to be annotated within the human genome assembly. As scientific tools became more advanced, new genes were continuously added to the list. RNA sequencing (RNA-Seq) is a tool used to profile the transcriptome by using next generation deep-sequencing technologies.¹³⁰ The purpose of RNA-Seq is to determine which transcripts—mRNA, ncRNA and small RNA—are found, and quantify the amount of each individual transcript in a particular sample. However, there are still limitations, especially when it comes to characterizing lncRNAs due to the fact that they are expressed at very low levels compared to protein coding genes; and since lncRNAs are mostly tissue-specific, RNA has to be extracted from the right cell type.

As discussed in Chapter 1, the 50-bp transcript found in the *HBS1L-MYB* intergenic region was discovered using ENCODE datasets from RNA-Seq analyses of K562 cells. While it was plausible to consider this transcript a small noncoding RNA, we decided to perform additional analyses to determine if the outcome could possibly be different. RNA-Seq of RNA derived from primary erythroblasts showed multiple reads from this same region.

2.2 Determining the Full Length of the Transcript

The first set of experiments done to address the question of whether the 50-bp transcript was part of a lncRNA, used reverse transcription polymerase chain reactions (RT-PCRs). First, multiple primers were developed consecutively downstream and upstream the genomic region of the 50-bp transcript, at 100-bp intervals, spanning a 1400 bp region. We used for PCR complementary DNA (cDNA) (DNA synthesized from RNA) of K562 cells, which are immortalized erythroid cells that can be induced to make HbF. The controls used for PCR were PCR mix without DNA input (negative control), PCR mix with genomic DNA (positive control), and PCR mix with cDNA generated without reverse transcriptase (negative control to ensure RNA was not contaminated with DNA). The PCR reactions were done with primers starting from the 50-bp region and extended further with the next consecutive primers until there was no amplification. The longest PCR product amplified was 1180 bp (chr6: 135096355-135097534; UCSC Genome Browser assembly ID: hg38) (**Figure 2.1A**), and therefore upheld our hypothesis that a lncRNA was expressed from the *HBS1L-MYB* intergenic region.

We next determined the entire length of the lncRNA by using rapid amplification of cDNA ends (RACE). RACE is used to determine the 5'- and 3'-ends of full length RNA by ligating adapters to ends of cDNA, and then amplifying the transcript of interest with adapter-specific and gene-specific primers.¹³¹ After confirmation of a PCR product by running the reaction on an agarose gel, the product is purified, cloned into a vector, and analyzed further to determine the exact location of the ends of the RNA (see subsection 2.8.6). Most RACE protocols use cDNA as the starting material, which can

produce non-specific amplification (especially for the 5'-end reactions) due to the lack of selectivity of specific RNAs. However, improvements have been made to produce a single band after PCR. Therefore, RNA Ligase Mediated (RLM) RACE was used, which, for the 5'-end, used RNA as the starting material; removed all degraded RNA, rRNA, tRNA and DNA; removed the 5'-cap (leaving only a monophosphate); ligated an adapter (will only ligate to RNA with the 5'-monophosphate); reverse transcribed RNA; and amplified 5'-end with adapter-specific and gene-specific primers. The RLM-RACE protocol for the 3'-end has fewer steps and higher success rate than the 5'-RACE. To determine the 3'-end, total RNA is reverse transcribed, adapter is ligated to 3'-end, and then PCR reactions are done with adapter-specific and gene-specific primers. RNA from K562 cells was initially used for both 5'- and 3'-RACE, however there was a huge DNA smear for the 5'-RACE, which could not be cloned into a vector for further analysis. The sequencing results for the 3'-RACE of K562 cells revealed that the 3'-end of the lncRNA was at location ch6: 135097644 (hg38) (Supplemental Figure 3), which was 110 bp downstream from what was found by RT-PCR (Figure 2.1B). As for the 5'-RACE, it was repeated using RNA from erythroblasts derived from cord blood CD34⁺ cells. This time there were multiple bands, and each band was cloned and analyzed further, however, none of the sequencing results were found to be from this region. During the analysis of the expression pattern of this lncRNA among various types of tissues, it was found that the thymus expressed the lncRNA at much higher levels than both K562 cells and $CD34^+$ cells (see section 2.3). Repeating 5' RLM-RACE using RNA from the thymus showed only one band. Sequencing results of that band revealed that the 5'-end of this lncRNA

was at location ch6: 135096362 (hg38) (**Supplemental Figure 4 and Figure 2.1B**). HUDEP-2 cells, another immortalized erythroid cell line that expresses HbA, expressed more of the transcript than both K562 cells and erythroblasts derived from CD34⁺ cells. Both the 5' and 3' RLM-RACE analyses gave the same results in these cells. The full length of this lncRNA, which we named the *HBS1L-MYB* intergenic long noncoding RNA (*HMI-LNCRNA*), is 1283 bp at chr6: 13509362-135097644 (hg38) (**Figure 2.1C**). *HMI-LNCRNA* is located 84.9 kb upstream from *MYB* and 41.5 kb upstream from *HBS1L*, and includes binding sites for erythroid transcript factors TAL1/E47, GATA1 and RUNX1, HbF-associated polymorphism rs66650371, and is transcribed from the enhancer for *MYB*. There is also no evidence of protein coding potential of *HMI-LNCRNA* based on the fact that it is absent in protein databases and contains no open reading frames longer than 300 nucleotides.



Figure 2.1. Determining the full length of the transcript found within the *HBS1L-MYB* intergenic region.

(A) PCR was done to amplify a 1180 bp region within the *HBS1L-MYB* intergenic region using cDNA from K652 cells. Samples were run on agarose gels and exposed under UV light. Lane 1: 100-bp DNA ladder; lane 2: negative control, PCR mix without DNA input; lane 3: positive control, PCR mix with genomic DNA; lane 4: PCR mix with cDNA generated without reverse transcriptase; and Lane 5: PCR mix with cDNA generated with reverse transcriptase. (B) Agarose gels show PCR products from the 5'-RACE reactions (using RNA from thymus) and 3'-RACE reactions (using RNA from K562 cells). For 5'-RACE, lane 1: 100-bp DNA ladder; lane 2: PCR mix with cDNA generated without Tobacco Acid Pyrophosphatase (TAP) treatment; and lane 3: PCR mix with cDNA generated with cDNA generated with the top treatment. For 3'-RACE, lane 1: 100-bp DNA ladder; lane 2: PCR mix with cDNA generated with cDNA generated without reverse transcriptase. (C) Illustration of genomic region between 134950000-135250000 (hg38 coordinates) of chromosome 6, showing

approximate locations of rs66650371 and rs9399137, and *HBS1L* and *MYB* (arrows represent transcription direction and approximate length of genes). Based on DNA sequencing of RACE products, the 5'- and 3'-ends of the transcript were revealed to determine the full length of the transcript, which is 1283 bp in length and named the *HBS1L-MYB* Intergenic Long Noncoding RNA (*HMI-LNCRNA*). Located in the genomic sequence for *HMI-LNCRNA* are binding sites for erythroid-specific transcription factors TAL1/E47, GATA and RUNX1, and the 3-bp polymorphism (rs66650371). *HMI-LNCRNA* does not include rs9399137.

2.3 Tissue Specificity of HMI-LNCRNA

Since lncRNAs are expressed in either one or a few tissue types (more tissuespecific than mRNA), we explored the expression pattern of HMI-LNCRNA among 25 different types of cells and tissues. RNA samples were either purchased or extracted, and HMI-LNCRNA expression was analyzed by quantitative PCR (qPCR). Not all tissue expressed *HMI-LNCRNA*. The fold change of expression by qPCR in multiple samples that included erythroblasts derived from cord blood CD34⁺ cells, bone marrow, fetal liver, brain, spleen, skeletal muscle, spinal cord, testis and thymus, had expression level of at least 1-fold above K562 cells; only three samples (erythroblasts derived from cord blood CD34⁺ cells, testis and thymus) had a fold change above 10; and only one sample (thymus) had a fold change above 1000 (Figure 2.2). T-cells found in the thymus and erythroid cells both are derived from hematopoietic stem cells (HSCs), and require MYB to regulate differentiation. MYB is low in both mature erythroid cells and mature Tcells.²⁷ This suggests that HMI-LNCRNA may work together with MYB to regulate differentiation of both cell types. If this is so, we might expect higher levels of HMI-LNCRNA in HSCs, and declining levels during the maturation of each cell type. However, it is unknown if the expression of *HMI-LNCRNA* in the thymus is from the Tcells, their precursors or a non-associated cell type. HMI-LNCRNA in peripheral blood mononuclear cells (PBMCs), which consists of 40-70% mature T-cells, was low, suggesting that the high levels of this lncRNA maybe coming from T-cell precursors.

The expression pattern of *MYB* in these same samples was evaluated to determine if it correlated with the expression pattern of *HMI-LNCRNA*. K562 cells, which express

very low levels of *HMI-LNCRNA* compared to erythroblasts derived from cord blood $CD34^+$ cells and the thymus, had the highest expression of *MYB* among all the samples tested (**Supplemental Figure 1**). However, *MYB* has 13 isoforms, and the primer used for qPCR only recognizes one isoform. If different isoforms for *MYB* were expressed among these samples, primers specific to the isoform expressed in each sample would be needed to evaluate *MYB* expression.

2.4 Cellular Location of HMI-LNCRNA

Unlike mRNAs, which translocate to the cytoplasm to undergo translation, lncRNAs can either stay in the nucleus or move to the cytoplasm after transcription, depending on their function. Most lncRNAs are located in the nucleus because they regulate gene expression. Hence, it was important to know if *HMI-LNCRNA* is expressed in the nucleus or the cytoplasm. Primary erythroblasts derived from cord blood CD34⁺ mononuclear cells were harvested and underwent nuclear and cytoplasmic fractionation. RNA was extracted from each fraction, and the relative transcript level of *HMI-LNCRNA* was analyzed by qPCR. *HMI-LNCRNA* was expressed almost entirely in the nucleus (**Figure 2.3**), suggesting that *HMI-LNCRNA* regulates gene expression. Regulation could affect *MYB*, *HBS1L* or another protein-coding gene.



HMI-LNCRNA

Figure 2.2. Expression pattern of *HMI-LNCRNA* among various human cells and tissue.

Relative quantity of *HMI-LNCRNA* was measured by qPCR in 25 different samples. $CD34^+$ cord blood day 11 is erythroblasts harvested at Day 11 of two-phase expansion and differentiation culture of $CD34^+$ mononuclear cells derived from cord blood. Means are shown for samples with two or more independent samples. *ACTB* was used as the endogenous control.



Figure 2.3. *HMI-LNCRNA* expression in nuclear and cytoplasmic fractions of primary erythroid cells derived from cord blood CD34⁺ mononuclear cells.

Nuclear and cytoplasmic fractions were extracted from erythroid cells derived from cord blood CD34⁺ mononuclear cells at Day 8 (n=1) and Day 12 (n=2) of differentiation. RNA extracted from each fraction was analyzed by qPCR analysis for expression of *HMI-LNCRNA*. The ratio of the relative quantity of nuclear to cytoplasmic *HMI-LNCRNA* transcripts was measured. *ACTB* was used as the endogenous control. p-value: ***<0.005. p-value obtained by Student T-test.

2.5 Comparing the Relative Expression Level of *HMI-LNCRNA* to a Protein-Coding Gene

LncRNAs have been shown to be less abundant than mRNA from protein-coding genes. To determine if *HMI-LNCRNA* shares this same characteristic, the relative expression level of *HMI-LNCRNA* was compared to the expression of *MYB* by qPCR. Using RNA extracted from HUDEP-2 cells and erythroblasts derived from peripheral blood CD34⁺ cells, *MYB* was expressed 300 to 500 times more than *HMI-LNCRNA* (Figure 2.4).





Figure 2.4. Ratio of *MYB* to *HMI-LNCRNA* expression in HUDEP-2 cells and erythroid cells derived from peripheral blood CD34⁺ mononuclear cells.

Relative quantity ratio of *MYB* to *HMI-LNCRNA* was determined in HUDEP-2 cells (n=2) and erythroid cells derived from primary peripheral blood mononuclear cells at Day 9 of differentiation (n=3) by qPCR. *ACTB* was used as the endogenous control.

2.6 Relative Expression Pattern of *HMI-LNCRNA* Between Erythroid Cells with Different Hemoglobin Phenotypes and During Erythroid Differentiation

MYB is known to be associated with low HBG and high HBB expression. We therefore sought to determine if HMI-LNCRNA was associated with the expression of fetal or adult globin genes. HMI-LNCRNA and MYB mRNA expression were compared in fetal-like erythroid cells expressing *HBG* and adult-like cells expressing *HBB*. The first set of cells analyzed were HUDEP-1 cells, which express 99% HBG and HUDEP-2 cells, which express 99% HBB. HMI-LNCRNA was significantly higher in HUDEP-2 cells compared to HUDEP-1 cells, however, there was no difference in expression of *MYB* (Figure 2.5A). The results for *MYB* do not mirror protein expression of c-MYB, which was found to be higher in HUDEP-2 cells compared to HUDEP-1 cells (see Chapter 3, subsection 3.2.4). This suggested that even though there may not be a difference transcriptionally, there could be differences in the translation of MYB mRNA, or degradation and/or stability of c-MYB between HUDEP-1 and HUDEP-2 cells. In primary erythroblasts derived from cord blood CD34⁺ cells that express 95% HBG and peripheral blood CD34⁺ cells that express 65% *HBG*, both *HMI-LNCRNA* and *MYB* were significantly higher in erythroblasts derived from peripheral blood compared to cells derived from cord blood (Figure 2.5B). These data suggest that high levels of MYB and HMI-LNCRNA expression are associated with increased HBB expression and reduced *HBG* expression.



Figure 2.5. *HMI-LNCRNA* and *MYB* expression pattern in HUDEP-1 and HUDEP-2 cells, and erythroid cells derived from cord blood and peripheral blood CD34⁺ mononuclear cells.

(A) Percent *HBG* to *HBB*, and the relative quantity of *HMI-LNCRNA* and *MYB* were determined by qPCR analysis in HUDEP-1 (n=3) and HUDEP-2 (n=2) cells (both maintained in expansion medium). (B) Primary erythroid cells derived from cord blood (n=3) and peripheral blood (n=3) CD34+ mononuclear cells (both expanded for 7 days and differentiated for 2 days) were analyzed by qPCR analysis to determine percent *HBG* to *HBB*, and the relative quantity of *MYB* and *HMI-LNCRNA*. *ACTB* was used as the endogenous control. p-values: *< 0.05; **< 0.005; ***<0.0005; NS (not significant). p-values were obtained by Student T-test.

Another characteristic of lncRNAs is that they are differentially expressed during development. Therefore, since *HMI-LNCRNA* is associated with erythroid cells that express more *HBB*, the expression pattern of this lncRNA was analyzed during differentiation of HUDEP-2 cells and peripheral blood CD34⁺ mononuclear cells. During the differentiation of both cell types, *HMI-LNCRNA* and *MYB* mRNA decreased (**Figure 2.6**).

Taken together, these data show that both *HMI-LNCRNA* and *MYB* have similar expression patterns in erythroid cells, and suggest that *HMI-LNCRNA* may regulate the transcription of *MYB*.



Figure 2.6. *HMI-LNCRNA* and *MYB* expression pattern during differentiation of HUDEP-2 cells and erythroid cells derived from peripheral blood CD34⁺ mononuclear cells.

(A) Fold change of *HMI-LNCRNA* and *MYB* were determined in HUDEP-2 cells at Days 0, 5 and 7 in cultures with differentiation medium. (B) Fold change of *HMI-LNCRNA* and *MYB* were determined during differentiation of CD34⁺ mononuclear cells derived from adult peripheral blood (n=3) at Days 7, 9, 11, 13 and 15. *ACTB* was used as the endogenous control. p-values: *<0.05; **<0.005; **<0.0005; NS (not significant). p-values were obtained by Student T-test.

2.7 Summary and Conclusions

The 50-bp transcript annotated by ENCODE and located within the HBS1L-MYB intergenic region, which is known to regulate erythroid-specific traits, became an interesting element in our mission to better understand hemoglobin regulation. Therefore, additional steps were taken to fully characterize this transcript. The first step was to determine the full length of this transcript. By RT-PCR, and 5'- and 3'-RACE, we discovered that the transcript is 1283 bp (chr 6: 135096362-135097644, hg38) and is a lncRNA; it contains the binding sites for erythroid-specific transcription factors; and it is transcribed from the enhancer region of *MYB*, which brands it an enhancer RNA (eRNA). This lncRNA was named the HBS1L-MYB intergenic region long noncoding RNA (HMI-LNCRNA). HMI-LNCRNA is not erythroid-specific, however, it was expressed in only a few cell/tissue types and could possibly be hematopoietic-specific. The expression pattern of HMI-LNCRNA in erythroid cells was mostly nuclear; it was less abundant than MYB mRNA; higher expression was associated with erythroid cells that express more HBB; and it was downregulated during erythroid differentiation. HMI-LNCRNA and MYB showed the same expression pattern in erythroid cells and during erythroid differentiation (Figure 2.7).



Figure 2.7. Summary of the characteristics of HMI-LNCRNA.
2.8 Materials and Methods

2.8.1 K562 cells

K562 cells are immortalized erythroid progenitors derived from pleural effusion of a patient with chronic myelogenous leukemia. They can be differentiated with arginine butyrate treatment toward erythroblasts that express only embryonic and fetal hemoglobin (**Supplemental Figure 2.4**).¹³² They were cultured at 37 degrees Celsius (°C) in RPMI medium containing 10% FBS and 2% penicillin/streptomycin. To induce hemoglobin expression, cells are seeded at a concentration of 1x10⁵ cells/ml with 1 mM arginine butyrate, and cultured at 37°C for seven days (medium changed every two days).

2.8.2 HUDEP cells

Human umbilical cord blood-derived erythroid progenitor (HUDEP) cells are an immortalized erythroid cell lines derived from cord blood CD34⁺ mononuclear cells.¹³³ HUDEP-1 and HUDEP-2 cells were cultured at 37°C maintained in StemSpan SFEM medium (StemCell Technologies) supplemented with SCF (50 ng/ml, Invitrogen), EPO (3 U/ml, Invitrogen), dexamethasone (1 μ M, Sigma), doxycycline (1 μ g/ml, Clontech), L-glutamine (1%, Life Technologies) and penicillin/streptomycin (2%, Life Technologies). For erythroid maturation, cells were cultured at 37°C in IMDM medium (Invitrogen) supplemented with heat inactivated human serum from human male AB plasma (5%, Sigma), EPO (3 U/ml, Invitrogen), insulin (10 μ g/ml, Sigma), doxycycline (1 μ g/ml, Clontech), holo-transferrin (500 μ g/ml, Sigma), heparin (3 U/ml, Sigma), SCF (100 ng/ml, Invitrogen), L-glutamine (1%, Life Technologies) and penicillin/streptomycin (3 U/ml, Sigma), SCF (100 ng/ml, Invitrogen), L-glutamine (1%, Life Technologies) and penicillin/streptomycin (3 U/ml, Sigma), SCF (100 ng/ml, Invitrogen), L-glutamine (1%, Life Technologies) and penicillin/streptomycin (3 U/ml, Sigma), SCF (100 ng/ml, Invitrogen), L-glutamine (1%, Life Technologies) and penicillin/streptomycin

(2%, Life Technologies) for 5 days. For further erythroid maturation, doxycycline was removed and cells were cultured for two more days.

2.8.3 Primary CD34⁺ mononuclear cells

Primary CD34⁺ mononuclear cells derived from cord blood and peripheral blood (StemCell Technologies) contain hematopoietic stem/progenitor cells that can be differentiated into multiple lineage-specific cells. To generate erythroid cells, CD34⁺ cells underwent a 2-phase culture system. The first phase is to expand the cells at 37°C in StemSpan SFEM II medium (StemCell Technologies) supplemented with StemSpan StemCell Technologies) and penicillin/streptomycin (2%, CC100 (1X, Life Technologies) for six or seven days. The second phase is to drive the cells to erythroid lineage. To induce erythroid differentiation, cells were cultured at 37°C in StemSpan SFEM II medium (StemCell Technologies) supplemented with SCF (10 ng/ml, Invitrogen), EPO (5 U/ml, Invitrogen), IL-6 (10 ng/ml, Sigma) and penicillin/streptomycin (2%, Life Technologies) for up to 10 days (Supplemental Figure 2).

2.8.4 RNA extraction

Cells taken from culture were centrifuged at 300 x g for 10 minutes at 4°C. Next, supernatant was removed without disturbing cell pellet, cells were resuspended in cold 1X Phosphate-buffered saline (PBS) and centrifuged again at 300 x g for 10 minutes at 4°C. After removal of supernatant, cell pellets were stored at -80°C or processed for RNA extraction. Total RNA was extracted using RNeasy Mini Kit (Qiagen), treated with DNase (RNase-Free DNase Set, Qiagen), and then purified by RNA cleanup using RNeasy Mini Kit. For tissue-specificity experiment, multiple human organ RNA panels (Invitrogen and Clontech) were treated with DNase, followed by RNA cleanup.

DNase treatment is necessary for qPCR analysis for *HMI-LNCRNA*. In my earlier experiments, there was non-specific amplification in reactions without reverse transcriptase, which suggest that the reaction was contaminated with DNA. However, the non-specificity was only a problem when I used the primer for *HMI-LNCRNA*, and not for the other primers, such as *ACTB* and *HBG*, which had no amplification in qPCR reactions without reverse transcriptase. I have heard from other investigators that this is problem when analyzing lncRNAs by qPCR, but they do not understand why this occurs. I tried multiple DNase treatment protocols, such as incubating the column with DNase before eluting the RNA, however there was still non-specific amplification. What did work is after extracting the total RNA, incubate the entire sample in DNase for 10 minutes, and then cleanup the RNA to remove the DNase and buffer.

2.8.5 Reverse transcription polymerase chain reaction (RT-PCR)

cDNA was synthesized from DNase-treated RNA using SuperScript III First-Strand Synthesis System for RT-PCR (Invitrogen). $Oligo(dT)_{20}$ was used for the annealing step, therefore, cDNA was synthesized from only $poly(A)^+$ RNAs.

PCR reactions were done using the Multiplex PCR kit (Qiagen). The following primers were used to amplify the 1180 bp product: 5'-ATCGCTCATGAGAAATGTGG-3' (forward) and 5'-GGAACCGCCCTGATAACATT-3' (reverse).

2.8.6 Rapid amplification of cDNA ends (RACE)

TTCACTCTGGACAGCAGATGTT-3'

5'- and 3'-RACE were done using the FirstChoice RLM-RACE Kit (Ambion), following the manufacturer's instructions, using SuperTaq Plus Polymerase (Life Technologies) for PCR reactions. For 5'-RACE, the negative control was cDNA synthesized from RNA that was not treated with Tobacco Acid Pyrophosphatase (TAP). TAP removes the 5'-cap from RNA and leaves only a monophosphate. After TAP treatment, the adapter is ligated to only RNA with the monophosphate at the 5'-end. Therefore, RNAs not treated with TAP cannot ligate to the adapter, and will not amplify during the PCR reactions because the primer specific for the adapter cannot bind to them.

The	following	gene-specific	primers	were	used:	5'-
GTCTA	AATGGTGTG	GCTCACAAA-3'		(5'-oute	r),	5'-
CCCCA	AGCTTCCTTA	ATCTGTAAA-3'		(5'-inner	r),	5' -

(3'-outer)

and

5'-

CGGTTCCCTCAGAAGACACTTA-3' (3'-inner). RACE PCR products were ligated to pCRII vector using TA Cloning Dual Promoter Kit (Invitrogen), transformed into One Shot INVaF chemically competent E. coli (Invitrogen), and grown on LB plates containing 100 µg/ml ampicillin and X-Gal. Insert-positive white colonies were picked and grown for DNA extraction. PCR reactions were done to amplify the regions where RACE ligated 5'the products were the vector (Forward: to TGTGGAATTGTGAGCGGATA-3' and Reverse: 5'-GTTTTCCCAGTCACGACGTT-3'), and then processed for DNA sequencing.

2.8.7 DNA sequencing

PCR products were purified using AccuPrep PCR Purification Kit, and prepared for sequencing using ABI Big Dye Terminator v3.1 Cycle Sequencing Kit, which uses only one primer (forward or reverse) that was previously used for PCR. The program used for PCR was based on the DNA sample. If sequencing genomic DNA or cDNA, the following program was used: 96°C for 1 min; 15 cycles (96°C for 10 sec, 50°C for 5 sec, 60°C for 1 min 15 sec; 5 cycles (96°C for 10 sec, 50°C for 5 sec, 60°C for 1 min 30 sec); 5 cycles (96°C for 10 sec, 50°C for 5 sec, 60°C for 2 min); 10°C hold. If sequencing plasmid DNA, the following program was used: 96°C for 1 min; 25 cycles (96°C for 10 sec, 50°C for 5 min, 60°C for 4 min); 4°C hold. ABI 3130xl Genetic Analyzer was used for sequencing. Sequence data was analyzed on FinchTV version 1.5.0. NCBI BLAST was used to determine location of sequence.

2.8.8 Nuclear and cytoplasmic fractionation

Cells taken from culture were centrifuged at 300 x g for 10 minutes at 4°C. Next, supernatant was removed without disturbing cell pellet, cells were resuspended in cold 1X Phosphate-buffered saline (PBS) and centrifuged again at 300 x g for 10 minutes at 4°C. After removal of supernatant, cells were processed using the Nuclear/Cytosol Fractionation Kit (BioVision), following manufacturer's instructions.

2.8.9 Quantitative PCR

qPCR—also known as real-time PCR—is a tool used to quantitate the relative transcript levels of a specific RNA. In the PCR phase, gene-specific primers are used to

amplify the target during a 40-cycle reaction period (1-40). More abundant transcripts are amplified at earlier cycle thresholds (Ct) values, while less abundant transcripts are amplified at later Ct values. To calculate the relative quantity of a transcript-of-interest (TOI), first take the difference between Ct of TOI and Ct of endogenous control, such as *ACTB* (β-actin), to get Δ Ct (Ct^{TOI} – Ct^{*ACTB*} = Δ Ct^{TOI}). Finally, the relative quantity of TOI is calculated using 2^{(-(Δ Ct))}. To calculate fold change between two samples—sample 1 (S1) and sample 2 (S2)—another approach is taken. For each sample, calculate Δ Ct^{TOI}. Next, calculate Δ ΔCt^{TOI} for each sample—for S1, [(Δ Ct^{TOI} for S1) – (Δ Ct^{TOI} for S1)] = (Δ ΔCt^{TOI} for S1) = 0; and for S2, [(Δ Ct^{TOI} for S2) – (Δ Ct^{TOI} for S1)] = (Δ ΔCt^{TOI} for S2). Last, the fold change of the TOI for each sample is calculated using 2^{(-(Δ ΔCt))}, in which S1 will equal 1.

Total RNA, TaqMan primers and reagents from the TaqMan RNA-to-CT 1-Step Kit (Applied Biosystems) were the components used to make 20 µl reaction mix for each sample (done in triplicate) that was loaded onto a 96-well plate, following manufacturer's instructions. The following TaqMan gene expression assays (Applied Biosystems) were used: *HBG1/2* (Hs00361131_g1), *HBB* (Hs00758889_s1), *MYB* (Hs00920556_m1), *HMI-LNCRNA* (custom TaqMan assay designed by Applied Biosystems to target genome position chr6: 135096354-135097644, hg38; assay ID number AJI1MTQ), and *ACTB* (Hs01060665_g1). QPCR reactions were ran on a StepOne Plus qPCR machine (Applied Biosystems). *ACTB* was used as the endogenous control.

CHAPTER 3. THE EFFECTS OF MODULATING *HMI-LNCRNA* EXPRESSION IN ERYTHROID CELLS

3.1 Introduction

Characterization of *HMI-LNCRNA* has given insight into how this lncRNA possibly functions in erythroid cells, possibly as an eRNA that promotes the expression of *MYB*, thereby regulating hemoglobin expression, and erythroid proliferation and maturation. To determine if *HMI-LNCRNA* regulated *MYB* and erythroid-specific traits, we employed various systems to modulate its expression in erythroid cells.

There are multiple techniques used to either knockdown or overexpress RNAs *in vitro*. To downregulate expression of RNAs, small interference RNAs (siRNAs), locked nucleic acids (LNAs) and short hairpin RNAs (shRNAs) can be used. siRNAs are synthetically made double-stranded RNAs (dsRNAs) of 20-25 nucleotides that are complementary to the RNA of interest. Cells are transfected with siRNAs using various methods that will allow the siRNAs to travel through the cell membrane and into the cytoplasm. Once in the cytoplasm, siRNAs are processed by Dicer to separate the two strands, and then binds to the RNA-induced silencing complex (RISC). The siRNA-RISC complex binds to the RNA of interest and cleaves it, reducing the amount of RNA in the cell. LNAs are a newer system that uses RNase H to cut the RNA of interest.¹³⁴ Cells are treated with a single-stranded RNase H activating DNA flanked by antisense oligonucleotides specific to the RNA of interest, and require no transfection reagent. After the LNA is in the cell, the antisense oligonucleotides binds to the RNA of interest and the RNA. Similar to siRNAs,

shRNAs use Dicer to separate the double-stranded DNA, and RISC to cleave RNA. However, they differ in the way they are made and introduced to the cell. An shRNA is a synthetically made double-stranded DNA template containing four specific regions sense, loop, antisense and terminator. The sense region is about 20 bp of the genomic region where the RNA of interest is transcribed from; the loop is 10 bp of nonsense nucleotides; antisense is complementary to the sense; and the terminator consists of nucleotides that will stop transcription. The shRNA template is ligated to a lentivector, which contains virus-specific genes that allow for efficient cell transduction and genome integration. Once integrated into the genome, shRNA is expressed, forms into a hairpin structure and undergoes the same processing as siRNAs.

There are also methods to completely stop transcription of RNAs. Gene editing uses nucleases such as zinc finger nucleases (ZFNs), transcription-activator like effector nucleases (TALENs) and clustered regularly interspaced short palindromic repeats (CRISPR) to edit the genome to inhibit gene transcription. One caveat to using gene editing, especially when studying intergenic lncRNAs, is that genomic regions with epigenetic regulatory elements such as enhancers and promoters cannot be targeted. The effects observed maybe due to the lack of such elements, not the lack of transcription of the lncRNA of interest.

The decision in determining which system to use to knockdown *HMI-LNCRNA* was based on one key characteristic—its expression is almost entirely in the nucleus. siRNAs will not be effective as they stay in the cytoplasm. Therefore, shRNAs were used due to the fact that they integrate into the genome, and therefore are expressed in the

nucleus. Overexpressing *HMI-LNCRNA* also required its expression in the nucleus. Hence, full length *HMI-LNCRNA* was cloned into a lentivirus and used to transduce erythroid cells.

3.2 Downregulation of HMI-LNCRNA in erythroid cells

3.2.1 Determining which shRNA most effectively reduces expression of HMI-LNCRNA

Three different shRNA templates (HMIT-1, HMIT-2 and HMIT-3) were developed to target HMI-LNCRNA, and cloned into the pGreenPuro lentivector. pGreenPuro is an HIV-based expression lentivector, and contains multiple components for cloning, lentiviral production and transduction, and shRNA expression (see subsection 3.5.1 for details). pGreenPuro also expresses green fluorescent protein (GFP), which can be detected by flow cytometry to determine the percentage of cells infected with lentivirus. To determine which shRNA template most efficiently downregulates HMI-LNCRNA, each shRNA plasmid was transduced into TF-1a cells and cultured in puromycin for up to two weeks to select for only cells infected with lentivirus. At the time cells were harvested for analysis for expression of HMI-LNCRNA, about 95% of the cells were GFP-positive (GFP⁺). HMI-LNCRNA was reduced with all shRNA templates, with HMIT-1 showing the greatest reduction at 70% (Supplemental Figure 3.1). This experiment was repeated in HUDEP-2 cells, however the cells were analyzed two days after transduction without puromycin treatment. When cells were harvested, about 30% were GFP⁺, and a significant decrease in HMI-LNCRNA was found only in cells transduced with HMIT-1 (Supplemental Figure 3.2). Based on the observations from

qPCR analysis for *HMI-LNCRNA*, HMIT-1 shRNA was used for all knockdown experiments, and was labeled as HMI-lncRNA shRNA.

3.2.2 Downregulation of HMI-LNCRNA in HUDEP-1 and HUDEP-2 cells

HUDEP-1 and HUDEP-2 cells were used for knockdown of *HMI-LNCRNA* expression. Each cell line was split into three groups—naïve (non-transduced), scrambled shRNA and HMI-lncRNA shRNA. Each group had three biological replicates. Two weeks after the second transduction, cells were collected for the following analyses: flow cytometry for GFP; qPCR for *HMI-LNCRNA*, *HBS1L*, *MYB*, *HBG*, *HBB*, and *ACTB*; Western blot analysis for HBG, HBB, c-MYB; and slide preparations for immunofluorescent staining for HBG and HBB (**Figure 3.1A**). The first two experiments that were done to ensure that almost all cells were infected and that *HMI-LNCRNA* was significantly downregulated, and therefore used flow cytometry for GFP and qPCR for *HMI-LNCRNA*, respectively.

Flow cytometry results in HUDEP-2 cells showed that there were no GFP⁺ cells in the naïve group, 84% GFP⁺ in the scrambled shRNA group and 90% GFP⁺ cells in the HMI-lncRNA shRNA group (**Supplemental Figure 3.3**). qPCR results for *HMI-LNCRNA* showed there was about 50% reduction in the HMI-lncRNA group compared to both naïve and scrambled shRNA groups (**Figure 3.1B**). Almost the entire cell population of HUDEP-2 cells transduced with either scrambled shRNA or HMI-lncRNA shRNA was infected, and there was a significant reduction in of *HMI-LNCRNA*. Transduction of HUDEP-2 cells was repeated but with the addition of allowing the cells to mature. Both GFP-positivity and reduction of *HMI-LNCRNA* was similar to the first experiment.



Figure 3.1. Downregulation of HMI-LNCRNA in HUDEP-2 cells.

(A) Illustration of timeline for transduction and culture of HUDEP-2 cells. Cells were maintained in expansion medium. (B) Expression level of *HMI-LNCRNA* was determined by qPCR in HUDEP-2 cells that were not transduced (naïve), and transduced with either scramble shRNA or HMI-lncRNA shRNA lentiviruses by qPCR. *ACTB* was used as the endogenous control. p-values: *< 0.05; **< 0.005; **< 0.005; **< 0.005; NS (not significant). p-values were obtained by Student T-test.

In HUDEP-1 cells, no GFP⁺ cells were found in the naïve control group, 89% GFP⁺ cells were found in the scrambled shRNA group and 92% GFP⁺ cells were present in the HMI-lncRNA shRNA group (**Supplemental Figure 3.4A**). This result showed that almost all cells that were transduced expressed the shRNAs. However, there was no change in *HMI-LNCRNA* expression (**Supplemental Figure 3.4B**). This was probably due to the fact that these cells expressed relatively low levels of *HMI-LNCRNA* compared with HUDEP-2 cells, reducing any effect of shRNA knockdown. In further experiments, knockdown of *HMI-LNCRNA* was only done in HUDEP-2 cells.

3.2.3 The effect of knocking down HMI-LNCRNA on the expression of HBS1L and MYB

Since *HBS1L* and *MYB* are the two genes in proximity to *HMI-LNCRNA*, the mRNA levels of both genes in HUDEP-2 cells infected with HMI-lncRNA shRNA was assessed. qPCR analysis showed that there was no change in *HBS1L* and a significant 30% reduction in *MYB* compared with HUDEP-2 cells transduced with scrambled shRNA control, and this was also reflected in the protein level of c-MYB (**Figures 3.2A and 3.3A**). This change in *MYB* expression was reflected in one of two experiments. Pending additional studies, the regulation of *MYB* by *HMI-LNCRNA* is inconclusive.

3.2.4 Effects of knocking down HMI-LNCRNA on the expression of HBG and HBB

The expression pattern of both *HBG* and *HBB* among the naïve, scrambled shRNA and HMI-lncRNA shRNA groups were investigated by qPCR. Knocking down *HMI-LNCRNA* had no effect on *HBB* expression in one experiment and a 4-fold increase in expression in the second experiment. In contrast, there was a 200-fold increase in

HBG mRNA in HUDEP-2 cells transduced with HMI-lncRNA shRNA compared to both naïve and scrambled shRNA controls (**Figure 3.2A**). The percent *HBG* increased from less than 1% in naïve and scramble shRNA cells to more than 20% of the total *HBG* and *HBB* expression in cells with HMI-lncRNA shRNA (**Figure 3.2B**). Modulation of γ globin expression was also seen at the protein level, with detectable HBG expression only found in cells with knockdown of *HMI-LNCRNA* (**Figure 3.3B**). These results were corroborated with immunofluorescent staining using anti-HBG antibody tagged with a secondary red fluorescent antibody. The high HBG expression appeared to be restricted to a small subpopulation of HUDEP-2 cells with knockdown of *HMI-LNCRNA*. There was no difference in protein levels of HBA and HBB between the three groups (**Supplemental Figure 3.5**). In another experiment, *HBG* mRNA was also significantly elevated 900-fold with knockdown of *HMI-LNCRNA* compared with naïve and scrambled shRNA controls, with about 20% *HBG* of total *HBG* and *HBB* mRNA. This effect was also present at the protein level.



Figure 3.2. The effects of knocking down *HMI-LNCRNA* on the expression of *HBG*, *HBB*, *MYB* and *HBS1L* in HUDEP-2 cells.

(A) *HBG*, *HBB*, *MYB* and *HBS1L* transcript levels, and (B) percent *HBG* and *HBB* out of the total of both transcripts were measured in these same samples For qPCR analyses, *ACTB* was used as the endogenous control. p-values: *<0.05; **<0.005; **<0.005; **<0.0005; NS (not significant). p-values were obtained by Student T-test.



Figure 3.3. The effects of knocking down *HMI-LNCRNA* on the protein expression of c-MYB, HBG and HBB.

(A) Protein expression of c-MYB, HBG and HBB were analyzed by Western blot in naïve HUDEP-2 cells, and cells transduced with scrambled shRNA and HMI-lncRNA shRNA (HUDEP-1 cells were used as control). GAPDH was used as loading control.
(B) Naïve HUDEP-2 cells, and cells transduced with scrambled and HMI-lncRNA shRNAs were stained with anti-HBG antibody, followed by secondary antibody labeled with Alexa Fluor-594 (in red) and DAPI to stain nuclei (in blue), and imaged with a fluorescent microscope at 40X magnification.

In addition to analyzing the effect of downregulating *HMI-LNCRNA* in nondifferentiated HUDEP-2 cells, after the two-week selection for transduced cells with puromycin treatment, the cells were allowed to undergo erythroid maturation for up to seven days (**Figure 3.4**). *HBG* was significantly higher at Days 0, 5 and 7 in HUDEP-2 cells with *HMI-lncRNA* shRNA, and *HBB* was significantly higher at Day 0 and 7 with knockdown of *HMI-LNCRNA* (**Figure 3.5A**). *HBG* expression remained at about 20% of total *HBG* and *HBB* transcripts at each timepoint (**Figure 3.5B**). These observations were also seen at the protein level (**Supplemental Figures 3.6 and 3.7**).



HUDEP-2 cell culture

Figure 3.4. Timeline for infection and culture of HUDEP-2 cells.

Cells were maintained in expansion medium for 2 weeks after transduction, and then placed in differentiation medium for up to 7 days. Doxycycline (DOX) was removed at Day 5 to promote erythroid maturation.



Figure 3.5. *HBG* and *HBB* expression during erythroid differentiation of HUDEP-2 cells with knockdown of *HMI-LNCRNA*.

Cells were maintained in expansion medium for 2 weeks after transduction, and then placed in differentiation medium for up to 7 days. Doxycycline (DOX) was removed at Day 5 to promote erythroid maturation. (A) Relative quantity for *HBG* and *HBB* transcripts were analyzed by qPCR in naïve HUDEP-2 cells (n=3), and cells transduced with scramble shRNA (n=3) and HMI-lncRNA shRNA (n=3) at Day 0, 5 and 7 of differentiation. (B) Percent hemoglobin of *HBG* to *HBB* was determined by qPCR. For all qPCR analyses, *ACTB* was used as the endogenous control. p-values: *< 0.05; **< 0.005; ***< 0.0005; NS (not significant). p-values were obtained by Student T-test.

3.2.5 The effect of knocking down HMI-LNCRNA on the transcription factors that regulate the expression hemoglobin

As mentioned in Chapter 1, various transcriptions factors regulate hemoglobin gene expression. Since a significant change in γ -globin expression is observed with downregulation of *HMI-LNCRNA*, the expression pattern of a few of these transcription factors were studied in HUDEP-2 cells transduced with naïve and scrambled shRNA controls and HMI-lncRNA shRNA. The expression of *KLF1*, *BCL11A*, *ZBTB7A*, *CHD4*, *NR2C1*, *NR2C2* and *KDM1a*, all of which were known to downregulate *HBG*, were analyzed by qPCR. Expression of these genes was either unchanged or slightly upregulated in HUDEP-2 cells transduced with HMI-lncRNA shRNA compared to controls (**Figure 3.6**). This suggests that the significant increase in γ -globin gene expression when *HMI-LNCRNA* is reduced in HUDEP-2 cells is independent of the expression of the transcription factors analyzed by qPCR.

3.2.6 The effect of knocking down HMI-LNCRNA on erythroid maturation

HUDEP-2 cells are characterized as basophilic erythroblasts expressing both transferrin receptor (CD71), a cell surface marker for immature erythroblasts, and glycophorin A (CD235), a cell surface marker for more mature erythroblasts.³¹ The percentage of cells positive for these markers were examined by flow cytometry in naïve cells, cells expressing scrambled shRNA and cells expressing HMI-lncRNA shRNA (**Figure 3.7A**). Within both control groups, approximately 55-65% of the cells were positive for both CD71 and CD235, while about 85% of cells transduced with HMI-lncRNA shRNA were positive for both markers, which was significantly higher than

control groups (**Figure 3.7B**). These observations suggest that downregulation of *HMI-LNCRNA* promotes maturation of HUDEP-2 cells.



Figure 3.6. The effects of knocking down *HMI-LNCRNA* on erythroid-regulating transcription factors in HUDEP-2 cells.

Relative expression of *KLF1*, *BCL11A*, *ZBTB7A*, *CHD4*, *NR2C1*, *NR2C2* and *KDM1a* transcripts were analyzed by qPCR in naïve HUDEP-2 cells, and cells infected with scrambled shRNA and HMI-lncRNA shRNA. Beta actin was used as the endogenous control. p-values: *< 0.05; **< 0.005; ***<0.0005; NS (not significant). p-values were obtained by Student T-test.



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Figure 3.7. The effects of knocking down *HMI-LNCRNA* on erythroid maturation of HUDEP-2 cells.

(A) Naïve HUDEP-2 cells, and cells transduced with scrambled shRNA and HMIlncRNA shRNA (cultured in expansion medium) were stained with PE-labeled transferrin receptor (CD71) and PerCP/Cy5.5-labeled glycophorin-A (CD235) antibodies, and analyzed by flow cytometry to discriminate between cells that are positive and negative for GFP, CD71 and CD235. Data was analyzed with FlowJo. (B) Average percent GFP+, CD71+/CD235- and CD71+/CD235+ for each group was plotted on a bar graph. p-values: *< 0.05; **< 0.005; ***<0.0005; NS (not significant). p-values were obtained by Student T-test.

3.3 Overexpression of HMI-LNCRNA in erythroid cells

Knocking down HMI-LNCRNA significantly increases expression of HBG. To ascertain the effects of overexpressing HMI-LNCRNA, full-length HMI-LNCRNA was cloned into pLVX-EF1a-IRES-ZsGreen1 expression lentivector (ZsGreen lentivector), which co-expresses ZsGreen1, a fluorescent protein that can be detected by flow After successfully cloning HMI-LNCRNA in this ZsGreen lentivector cytometry. (ZsGreen HMI-lncRNA), it was used to transduce HUDEP-1 cells, which, unlike HUDEP-2 cells, express mostly HBG and have lower expression of HMI-LNCRNA. Unlike pGreenPuro lentivector, ZsGreen lentivector contains no genomic cassette for antibiotic selection in cell culture. Therefore, two days after transduction, cells were sorted for ZsGreen-positive (ZsGreen⁺) cells at the Flow Cytometry Core and cultured for two more weeks for analyses. Naïve cells and cells transduced with vector alone (ZsGreen) were used as controls. Similar to the knockdown experiments, the percent ZsGreen⁺ cells was checked by flow cytometry and showed 0% for naïve, 84% for ZsGreen and 88% for ZsGreen HMI-lncRNA (Supplemental Figure 3.7). qPCR analysis for expression of HMI-LNCRNA showed over a 2000-fold increase in expression compared to naïve and ZsGreen cells allowing further analyses as most of the cells were transduced with either ZsGreen or ZsGreen HMI-lncRNA, and there was significant upregulation of HMI-LNCRNA (Figure 3.8A).

We hypothesized that when *HMI-LNCRNA* is overexpressed in HUDEP-1 cells, *MYB* would increase and *HBG* will decrease. However, among the three groups of cells—naïve, ZsGreen and ZsGreen HMI-lncRNA—there was no difference in *HBG*, *HBB, HBS1L* or *MYB* (Figure 3.8B). It is not necessarily true that overexpressing a gene will give the opposite outcome to knocking down a gene. It is possible that other components that work with *HMI-LNCRNA* to regulate *HBG* expression are not expressed in HUDEP-1 cells, and therefore overexpressing *HMI-LNCRNA* will not show any effect.



Figure 3.8. The effects of overexpressing *HMI-LNCRNA* on the expression of *HBG*, *HBB*, *MYB* and *HBS1L* in HUDEP-1 cells.

Expression levels of (**A**) *HMI-LNCRNA* and (**B**) *HBG, HBB, MYB* and *HBS1L* were determined by qPCR in HUDEP-1 cells that were not transduced (naïve), and transduced with either ZsGreen or ZsGreen HMI-lncRNA lentiviruses by qPCR. *ACTB* was used as the endogenous control. p-values: *<0.05; **<0.005; ***<0.005; NS (not significant). p-values were obtained by Student T-test.

3.4 Summary and Conclusions

To determine the effect of modulating HMI-LNCRNA on neighboring proteincoding genes, HBS1L and MYB, and HBG and HBB in erythroid cells, we knocked down and overexpressed this lncRNA in HUDEP-2 and HUDEP-1 cells that produce primarily adult and fetal globins, respectively. To downregulate HMI-LNCRNA, HUDEP-2 cells were transduced with lentiviral particles that express an shRNA template targeting the IncRNA. After confirming successful transduction of cells and significant reduction of *HMI-LNCRNA*, we observed that these cells also displayed high upregulation of γ -globin at both the mRNA and protein levels, and a higher population of mature erythroid cells compared to cells in the naïve and scrambled shRNA control groups. Furthermore, there was consistent upregulation and percent HBG of total HBG and HBB during differentiation of these cells. Thus, these results suggest that HMI-LNCRNA regulates the expression of HBG and erythroid maturation (Figure 3.9). Overexpression HMI-LNCRNA in HUDEP-1 cells, despite a high level of cell transduction and increase in HMI-LNCRNA, was not accompanied by a change in the expression of MYB and globin genes.



Figure 3.9. Model for the function of *HMI-LNCRNA*.

HMI-LNCRNA is expressed from the enhancer region of *MYB*. Downregulation of *HMI-LNCRNA* increased the expression of *HBG* and promoted erythroid maturation. Further investigation is required to determine if *HMI-LNCRNA* also regulates the expression of *MYB*.

3.5 Materials and Methods

3.5.1 Development of shRNA plasmids for knockdown of HMI-LNCRNA

Three different shRNA templates were made to target three different areas of *HMI-LNCRNA*. This was accomplished by using Block-iT[™] RNAi Designer (Life Technologies), which analyzed the genomic region of *HMI-LNCRNA* and designed shRNA templates that are ranked based on the likelihood that it is effective. I used the top three ranking shRNA templates and labeled them *HBS1L-MYB* Intergenic Transcript 1 (HMIT-1), HMIT-2 and HMIT-3. Top and bottom strands of each shRNA were synthetically made (**Table 3.2**).

shRNA Template		
	Top strand	5'- GATCCGCTAGTATGTGAAGCACTTAGCTTCCTGTC AGACTAAGTGCTTCACATACTAGCTTTTTG-3'
HMIT-1	Bottom Strand	5'- AATTCAAAAAGCTAGTATGTGAAGCACTTAGTCTG ACAGGAAGCTAAGTGCTTCACATACTAGCG-3'
	Location (ch6, hg38)	135096650-135096670
	Top strand	5'- GATCCGCCGCTGCCTTTAATTGATGTCTTCCTGTCA GAACATCAATTAAAGGCAGCGGCTTTTTG-3'
HMIT-2	Bottom Strand	5'- AATTCAAAAAGCCGCTGCCTTTAATTGATGTTCTG ACAGGAAGACATCAATTAAAGGCAGCGGCG-3'
	Location (ch6, hg38)	135096889-135096909
	Top strand	5'- GATCCGCACCATCATGGTTTCAGTGCCTTCCTGTCA GAGCACTGAAACCATGATGGTGCTTTTTG-3'
HMIT-3	Bottom Strand	5'- AATTCAAAAAGCACCATCATGGTTTCAGTGCTCTG ACAGGAAGGCACTGAAACCATGATGGTGCG-3'
	Location (ch6, hg38)	135097331-135097351

TADIC J.1. SHIVIA COMPLETE IN THEFT LIVENIA	Table 3.1.	shRNA	templates	for	HMI-	LNCRNA
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Each shRNA template contained the following regions: sense, loop, antisense and terminator. Also, restriction sites for BamHI and EcoRI were placed on the 5'-end of the top strand and the bottom strand, respectively, to ligate shRNA templates to the

pGreenPuro expression lentivector (System Biosciences). Next, the top strand and bottom strand of each shRNA template were annealed to make double-stranded oligonucleotides by incubating both strands at 95°C in annealing buffer (100 mM Tris-HCL, 10 mM EDTA, 1 M NaCl and H₂O) for 4 minutes. The end products are doublestranded shRNA templates with restriction sites overhangs.

We next ligated each shRNA individually to pGreenPuro shRNA expression pGreenPuro is an HIV-based expression lentivector with the following lentivector. H1 promoter that is bound by RNA polymerase III for transcription of attributes: shRNA; EF1 α promoter that expresses both GFP and puromycin-N-acetyl transferase in transduced cells; RSV-5'LTR to express high levels of lentiviral particles in 293T cells; cPPT, GAG and LTRs, which are required for packaging, transduction and integration of viral components; ampicillin-resistance gene for selection in competent cells; and SV40 polyadenylation, which ensures termination of transcription, pGreenPuro lentivector was linearized by restriction digest with both BamHI and EcoRI, and then ligated to each of the double-stranded shRNA templates. After ligation, products were transformed into competent E. coli cells, grown overnight on LB plates treated with ampicillin, and colonies were selected and screened further by PCR to identify which clones had the The following primers were used for PCR: Forward (5'shRNA inserts. AATGTCTTTGGATTTGGGAATCTTAT-3') (5' and Reverse TGGTCTAACCAGAGAGACCCAGTA-3'), which would amplify about a 180 bp product if shRNA insert is present, and a 110 bp product is insert is not present. After positive clones were identified, each shRNA insert was confirmed by DNA sequencing.

pGreenPuro scramble shRNA (purchased from vendor, System Biosciences) was used as the negative control.

3.5.2 Development of plasmid to overexpress HMI-LNCRNA

Full-length HMI-LNCRNA was synthesized by IDT with an XbaI restriction site on the 5'-end and a BamHI restriction site on the 3'-end, and inserted into their pUCIDT-AMP vector. We next cloned HMI-LNCRNA into the pLVX-ER1a-IRES-ZsGreen1 (ZsGreen) expression lentivector. ZsGreen is an HIV-based expression lentivector that co-expresses ZsGreen, which is another green fluorescent protein. ZsGreen have the same components as pGreenPuro that are necessary to cloning, lentivirus production and integration, except there is so antibiotic cassette to select for infected cells in culture. Next, both pUCIDT-AMP with HMI-LNCRNA insert and the ZsGreen underwent restriction digest with XbaI and BamHI to isolate and purify HMI-LNCRNA insert and to linearize ZsGreen. Following digest, HMI-LNCRNA was ligated to ZsGreen, transformed into competent E. coli cells and cultured overnight on LB plates treated with ampicillin, and colonies were selected and screened further by restriction digest to identify which clones had the HMI-LNCRNA insert. DNA sequencing was utilized to confirm insertpositive clones, which was labeled as ZsGreen HMI-lncRNA. The ZsGreen vector alone was used as a negative control.

3.5.3 TF-1a cells

TF-1a cells are immortalized erythroblasts derived from the bone marrow of a patient with erythroleukemia.¹³⁵ Cells are cultured at 37°C in RPMI medium containing 10% FBS and 2% penicillin/streptomycin.

3.5.4 HUDEP cells

Human umbilical cord blood-derived erythroid progenitor (HUDEP) cells are immortalized erythroid cell lines derived from cord blood CD34⁺ mononuclear cells.¹³³ HUDEP-1 and HUDEP-2 cells were cultured at 37°C maintained in StemSpan SFEM medium (StemCell Technologies) supplemented with SCF (50 ng/ml, Invitrogen), EPO (3 U/ml, Invitrogen), dexamethasone (1 μ M, Sigma), doxycycline (1 μ g/ml, Clontech), Lglutamine (1%, Life Technologies) and penicillin/streptomycin (2%, Life Technologies). For erythroid maturation, cells were cultured at 37°C in IMDM medium (Invitrogen) supplemented with heat inactivated human serum from human male AB plasma (5%, Sigma), EPO (3 U/ml, Invitrogen), insulin (10 μ g/ml, Sigma), doxycycline (1 μ g/ml, Clontech), holo-transferrin (500 μ g/ml, Sigma), heparin (3 U/ml, Sigma), SCF (100 ng/ml, Invitrogen), L-glutamine (1%, Life Technologies) and penicillin/streptomycin (2%, Life Technologies) for 5 days. For further erythroid maturation, doxycycline was removed and cells were cultured for two more days.

3.5.5 Quantitative PCR

qPCR, also known as real-time PCR, is a tool used to quantitate the relative transcript levels of a specific RNA. In the PCR phase, gene-specific primers are used to

amplify the target during a 40-cycle reaction period (1-40). More abundant transcripts are amplified at earlier cycle thresholds (Ct) values, while less abundant transcripts are amplified at later Ct values. To calculate the relative quantity of a transcript-of-interest (TOI), we take difference between Ct of TOI and Ct of endogenous control, such as *ACTB* (β-actin), to get Δ Ct (Ct^{TOI} – Ct^{*ACTB*} = Δ Ct^{TOI}). Finally, the relative quantity of TOI is calculated using 2^{(-(Δ Ct))}. To calculate fold change between two samples—sample 1 (S1) and sample 2 (S2)—another approach is taken. For each sample, we calculate Δ Ct^{TOI} followed by calculating Δ ΔCt^{TOI} for each sample—for S1, [(Δ Ct^{TOI} for S1) – (Δ Ct^{TOI} for S1)] = (Δ \DeltaCt^{TOI} for S1) = 0; and for S2, [(Δ Ct^{TOI} for S2) – (Δ Ct^{TOI} for S1)] = (Δ ΔCt^{TOI} for S2). The fold change of the TOI for each sample is calculated using 2^{(-(Δ Ct))}, in which S1 will equal 1.

Total RNA, TaqMan primers and reagents from the TaqMan RNA-to-CT 1-Step Kit (Applied Biosystems) were used to make 20 μl reaction mix for each sample (done in triplicate) that was loaded onto a 96-well plate, following manufacturer's instructions. The following TaqMan gene expression assays (Applied Biosystems) were used: *HBG1/2* (Hs00361131_g1), *HBB* (Hs00758889_s1), *MYB* (Hs00920556_m1), *HBS1L* (Hs04188641_g1), *HMI-LNCRNA* (custom TaqMan assay designed by Applied Biosystems to target genome position chr6: 135096354-135097644, hg38; assay ID number AJI1MTQ), *BCL11A* (Hs01093197_m1), *CHD4* (Hs00172349_m1), *KLF1* (Hs00610592_m1), *ZBTB7A* (Hs00252415_s1), *NR2C1* (Hs00915957_m1), *NR2C2* (Hs00991824_m1), *KDM1a* (Hs01002741_m1) and *ACTB* (Hs01060665_g1). QPCR

reactions were ran on a StepOne Plus qPCR machine (Applied Biosystems). *ACTB* was used as the endogenous control.

3.5.6 293T cells

293T cells are a sub-line of 293 cells, which are immortalized human embryonic kidney cells. 293T cells are competent for replicating plasmids carrying SV40 origin of replication, and are used to produce high titers of pseudoviral particles such as lentivirus. They are cultured at 37°C in DMEM supplemented with 10% FBS and 2% penicillin/streptomycin.

3.5.7 HT1080 cells

HT1080 cells are derived from human fibrosarcoma, and are cultured at 37°C in DMEM supplemented with 10% FBS and 2% penicillin/streptomycin. HT1080 cells can efficiently be transduced with lentiviruses, and therefore are used to determine viral titer.

3.5.8 Generation of lentiviral particles

For shRNA experiments, 293T cells were transfected with scrambled shRNA and HMI-lncRNA shRNA using the LentiStarter 2.0 kit (System Biosciences), which uses pPACKH1 lentivector packaging system, following manufacturer's instructions, to generate lentiviral particles. For overexpression experiments, 293 T cells were transfected with either ZsGreen vector or ZsGreen HMI-lncRNA plasmid using the Lenti-X Packaging Single Shots, following manufacturer's instructions, to generate lentiviral particles.

Viral titer was determined using HT1080 cells— 1×10^5 cells were transduced with various amount of lentivirus using polybrene, and GFP-positivity was determined 48 hours later by flow cytometry. Transducing units per mL were determined using the following equation: (($1 \times 10^5 \times \%$ GFP-positive cells) x 1000)/Lentivirus volume (µL).

3.5.9 Lentivirus transduction

For shRNA experiment, HUDEP-1 and HUDEP-2 cells were transduced at an multiplicity of infection (MOI) 50 with 5 μ g/mL polybrene, and centrifuged at room temperature for 30 minutes at 1250 x g, and repeated again 48 hours later. 48 hours post-transduction, cells were cultured in expansion medium with 1 ug/mL puromycin for up to two weeks to select for transduced cells. After 2 weeks, FACS for GFP expression was done to ensure most cells were GFP-positive, and therefore express the shRNA plasmids. For overexpression experiments, HUDEP-1 cells were transduced with either ZsGreen or ZsGreen HMI-lncRNA at an MOI of 50 with 5 μ g/mL polybrene, and centrifuge at room temperature for 30 minutes at 1250 x g. Two days after transduction, cells were collected and sorted for GFP⁺ cells using flow cytometry. After sorting, cells were cultured in expansion medium for up to two weeks to provide sufficient cells for analyses. After 2 weeks, FACS for ZsGreen expression was done to ensure most cells were to many the provide sufficient cells for analyses.

3.5.10 Cell surface staining for FACS

 $5x10^5$ cells were harvested and washed in FACS buffer. Cells were resuspended in 100 µl of FACS buffer with antibody, and incubated on ice for 30 minutes. Next, cells were washed twice in FACS buffer by centrifugation at 300 x g for 5 minutes each. Finally, cells were fixed in IC Fixation buffer (eBioscience) for 30 minutes before analyzing the cells using the BD FACScan. The following antibodies were used: CD71-PE (334105, BioLegend) and CD235-PerCP/Cy5.5 (306613, BioLegend).

3.5.11 Western blot analysis

Culture cells were centrifuged at 300 x g for 10 minutes at 4°C, washed in cold 1X PBS and centrifuged again at 300 x g for 10 minutes at 4°C. Cell pellets were suspended in Roche lysis buffer (protease inhibitor, 0.3% NP40, 10% glycerine, 2 mM EDTA, 246 mM NaCl, 10% phosphatase inhibitor, PBS and water), placed on ice for 1 hour and centrifuged at 14,500 rpm for 15 minutes at 4°C to extract protein.

For Western blot analysis, protein lysates (10 µg) were denatured with lithium dodecyl sulfate (LDS; Life Technologies), reduced in sample reducing agent (Life Technologies) and incubated at 70°C for 10 minutes. Samples were ran on 4-12% Bis-Tris protein gel (Life Technologies) in 1X MED SDS running buffer and 1 ml NuPAGE antioxidant (Life Technologies) at 200 volts for 40 minutes. Protein from the Bis-Tris gel was transferred to a polyvinylidene difluoride (PVDF) membrane (Bio-Rad) in transfer buffer (2X NuPAGE transfer buffer, 10% methanol, 1 ml NuPAGE antioxidant and H₂O) at 100 volts for 1 hour in a transfer cell kept cold with an ice package and ice. After transfer, membranes were immersed in Ponceau S solution (Sigma) to determine if transfer was successful and to cut membrane evenly to probe with different antibodies that recognize proteins of different molecular weights. The following steps were taken to probe membrane with antibodies (all done on a rocker at 5-6 rpm at room temperature): wash in 1X PBS-tween 3 times for 5 minutes each; block in 1X PBS-tween with 5% milk

for 1 hour; wash in 1X PBS-tween; probe in primary antibody in 1X PBS with 5% milk for 2 hours; wash in 1X PBS-tween 3 times for 5 minutes each; probe in secondary antibody in 1X PBS-tween for 1 hour; and wash in 1X PBS-tween 3 times for 5 minutes each. Last, the membranes were treated with enhanced chemiluminescence (ECL) substrate for 5 minutes, and then exposed by chemiluminescence using a LAS-4000 Luminescent Image Analyzer at various exposure times. The following antibodies were used: c-Myb (ab109127, Abcam), hemoglobin β (16216-1-AP, Proteintech), hemoglobin γ (sc-21756, Santa Cruz Biotechnology) and GAPDH (sc-47724, Santa Cruz Biotechnology). For densitometry, ImageJ software was used to analyze western blots to determine percent intensity of each band, which was used to calculate the relative densities. Relative densities for c-Myb, hemoglobin beta and hemoglobin gamma were adjusted by GAPDH.

3.5.12 Slide preparation for microscopy

Cells were rinsed in PBS and resuspended in 2% BSA (in PBS). 200 μ L of cell suspension (5-7x10⁴ cells) were added to a cytology funnel in a slide holder and centrifuged in a cytospin at 800 rpm for 5 minutes. Cells were allowed to dry on slide for at least 30 minutes, and then fixed in 100% methanol for 10 minutes. Slides were stored at 4°C.

3.5.13 Wright-Giemsa staining

Fixed cells on slides were placed in diluted (1:3) Wright-Giemsa Stain (RICCA Chemical) for up to 1 hour. Slides were rinsed, dried and mounted with Permount

(Fisher). Images were taken in bright field with the Eclipse Ci (Nikon) microscope with a DS-Fi-U3 color-type (Nikon) camera, using NIS-Elements Br Imaging software (Nikon).

3.5.14 Immunofluorescence

Fixed cells were washed in PBS, blocked in 5% BSA (in PBS) and incubated in primary antibody for 1 hour. Cells were washed, blocked in 5% normal goat serum and incubated in secondary fluorescent-labeled antibody for 30 minutes. Finally, cells were washed and covered with mounting medium containing DAPI. Images were taken with the Eclipse Ci (Nikon) microscope with a DS-Qi1Mc-03 mono-type (Nikon) camera, using NIS-Elements Br Imaging software (Nikon). The following antibodies were used: Hemoglobin γ (sc-21756, Santa Cruz), Monoclonal Mouse Anti-human β -globin (a gift from IsoLab), Anti-human alpha globin (Wallac Inc.), and Alexa Fluor-594-conjugated AffiniPure F(ab)2 Fragment Goat Anti-mouse IgG (H+L) (115-586-062, Jackson ImmunoResearch).

CHAPTER 4. DISCUSSION

4.1 The *HBS1L-MYB* Intergenic Long Noncoding RNA (*HMI-LNCRNA*)

The *HBS1L-MYB* intergenic region on chr6q23 is important for regulating erythroid-specific traits. It contains an enhancer for the downstream gene MYB and binding sites for erythroid-specific transcription factors, which all bind to the promoter of MYB to increase its transcription. Furthermore, it is also known that upregulation of MYB increases erythroid cell proliferation, inhibits erythroid cell maturation, decreases HBG and increases *HBB* expression. Consequently, polymorphisms found within this region are associated with high HbF because they disrupt the enhancer-promoter interaction, thus reducing MYB expression and increasing expression of HBG and HbF. This doctoral work adds another element to this intergenic region, a novel 1283 bp lncRNA (chr6: 135096362-135097644, hg38) that is transcribed from the enhancer of MYB, and contains the HbF-associated polymorphic region rs66650371. Further characterization of HMI-LNCRNA revealed that it is not erythroid-specific but might be hematopoietic-specific; is located in the nucleus; is expressed at very low levels compared to the protein-coding gene *MYB*, and it is expressed at significantly higher levels in erythroid cells expressing HBB compared to cells that expressing HBG. The expression pattern of HMI-LNCRNA in these cells is similar to that of *MYB*.

Since *HMI-LNCRNA* is transcribed from an enhancer regulatory element for *MYB*, this suggests that it functions as an eRNA to support and promote binding of the erythroid-specific transcription factors to the promoter of *MYB*. However, additional experimentation will have to be done to determine if *HMI-LNCRNA* does work in

complex with these transcription factors. One method that can be used is chromatin isolation by RNA purification sequencing (ChIRP-seq).¹³⁶ ChIRP-seq is a relatively new approach to determine which genomic regions and proteins bind to a specific lncRNA. It crosslinks genomic DNA and proteins that are bound to lncRNA, adds biotinylated tilling oligonucleotides to bind to the lncRNA of interest, and then purifies out the complex by using magnetic beads to pull out the biotinylated oligonucleotides that are attached to the lncRNA. After purification, the complex can be dissociated and further analyses can be done to determine which genomic region and proteins bound to the lncRNA. If *HMI-LNCRNA* does function as an eRNA to promote the expression of *MYB*, we should find that it is associated with the promoter of *MYB*, and binds to TAL1/E47, RUNX1, GATA, LDB1 and KLF1. It is also possible for *HMI-LNCRNA* to regulate other protein-coding genes either in *cis* or *trans*, which ChIRP-seq will be able to elucidate.

We observed that the expression pattern of *HMI-LNCRNA* decreases during erythroid differentiation, as does the expression pattern of *MYB*. It will also be informative to determine the expression pattern of *HMI-LNCRNA* during the earlier phase of hematopoiesis, such as the HSCs, the common myeloid progenitors (CMPs) and the common lymphoid progenitors (CLPs) to determine if expression differs from that of erythroblasts. Since the expression of *MYB* is higher in HSCs, CMPs and CLPs, I would hypothesize that *HMI-LNCRNA* will be higher in these cells as well.²⁷ Additionally, *MYB* is not expressed during primitive erythropoiesis.² Therefore, it will also be of interest to determine if *HMI-LNCRNA* is not expressed in primitive erythroid cells.
4.2 HMI-LNCRNA and HbF

Increasing HbF levels in individuals with SCD is a major goal of clinical Hydroxyurea, which was the first FDA-approved drug for SCD, blood treatment. transfusions and gene therapy, are either not effective in all patients or financially and physically taxing. Therefore, it is vital to better understand how hemoglobin expression can be regulated in order to lead the field to more effective therapeutic treatments that will promote the expression of HbF. Various transcription factors such as BCL11A, KLF1 and LRF have been discovered to regulate HbF, and are considered possible therapeutic targets.^{31,137,138} Once thought of as "junk" RNA, lncRNAs are now considered to be important regulators of cellular processes, biomarkers for disease prognosis and possible therapeutic targets for disease treatment. This provides a rationale for further study of lncRNAs and erythroid ontogeny. Published data implicate lncRNAs as important regulators of hematopoiesis and erythropoiesis, however, before my studies, none have been implicated as regulators of hemoglobin expression. Therefore, it was very interesting and exciting to see that when HMI-LNCRNA was downregulated in HUDEP-2 cells, γ -globin expression increased both at the mRNA and protein levels. HUDEP-2 cells have been used to study the regulation of hemoglobin expression and erythroid maturation.^{31,137} However, additional functional analyses will be necessary in other cell culture systems, such as induced pluripotent stem cells (iPSCs)¹³⁹ and primary erythroid cells derived from CD34⁺ mononuclear cells, to ensure these results can be replicated in cells of different genetic backgrounds and normal cells directly from human tissue. Also, we can determine if the upregulation of HBG provides sufficient HbF to

inhibit polymerization of sickle hemoglobin after knockdown of *HMI-LNCRNA*, which will clinically benefit patients with SCD.

As mentioned in Chapter 3, gene editing is a tool used to stop transcription by removing genomic regions where genes are transcribed. In early work, investigators used gene editing to remove lncRNAs in order to determine their function. However, it has been discovered that removal of these genomic regions, which are mostly in intergenic regions, not only inhibits transcription of the lncRNA, but also removes regulatory regions that regulate the expression of neighboring genes. Paralkar et al determined if the lncRNA, Lockd, actually does regulate the transcription of its neighboring gene, Cdkn1b, as was suggested in the literature.¹⁴⁰ Therefore, gene editing was used in two different experiments to delete the entire *Lockd* gene, and to stop transcription of *Lockd* by placing a polyadenylation signal at its transcription start site. When *Lockd* is deleted from the genome, there was low expression of *Cdkn1b*. However, when a polyadenylation signal was inserted at the transcription start site of Lockd, thus inhibiting its transcription, *Cdkn1b* was transcribed normally. They further showed that *Lockd* resided within an enhancer region for *Cdkn1b*, and when the region was deleted, it reduced expression of However, if the genomic region was left basically intact and only the this gene. transcription of the lncRNA was inhibited by inserting a polyadenylation signal, there was no change in *Cdkn1b* expression. This suggested that *Lockd* did not regulate *Cdkn1b* and that previously published work was likely to be incorrect. A similar approach would have to be taken if we stop transcription of HMI-LNCRNA. Since shRNA does not

entirely knockdown *HMI-LNCRNA*, and since gene editing can fully curtail its expression, without residual lncRNA, a significant downregulation of *MYB* might occur.

4.3 Generating a Mouse Model for HMI-LNCRNA

If downregulation of *HMI-LNCRNA* can be confirmed to significantly increase HbF in primary erythroid cells to levels that could be therapeutically useful, the next step to consider is developing a mouse model for this lncRNA. Investigators have used various methods to determine the functionality of lncRNAs via mouse models that are similar to mouse models of protein coding genes. Xenograft models have been useful in determining if neoplastic cells with a specific genetic modification have the ability to grow and/or metastasize in immunocompromised mice.¹⁰⁷ Many lncRNAs have been discovered to be involved in the pathophysiology of specific cancers, and therefore this approach is frequently used. Another methodology is treating mice with antisense oligonucleotides (ASOs) that will bind to and thus reduce the amount of the specific lncRNA targeted.¹⁴¹ If ASOs were used for *HMI-LNCRNA*, this could be a preclinical model, which could possibly translate to the clinic as an approach for drug development for patients with SCD. Transgenic mouse models also have been useful for studying disease pathophysiology and treatment.¹⁴²

To study *HMI-LNCRNA in vivo*, there are two approaches to consider, and both include using the sickle cell transgenic mouse model.¹⁴³ The first approach is to treat the SCD mice with ASOs for *hmi-lncrna*, and determine if HbF increases. The second approach is to make an *hmi-lncrna* knockout mouse and cross it with an SCD mouse, and

determine if HbF increases in these mice, and if it is enough to inhibit polymerization of HbS.

There are a couple factors to consider before developing a mouse model for *hmi-lncrna*, and the first requires that it be expressed in the mouse. There is a lncRNA annotated in the mouse genome (Gm33728; chr10: 21228529-21252242; mm10) that is 23.7 kb in length. Human *HMI-LNCRNA* maps to an intron of Gm33728, and therefore they do not correspond to each other. However, it does not mean they do not have similar functions. There is no published data on Gm33728, therefore it will be of interest to determine if Gm33728 can regulate the expression of *myb* and hemoglobin as well. Another factor to consider is murine hemoglobin expression. Mice do not express HbF, but they do have embryonic hemoglobin, and it will be interesting to determine if Gm33728 regulates its expression.¹⁴³

4.4 HMI-LNCRNA as a Therapeutic Target

The severity of symptoms related to SCD is correlated not only to whether a patient express high levels of HbF, but also the HbF concentration in sickle erythrocytes and the distribution of HbF amongst F-cells.¹⁴⁴ Therefore, it is important to know if reduction of *HMI-LNCRNA* will increase HbF to levels that are beneficial to SCD patients. Knockdown of *HMI-LNCRNA* in HUDEP-2 cells resulted in an increase of *HBG* to 20% of the total *HBG* and *HBB* transcripts. Furthermore, this led to detectable protein expression of HBG. But will that ameliorate symptoms of SCD? HUDEP-2 cells also displayed a heterocellular expression pattern of HBG when *HMI-LNCRNA* was downregulated, which reflected a phenotype found in patients with $(\delta\beta)^0$ -thalassemia. As

mentioned in Chapter 1 (subsection 1.3.1), $(\delta\beta)^0$ -thalassemia and HPFH vary both in the concentration and distribution of HbF among RBCs.⁷⁰ These variations are the reason why patients with $(\delta\beta)^0$ -thalassemia are symptomatic and patients with HPFH are not. Therefore, the heterocellular distribution of HbF found in HUDEP-2 cells when *HMI-LNCRNA* was downregulated is not ideal when considering it as a therapeutic target. However, there were factors in the experiment that would have caused this uneven expression pattern of HbF. Transduction of HUDEP-2 cells with lentiviral particles is not homogenous. Therefore, not every cell expressed the same copy number of shRNA template. This suggests that cells with more copies of HMI-IncRNA shRNA expressed more HBG than cells with fewer copies. One way to address this issue is to undertake clonal expansion from a single cell after transduction.

4.5 Summary

The specific aims of my doctoral thesis was to characterize a transcript found within the *HBS1L-MYB* intergenic region on chr6q23, and to determine its function as it relates to the expression of *MYB* and hemoglobin. I discovered a novel 1283 bp lncRNA, named the *HBS1L-MYB* intergenic lncRNA (*HMI-LNCRNA*), which is transcribed from an intergenic region important for the regulation of erythroid-associated traits, and has been found to significantly increase γ -globin expression at both the mRNA and protein levels when downregulated by shRNA. Furthermore, due to this remarkable increase in γ -globin expression, *HMI-LNCRNA* can be considered as a possible therapeutic target for individuals with SCD and β -thalassemia where increased HbF is clinically beneficial.

SUPPLEMENTAL DATA

<u>3'-RACE Sequencing Results</u>



Supplemental Figure 2.1. Sequencing results for 3'-RACE PCR product.

Position 571 of sequencing results was the last nucleotide aligned on chr6q23 of NCBI Blast results, which is 135097644 (hg38).



Supplemental Figure 2.2. Sequencing results for 5'-RACE PCR product.

Position 32 of sequencing results was the last nucleotide aligned on chr6q23 of NCBI Blast results, which is 135096362 (hg38).



Supplemental Figure 2.3. Expression pattern of *MYB* among various human cells and tissue

Relative quantity of *MYB* was measured my qPCR in 25 different samples. Means are shown for samples with two or more independent samples. *ACTB* was used as the endogenous control.



Supplemental Figure 2.4. Hemoglobin expression in K562 cells treated with arginine butyrate.

K562 cells were either untreated (n=1) or treated (n=3) with 1 mM arginine butyrate for seven days. At day 7, cells were harvested to determine (**A**) cell color, (**B**) *HBG* mRNA expression by qPCR, and (**C**) HBG protein expression by Western blot analysis (left panel) and immunofluorescence staining (blue is DAPI, green is HBG). *ACTB* was used as endogenous control for qPCR analysis.



CD34⁺ Mononuclear Cell Culture

Supplemental Figure 2.5. Timeline for the expansion and differentiation of CD34+ cells derived cord blood and peripheral blood.



Supplemental Figure 3.1. *HMI-LNCRNA* expression in TF-1a cells transduced with three different HMI-lncRNA shRNA templates.

TF-1a cells were transduced with pGreenPuro (pGP) (n=1), and HMIT-1 (n=1), HMIT-2 (n=1) and HMIT-3 (n=1) shRNAs and cultured for two weeks in 8 μ g/ml puromycin for selection. qPCR analysis for *HMI-LNCRNA* was analyzed to determine fold change for each sample. *ACTB* was used as endogenous control.



Supplemental Figure 3.2. *HMI-LNCRNA* expression in HUDEP-2 cells transduced with three different HMI-lncRNA shRNA templates.

HUDEP-2 cells were transduced with pGreenPuro (pGP) (n=3), and HMIT-1 (n=3), HMIT-2 (n=3) and HMIT-3 (n=2) shRNAs and cultured for one week in 1 μ g/ml puromycin for selection. qPCR analysis for *HMI-LNCRNA* was analyzed to determine fold change for each sample. *ACTB* was used as endogenous control. p4 sample is non-transduced HUDEP-2 cells.



Supplemental Figure 3.3. FACS analysis for GFP expression in HUDEP-2 cells transduced with scramble shRNA and HMI-lncRNA shRNA.

Naïve HUDEP-2 cells (n=3), and cells transduced with scramble shRNA (n=3) and HMIlncRNA shRNA (n=3) were fix, and analyzed by flow cytometry based on SSC-H (side scatter for granularity) and expression of GFP^+ .



Supplemental Figure 3.4. GFP and *HMI-LNCRNA* expression in HUDEP-1 cells transduced with scramble shRNA and HMI-lncRNA shRNA.

(A) Naïve HUDEP-1 cells (n=3), and cells transduced with scramble shRNA (n=3) and HMI-lncRNA shRNA (n=3) were fix, and analyzed by flow cytometry based on SSC-H (side scatter for granularity) and expression of GFP^+ . (B) qPCR analysis for *HMI-LNCRNA* of HUDEP-1 cells from each group. *ACTB* was used as endogenous control. NS means not significant.



 VDEP-2 cells

 Vor
 Scramble shRNA
 HMI-IncRNA shRNA

 Image: Image:

Supplemental Figure 3.5. HBB and HBA expression.

В

Naïve HUDEP-2 cells, and cells transduced with scramble and HMI-lncRNA shRNAs were stained with (**A**) anti-HBB antibody and (**B**) anti-HBA antibody, followed by secondary antibody labeled with Alexa Fluor-594 (in red), and DAPI to stain nuclei (in blue), and imaged with a fluorescent microscope at 40X magnification.



Supplemental Figure 3.6. Protein expression of c-MYB and hemoglobin with knockdown of *HMI-LNCRNA* during erythroid differentiation of HUDEP-2 cells.

Naïve HUDEP-2 cells, and cells transduced with scramble shRNA and HMI-lncRNA shRNA were placed in differentiation medium for up to 7 days. Protein expression of c-MYB, HBG and HBB were analyzed in cells from (**A**) Day 0, (**B**) Day 5 and (**C**) Day 7 (HUDEP-1 cells were used as control). GAPDH was used as loading control. Densitometry was used to quantify western blot results (measured by ImageJ). p-values: *< 0.05; **< 0.005; **< 0.005; NS (not significant). p-values were obtained by Student T-test.



HUDEP-2 cells (Day 7 of differentiation)

Supplemental Figure 3.7. Protein expression of HBG with knockdown of *HMI-LNCRNA* at Day 7 of erythroid differentiation of HUDEP-2 cells.

Naïve HUDEP-2 cells, and cells transduced with scrambled and HMI-lncRNA shRNAs were placed in differentiation medium for 7 days. Cells were stained with anti-HBG antibody, followed by secondary antibody labeled with Alexa Fluor-594 (in red) and DAPI to stain nuclei (in blue), and imaged with a fluorescent microscope at 40X magnification.

LIST OF ABBREVIATED JOURNAL TITLES

Acta Biochim Pol	Acta Biochimica Polonica
Am J Clin Pathol	American Journal of Clinical Pathology
Am J Pathol	American Journal of Pathology
Ann Hematol	Annals of Hematology
Ann Hum Genet	Annals of Human Genetics
Ann N Y Acad Sci	Annals of the New York Academy of Sciences
Annu Rev Genet	Annual Review of Genetics
Arter Thromb Vasc Biol	Arteriosclerosis, Thrombosis, and Vascular Biology

Biochim Biophys Acta	Biochimica et Biophysica Acta
Biophys Chem	Biophysical Chemistry
Blood Cells, Mol Dis	Blood Cells, Molecules and Diseases
BMC Blood Disord	BMC Blood Disorders
Br J Haematol	British Journal of Haematology
Bull N Y Acad Med	Bulletin of the New York Academy of Medicine

Cell Death Differ	Cell Death and Differentiation
Cell Rep	
Cold Spring Harb Perspect Biol	Cold Spring Harbor Perspectives in Biology
Cold Spring Harb Perspect Med	Cold Spring Harbor Perspectives in Medicine

Eur J Biochem	European Journal of Biochemistry

Front Med	Frontiers in Medicine
Front Physiol	Frontiers in Physiology

Genes Dev	Genes and Development
Gene Ther	Gene Therapy
Genome Biol	Genome Biology
Genome Res	Genome Research

Hum Genet	Human Genetics
Hum Mol Genet	

Int J Hematol	International Journal of Hematology
Int J Mol Sci	International Journal of Molecular Sciences

JAMA	Journal of the American Medical Association
J Blood Med	Journal of Blood Medicine
J Clin Invest	Journal of Clinical Investigation
J Clin Lab Anal	Journal of Clinical Laboratory Analysis

Leuk Res	 Leukemia Research

Med Res Rev	
Methods Mol Biol	Methods of Molecular Biology
Mol Cell	Molecular Cell
Mol Cell Biol	Molecular and Cellular Biology
Mutat Res - Rev Mutat Res	Mutation Research/Reviews in Mutation Research

Nat Commun	Nature Communications
Nat Genet	Nature Genetics
Nat Med	Nature Medicine
Nat Methods	Nature Methods
Nat Rev Genet	Nature Reviews Genetics
N Engl J Med	

PLoS Genet	PLOS Genetics
PLOS Med	PLOS Medicine
PNAS	Proceedings of the National Academy of Sciences

Proc Nat Acad Sci USA	
Proceedings of the National Academy of Sciences of the United States of Americ	ca

Proc Natl Acad Sci	Proceedings of the National Academy of Sciences
RNA Biol	RNA Biology
Sci Signal	Science Signaling
Semin Immunol	Seminars in Immunology
Trends Biochem Sci	Trends in Biochemical Sciences
Trends Genet	Trends in Genetics
Trends Pharmacol Sci	Trends in Pharmacological Sciences

YALE J Biol Med	Yale Journal of Biology and Medicine
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CURRICULUM VITAE












