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# A therapeutic approach to chronic myeloid leukaemia using short hairpin RNA molecules

Al-Mazedi, Maryam

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# A Therapeutic Approach to Chronic Myeloid Leukaemia Using Short Hairpin RNA Molecules

Ву

Maryam S. Al-Mazedi

A thesis submitted in partial fulfilment of the requirements for the degree of Doctor of Philosophy at King's College London

The Institute of Pharmaceutical Science, School of Biomedical and Health Sciences, King's College London, Hodgkin's Building, Guy's Campus, London SE1 1UL

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### Abstract

Chronic myeloid leukaemia (CML) was one of the first cancers to be linked to a chromosomal abnormality, the Philadelphia chromosome. This chromosome results in a translocation between chromosomes 9 and 22, where the ABL gene on chromosome 9, a tyrosine kinase, is translocated to the BCR gene region on chromosome 22 giving rise to an abnormal BCR/ABL fusion gene. The resultant fusion gene has an abnormally upregulated tyrosine kinase activity that results in an increase in the proliferation of immature white blood cells, thus leading to the development of CML. There are several breakpoints that can occur in the BCR gene two of which give rise to 95% of CML cases. These fusion points are called the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  depending on where the chromosomal breakages occur in the BCR gene. The aim of the project was to establish a new method of treatment for CML through the use of RNAi to abolish the increase in tyrosine kinase activity of the abnormal fusion gene product.

The K562 and KCL22 cell lines incorporating the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  fusion points respectively were used in this project. The fusion points were cloned and sequenced. The human U6 and H1 promoters, were selected for the production of the antisense molecules and were obtained by PCR of K562 cDNA. Short hairpin RNA molecules were designed to the sequences of the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  fusion points. These designed shRNA molecules were synthesized as oligonucleotides and were incorporated into a reverse PCR primer. Cassettes containing shRNA molecules and a respective promoter were produced by means of PCR and the products cloned into pB12mcs-eGFP vector, which was used as a GFP reporter system. Constructs were then transfected into the appropriate cell lines, and expression studies including qPCR and Western blot analysis were conducted, to examine the effects of the designed shRNA constructs to their target sites on both mRNA and protein levels. In addition, these experiments also indicate the efficiency of the construct and also their specificity to their targets. qPCR and Western blot analysis, show that both the shRNA molecules designed against the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  fusion points, efficiently induced RNAi based gene silencing to their target sites. In addition, the designed constructs show high specificity to only its target sites and not to other unrelated or related genes. These results point towards the use of molecular modulation of gene expression as a promising strategy for potential CML therapy.

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# List of Abbreviations

Α	Adenine
AAV	Adeno-associated viral
A <sub>260nm</sub>	Absorbance at 260nm
Ab	Antibody/s
ALL	Acute lymphoid leukaemia
AML	Acute myeloid leukaemia
АТР	Adenosine triphosphate
bcr	Breakpoint cluster region
ВМ	Bone marrow
BSA	Bovine serum albumin
c	Cytosine
cDNA	Complimentary DNA
CLL	Chronic lymphoid leukaemia
CML	Chronic myeloid leukaemia
CrkL	Crk Like
DEPC	Diethylpyrocarbonate
cfu	Colony forming Unit
CML	Chronic Myeloid Leukaemia
CMV	Cytomegalovirus
DNA	Deoxyribonucleic acid
dNTP	Deoxyribonucleotide triphosphate
DMSO	Dimethylsulfoxide
dsRNA	Double stranded RNA
DTT	Dithiothreitol
E.Coli	Escherichia coli
ECL	Enhanced chemiluminescence
EDTA	Ethylenediaminetetraacetic acid
eGFP	Enhanced Green Fluorescent protein

FACS	Fluorescence-Activated Cell Sorter
FBS	Foetal bovine serum
G	Guanine
GAP	GTPase activating protein
GAPDH	Glyceraldehyde-3-phosphate dehydrogenase
GDP	Guanosine diphosphate
GEFs	Guanine nucleotide exchange factors
GFP	Green fluorescent protein
GMCSFR	Granulocyte monocyte colony stimulating factor
	receptor
Grb2	Growth factor receptor-bound protein 2
GTP	Guanosine triphosphate
H <sub>2</sub> O	Water
HEPES	4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid
hr	hour/s
INF	Interferon
IPTG	Isopropyl-β-D-thio-galactoside
IRS1	Insulin receptor substrate 1
JAK	Janus kinase
KDa	Kilo-Daltons
Kb	Kilobase
LB	Lysogeny broth
Max	Maximum
M-bcr	Major breakpoint cluster region
m-bcr	minor breakpoint cluster region
μ-bcr	micro breakpoint cluster region
MCR	Multiple Cloning Region
μg	Micro-gram
miRNA	MicroRNA
miRNP	MicroRNA protein complex
Min	Minutes
μΙ	Micro-Litre

M-MLV	Moloney Murine Leukaemia Virus
ng	Nanograms
nt	Nucleotides
OD	Optical density
PBS	Phosphate buffered saline
PCR	Polymerase chain reaction
PDK	Phosphoinositide-dependent kinase
PFA	Paraformaldehyde
pg	Picogram
Ph	Philadelphia chromosome
Ph⁺	Philadelphia chromosome positive
PIP2	Phosphoinositide-4, 5-triphosphate
PIP3	Phosphoinositide-3, 4, 5-triphosphate
РІЗК	Phosphatidylinositol 3-kinases
Pol III	Polymerase III
Pri-miRNA	primary miRNA
PRL13	Ribosomal protein large 13
PVDF	Polyvinylidene fluoride
qPCR	quantitative PCR
RISC	RNA-induced silencing complex
RNA	Ribonucleic acid
RNAi	RNA interference
Rpm	Revolutions per minute
RT-PCR	Reverse transcription polymerase chain reaction
SCID	Severe Combined Immunodeficiency
SD	Standard deviation
SDS	Sodium dodecyl sulfate
SDS-PAGE	Sodium dodecyl sulfate polyacrylamide gel
	electrophoresis
SH3	Src-homology 3
SHIP1	SH2 domain-containing inositol 5-phosphatase
shRNA	Short hairpin RNA

siRNA	Short interfering RNA
SOS	Son of sevenless protein
STAT	Signal transducer and activator of transcription
т	Thymine
TAE	Tris-acetic acid EDTA
TBS	Tris buffer saline
ТЕ	Tris-Ethylenediaminetetraacetic acid
TFO	Triplex forming oligonucleotides
ткі	Tyrosine kinase inhibitor
UPL	Universal probe library
UV	Ultra violet
WBC	White blood cell
w/v	Weight per volume
X-gal	5-bromo-4-chloro-3-indolyl-b-D-galactoside

### Chapter 1 : Introduction

### 1.1 Haematopoiesis and Leukaemia

Haematopoiesis is defined as the process of white blood cell (WBC) production. The process starts with the haematopoietic stem cell, then goes through several steps of differentiation and proliferation, involving several haemopoietic growth factors and cytokines. Mature WBC are then produced.

There are two main types of WBC, the myeloid or the phagocytic lineage and the lymphoid or the immunocytic lineage. The myeloid lineage gives rise to the monocytes and granulocytes, the latter includes the neutrophils, basophils and eosinophils, whereas the lymphoid lineage gives rise to the B- and Tlymphocytes (Hoffbrand *et al.*, 2006). Figure 1.1

Leukaemias or blood cancers are a group of malignancies that involve the haemopoietic system (Sobecks and Theil, 2003) in particular the WBC. In all leukaemias the WBC proliferation dramatically increases; however, differentiation decreases to a point where the bone marrow (BM) becomes hypercellular and immature malignant cells are seen in the peripheral blood.

Four types of leukaemia exist depending on the cell lineages affected, the types of malignant cells seen and the disease progression. These are the acute myeloid leukaemia (AML), the chronic myeloid leukaemia (CML), the acute lymphoid leukaemia (ALL) and the chronic lymphoid leukaemia (CLL) (Hoffbrand *et al.*, 2006). This project considers CML, therefore only this type of leukaemia will be discussed in detail.



Figure 1.1 Classification of the White Blood Cells (Hoffbrand et al., 2006).

A normal adult has 4 -11x10<sup>9</sup>/L WBC in their peripheral blood. WBC can be mainly divided into phagocytes and immunocytes. The phagocytes are further divided into the monocytes and the granulocytes. Monocytes are agranulocytic cells making up 4-6% of the total WBC count, while the granulocytes consist of neutrophils, eosinophils and basophils that make up 50-70%, 1-3% and 0.4-1% of the total WBC count respectively. The immunocytes, on the other hand, consist of the T- and B- lymphocytes that make up 25-35% of the total WBC count.

### 1.2 Chronic Myeloid Leukaemia (CML)

In 1845 Chronic Myeloid leukaemia (CML) was independently described and recognized by three German pathologists Virchow, Craigie and Bennett in Edinburgh, (Chopra *et al.*, 1999). However, it was not until the 1920s that the epidemiology of the disease was recognized (Chopra *et al.*, 1999). The incidence of CML is approximately 1 in 100,000 (Jemal A. *et al.*, 2002) with the disease affecting both sexes. However, it is slightly higher in males (Sobecks and Theil, 2003) and increases with age (Michor *et al.*, 2006). The median age of presentation of CML is between 50 and 60 years, nevertheless, 12-30% of cases are diagnosed above 60 (Kalidas *et al.*, 2001).

### 1.2.1 Definition of CML

CML is a myeloproliferative disorder that arises due to a clonal process involving the haematopoietic stem cells. The disease primarily affects the cells of the myeloid lineage, however, as the disease progresses it also affects the erythroid, megakaryocytic and lymphoid lineages (Faderl *et al.*, 1999).

The disease is characterised by having a severe increase in the WBC count, a mild to moderate anaemia and a hypercellular BM with a marked proliferation of all the granulocytic elements. The latter, leads to BM failure and cytopenias that will in turn lead to haemorrhage, recurrent infections and organ failure as the disease progresses to its advanced stages (Hoffbrand *et al.,* 2006).

### 1.2.2 Clinical Staging of CML

CML is known to be a triphasic disorder, and the chronic phase is the first phase of the disease. 50% of Patients are usually asymptomatic at this stage of the disorder. However, if symptoms persist, they can include fatigue, weight loss, abdominal fullness, bleeding, sweats, splenomegaly, hepatomegaly (Advani and Pendergast, 2002), anaemia, leukocytosis, and thrombocytosis (Faderl *et al.*, 1999). It is then followed by the accelerated phase and subsequently the blastic phase (Sawyers, 1999). The last two phases are sometimes joined together and considered to be advanced-phase CML. The duration of the three phases together, if untreated, has a median span of 2

to 5 years. Usually the chronic phase lasts from 2-5 years; the accelerated phase can last up to one year, whereas the blastic phase may lasts for 3-6 months (Sessions, 2007). Patients may be diagnosed at any of the three stages. Table 1.1 summarizes the main characteristics of each phase. 85% of Patients are usually diagnosed in the chronic phase and 50% of them are diagnosed from routine tests where patients are asymptomatic (Faderl *et al.*, 1999): they visit their physicians for other unrelated issues, and end up being diagnosed through an abnormally elevated WBC (Sessions, 2007).

Name of CML clinical phase	Main characteristics
Chronic phase	<ul> <li>Usually an asymptomatic phase. If symptoms persist, they may include splenomegaly, anaemia and hyperviscosity.</li> <li>This phase is easily controlled by conventional therapy.</li> <li>Lasts for an average of 3 years.</li> </ul>
Accelerated phase	<ul> <li>WBC doubling every five days, hard to control by conventional therapy.</li> <li>20% of blasts and promyelocytes are found in the blood and BM.</li> <li>30% basophils and eosinophils are seen in the blood.</li> <li>Splenomegaly in consistently increasing.</li> <li>Anaemia and thrombocytopenia are non- respondent.</li> </ul>
Blastic phase	<ul> <li>30% of blasts and promyelocytes are found in the blood and BM.</li> <li>Patients experience splenic and bone pain in addition to lymphoadenopathy.</li> <li>Median survival rate 3-6 months.</li> </ul>

Table 1.1 The characteristics of CML clinical phases. (Hoffbran et al., 2006).

#### **1.3** The Molecular Biology of CML

CML was one of the first cancers to be associated with a chromosomal abnormality (Sawyers, 1999). In 1960 Nowell and Hungerford noticed an abnormally small chromosome in patients with CML (Nowell, 2007). Further studies by Rowley J.D in 1973, using quinacrine fluorescent and Giemsa staining band technique, revealed that the abnormal chromosome in these patients was produced due to a balanced translocation between chromosomes 9 and 22. This new chromosome came to be known as the Philadelphia chromosome (Ph): named after the city where it was first discovered. The chromosome plays a significant role in the development of the disease and it is found to be present in 95% of all CML cases (Evans and Sillibourne, 1996 and Wertheim *et al.*, 2002). Patients with t(9;22) translocation are termed Ph positive (Ph<sup>+</sup>).

In 1982 the first aspects of the molecular biology of CML were subsequently discovered. In that year a research group in the USA showed that the gene on chromosome 9, which was translocated to chromosome 22, was the human *ABL* (Heisterkamp *et al.*, 1982 and Heisterkamp and Groffen, 2002) proto-oncogene (Advani and Pendergast, 2002). It was then found that this gene was translocated into a limited region, around 5.8 kilo bases (KB), on chromosome 22. This region was termed the breakpoint cluster region (bcr) (Groffen *et al.*, 1984). Hence, the name of the gene on chromosome 22 is now known as the *BCR* gene. Figure 1.2 shows the translocation between chromosomes 9 and 22.





The *ABL* gene located in chromosome 9 is translocated on chromosome 22, where the *BCR* gene is located via a reciprocal translocation. This results in the Philadelphia chromosome that bears the abnormal *ABL-BCR* fusion gene.

### 1.3.1 The ABL Protein

The normal *ABL* gene is composed of 11 exons (Laurent *et al.*, 2001), and transcribes the c-ABL protein. This 145 Kilo-Daltons (KDa) protein is a non-receptor tyrosine kinase. c-ABL has roles in assembling signalling proteins,

growth factor signalling, cytoskeletal organization, lipid metabolism, and transcriptional regulation (Chopra *et al.*, 1999). Two alternative first exons are present (1a and 1b) and depending on the alternative splicing of the transcribed RNA of the human *c-ABL* gene, two splice variants are present (Shtivelman *et al.*, 1986). Furthermore, the c-ABL type 1a is the spliced form of the protein and does not include a myristolation signal. In both variants, at the amino-terminus, a Cap region is present, then an Src-homology 3 (SH3) domain, followed by an SH2 domain, a tyrosine kinase domain, and finally a long carboxyl-terminal extension, which is also called the last exon region (Hantschel, 2004). The c-ABL protein also contains both nuclear localization and F-actin binding motifs. This explains the observations that the c-ABL protein can be seen in both the nucleus and the cytoplasm (Laurent *et al.*, 2001).

In addition, Pluk *et al.*, 2002, showed that the c-ABL protein is autoregulated, and the Cap, SH3, SH2 and the SH3-SH2 linker catalytic domain, play a role in autoinhibition of the normal c-ABL protein. The structural organization of this protein can be seen in Figure 1.3.



Figure 1.3 The Structural Organization of the c-ABL protein.

The c-ABL protein is 145 KDa, and consists of several structural motifs. These motifs include the Cap, the SH3, the SH2, and the tyrosine kinase domain. The protein is autoregulated and the kinase domain is autoinhibited in the normal protein.

#### 1.3.2 The BCR Protein

The *BCR* gene spans 130kb and is composed of 23 exons. The gene is found to have two alternative exons; an alternative exon 1 and an alternative exon 2, which are usually spliced out (Laurent *et al.*, 2001). The gene encodes a 160 KDa protein. The protein has several structural domains that are illustrated in Figure 1.4. From the N-terminus of the BCR protein the following domains are present; an oligomerization domain (Zhao *et al.*, 2002), a serine/threonine kinase domain (Maru, 1991), SH2-binding domains (Laurent *et al.*, 2001), and a GDP-GTP exchange domain (Chopra *et al.*, 1999). At the C-terminus, a domain showing homology to the GTPase activating protein, GAP domain, is present (Diekmann *et al.*, 1991). The first two domains are encoded from the first exon of the gene and are preserved in all BCR-ABL variants (Laurent *et al.*, 2001), as all breakages in the *BCR* gene occur downstream to exon 1.



Figure 1.4 The Structural Organization of the BCR Protein.

The BCR protein is composed of several structural motifs. These include the oligomerization domain, the serine/threonine domain and the GDP-GTP exchange domain on the N-terminus of the protein. At the C-terminus, the GAP domain is present.

The oligomerization domain (a coiled-coil motif) facilitates heterooligomerization with the BCR protein. In the abnormal CML fusion protein this

motif allows homo-dimerization of the fusion protein that is necessary for the activity (Faderl et al., 1999) and localization of the abnormal BCR-ABL protein. Deletions of this portion of the protein were found to decrease its binding to Factin, because a F-actin binding motif is found within the ABL portion of the protein. It is found that the oligomerization domain is responsible for the cytoplasmic localization of the fusion protein (Laurent et al., 2001) to one of the SH2-binding domains of the c-ABL protein. This interaction is thought to be essential for the oncogenic activation of the BCR-ABL protein (Laurent et al., 2001). The GDP-GTP exchange domain serves as an important control element for the RAS family, and is a GDP-GTP exchange factor for other GTP binding proteins, whereas the GAP domain is associated with the RAS-related protein family, RAC and RHO. The GAP domain controls the rate of GTP hydrolysis of the active RAS proteins to their inactive form, bound to GDP (Faderl et al., 1999). The structure of the BCR protein, therefore, makes it a multifunctional signal transducting molecule (Chopra et al., 1999 and Laurent et al., 2001). And because of the presence of the GDP-GTP exchange factor and GDP domains, see Figure 1.4, the protein seems to be especially concerned with phosphorylation, GTP binding of various proteins in several pathways that are involved in cytoskeletal organization, cell growth, and normal development (Laurent et al., 2001). These pathways are further discussed in sections 1.3.5.1-1.3.5.4.

In terms of BCR expression, the protein is constantly expressed in various tissues and cells, indicating its role in general cell metabolism rather than being specific to a particular tissue (Maru, 1991). The protein's mRNA levels were found to be highest in brain and hematopoietic cells. Interestingly, the BCR protein is expressed in the first stages of myeloid differentiation, and the levels of the protein decreases significantly with leukocyte maturation (Laurent *et al.*, 2001). Finally, the protein is usually cytoplasmic. However, it has also been found to be associated with condensed DNA.

#### 1.3.3 The BCR-ABL Fusion Gene

When the *c*-ABL gene is translocated from chromosome 9 to chromosome 22, the gene will be under the regulation of the BCR promoter (Shah et al., 1991 and Zhu et al., 1990). This translocation gives rise to an abnormal upregulated tyrosine kinase activity of the BCR/ABL fusion gene. The breakpoint within the *c*-ABL gene occurs within the first intron of this gene, leading to the fact that the resultant fusion transcripts always include the portion of the ABL gene from exon 2 onwards (Shtivelman et al., 1985). The BCR part of the fusion gene has three different points at which a chromosomal breakage can occur. Three breakpoints exist, and are called the Major breakpoint cluster region (M-bcr), the minor breakpoint cluster region (m-bcr) and the micro breakpoint cluster region ( $\mu$ -bcr) (Chopra *et al.*, 1999 and Melo, 1997); this is how the BCR gene got its name. Figure 1.5 demonstrates these breakpoints. The M-bcr spans a 5.8kb region within the BCR gene (Groffen et al., 1984). The breakpoints here span 5 exons historically named b1-b5, now known as exons 12 to 16 (Melo, 1996), and all breakpoints within this region are found to be intronic (Heisterkamp et al., 1985).



Figure 1.5 The Breakpoints within the BCR gene found on chromosome 22.

The position of the *BCR* gene is indicated by the dark transverse band shown on the long arm of chromosome 22. The gene is then further amplified to show the three points at which breakages may occur in the chromosome. These are the minor-bcr, the major-bcr and the micro-bcr breakpoints. The minor-bcr is located between exons 1 and 2, fusions to the *ABL* gene at this point give rise to the p190 fusion protein that is responsible for 30% of ALL cases. The major-bcr is between exons 13 and 15 and when fused to the *ABL* gene the p210 fusion protein results. This protein is responsible for 95% of CML cases. Finally, the micro-bcr is between exons 19 and 20, and fusions to the *ABL* gene at this point give rise to the protein ceutrophilic leukaemia. Note that exons 13 and 14 are also known as  $\beta$ 2 and  $\beta$ 3 respectively.

The M-bcr occurs between exons 13 and 14 or 14 and 15 giving rise to two variants of the p210 protein that are associated with 95% of CML cases. The mbcr occurs between the alternative exons 1 (1b and 1a) and exon 2. Breaks at this point give rise to the p190 protein that is associated with 15-30% of adult ALL and around 3-5% of childhood ALL. Finally, the  $\mu$ -bcr occurs between exons 19 and 20 of the *BCR* gene. The resultant protein is the p230 (Melo, 1997 and Van Der Burg *et al.,* 2004) and it is found to be associated with chronic neutrophilic leukaemia (Advani and Pendergast, 2002 and Quackenbush *et al.,* 2000). Figure 1.6 demonstrates the different BCR-ABL fusion proteins. Note that the numbering of each protein corresponds to their size in kDa (Pane *et al.,* 2002).



Figure 1.6 The BCR and ABL genes, breakpoints, and transcripts.

Depending on where the chromosomal breakage occurs in the *BCR* gene, four transcriptional variants can result. Breakages in the m-bcr region give rise to the e1a2 variant that is transcribed into a 190 kDa protein, p190. Breaks in the M-bcr region may give rise to the b3a2 or the b2a2 variants both of which transcribe a 210 kDa protein, p210. Finally, breakages in the  $\mu$ -bcr region give rise to the e19a2 variant that transcribes a 230 kDa protein, p230.

Each of the different BCR-ABL proteins is associated with a different type of leukaemia, as demonstrated above. This strongly suggests that the type of leukaemia, phenotype outcome, depends highly on the domains of the *BCR* 

gene included. The p190 is the smallest of the BCR-ABL proteins and is associated with cases of ALL whereas the p230, the largest of the BCR-ABL proteins, is associated with chronic neutrophilic leukaemia, which is considered to be a less aggressive form of leukaemia in comparison with ALL and CML (Melo, 1996 and Advani and Pendergast, 2002). Hence, the less the amount of the *BCR* gene included, the more aggressive the leukaemia.

The amount of the *BCR* gene being encoded in the BCR-ABL fusion protein correlates with the BCR domains present in the latter. Thus, in p190 protein only the oligomerization domain and the serine/threonine domains are present, whereas in the p210 protein, the previous domains in addition to the GDP-GTP exchange domain are present. Finally, the p230 protein includes all the domains present in the p210 protein and part of the GAP domain. This suggests the presence of a correlation between the structure of the BCR-ABL fusion protein and the disease phenotype (Melo, 1996). Figure 1.7 illustrates the relation between breakpoint cluster regions and the BCR domains. This project will only take into account the p210 BCR/ABL fusion protein due to its significant association with CML. Therefore only the M-bcr region will be further discussed.



Figure 1.7 Schematic representation of the structural motifs of the normal BCR protein and their association with the breakpoint cluster regions.

The red arrows show the positions of the three breakpoints. The first arrow represents the mbcr. Breakpoints at this position results in a p190 bcr-abl fusion protein that retains the oligomerization domain and the serine/threonine domain. The second arrow indicates the Mbcr. Breakpoints here give rise to the p210 BCR-ABL fusion proteins. These proteins include the domains present in the p190 protein and the GDP-GTP exchange domain. The p210 fusion protein is associated with CML and has two subsequent variants ( $\beta 2\alpha 2$  and  $\beta 3\alpha 2$ ). Finally, the third arrow labelled  $\mu$ -bcr points to the last breakpoint. Breakpoints at this position results in the p230 BCR-ABL protein. This protein has all the domains with the exception of the GAP domain, where only a part of it is present.

### 1.3.4 The p210<sup>BCR-ABL</sup> Fusion Protein

The M-bcr is around 5.8 kb in size (Pane *et al.,* 2002) and as figure 1.5 demonstrated the breakage occurs between exons 13/14 or 14/15, thus

retaining the coding region of the *BCR* gene from exon 1 to exon 13 or 14; the last two are also known as exons b2 and b3 respectively (Melo, 1996 and Chopra *et al.*, 1999). Moreover, breakages on the *ABL* gene occur between the alternative exons 1b and 1a, which are then spliced out.

The net result of the t(9;22) translocation is that the second exon of the *ABL* gene will fuse (at the gene's 3' end) to either exon 13 or 14 of the *BCR* gene (at it's 5' end) (Pane *et al.*, 2002 and Laurent *et al.*, 2001). Hence, giving raise to the two BCR-ABL alternative transcripts, the  $\beta 2\alpha 2$  and the  $\beta 3\alpha 2$  variants, with the second variant having a longer transcript. (Advani and Pendergast, 2002) Both variants encode the p210 BCR-ABL protein in which the ABL tyrosine kinase is constitutively active. This constitutive activation is the result of oligomerization domain of the *BCR* gene that promotes tetramerization of the fusion protein (Faderl *et al.*, 1999), thus allowing the Serine/threonine domain of the BCR protein to bind to the SH2 domain of the ABL protein leading to activation of it's the tyrosine kinase domain (Laurent *et al.*, 2001). Figure 1.8 demonstrates the BCR and ABL domains included in the p210 BCR-ABL protein.

On the basis of the more *BCR* sequence included in the protein, the less aggressive the leukaemia, mentioned earlier, scientists were interested to examine if this was true for the p210 variants (Mills *et al.*, 1988, Leibowitz *et al.*, 1991, and Shepherd *et al.*, 1995). The  $\beta 3\alpha 2$  being initiated from a breakage in the 3' end of the M-bcr is the longer transcript, whereas the  $\beta 2\alpha 2$  was produced due to breakages in the 5' end of the M-bcr yielding shorter transcripts (Melo, 1997). Studies that examined the presence of a correlation between the p210 BCR-ABL variants and CML disease prognosis had different results. Some studies found that a correlation does exists; those studies implied that the  $\beta 3\alpha 2$  variant had a better prognosis and a longer chronic phase than  $\beta 2\alpha 2$  (Mills *et al.*, 1988 and Leibowitz *et al.*, 1991), others found no evidence (Shepherd *et al.*, 1995). From this point onward the BCR-ABL protein will only refer to p210 variant.



Figure 1.8 Schematic representation of the structural motifs of the p210 BCR-ABL protein.

The BCR part of the p210 fusion protein is indicated by the red arrows and include the oligomerization, the serine/threonine, and the GEF domains. The ABL part of the fusion protein is indicated by the black arrows and include the SH2, the SH3, and the tyrosine kinase domains.

#### 1.3.5 BCR-ABL Activity in CML

How the BCR-ABL protein's activity leads to leukaemogenesis is of great interest. There is no single protein, or one signalling pathway that has been identified to cause leukaemogenesis. In fact a multi-network of signalling pathways and different proteins are involved in this process (Heisterkamp and Groffen, 2002). Therefore, the main pathways involved in leukaemogenesis of CML due to the deregulation of the *BCR-ABL* gene will be described below.

### 1.3.5.1 The RAS Pathway

This pathway is a central signal transduction pathway that transmits signals from several cell surface receptors to several transcription factors in the nucleus. Ras protein is a small protein that is a common molecule for several other signalling pathways. The protein is in its active form when bond to GTP, and is inactive when bound to GDP. Thus, the regulation of the RAS protein can be achieved by GTPase activating proteins and GEFs (guanine nucleotide exchange factors). Mutations that result in the continuous expression of the active RAS protein are observed in around 30% of cancers (Steelman *et al.*, 2004). The Ras pathway is demonstrated in Figure 1.9.



Figure 1.9 A Schematic diagram of the RAS pathway and the effects of the BCR-ABL on it.

Illustrated is the RAS pathway. Under normal conditions the RAS pathway will be activated through the activation of Shc via a stimulated cell surface receptor (not shown here). The Shc+ will recruit Grb2, activate it, and in turn Grb2+ activates the downstream GDP bound RAS to the GTP bound RAS. RAS+ will activate RAF that in turns activates MEK that will activate ERK. Finally ERK+ activates p90<sup>Rsk-1</sup> that will make its way to the nucleus and cause activation of various transcriptional factors. It should also be noted that RAF can translocate to the mitochondrial membrane and through the interactions with Bcl2 family, especially BAD, cause anti-apoptosis effects by inactivation of the pro-apoptotic protein BAD. Finally, the red arrows from the fusion protein itself demonstrate the points at which the BCR-ABL protein activates the RAS pathway.
For activation of RAS, a cell surface receptor that can be a growth factor, mitogen, or cytokine receptor, should be first simulated. This will lead to the Shc adaptor protein to be associated with the simulated receptor. The Shc adaptor protein will recruit the Grb2 (growth factor receptor-bound protein 2) and SOS (son of sevenless protein). This interaction leads to loading the membrane bound RAS with a GTP, hence its activation (Steelman *et al.*, 2011). The activated RAS protein undergoes conformational change that leads to the translocation of the RAF protein to the cell's membrane where it binds to the active RAS and will be activated by phosphorylation. The RAF protein is involved in proliferation, differentiation, apoptosis induction, and further to cytokine stimulation. In addition, over expression of this protein abolishes cytokine dependence in the haematopoietic cells.

Furthermore, the activated RAF will in turn activate MEK1 that in turn activates ERK1 and ERK2 through phosphorylation (Steelman *et al.*, 2011 and Steelman *et al.*, 2004). This results in phosphorylation of transcription factors, proliferation, and apoptosis inhibition (Steelman *et al.*, 2004), which is achieved directly through the activated ERK where it translocate to the nucleus phosphorylation several transcription factors that in turn bind to promoters of growth factors and cytokines. The activated ERK can further activate downstream p90<sup>Rsk-1</sup> that will influence gene expression (Steelman *et al.*, 2011).

Activated RAF protein can be translocated to the mitochondria (Hindley and Kolch, 2002) leading to the alteration of the mitochondrial distribution of another downstream protein called BAD (Steelman *et al.*, 2004), a pro-apoptotic protein that will be inactivated (Henson and Gibson, 2006). This alteration has been seen to prevent apoptosis in hematopoietic cells (Steelman *et al.*, 2004). Thus, the activation of this pathway plays a role in neoplasia and to be more precise in leukaemogenesis. In CML the BCR-ABL protein has direct effects on many of the proteins within the RAS pathway. The effects of the BCR-ABL protein is through direct activation of Grb2 (Puil *et al.*, 1994, Chopra *et al.*, 2011), RAS and RAF (Steelman *et al.*, 2004 and Steelman *et al.*, 2011).

See Figure 1.9. In summary, it can be concluded that the BCR-ABL fusion protein can play a role in CML development through the activation of the RAS pathway.

#### 1.3.5.2 The Phosphatidylinositol 3' Kinase Pathway

In CML, BCR-ABL protein is seen to activate the PI3K pathway (Zhu *et al.*, 2011, Naughton *et al.*, 2009, Steelman *et al.*, 2004, Chopra *et al.*, 1999, and Faderl *et al.*, 1999). Phosphatidylinositol 3' kinase (PI3K) is a protein made of 2 subunits a catalytic subunit (p110) and a regulatory subunit (p85) (Chopra *et al.*, 1999). The protein's activity is involved with cell division, cytoskeletal organization, and inhibition of apoptosis (Steelman *et al.*, 2004). The catalytic subunit of the protein is the part that interacts with receptor and non-receptor protein tyrosine kinases leading to the activation of PI3K (Chopra *et al.*, 1999). In addition PI3K can also be activated through RAS (Steelman *el al.*, 2004). Once activated, the active PI3K will act on its substrate PIP2 (phosphoinositide-4, 5-triphosphate) generating PIP3 (phosphoinositide-3, 4,5-triphosphate) that will lead to the activation of PDK (Phosphoinositide-dependent kinase) and Akt. In addition, Akt is also a substrate for the active PDK, hence it will be activated by that protein (Georgescu, 2010).

From this point the activated Akt protein will lead to the phosphorylation of several target sites, these will include the TSC2 and PRS40, GSK3, and FOXO. TSC2 and PRAS40 are two upstream inhibitors of RehB. RehB is a protein that activates mTORC1, which is responsible for protein synthesis in cells. When Akt phosphorylates TAC2 and PRAS40, it will lead to their inactivation, hence there will be no negative regulation of RehB and an increase in the activation of mTORC1 is established. The net result here is an increase in protein synthesis (Georgescu, 2010). Akt also negatively regulates GSK3, a protein that usually signals apoptosis. When Akt phosphorylates GSK3, the latter is inhibited leading to cell proliferation (Henson and Gibson, 2006 and Georgescu, 2010). Finally, FOXO proteins induce cell cycle arrest and apoptosis. They achieve this

through regulating the expression of various cell cycle regulators, and through up-regulation of pro-apoptotic members of the Bcl2 family. The active Akt protein again has a negative inhibitory effect on FOXO, thus leading to cell survival, proliferation, and apoptosis resistance (Georgescu, 2010). Figure 1.10 summarises the PI3K pathway.



Figure 1.10 Schematic Diagram of the PI3K pathway.

The PI3K pathway as demonstrated is rather a complex pathway. It is initiated by the activation of PI3K that in turn leads to a series of lipid-protein and protein-protein interactions. These interactions lead to the activation of the protein Akt. From here the protein negatively regulates further downstream substrates that usually inhibit cell growth, proliferation and protein synthesis. Akt negatively regulates downstream proteins leading to the net result of cell growth and anti-apoptosis. This complex pathway is activated in Philadelphia positive cells, hence, the proliferation and apoptotic resistance of these cells.

## 1.3.5.3 The JAK/STAT Pathway

CML also affects the JAK (janus kinase) family of kinases which are activated by signal transduction from cytokine receptors such IL-3 and GMCSFR (granulocyte monocyte colony stimulating factor receptor). Activated JAK protein will lead to activation of the STAT (signal transducer and activator of transcription) protein through phosphorylation. In turn the activated STAT proteins dimerize and move into the nucleus where they direct transcription and regulate gene expression (Constantinescu *et al.*, 2008) through binding to specific DNA elements (Henson and Gibson, 2006 and Chopra *et al.*, 1999). The down-regulation of the JAK/STAT pathway is achieved through the CIS/SOCS proteins. Where the CIS proteins inhibit STATs and the SOCS proteins bind and inhibit JAKs (Steelman *et al.*, 2004). The activity of the JAK/STAT pathway is highly increased in CML (Laurent *et al.*, 2001), and the BCR-ABL protein can directly activate STAT (Steelman *et al.*, 2004, Laurent *et al.*, 2001 and Noor *et al.*, 2011). This results in the growth factor independence observed in CML (Laurent *et al.*, 2001). Figure 1.11 gives a schematic diagram of this pathway.



Figure 1.11 A Schematic diagram of the JAK/STAT pathway.

The JAK/STAT pathway is activated in CML, this is due to the direct activation effects of the BCR-ABL fusion protein on STAT. giving a net results of increased transcription and growth factor independence observed in CML cells. CIS/SOCS proteins negatively regulate the pathway.

The JAK/STAT pathway has also been found to be activated in BCR-ABL negative myeloproliferative leukaemias, thus emphasising the important role of this pathway in leukaemogenesis. In these leukaemias, the activation of the JAK/STAT pathway is initiated through a mutated JAK, where the protein is constitutively activated and bypasses the need for cytokine receptor activation (Constantinescu *et al.*, 2008). Consequently, scientists have taken an interest in developing Anti-cancer drugs targeting this mutated protein (Knoops and Constantinescu, 2011 and Noor *et al.*, 2011). In summary, the JAK/STAT pathway is involved in leukaemogenesis, and due to the effects of the *BCR-ABL* gene on STAT, the pathway is continuously activate in CML.

#### **1.3.5.4 Other Proteins Associated with BCR-ABL Activity**

Other proteins that are associated with the activity of BCR-ABL can include Crk Like (CrkL) and SH2 domain-containing inositol 5-phosphatase (SHIP1) proteins. CrkL protein has been noted to be associated with the fusion BCR-ABL protein, however, how this protein plays a role in haematopoietic diseases and CML is still not understood (Sriram and Birge, 2010). In spite of this, the interaction between CrkL and the BCR-ABL protein is confirmed and the phosphorylation levels of CrkL are considered as a prognostic parameter for CML patients (La Rosee *et al.*, 2008 and Xu *et al*, 2011). A study done by Siriram *et al.* in 2011 has revealed that a tyrosine phosphorylation site on the CrkL is constitutively phosphorylated by BCR-ABL, this further confirms that the CrkL plays a role in CML; still more light is yet to be shed in this area.

SHIP1 protein has been shown to regulate haematopoiesis in mice and when disrupted leads to the development of myeloproliferative disorders (Laurent *et al.*, 2001). It is a negative regulator of the PI3K pathway by hydrolysing 5' phosphate of the PI3K substrate, PIP3, to its inactive form PIP2 (Ruschmann *et al.*, 2010). It has also been documented that the BCR-ABL protein inhibits SHIP1 (Steelman *et al.*, 2004) by tyrosine phosphorylation. This phosphorylation signals the proteasomal degradation of SHIP1 (Ruschmann *et al.*, 2010), thus abolishing its negative control on the PI3K pathway.

# 1.3.5.5 BCR-ABL and Protein Pathways: A Conclusion

In conclusion, the BCR-ABL has an upregulated tyrosine kinase activity. The BCR-ABL protein interacts with many other proteins that are involved in different pathways (Brehme *et al.*, 2009) leading to increased gene expression, protein synthesis, cell proliferation, differentiation, growth factor independence, and resistance to apoptosis. These effects all lead to the developments of leukaemic transformation and the development of CML.

The BCR-ABL activated pathways involve the RAS pathway, the PI3K pathway, the JAK/STAT pathway and other proteins such as CrkL and SHIP. These pathways are non-linear and are very complex as discussed above. The BCR-ABL protein can cause activation of these pathways at different levels and can also deactivate some negative regulatory events. This makes targeting the BCR-ABL fusion protein a highly appealing approach for CML therapy.

#### 1.4 CML and Therapy

Like all cancers, the treatment approach in CML is multiple sessions of chemotherapy via the usage of drug cocktails (Hoffbrand *et al.,* 2006). Drugs for the disease have been developed over the years, differing in their actions and target sites. Historically, drugs such as busulfan and hydroxyurea, which fall under the name of cytoreductive chemotherapy, were used to control the WBC of patients with CML. However, this kind of treatment has not stopped the disease progressing to the accelerated phase and the blastic phase (Fausel, 2007). Then interferon (IFN) was developed and used. This drug showed a complete haematologic response in 50% of patients. However, IFN's adverse effects, toxicity, cost effectiveness, and inconvenience limited its benefits. Therefore, new strategies of treatment were required (Fausel, 2007).

# 1.4.1.1 Current Drug Treatment for CML

With further understanding of the molecular pathology of CML, new drugs have been designed and developed. These drugs aimed at inhibiting the ABL tyrosine kinase activity (Milojkovic and Apperiey, 2008, Puttini *et al.*, 2008) therefore, are named tyrosine kinase inhibitors (TKI). Three generations of TKI exist: first generation TKI include Imatinib, which to date is the most popular drug for CML. Second generation TKI include Dasatinib and Nilotinib. Finally, the third generation TKI include Ponatinib and Bosutinib. Ponatinib is in clinical phase II (Vaidya *et al.*, 2011), while Bosutinib has just finished phase III clinical trials (Shieh *et al.*, 2011).

Imatinib, also known as STI571, is a drug made by Novartis and has a commercial name of Gleevec or Glivec (Druker, 2002). The drug works by competing with adenosine triphosphate (ATP) for its binding site to the *BCR/ABL* gene product. This inhibits the tyrosine kinase activity of the mutated gene, hence inhibiting its autophosphorylation, and subsequent activation of all its protein substrates that are involved in the various pathways discussed earlier. This will finally lead to the decrease in WBC proliferation and cell survival (Fausel, 2007 and Stentoft *et al.,* 2001). See Figure 1.12. Chronic phase patients on imatinib showed a 90% complete haematologic response. The drug initially showed promising outcomes, however, cases of drug resistance quickly emerged (Steinberg, 2007 and Talpaz *et al.,* 2006).



Figure 1.12 The mechanism of BCR-ABL inhibition by Imatinib.

In CML, BCR-ABL is able to phosphorylate downstream proteins when bound to ATP, and is therefore termed "active" in this state. The BCR-ABL protein switches from being active and inactive. When BCR-ABL is in its inactive state, Imatinib binds to the protein at its ATP binding site, locking the protein in its inactivated form, consequently, inactivating the protein pathways responsible for leukaemogenesis.

# 1.4.1.2 Mechanisms of Imatinib Drug Resistance in CML

Drug resistance to Imatinib can be classified into primary and secondary resistance. Patients with primary resistance illustrate a lack of effectiveness to the drug from the start of treatment. Those with secondary/acquired resistance show a loss in haematologic, cytogenetic and molecular response. The mechanisms by which resistance to Imatinib develop can be either BCR-ABL dependent or independent (Vaidya *et al.*, 2011).

#### **1.4.1.2.1 BCR-ABL Independent Mechanisms of Drug Resistance**

BCR-ABL independent mechanisms include the persistence of insensitive haematopoietic stem cells; these cells seem to be persistent due to their over expression of the BCR-ABL protein (Vaidya et al., 2011). Plasma enzymes inactivate Imatinib by binding and metabolizing it. This also demonstrates that the drug's plasma levels do not reflect the actual concentrations of the drug within the CML cells (Gambacorti-Passerini et al., 2003). Another BCR-ABL independent mechanism includes drug efflux. Drug efflux alters the intracellular availability of the drug. It has been shown that some CML patients resistant to Imatinib over express proteins, such as P-glycoprotein, responsible for imatinib efflux. When those patients were treated with P-glycoprotein inhibitor their sensitivity to Imatinib was increased (Jabbour et al., 2011). Finally, Imatinib resistance independent from BCR-ABL can also be due to the activation of the pathways implicated in CML due to mutations in the different proteins involved in these pathways. Many studies have demonstrated that mutations in proteins involved in any of the three main pathways involved in leukaemogenesis can independently lead to CML or a myeloproliferative disorder. Nambu et al., 2010, showed that one mechanism of independent resistance was the phosphorylation of ERK1 and 2. Scientists here developed a K562R cell line that is highly resistant to imatinib and had phosphorylated ERK1/2, STAT 5 and Akt proteins. When treated with Imatinib, the drug reduced the phosphorylation of all proteins with the exception of ERK1 and ERK2. A study done by Esposito et al., 2011, found that the levels of SHIP1 were decreased in cell lines resistant to imatinib, suggesting it as another mechanism for independent resistance. Another study showed that five cell lines, all resistance to Imatinib, had an increase in activity of the PI3K pathway due to a mutation in PI3K making the protein continuously active (Quentmeier et al., 2011).

#### 1.4.1.2.2 BCR-ABL Dependent Mechanisms of Drug Resistance

The BCR-ABL dependent resistance to Imatinib can be classified into three categories. Resistance due to mutations within the tyrosine kinase domain, outside the tyrosine kinase domain, and resistance due to over expression of BCR-ABL. Mutations within the tyrosine kinase domain all result in the inability of the drug to bind to its target site on the BCR-ABL protein (Vaidya *et al.*, 2011). An example is the T315I mutation that is the first BCR-ABL mutation to be discovered leading to Imatinib resistance (Gorre *et al.*, 2001). Mutations outside the kinase domain include mutations in the Cap, SH3, SH2, and SH3-SH2 linker domains. These domains are usually responsible for autoinhibition of the tyrosine kinase (Sherbenou *et al.*, 2010 and Vaidya *et al.*, 2011). Resistance due to over-expression of the BCR-ABL protein develops because the BCR-ABL protein is over expressed to a point where it can no longer be inhibited by the therapeutic dosage (Vaidya *et al.*, 2011).

As demonstrated a wide range of resistant mechanisms exist against the first generation TKI Imatinib leading to drug resistance in CML. Mutations leading to drug resistance can occur at any stage of the disease and is not limited to the accelerated or blast phase of the disorder (Roche-Lestienne *et al.,* 2002). Therefore, other TKI were developed. These include second and third generation TKI (Vaidya *et al.,* 2011).

## 1.4.2 Second and third generation TKI

New second generation TKIs for relapsed patients, resistant to Imatinib, have been developed. These include a drug called Dasatinib (Steinberg, 2007 and Talpaz *et al.*, 2006), developed by Bristol Pharmaceuticals in Switzerland, and FDA approved in October 2010 (Shieh *et al.*, 2011), and another called Nilotinib. The latter is developed by Novartis in the USA (Vaidya *et al.*, 2011),

and was FDA approved in June 2010 (Eskazan *et al.*, 2011 and Faber *et al.*, 2010 and Shieh *et al.*, 2011). These drugs were developed due to the cases of CML patient relapse. Nilotinib shows 10-25 fold more potency then the drug Imatinib, whereas Dasatinib shows between 100-300 fold greater activity. CML drug resistance to Nilotinib has also developed, and in some cases can be overcome by treatment with Dasatinib (Okabe *et al.*, 2011). Nevertheless, the first BCR-ABL mutation to give rise to imatinib drug resistance remains resistant to all three drugs discussed above (Skaggs *et al.*, 2006).

Third generation drugs, Ponatinib and Bosutinib, are in clinical phases with Bosutinib recently just completed phase III clinical trials (Shieh *et al.*, 2011). Data from the third generation TKI are yet to be collected and evaluated. Meanwhile, TKI available for CML treatment can be used for CML patients in any phase of the disease, with always a better prognosis when used in the chronic phase.

However, these drugs are only treatments not cures for CML. Patients tend to relapse, and the issue of drug side effects still remain. These will include toxicity, fluid retention, cardiac toxicity, platelet dysfunction and bleeding, immunosuppression, proinflammatory effects, and gastrointestinal side effects (Shieh *et al.*, 2011). The only cure available for this disease to date is allogeneic BM transplants (Fausel, 2007). Nevertheless, BM transplants are limited to younger patients (patients with CML are usually diagnosed between ages 50 – 60) and the availability of an appropriate HLA matching donor (Advani and Pendergast, 2002). Thus, making modulation of gene expression a promising strategy for potential CML treatment.

# 1.5 Modulation of gene expression

Various strategies exist for modulating gene expression, and expression can be increased, decreased or even abolished (Lavrovsky *et al.*, 1997). For the purpose of this study, only those that decrease or abolish gene expression will be considered.

Strategies that abolish gene expression can act at different molecular levels. These will include intervention at pre-transcription (DNA), post transcription (RNA), or post translational levels (protein). In past years, studies in down-regulating gene expression have focused on specific areas. These are, the formation of DNA triple helices, antisense, ribozyme, and RNA interference (RNAi) technologies. Each approach has its own advantages and disadvantages, depending on the nature of the pathological condition (Karagiannis and El-Osta, 2005 and Lavrovsky *et al.*, 1997).

# 1.5.1 DNA triple helices

DNA triple helices, also known as triplex forming oligonucleotides (TFO), are short oligonucleotides designed to be complementary to a regulatory region in a gene (Duca *et al.*, 2008). Hence the binding of the TFO to their specific target sites in the major grove of the double-stranded DNA, results in the formation of a triple helix (Arya, 2011). The resultant triple helix will block the binding of various DNA binding proteins, such as endonucleases, transcription factors, DNA methylating enzymes and polymerases, resulting in down regulation of gene expression (Lavrovsky *et al.*, 1997). One example of the use of TFO technology to target the fusion gene in CML is a study done by Rapozzi *et al.* 2002. In this study a 13-mer TFO targeting the BCR-ABL promoter region was designed. The TFO<sup>BCR-ABL</sup> showed down regulation of the BCR-ABL mRNA expression in K562 cell lines.

Although TFO are attractive tools for modulating gene expression, limitations do exist. These include the dependency of the presence of an oligopyrimidine/oligopurine in the DNA target site for binding, stability of the triple helix (Duca *et al.*, 2008), inefficient cellular uptake, insufficient nuclear internalization and oligonucleotide aggregation (Rapozzi *et al.*, 2002).

## 1.5.2 Antisense oligonucleotides

This technology involves molecules that modulate gene expression by targeting mRNA sequences, and uses antisense oligonucleotides of singlestranded DNA or RNA molecules. These sequences are 13-25 nt long, complementary to a specific site on the mRNA molecule of the gene of interest, and the binding results in the formation of sequence specific double helices. The use of synthesized phosphodiester oligonucleotides have proven to be of limited use. This is because these oligonucleotides are easily degraded by intracellular endo- and exo-nucleases. Therefore, attempts to chemically modify these synthesized oligonucleotides have been made with the overall purpose to gain more stable, nuclease resistant, high mRNA target hybridization affinity oligonucleotides. Dias and Stein reviewed these chemical modifications, in 2002 (Dias and Stein, 2002).

The double-helix formation may then down-regulate gene expression in two ways: one is by blocking the translation of the bonded mRNA molecule, and the other, is through the activation of cellular ribonuclease called RNase-H. This ribonuclease specifically cleaves RNA that is in a RNA-DNA hybrid. Theoretically, only the desired gene will be affected, making this technology an effective and powerful tool in gene expression modulation of various diseases and disorders (Dias and Stein, 2002 and Lavrovsky *et al.*, 1997).

The use of antisense oligonucleotides has been applied to target the mRNA of the *BCR-ABL* gene in CML. Though it was reported to be successful in reducing the mRNA expression in SCID mouse tissue, previously injected with Ph+ CML blast crisis cell line, and increased the duration of survival in treated mice (Agarwal and Gewirtz, 1999). Its clinical applications have been held back due to lack of stability, and the requirements of high dosages to achieve an anti-leukaemic effect. These dosages can also lead to unspecific binding and off-target toxicity (Quintas-Cardama, 2008).

#### 1.5.3 Ribozymes

Ribozymes are RNA structures with a specific catalytic activity. They are able to bind to other RNA molecules; thus breaking and/or forming covalent bonds with extraordinary specificity, thereby accelerating the rate of these reactions (Castanotto *et al.*, 2002).

Naturally occurring ribozymes act in a *cis* manner. These include hammerhead and hairpin ribozymes, that are derived from plants, RNase P (Kijima *et al.*, 1995), and group I and group II ribozymes that are found as introns in organelles and bacteria (James and Gibson, 1998). However, ribozymes can be also engineered to act in a *trans*-acting manner (Castanotto *et al.*, 2002 and Kijima *et al.*, 1995).

In the hammerheads, hairpins and group I introns, the specificity of a ribozyme to a particular target site depends on base-pairing between the ribozyme and its RNA target (Kijma *et al.*, 1995). In the RNase P ribozymes, it depends on the pairing of a guide RNA with the RNA target (Castanotto *et al.*, 2002), while in cases of group II introns it is contingent on the pairing of the ribozyme to its target DNA (Lewin and Hauswirth, 2001).

Once the ribozyme is attached to its target site, it will form a specific complex that will catalyse the hydrolysis of the phosphodiester bonds, hence cleaving the target RNA (Lavrovsky *et al.*, 1997). These properties make ribozymes useful tools in gene therapy (Kijma *et al.*, 1995) especially against HIV and cancer (Mulhbacher *et al.*, 2010). Ribozymes have been also previously used against the mRNA of the *BCR-ABL* gene. A reduction in the mRNA levels was observed in several studies (Lange *et al.*, 1994, Mendoza-Maldonado *et al.*, 2002), however, other studies didn't replicate these findings but showed nonspecific cleavage of the normal c-ABL mRNA (James *et al.*, 1996). Limitations of ribozymes exist, these include ribozyme stability, efficient target binding, co-localization of target and ribozyme, and specificity of delivery (Castanotto *et al.*, 2002 and James and Gibson, 1998).

#### 1.5.4 RNA interference

In 1998, Fire *et al.* observed that double stranded RNA (dsRNA) in *C.elegans* caused a substantially more effective RNA interference than using either an individual sense or anti-sense strand to modify gene expression. From that point on the process of RNA interference (RNAi) was discovered. RNAi quickly become evident as an ancient process. It is now known to be the oldest and most ubiquitous antiviral system in plants and animals, where RNAi is seen to be linked with post-transcriptional gene silencing in the above organisms (Sharp, 2001 and Elbashir *et al.*, 2001).

# 1.5.4.1 The Mechanism of RNAi

Basically, RNAi is the process whereby a double-stranded RNA (dsRNA) induces sequence specific post-transcriptional gene silencing. The dsRNA (>30 bp) does this by generating specific 21- and 22- nucleotide (nt) RNA fragments (Elbashir *et al.*, 2001). These fragments are generated by the cleaving action of a RNase-III family nuclease called "dicer", which is believed to be the first initiating step in RNAi (Bernstein *et al.*, 2001) and the fragments are called short interfering RNAs (siRNAs) (Cullen, 2002, and Elbashir *et al.*, 2001).

Dicer is associated with another protein called TRBP. The last recruits the RNA endonuclease Ago2 (Chendrimada *et al.*, 2005). Ago2 is also the catalytic component of the RNA-induced silencing complex (RISC) (Matranga *et al.*, 2005). The double stranded siRNA gets loaded onto Ago2 nuclease, and the passenger strand (sense strand) of the siRNA molecule is cleaved by the action of this protein. This cleavage occurs 9-10 nt counting from the 5' end of this strand (Rand *et al.*, 2005). This leads to the activation of the RISC complex that is guided by the siRNA guide strand (anti-sense strand) to its complementary target mRNA causing its degradation by the endonuclease activities of Ago2 (Matranga *et al.*, 2005).

In summary, the Dicer generated siRNA fragments are the mediators for RNAi and they do not exist independently. They are found associated with a number of proteins, which are still being studied in order to shed more light on their structure and how they contribute in the RNAi pathway (Wang et al., 2009). Together these molecules form complexes called silencing complexes that recognize their single-stranded mRNA targets in a highly sequence specific manner. The silencing complexes will then cause direct cleavage of their target sites, thus destroying any mRNAs that perfectly match the guiding siRNA strands. This will prevent the mRNA molecule(s) from being translated into protein. It should also be noted that even with a single nucleotide mismatch between the siRNA and its target, the effect is greatly diminished and may be even eliminated entirely (Dykxhoorn et al., 2003), making RNAi a highly precise and very specific tool. Figure 1.13 illustrates this process. Since the discovery of RNAi, this process has been used for various loss of function studies and other therapeutic approaches, which will be discussed later on. To date the application of RNAi has been mediated through synthetic siRNA, microRNA, and short hairpin RNA (shRNA) molecules.



Figure 1.13 Schematic diagram of RNAi mechanism.

dsRNA is recognised by the protein Dicer to which it binds. Once bound, the TRBP protein associated with Dicer recruits the RSIC complex and the dsRNA gets loaded onto the Ago2 protein within the complex. The endonuclease action of Ago2 cleaves the passenger strand, shown in blue, and the RISC is then activated. The guide strand, shown in red, directs the activated complex to its complementary mRNA. The antisense strand binds to its target mRNA and through the nuclease action of Ago2, the target mRNA will be cleaved 10-12 nt from the 5' end of the guide strand, thus RNAi is achieved.

#### 1.5.4.2 MicroRNAs

MicroRNAs (miRNAs) are a group of endogenously expressed small RNA molecules that are encoded by genes present in the host's genome. miRNAs do not lead to protein production, instead they act as regulators for mRNA expression. miRNAs are single stranded 21-23 nt long molecules that are produced by the cleavage of 70-90 nt shRNA precursor by the enzyme Dicer. This shRNA precursor is also known as pre-miRNA (Terasawa *et al.*, 2011). The miRNA molecules go on to associate with RISC that in turn induces translational repression by blocking the ribosomal translation (Karagiannis and El-Osta, 2005), and leading to inhibition of the translation of the target mRNA. Translational inhibition occurs when the miRNAs are partially complementary to the 3' untranslated region of their target sites. However, if the miRNAs are completely complementary to their targets, they will act in a way similar to siRNA molecules. Hence, by causing cleavage and degradation of their targets, see figure 1.14.

miRNA molecules have been identified in many multi-cellular organisms such as fruit flies, plants and humans (Pushparaj and Melendez, 2006 and Karagiannis and El-Osta, 2005). A study by Terasawa *et al.*, 2011, demonstrated that synthetic pre-miRNA based-shRNA had more potent RNAi activity than their corresponding siRNA molecules. In addition, they also observed that the guide strand of these pre-miRNA molecules was more efficiently incorporated into RISC. Thus, giving scientists further options in designing RNA molecules with more potent RNAi.

# 1.5.4.3 siRNAs

As stated earlier, siRNA molecules are the first discovered naturally occurring mediators of an RNAi response. They are 21-22 nt long and are the products of dsRNA precursors. Synthetic siRNA molecules have been

developed and used for loss of function studies or as therapeutic tools to modulate gene expression, of either an abnormal or an over expressed wild type gene (Dykxhoorn et al., 2003). These synthetic siRNA can be introduced exogenously or by the expression of viral or non-viral vectors (Pushparaj and Melendez, 2006), see figure 1.14. The classical siRNA structures proposed by Elbashir et al. in 2001 are siRNA molecules that are 19 nt in length with 3' overhang of 2 nt. A fixed length, symmetry in structure and the 3' overhang, were structural design rules to achieve optimum RNAi. However, these rules were observed in *Drosophila* embryonic extract. However, a recent review by Chang et al., 2011, shed light that siRNA structures used in mammalian cell lines can undergo more flexibility in design and chemical modifications based on the different siRNA structures that have been used with successful induction of RNAi to their target genes. Hence, the previously proposed fixed rules can be altered in mammalian cell lines, giving the advantage of developing siRNA molecules with more potent RNAi, reduced off target responses, and enhanced cellular delivery.

# 1.5.4.4 shRNAs

These RNA molecules are characterized by base-paired stems and a loop region, hence their name hairpin. shRNA is also processed by Dicer into siRNA that are 21 - 23 nt duplexes with a two base overhang at the 3' ends. The processing of such molecules yields mature products similar to the naturally occurring miRNA. Thus, competition between synthetic shRNA and naturally occurring miRNA for the endogenous RNAi machinery may result. However, to bypass this, the best approach is to select potent shRNA molecules capable of down-regulation of target gene when expressed in low levels (Rossi, 2008).

The use of synthetic siRNA is appropriate when transient inhibition of gene expression is satisfactory. However, in cases of prolonged gene silencing, several or continuous siRNA administrations will be required in order to maintain a long-term inhibition of gene expression. Expression vector systems have been developed where the DNA is transcribed and processed into short RNA within the cells, see Figure 1.14. In these systems the expression of shRNA molecules are favourable (Karagiannis and El-Osta, 2005). These systems usually use RNA polymerase III promoters (Pol III), especially the U6 small nuclear RNA and the H1 RNase P RNA promoters. These promoters are known for their simplicity, and strength of expression (Rossi, 2008). They do not add any extra bases that might affect the functional activity of the transcript, and their expression is well known to be constitutive and ubiquitous (Ohkawa and Taira, 2000). Hence, the combination of an expression vector system that uses a Pol III promoter to induce a high RNAi through the expression of shRNA molecules is a recommended system when long term gene silencing is required.



# Figure 1.14 The Mechanisms of Endogenous and Exogenous expression of miRNA, shRNA, and siRNA.

The process of RNAi can be achieved from either endogenous or exogenous triggers. The endogenous triggers of RNAi are the miRNA molecules. These are expressed from non-protein genes. miRNA are initially expressed as a long primary miRNA (pri-miRNA) that is further processed by an RNase-III type endonuclease enzyme called Drosha and its cofactor DGCR8 (Han *et al.*, 2004) into the miRNA precursors known as pre-miRNA. The pre-miRNA than exits the nucleus and gets processed by Dicer to imperfect 21- 23 nt long duplexes. These duplexes are taken up by RISC that cleaves the passenger strand and the mature miRNA directs the activated complex to its target mRNA where translational repression occurs. It should be noted that if the miRNA perfectly binds to its target mRNA it will leads to nuclease cleavage of its target. Exogenous RNAi is induced through the expression of shRNA constructs or the exogenous administration of synthetic siRNA. shRNA can be also administrated into the cells via the use of viral vectors. Dicer processes the shRNA molecules into siRNA duplexes that are loaded into RISC. Within RISC the passenger strand is cleaved and the active complex is directed to the target mRNA. Complementary binding between the guide strand and target leads to the cleavage of the latter and the achievement of RNAi.

## 1.5.5 The use of RNAi as a therapeutic tool

With the developments of siRNA and shRNA based technologies, several research groups have directed their work and interests in using the RNAi technology for therapeutic applications. These applications included viral infections, neurodegenerative disorders, cancer (Karagiannis and El-Osta, 2005), dominant gene disorders (Seyhan, 2011) and targeting downstream modulators of disease (Pushparaj and Melendez, 2006). Nevertheless, several limitations do exist for the use of RNAi as a therapeutic agent. These limitations include the stability of the siRNA molecules, and the major drawback of delivery into tissues in addition to the specificity of delivery.

To over come these limitations, various chemical modifications to the backbone of the siRNA molecules can be introduced. These modifications have been proven to increase the time life of the short RNA molecules. For the issues of delivery and specificity of siRNA/shRNA molecules, several delivery systems have been developed these can be mainly divided into viral and non-viral systems. Non-viral systems include polymer and lipid based systems. Here the short RNA molecules complexes with the transfecting reagent to form lipoplexes, dendriplexes, and polyplexes depending on the vector used, cationic lipids, dendrimer, or cationic polymers respectively (Shegokar *et al.*, 2011).

Viral vectors, on the other hand, have also been developed. These vectors provide sustained expression of their transcripts and have higher transfection efficiencies. Many vectors exist and the choice of the most suitable vector to be used highly depends on the therapeutic aim and tissue to be targeted. The most frequently used viral vectors will include adenoviral vectors, adenoassociated viral (AAV) vectors, retroviral vectors, and lentiviral vectors (Davidson and McCary, 2011). These vectors have the advantage of transducing both dividing and non-dividing cells. The adenoviral vectors are usually used for the delivery of shRNA or miRNA *in vitro*. However, they do not usually integrate into their target cell's genome. Thus, are used for short-term and single gene expressions. Three generations of adenoviral vectors exists. AAV vectors offer stable and high-level gene expression. They don't usually integrate into their host genome but exists as episomal forms. There are several stereotypes of AAV, thus giving a choice for selection of the most appropriate type depending on the cells of interest. Retroviral vectors on the other hand, yields long-term expression of the transcript of interest. This is because the viral RNA is reverse transcribed into double stranded DNA upon entry into the host cells. The transcribed DNA will then integrate into the host's chromosome. Nevertheless, with time, persistent gene expression may not be sustained; this maybe due the occurrence of silencing transcriptional units. Another drawback in retroviral vectors is the risk of T-cell leukaemia development that has been previously observed in patients. Finally, lentiviral vectors, a subgroup of retroviruses, can produce persistent transgene expression, which can also be silenced over time. These vectors permit the integration in their host's genome within intronic regions of active transcriptional units. (Liu et al., 2011).

With all the recent developments in RNAi, the technology has been rapidly improving since its discovery in 1998. Over the years many studies have been now approved and are in pre-clinical and clinical trials. In addition some RNAi therapeutic approaches have already completed phase II. These studies cover a wide range of diseases and disorders. Their clinical setting for RNAi therapy include 7 drugs for ocular and retinal disorders, 8 drugs for different types of cancers, 3 drugs for kidney disorders, 2 drugs for LDL lowering, and 5 drugs for antiviral infections (reviewed by Davidson and McCary, 2011). These are promising developments for RNAi as a constantly evolving technology to become the therapeutic tool of the future.

#### 1.5.6 RNAi and CML

CML was characterized as the first cancer to be linked with a chromosomal abnormality that is constant in 95% of CML cases. The disease results mainly from overexpression of the c-ABL tyrosine kinase activity. For many years the BCR-ABL abnormal protein have been an attractive site for modulators of gene expression. These include antisense oligonucleotides, ribozymes and RNAi, including both siRNA and shRNA molecules.

In this project, gene modulation of the expression of the abnormal BCR-ABL gene through RNAi is of interest. Several studies have attempted to target the BCR-ABL chimeric protein through RNAi by the use of either synthetic siRNA or shRNA molecules. These include studies done by Mahmodabady et al., 2010, Myssina et al., 2009, Li et al., 2003, and Arthanari et al., 2010 for shRNA molecules, and Rapozzi and Xodo, 2004, Wohlbold et al., 2003, and Bártová et al. 2005 for siRNA molecules. The studies using shRNA molecules targeted both fusion points,  $\beta 3\alpha 2$  (Mahmodabady *et al.*, 2010, Myssina *et al.*, 2009, Li *et al.*, 2003, and Arthanari et al., 2010) and the  $\beta 2\alpha 2$  (Li et al., 2003). The studies used chemical and physical methods of transfection, and different shRNA sequences that were regulated and expressed from different promoters and vector systems. However, all publications reported a decrease in both mRNA and protein levels after shRNA treatment of only the  $\beta 3\alpha 2$  variant. The mRNA and protein levels of the  $\beta 2\alpha 2$  variant, however, were not affected (Li *et al.*, 2003). The findings of these publications will further be discussed in detail in chapter 5.

Other studies have also considered RNAi technology either as a therapeutic tool for CML, through targeting proteins down stream BCR-ABL, or to understand the many pathways involved in leukaemogenesis by loss of function analysis. Examples include a study done by Machado-Neto *et al.* 2011, that used an shRNA-lentiviral delivery system to study the knockdown effects of a protein called IRS1 (insulin receptor substrate 1) on the PI3K-Akt/mTOR and MAPK pathways in K562 cell lines. This study was done on the basis that IRS1 is continuously phosphorylated and associated with BCR-ABL in K562 cell line (Traina *et al.*, 2003). Silencing of IRS1 did down-regulate the associated pathways, causing a decrease in cell proliferation and colony formation, but apoptosis was not affected.

Another study conducted by Albers et al. 2011, demonstrated the use of RNAi based systems for loss of function analysis. The group found that Raf1 protein is a crucial mediator of BCR-ABL dependent CML, through activation of the MAPK/ERK cascade. In this study the shRNA was expressed from a retroviral vector in a coupled expression system. Another study done by Scherr et al. 2006, have also demonstrated that SHP2, STAT5 and Grb2 proteins can be therapeutic targets in cases of drug resistance. They have identified these target proteins through the use of a lentiviral mediated RNAi. The study also demonstrated that co-expression of shRNA against the BCR-ABL and SHP2 proteins from a single lentiviral vector induces stronger colony formation inhibition when compared to using shRNA alone. Elmaagacli et al. 2005, found that the use of two siRNA molecules had additive effects in the induction of apoptosis in CML cells, in comparison to when each siRNA was used individually. In this study two synthetic siRNAs were co-transfected. One siRNA was against the BCR-ABL, while the other targeted the Wilms' tumor genes, a gene found to be abnormally over-expressed in leukemic cells.

Other studies have found that the use of RNAi technology can sensitize the drug resistant leukaemic cells to TKI, such as Imatinib and Nilotinib. These studies include those done by Wohlbold *et al.*, 2003, Rumpold *et al.*, 2005, and Koldehoff *et al.* 2010, respectively. In these studies the RNAi mediated knockdown was against the BCR-ABL transcripts, giving a basis to combining both RNAi technology and TKI, thus targeting the *BCR-ABL* fusion gene molecularly at both mRNA and protein levels. A study conducted by Mendonca *et al.* 2010, showed that siRNA against the *BCR-ABL* gene could be co-encapsulated with Imatinib giving a new therapeutic approach for CML and at the same time bypassing the effects of drug resistance.

In conclusion, since the discovery of RNAi by Fire *et al.* in 1998, the technology has been advancing at a high speed. A lot has been unravelled

about RNAi mediated gene silencing, the proteins involved in the process, and the different mechanisms of the different triggers of RNAi. The technology is rapidly becoming a standard tool for loss of function studies and an appealing therapeutic approach for a wide range of diseases and disorders including cancer in general and targeting leukemic fusion proteins in particular (Thomas *et al.*, 2006).

#### 1.6 Cell Lines as a Model for CML

Since the establishment of the first CML cell line in 1975 by Lozzio and Lozzio, many other CML cell lines have been established. There are around 30 CML cell lines as listed in a review by Drexler in 1994, some of which express the  $\beta 3\alpha 2$  BCR-ABL variant, whereas others express the  $\beta 2\alpha 2$  fusion protein. This wide range of CML cell lines differ morphologically and phenotypically, giving scientists the opportunity to study the pathobiology of CML and the role of the BCR-ABL protein in the development of leukaemogenesis (Drexler *et al.*, 1999).

In 1975 the first CML cell line, the K562 cell line, was established (Lozzio and Lozzio, 1975). These cells express the  $\beta 3\alpha 2$  BCR-ABL variant that is a free-floating suspension cell line (Lozzio and Lozzio, 1975). Cytologically, K562 cells are composed of undifferentiated myeloid cells (lozzio and Lozzio, 1979) and molecularly, are a triploid cell line (Drexler *et al.*, 1999). K562 cells were used over the years since its establishment for various studies including those that shed light on the nature of *BCR-ABL* fusion gene (Heisterkamp *et al.*, 1983), or those that used the cell line for gene modulation (Wilda *et al.*, 2002, Rapoozi *et al.*, 2002, Mahmodabady *et al.*, 2010, and Arthanari *et al.*, 2010). Because the cell line is easy to grow and maintain, in addition, molecularly they express the *BCR-ABL* gene in an oncogenic amplification manner (20 – 30 fold), this cell line provides a suitable CML model for gene modulation studies; therefore, the K562 cell line was selected for the purposes of the current study.

The KCL-22 cell line, on the other hand, expresses the  $\beta 2\alpha 2$  BCR-ABL variant. Kubonishi and Miyoshi, 1983, first established this cell line in 1983. KCL-22 cells are also free-floating in suspension, cytologically the cells are composed of immature undifferentiated haematopoietic cells; and molecularly it is a diploid cell line with double Ph chromosome, in addition to a normal chromosome 9 and 22 (Kubonishi and Miyoshi, 1983). The double Ph chromosomes are not surprising as this cell line (and K562) was established from a CML patient in blast phase. In the CML blast phase/crisis other chromosomal abnormalities occur including trisomy #8, isochromosome of the long arm of chromosome  $\pm 17$ , and double Ph chromosomes are observed (Kubonishi and Miyoshi, 1983, Drexler, 1994, and Drexler et al., 1999), hence, actually giving evidence of the leukaemic origin of the KCL-22 cells. Rapozzi and Xodo, in 2004 used the KCL-22 cell line as a leukaemic model in attempt to down-regulate the expression of the  $\beta 2\alpha 2$  BCR-ABL by siRNA molecules, and Ohmine et al., 2003, used these cells to establish a new Imatinib-resistant cell line, KCL-22/SR in 2003. A publication by Esposito et al., in 2011 used a KCL-22/R (imatinib resistant) cell line to study the mechanisms of imatinib resistant in CML. The cells are also easy to grow and maintain, and have a rate of doubling every 24hr (Kubonishi and Miyoshi, 1983), thus, for the purpose of this study, this cell line was chosen as it serves as an appropriate model for CML expressing the  $\beta 2\alpha 2$  BCR-ABL fusion protein.

# **1.7** Aim of the project

As previously mentioned, RNAi is becoming a tool for therapeutic approaches against several diseases and disorders, including cancer. CML was the first cancer to be characterized by the presence of an abnormal Philadelphia chromosome that transcribes an abnormal protein through as the *BCR-ABL* oncogene. This abnormal fusion gene has two variants; the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$ , both playing an important role in leukaemogenesis of CML. TKI are currently the first line of treatment for CML. However, first and second generations of TKI have failed to overcome the development of leukaemic drug resistance in relapsed patients, making modulation of the expression of the *BCR-ABL* oncogene a favourable approach. Using RNAi mediated gene silencing of the *BCR-ABL* oncogene through stable expression of shRNA<sup>BCR-ABL</sup> would be an excellent therapeutic approach for CML. The combination of two or more shRNA, targeted against one fusion variant or both together (in some cases of CML both fusion variants are expressed simultaneously (Adler *et al.*, 2009)), would be another therapeutic approach. This is based on the studies mentioned earlier where siRNA molecules had additive effects on CML. Finally, shRNA molecules against CML can also be used to sensitize leukemic cells to TKI, or the shRNA constructs can be administrated with TKI to establish complete hematologic response for patients with CML.

This project aims to decrease or abolish the expression of the two BCR/ABL variants. It aims to accomplish this by inducing RNAi through designing shRNA molecules against the two BCR-ABL transcripts. The designed molecules were then expressed and delivered via a non-viral system in two CML cell lines; one for each variant. The shRNA molecules to be expressed were designed to be as symmetrical as possible, with an 8 nt long loop, and were under the regulation of the human Pol III promoters, U6 or H1. The efficacy and specificity of the shRNA constructs were established as a prelude to these molecules being incorporated in a lentiviral system that would be tested on cell lines, and on primary CML cells from patient samples *in vitro*.

# Chapter 2 : Materials and Methods

# 2.1 Molecular Biology

#### 2.1.1 RNA Extraction and Purification

Cell lines were grown and  $1 \times 10^6$  cells were used for RNA extraction. RNA extraction was done using the RNeasy Mini Kit (Qiagen, UK). The extraction was done according to the manufacturer's instructions. The extracted RNA was further treated with DNase in order to remove any traces of contaminating DNA. This was done using the TURBO DNA-free<sup>TM</sup> Kit (Ambion Applied Biosystems, UK), following the manufacturer's protocol. The amount of the extracted RNA and the purity of the sample obtained are described in section 2.1.3.

# 2.1.2 DNA Extraction and Precipitation

To the DNA containing solution, an equal volume of phenol/chloroform mixture was added. The mixture was vortexed for a couple of seconds, kept at room temperature for 15 minutes, then centrifuged for 5 minutes at 14,000g. The layer containing the DNA, the top layer, was transferred to a fresh eppendorf. To that 1/10 the volume of 10M ammonium acetate and 2.5 the volume of 100% ethanol was added. The solution was vortexed for a couple of seconds, incubated on ice for 30 minutes, and then centrifuged for 5 minutes at 14,000g. The supernatant was removed, and the pellet was washed with 70% ethanol. The DNA pellet was air dried for 20 minutes, and then resuspended in an appropriate volume of TE buffer or DEPC H<sub>2</sub>O. The amount of the extracted and purified DNA, and the purity of the sample, is described in section 2.1.3.

#### 2.1.3 Quantification of Nucleic Acid Concentration

The concentration of nucleic acids was quantified by measuring the diluted sample's absorbance at 260nm. This was done by the use of a GeneQuant spectrophotometric measurement (Pharmacia Biotech, Sweden), and depending on what the nucleic acid pellet was resuspended in, TE buffer or DEPC H<sub>2</sub>O was used as a blank. The following formula was used to estimate the nucleic acid concentration:

1.0 OD<sub>260</sub> = 44ng/ μl for RNA

1.0 OD<sub>260</sub> = 50ng/ μl for DNA

The purity of the samples was estimated through the  $OD_{260}/OD_{280}$  ratio. Ratios of 1.8 for DNA and 2.0 for RNA were considered to be highly pure.

#### 2.1.4 cDNA Synthesis

cDNA synthesis is the production of a DNA complementary strand from an existing RNA strand. This process can be obtained by a reverse transcription (RT) reaction. The reaction is accomplished by the use of a reverse transcriptase enzyme that extends an RNA hybridised primer. Since, eukaryotic mRNA is 3' polyadenylated, a polydeoxythymidine oligomer (16-18 bases) can be used as a primer for this reaction, consequently giving rise to RNA-DNA hybrids (Gerard *et al.*, 1997).

For the RT reaction the following mix was prepared: a volume equivalent to  $1-2\mu g$  of RNA,  $1\mu l$  of oligo  $dT_{16-18} (2\mu g/\mu l$ , Synthesized by the Molecular Biology Unit at Hodgkin Building, King's College London), and RNase free water (Qiagen, UK) to a total volume of  $45\mu l$  was added. The mixture was incubated at 70°C for 10 minutes (Thermal Reactor- HYBAID Omnigene, UK). This was done to denature the RNA secondary structure.

The 45µl mixture was left on ice for 5 minutes and the following were added: 5µl of 10mM deoxynucleotide triphosphate (dNTP) (Promega, UK), 20µl of first strand 5x Buffer (250mM Tris-HCl: pH 8.3, 375mM KCl, 15mM MgCl<sub>2</sub>:

Invitrogen, UK), 10µl of 1M DTT (Invitrogen, UK), 5µl of RNasin Ribonuclease Inhibitor, 20-40u/µl (Promega, UK), and 5µl of M-MLV reverse transcriptase, 200u/µl (Promega, UK). The final volume of the RT mixture was 100µl which was incubated at 37°C for one hour (Techne DB, DRI-Block<sup>®</sup>, U.S.A).

# 2.1.5 Polymerase Chain Reaction

PCR involves the amplification of a DNA segment at high temperature by the use of sequence specific primers. The DNA template is initially denatured, primers then anneal to the template and a heat stable DNA polymerase extends them. This temperature cycle is repeated for a number of times, usually 30-40 cycles. The product of the first amplification becomes the template for the next one, and the amount of product increases exponentially, theoretically, two fold for each cycle.

All PCR reactions were carried out either with PCR master mix (Promega, UK) or Fast Start High Fidelity PCR system (Roche, UK). Tables 2.1 and 2.2 list all components that were used with the above PCR polymerases respectively.

Component	Volume per 25µl	
	reaction	
20ng of DNA	xμl	
10μM Forward Primer	1µl	
10μM Reverse Primer	1µl	
PCR Master Mix	12.5µl	
H <sub>2</sub> O	Add to 25µl	

Table 2.1 Components of the PCR Master Mix (25µl reaction mixture).

Component	Volume per 25µl	
	reaction	
20ng of DNA	xμl	
10µM ForwardPrimer	1µl	
10μM Reverse primer	1µl	
10mM dNTP	1.25µl	
10x fast start high	2.5µl	
fidelity buffer		
Fast start high fidelity	1.25 U	
<i>Taq</i> polymerase		
enzyme		
H <sub>2</sub> O	Add to 25µl	

Table 2.2 Components of the Fast Start High Fidelity PCR system (25 $\mu$ l reaction mixture).

The preparation of the reaction master mix was performed on ice, and the high fidelity *Taq* polymerase was added just before the initiation of the reaction.

The parameters set for the PCR systems are shown below and all sets of primers used for this project can be seen in table 2.3.

The following PCR conditions were applied for the reactions done by the use of the PCR Master Mix and the Fast Start high Fidelity PCR system:

Initial denaturation at 96°C for 5 minutes			
30 cycles of $\rightarrow$	Denaturation at 95°C for 1 minute		
	Annealing (temperature depending on the primer set, see table		
	2.3) for 50 seconds		
	Elongation at 72°C for 15 seconds x length of the amplicon/Kb		
Final extension at 72°C for 5 minutes			

The PCR products were analysed by agarose gel electrophoresis.

PCR Primer Set		Annealing
	Primer Set Sequence	temperature
		used
β3α2	Forward 5' d-AGCTTCTCCCTGACATCCGTGGA'3	60°C
primers*	Reverse 5' d-CCCATTGTGATTATAGCCTAAGA'3	
$\beta 2\alpha 2$ primers*	Forward 5' d-GCTACGGAGAGGCTGAAGAA'3	60°C
	Reverse 5' d-CGTGATGTAGTTGCTTGGGA'3	
U6 primers	Forward 5' d-CCCAAGCTTCAAGGTCGGGCAGGAAGAGGGCCTA'3	64°C
	Reverse 5' d-AT <mark>GGATCC</mark> TAGTATATGTGCTGCCGAAGCGAGCAC'3	
H1 primers	Forward 5' d-CCCAAGCTTCGAACGCTGACGTCATCAAC'3	64°C
	Reverse 5' d-AT <mark>GGATCC</mark> GAAAGAGTGGTCTCATACAGAACTT'3	
CMV-340	Forward 5' d-TTCCAAGTCTCCACCCCATTGACG T'3	60°C
CMV-207	Forward 5' d-CCACTTGGCAGTACATCAAG'3	60°C
EGFPN1	Reverse 5' d-GTCCAGCTCGACCAGGATG'3	60°C
Sp6	5' d-ATTTAGGTGACACTATAG'3	50°C
pEGFP-N1	Forward 5' d-AACGCCAATAGGGACTTTCC'3	57°C
	Reverse 5' d-AAGTCGTGCTGCTTCATGTG'3	
B-actin primer N	5' d-CTGAGTCTCCTTTGGAACTCTGCAG '3	60°C
B-actin primer K	5' d-GAGCACAGAGCCTCGCCTTTGC'3	60°C
B-actin primer L	5' d-GGATCTTCATGAGGTAGTCAGTCAGG'3	60°C
PRL13 Primers	Forward 5' d-TGAAGGAGTACCGCTCCAAA'3	60°C
	Reverse 5' d-GGGTGGCCAGTTTCAGTTC '3	
T3 reverse	5' d-GCAATTAACCCTCACTAAAGG'3	50°C
T7 forward	5' d-TAATACGACTCACTATAGGG'3	50°C

**Table 2.3 Primer sets used within the project.** Primers were aimed to be 21-23bp with a GC content of 48-60%. SGD web primer design software was used to design the primers. Red coloured sequences in U6 and H1 primer sets, indicate added restriction sites. These are HindIII and BamHI restriction sites located in the forward and reverse primers respectively. Primers indicated by (\*) were taken from the following publications, (Campanini F *et al.*, 2001) for the  $\beta 3\alpha 2$  primer set and (Goh H.G *et al.*, 2006) for the  $\beta 2\alpha 2$  primer set.

#### 2.1.6 Agarose Gel Electrophoresis

DNA molecules carry a negative charge as a consequence of their phosphate backbone. Therefore, if DNA molecules are placed in an electrical field they will migrate towards the positive pole. An agarose gel is used for support, and comprises a network of pores, through which DNA molecules can travel. The pore size of a gel is determined by the agarose concentration, the greater the concentration of the gel, the smaller its pores.

To make a 1% gel, 0.5g of agarose powder (Fisher Bioreagents, UK) was weighed (Mettler R1210) and transferred into a clean conical flask that was then filled with 50ml of 1x TAE buffer (40mM Tris-Acetate-EDTA, 1mM EDTA pH 8.5, Sigma, Germany). The mixture was then heated on maximum power for 2 minutes in a microwave oven (Hotline, 930W Bejam, Korea). 10µl of ethidium bromide (1mg/ml, MBL International, USA) was then added and the solution was mixed.

An electrophoretic boat was placed horizontally into an electrophoretic tank (Max fill, Fisher brand, UK). A comb with the desired number of wells, was placed into the boat. The agarose was then poured into the boat and the gel was allowed to set. Once set, the comb was removed and the electrophoretic tank was then filled with 1x TAE buffer to a point where all the wells were covered. Samples were then mixed with 6x loading buffer (0.25% bromophenol blue, and 30% glycerol in 1x TAE buffer, BioLabs inc, England) and loaded into the wells. 300-450ng of the appropriate molecular weight marker was also loaded onto the gel. See Appendix 1 for all molecular weight markers used in this project.

The tank was then connected to an electrophoresis 3500 power supply (Pharmacia, UK) and a current between 90-100 volts was applied. Once the samples had migrated three-quarters of the way through the gel, the current was stopped and the gel was viewed under UV light (Gene flash, Syngene Bio Imaging, USA), and a picture was taken (Video graphic printer, UP-850, Sony, Japan). The DNA fragments of the size of interest were cut out of the agarose gel
and were purified by the use of the Wizard<sup>®</sup> SV Gel and PCR Clean-Up System according to the manufacturer's instructions (Promega, UK). The concentration of the purified DNA fragments was then estimated my means of OD<sub>260nm</sub> measurement.

## 2.1.7 Restriction Enzyme Digest

DNA digestions were carried out on PCR products or plasmids. According to the manufacturer's protocol, 1 unit of the enzyme is required to digest 1µg of DNA in one hour under the optimum temperature and buffer conditions (Promega and New England Biolabs, UK). In this project, DNA fragments of interest were digested for a duration of 3 hours at 37°C with the appropriate restriction enzyme at its optimum 10x buffer. All restriction enzymes and their buffers used in this study are listed in table 2.4.

Restriction Enzyme	Restriction Buffer (100% suitable)	Manufacturer
BamHI	Buffer B: 6mM Tris-Cl pH 7.5, 6mM MgCl <sub>2</sub> , 50mM	Promega
	NaCl, 1mM DTT	
	Or Multicore buffer: 25mM Tris-acetate pH 7.8,	
	100mM K(Ac), 10mM Mg(Ac) <sub>2</sub> , 1mM DTT	
EcoRI	Buffer H: 90mM Tris-Cl pH 7.9, 6mM MgCl <sub>2</sub> , 50mM	Promega
	NaCl	
	0r Multicore buffer	
HindIII	Buffer E: 6mM Tris-Cl pH 7.5, 6mM MgCl <sub>2</sub> , 100mM	Promega
	NaCl, 1mM DTT	
Sacl	Buffer J: 10mM Tris-Cl pH 7.5, 7mM MgCl <sub>2</sub> , 50mM KCl,	Promega
	1mM DTT	
SnaBl	Buffer 4: 20mM Tris-acetate pH 7.9, 50mM K(Ac),	New England
	10mM Mg(Ac) <sub>2</sub> , 1mM DTT	Biolabs
Xhol	Buffer B: 6mM Tris-Cl pH 7.5, 6mM MgCl <sub>2</sub> , 150mM	Promega
	NaCl, 1mM DTT	

Table 2.4 Restriction Enzymes used in this project and their incubation buffers.

# 2.1.8 DNA Ligation

Purified and digested PCR products were ligated into the vector pB12mcseGFP, that was kindly given by Dr Adam Rodaway (King's College London, Cardiovascular Division). The plasmid was also digested with the same restriction enzymes as that used for digesting the PCR products to be inserted. Following gel purification of DNA and estimation of its concentration by means of OD<sub>260nm</sub>, a molar ratio 10:1 insert to vector was used to establish ligation. To do so, the following equation was used:

ng of insert = (ng of vector x kb of insert size / kb of vector size) x (molar ratio of insert : vector)

The ligation reactions were done in a total volume of  $10\mu$ l, where 200ng of the vector, xng of the insert,  $1\mu$ l of T4 DNA ligase enzyme ( $3U/\mu$ l),  $1\mu$ l of 10x ligase buffer (Promega, UK) and a volume of  $x\mu$ l of DEPC treated water were added. The reaction was left for overnight ligation at 4°C. The ligation reaction was further assessed by means of PCR, where the ligation mixture was diluted by 1000 and used as a DNA template. Primer sets used were either both located within the plasmid or at least one was within the plasmid while the other was within the insert. In addition, a negative control was established by the use of the same set of primers to PCR the wild type plasmid.

## 2.1.9 Transformations

 $50\mu$ l of JM109 competent cells (> $10^7$  cfu/µg, Promega, UK) were thawed on ice for 5 minutes.  $5\mu$ l of the ligation mixture was added and the cells were kept in a 1.5ml Eppendorf tube on ice for another 20 minutes. To heat shock the cells, the tube was then transferred to a heat block at 42°C (Grant BT1 Block Thermostat, UK) for 45-55 seconds then back on ice for another 2 minutes.  $350\mu$ l of SOC medium (BioLine Ltd, UK) was then added to the cells, and the cells were incubated at  $37^\circ$ C for 45-90 minutes.

 $200\mu$ l of transformed competent cells were then spread on prepared LB agar plates. These plates were prepared from 16g of LB agar (Sigma, UK) that was dissolved in 500ml of Milli Q H<sub>2</sub>O (ELGA LA620, UK), autoclaved, then further boiled, left to cool down, and  $60\mu$ g/ml of ampicillin was added. LB agar plates were stored at 4°C for a month from the date of preparation. The plates were placed overnight in a 37°C incubator (Luckham R300), and were examined for colony growth the next day.

By the use of the appropriate set of primers, the grown colonies were screened by means of PCR. A negative control, wild type plasmid, was also included. Positive colonies that contain the inserted DNA fragment were picked up from the agar plate and further grown in 20ml of LB broth medium (Fisher Scientific, UK). The LB medium was prepared by dissolving 12.5g of LB broth in 500ml of Milli Q H<sub>2</sub>O, autoclaved, and  $60\mu$ g/ml of ampicillin was added just prior to bacterial colony addition. The 50ml tubes containing the LB medium were left overnight at 37°C in a shaker incubator (Gallenkamp Orbital incubator).

## 2.1.10 Plasmid DNA Isolation and Purification

Solutions used include:

- Cell suspension solution (P1): 25mM Tris-HCL, pH 8, 10mM EDTA with 50mM glucose and DNase-free RNase (100µg/µl).
- Cell lysis solution (P2): 0.2N NaOH, 1% SDS.
- Potassium acetate solution (P3): 20 Volumes of 5M potassium acetate, 3
  volume of Glacial acetic acid and 2 volumes of H<sub>2</sub>O.

Mini and Maxipreps were conducted on transformed bacterial cells, which were incubated in 20ml or 500ml of LB media respectively containing the appropriate antibiotic. The cells were grown overnight in a shaking incubator at 37°C and were harvested by centrifugation at 5000rpm at 4°C for 10 min (Mini) or 20 min (Maxi). The supernatant was discharged and the DNA isolation was performed by the use of the Qiagen Plasmid Mini or Endofree Plasmid Maxi kit that was conducted according to the manufacturer's protocol. The purity and the amount of the isolated DNA were determined as described earlier in section 2.1.3.

## 2.1.11 Glycerol Stocks

Upon confirmation of clones of interest by means of PCR colony screening, each clone was picked up from a positive single colony and grown overnight in LB media containing a suitable antibiotic. 900µl of the overnight grown bacterial colony was pipetted into a 1.5ml Eppendorf tube containing 100µl of glycerol (Sigma, UK). The tube was inverted several times and the glycerol stock was stored at -80°C.

#### 2.1.12 DNA sequencing

 $5\mu$ l of plasmid DNA, at a concentration of 100ng/ $\mu$ l, was enclosed in a 0.5ml Eppendorf tube and sent to Geneservice Ltd at UCL London for sequencing. Geneservice Ltd provided common primers for sequencing such as those that bind to T7 or Sp6 regions. However, for sequencing with custom made primers,  $5\mu$ l at 3.2pmol/ $\mu$ l of each primer per reaction was sent separately with the DNA samples to be sequenced. Sequencing results were analysed using the Sequence Scanner software v1.0 (Applied Biosystems, UK).

## 2.1.13 Generation of Blunt Ends

Some restriction enzymes generate blunt ends directly, such as Stul and SnaBI. However, other enzymes generate sticky ends with some base overhangs. To achieve blunt end cloning or even vector self-ligation through blunt ends, these overhangs must be converted to blunt ends. This can be achieved by either filling up the overhangs with complimentary bases or by removal of the excess bases. In this study, the Klenow fragment (3'  $\rightarrow$  5' exo-), kindly given by Dr. Cathy Joyce, the Department of Molecular Biophysics and Biochemistry, Yale University, New Haven, CT., USA, was used to create blunt ends by removal of the 3' overhangs.

To do so, 1µg of digested vector, was incubated with 1µl of Klenow fragment in 3-4µl of 10x buffer 2 (1x buffer 2: 10mM Tris-HCL pH 7.9, 50mM NaCl, 10mM MgCl<sub>2</sub>, 1mM DTT), 33µM of each dNTP and xµl of DEPC H<sub>2</sub>O to a total of 30-40µl. The reaction was left at room temperature for 15 minutes then stopped by heating at 75°C for 25 minutes. The Klenow treated vector, was then run on an agarose gel, were the band representing the vector was

purified by the use of the Wizard<sup>®</sup> SV Gel and PCR Clean-Up System according to the manufacturer's protocol. The concentration of the purified DNA fragments was then estimated by means of OD<sub>260nm</sub> measurement.

## 2.2 AT Cloning

PCR products generated by the use of *Taq* polymerase enzyme have an "A" residue overhang on the 3' ends. Therefore, unmodified PCR products can be cloned into a linearised vector having a "T" overhang of the 3' end. Based on these facts, unmodified PCR products were cloned into the pGEM-T vector (Promega, UK). The AT cloning was done by the use of the pGEM-T easy vector kit (Promega, UK) and was conducted according to the manufacturer's protocol. The pGEM®T easy vector system is shown in Figure 2.1. This vector has a *LacZ* gene within its multiple cloning region (MCR). The *LacZ* gene expresses  $\beta$ -galactosidase enzyme upon induction with IPTG. This means that bacterial colonies containing the pGEM®T vector when grown on agar plates containing both IPTG and X-gal will produce  $\beta$ -galactosidase, and these bacterial colonies will be blue in colour. However, when a DNA fragment is inserted into the MCR, the *LacZ* gene is disrupted and bacterial colonies, hence, allowing blue-white colony screening.

Pre-prepared LB agar plates with 100µg/ml ampicillin, were further plated with a 100µl of 100mM IPTG (promega, UK) and 20µl 50µg/ml X-Gal (Promega, UK) 30 minutes prior to plating the transformed bacterial cells. Colonies were left to grow overnight at 37°C, and the white colonies were screened by means of PCR.

Positive PCR screened colonies were further grown overnight in LB broth containing ampicillin and the plasmids were then extracted. The purified extracted plasmids were quantified and were then sent to Geneservice Ltd at UCL London for sequencing. The primers T7 forward and Sp6 reverse were used to conduct this sequencing.



Figure 2.1 Map of the pGEM®T Easy Vector system. The diagram was obtained from the Promega website.

The pGEM®T easy vector system was used in this study for the purposes of AT cloning. As the Map indicates, the vector has a "T" overhang within the MCR in which the DNA fragment of interest, having an "A" overhang, can be inserted.

# 2.3 Generation of Plasmids

Two plasmids were used for the purpose of incorporating shRNA molecules, pEGFP-N1 plasmid and pB12mcs-eGFP plasmid. The pEGFP-N1 vector (Clonetech, UK) has an enhanced GFP, and includes a human cytomegalovirus (CMV) promoter. For the purposes of this project the CMV promoter was not used. Instead either the human U6 or the human H1 promoter was used. The pEGFP-N1 plasmid with the CMV promoter removed was used as a negative control during expression experiments. The pEGFP-N1 vector, where either the U6 or H1 promoter had replaced the CMV promoter, would be used as the positive control. The pEGFP-N1 vector is shown in figure 2.2.

JM109 bacterial cells were transformed with the pEGFP-N1 vector. The cells were then grown overnight in 500ml of LB with the appropriate antibiotic.

1ml of the grown bacteria was stored at -80°C as glycerol stock, and a maxi prep was then conducted on the transformed bacterial cells. The extracted pEGFP-N1 plasmid was resuspended in an appropriate volume of Milli Q water, quantified and stored at -20°C.

The pB12mcs-eGFP vector was kindly given by Dr. Adam Rodaway. This vector also contains an enhanced GFP, however, it has no promoter. This is advantageous, as the wild type vector can be used as a negative control. Upstream of the eGFP is the MCR, in which the U6, the H1 human promoters, and the shRNA cassettes can be individually cloned. The vector with only the U6 or the H1 promoter can then serve as positive control. A map of the pB12mcs-eGFP plasmid is shown in Figure 2.3. Before deciding which restriction enzymes were to be used, the vector was sequenced to confirm its identity.



Figure 2.2 pEGFP-N1 Vector Map. The diagram was obtained from the Clontech website.

Top: The map of the pEGFP-N1 vector.

Bottom: The sequence and the restriction sites of the MCR of the pEGFP-N1 vector.

The pEGFP-N1 vector was selected to clone the shRNA cassettes. The vector has an eGFP gene that gives its high expression and bright fluorescence in mammalian cells.



Figure 2.3 The pB12mcs-eGFP Vector Map. The plasmid was constructed and given by Dr. Adam Rodaway

The pB12mcs-eGFP vector is 3029bp in size, has an ampicillin resisted gene, and an eGFP gene that gives a higher and brighter expression of this protein in mammalian cells. The wild type vector can be used as a negative control. Positive controls can be constructed through the insertion of a U6 or an H1 human promoter into the MCR.

## 2.3.1 Construction of the shRNA Cassettes

Short hairpin RNA (shRNA) molecules were designed according to the Cold Spring Harbor Lab guidelines. shRNA molecules were designed to be of 50-52 nt long and were to be cloned with, and downstream of, the U6 or H1 human promoters. The U6-shRNA primer transcripts were initialised with a "G" nucleotide, of the U6/H1-small nuclear RNA Pol III transcript, and termination was mediated by a stretch of 6 "T" nt at the end of the transcript.

Each of the designed shRNA molecules can be divided into four regions. These are the anti-sense region, the loop, and sense and termination regions. The sense region corresponds to 21-22 nt and it's sequence is the same as the target DNA, this region was designed to be complementary to the fusion points of the *BCR-ABL* gene. Thus, aiming to target the fusion gene only and not the normal *BCR* or *ABL* genes. The anti-sense region is complimentary to the sense part. Both the sense and anti-sense parts of the shRNA transcript are separated by an 8 nt long loop. The final region is made up of a stretch of "Ts", because the U6 and H1 promoters require an oligo T termination signal. The sequences of the designed shRNA transcripts can be found in table 2.5.

Brummelkamp and colleagues in 2002 studies the effects of the loop size on the RNAi potency of shRNA molecules that had the same sense and antisense sequences, but either had five, seven, or nine long nt loop sequences. Their study showed that the differences in loop size and sequence had different RNAi effects, where shRNA molecules with a five nt loop were inactive, those with seven were moderately active, and shRNAs with 9 nt long gave around 90% knockdown of the gene of interest (Brummelkamp *et al.*, 2002). Furthermore, in 2010, Schopman and colleagues demonstrated that moderate or even weak shRNA inhibitors could be transformed into strong potent inhibitors by simply changing the loop size and structure. In this study scientists found that shRNA molecules with an open loop structure (no sequences of the loop are complementary to each other at the sense and antisense part of the shRNA) exhibit stronger RNAi gene silencing (Schopman *et*  *al.*, 2010). On this basis, an 8 nt long open loop structure was used for the shRNA molecules designed in the current work.

Once the designing of the shRNA molecules was complete, the secondary structures of these molecules were predicted by the use of the program 'mfold'. These secondary structures can be seen in Appendix 2, except for  $\beta 3\alpha 2$  shRNA1 structure that is demonstrated in Figure 3.13.

Fusion point	shRNA sequences 5'→3'	
β3α2	shRNA1 <sub>(21nt)</sub>	gcagaguuca <mark>aa</mark> agcccuucagaagcuugugaagggcu <mark>uu</mark> ugaacucugctttttt
	shRNA2 <sub>(21nt)</sub>	cagaguucaaaagcccuucaggaagcuugcugaagggcuuuuugaacucugtttttt
	shRNA3 <sub>(22nt)</sub>	gcagaguuca <mark>aa</mark> agcccuucaggaagcuugcugaagggcu <mark>uu</mark> ugaacucugctttttt
β2α2	shRNA1 <sub>(21nt)</sub>	caauaaggaa <mark>ga</mark> agcccuucagaagcuugugaagggcu <mark>uc</mark> uuccuuauugtttttt
	shRNA2 <sub>(21nt)</sub>	aauaaggaagaagcccuucacgaagcuuggugaagggcuucuuauutttttt
	shRNA3 <sub>(22nt)</sub>	caauaaggaaggaaggeccuucacgaagcuuggugaagggcuucuuauugtttttt

#### Table 2.5 The designed shRNA transcripts.

The above table demonstrates that three different shRNA molecules were designed for each fusion point. shRNA1 and 2 are of 21 nt long, whereas shRNA3 is 22 nt long. shRNA1 has 11 nt upstream the fusion point and 10 nt downstream, shRNA2 had 10 nt upstream the fusion point and 11 nt downstream, whereas shRNA3 has 11 nt on either sides of the fusion point. The sequences shown in blue represent the anti-sense part of the molecule and the fusion points within this part are underlined and in bold. Any sequence differences between the three shRNA molecules are shown in green. The sequences in red represent the sense part. The sequences in black and underlined correspond to the hairpin loops, and the oligo T termination signal is in black.

## 2.3.2 Cloning of Promoter Constructs

PCR was used to produce the U6/shRNA and H1/shRNA cassettes. These were established by incorporating the shRNA sequence within a reverse primer

that had 28 nt of its sequence complimentary to the U6 or H1 promoters. HindIII and BamHI restriction sites were also incorporated into the forward and reverse primers respectively. Finally, PCR, conducted on K562 cDNA, was used to produce the whole cassettes. The sequences for these primers are shown in table 2.6.

The U6/shRNA and the H1/shRNA cassettes were cloned into the pB12mcseGFP plasmid. The plasmids were then sent for DNA sequencing to confirm the correct sequence. A schematic diagram for the steps involved in the synthesis of the shRNA cassette is shown in Figure 2.4. A flow diagram showing the overall procedures undertaken to generate the pB12mcs-eGFP containing the shRNA cassettes can be seen in Figure 2.5.



Figure 2.4 Schematic Diagram of the Synthesis of the shRNA cassettes.

The diagram illustrates how a shRNA cassette under the regulation of the U6 promoter was synthesised. cDNA extracted from the K562 cell line was used as a DNA template to amplify the human U6 promoter. To synthesise the shRNA cassette (top part of the diagram) the shRNA molecule was synthesised as an oligonucleotide that had 27bp complimentary to the U6 promoter. This oligo was part of a U6 reverse primer, which could be used together with the U6 forward primer in a PCR reaction. The resultant PCR product contained the U6 promoter linked to the shRNA (bottom part of the diagram). The reaction was conducted for every designed shRNA molecule, and the same technique was used to produce shRNA cassettes regulated under the human H1 promoter.

Primer Name	Primer sequence	Melting
		temperature
U6 Forward	5' d-CCCAAGCTTCAAGGTCGGGCAGGAAGAGGGCCTA'3	64°C
β3α2 shRNA1 U6 reverse	5' d-ATGGATCCAAAAAAGCAGAGTTCAAAAGCCCTTCACAAGCTT CTGAAGGGCTTTTGAACTCTGCGTAGTATATGTGCTGCCGAAGCGAGCAC'3	68.1°C
β3α2 shRNA2 U6 reverse	5' d-AT <mark>GGATCC</mark> AAAAAACAGAGTTCAAAAGCCCTTCAGCAAGCT TCCTGAAGGGCTTTTGAACTCTGGTAGTATATGTGCTGCCGAAGCGAGCAC'3	68.3°C
β3α2 shRNA3 U6 reverse	5' d-ATGGATCCAAAAAAGCAGAGTTCAAAAGCCCTTCAGCAAGC TTCCTGAAGGGCTTTTGAACTCTGCGTAGTATATGTGCTGCCGAAGCGAGCA C'3	68.9°C
β2α2 shRNA1 U6 reverse	5' d-ATGGATCCAAAAAACAATAAGGAAGAAGCCCTTCACAAGCT TCTGAAGGGCTTCTTCCTTATTGGTAGTATATGTGCTGCCGAAGCGAGCAC'3	67°C
β2α2 shRNA2 U6 reverse	5' d-AT <mark>GGATCC</mark> AAAAAAAAAAAGGAAGAAGCCCTTCACCAAGCTT CGTGAAGGGCTTCTTCCTTATTGTAGTATATGTGCTGCCGAAGCGAGCAC'3	67.1°C
β2α2 shRNA3 U6 reverse	5' d-ATGGATCCAAAAAAACAATAAGGAAGAAGCCCTTCACCAAGCT TCGTGAAGGGCTTCTTCCTTATTGGTAGTATATGTGCTGCCGAAGCGAGCAC' 3	68.2°C
H1 forward	5' d-CCCAAGCTTCGAACGCTGACGTCATCAAC'3	64°C
β2α2 shRNA1 H1 reverse	5' d-ATGGATCCAAAAAACAATAAGGAAGAAGCCCTTCACAAGCTT CTGAAGGGCTTCTTCCTTATTGGAAAGAGTGGTCTCATACAGAACTT'3	66.8°C
β2α2 shRNA2 H1 reverse	5' d-AT <mark>GGATCC</mark> AAAAAAAAAAAGGAAGAAGCCCTTCACCAAGC TTCGTG AAGGGCTTCTTCCTTATTGAAAGAGTGGTCTCATACA GAACTT'3	66.9°C
β2α2 shRNA3 H1 reverse	5' d-AT <mark>GGATCC</mark> AAAAAACAATAAGGAAGAAGCCCTTCACCAAGCT TCGTGAAGGGCTTCTTCCTTATTGGAAAGAGTGGTCTCATACAGA ACTT'3	68.2°C

**Table 2.6 Primer sequences used to construct the shRNA cassettes.** This table represents all the primers used to produce the shRNA cassettes. The sequences within the primers in red show the incorporated restriction enzyme sites, HindIII and BamHI, located in the forward and reverse primers respectively.



#### Containing the shRNA Cassettes.

This diagram summarises the steps and experiments conducted to construct the shRNA cassettes, directed against the *BCR-ABL* fusion genes, and to incorporate these cassettes into a suitable vector.

### 2.4 Tissue Culture

Two types of cell lines were used for the purposes of this study. These are the K562 cell line (kindly given by Dr. David Grimwade, Guy's tower, King's College London) and the KCL-22 cell line (DSMZ, Germany). These are CML cell lines, and as explained in chapter 1, the K562 cells are CML cells with the  $\beta 3\alpha 2$ fusion point, whereas the KCL-22 cells have the  $\beta 2\alpha 2$  fusion point.

# 2.4.1 Cell line Maintenance

## 2.4.1.1 Thawing and recovery of cell lines

A vial of CML cell lines was removed from the liquid nitrogen freezer (RS series, Jencons pls) and placed immediately in a 37°C water bath (Grant Instrument Ltd, England). The cells were washed with 2ml warm RPMI 1640 medium with L-glutamine (PAA, Austria), 10% Foetal Bovin Serum (FBS) (Sigma, UK) and 550µl of penicillin/streptomycin antibiotic (10,000iu PEN/ml, 10,000ug STREP/ml, MP Biomedicals, France), and centrifuged for 5 minutes at 1000 rpm (Beckman CS-R6 centrifuge, UK) at room temperature. The supernatant was discharged and the pellet resuspended in 1ml of medium. Cells were then transferred to a 75cm<sup>2</sup> culture flask that contained 30ml of medium and incubated in a humidified 37°C, 5% CO<sub>2</sub> incubator (Binder, USA).

#### 2.4.1.2 Feeding and Passageing of cell lines

For the cells to be passaged they must be confluent, that is their count should be about  $2.5 \times 10^6$  cells/ml in a 250ml flask; cells were fed every 2-3 days till confluency was reached.

#### - Feeding of Cell lines

Cells were counted and if non-confluent, one third of the medium in the flask was aseptically removed and replaced by an equal volume of fresh warm media. The flask was gently swirled and returned to the incubator. On the days cells were not fed, they were checked by gently swirling the flask, to resuspend them, and by observing any media colour changes which reflected their metabolic growth.

#### Passaging of Cell lines

When cells were confluent, half the cells in suspension were aseptically transferred to a new culture flask. To each flask an equal amount of warm medium was added and flasks were returned to the incubator.

### 2.4.1.3 Counting of cell lines

The process of cell counting is an important procedure through which optimisation of culture conditions and the performance of accurate quantification experiments can be conducted. To count the cells a haemacytometer (Bright-Line, Hausser Scientific, USA) was used. The device is a thick glass slide with a central counting chamber. The chamber is divided into two portions each made up of a 3 x 3 mm etched grid that is further divided into nine secondary squares, 1 x 1 mm each. The four corner squares and the central one are used for counting; they are divided into 16 and 25 tertiary squares respectively. A cover slip is also included with the device. See figure 2.6.



Figure 2.6 The Haemacytometer. (The diagram was taken from the Klinik Andrologi website)

(Top) The central portion of the haemacytometer, the counting chamber, is divided into two portions. Each portion has its counting grid. (Bottom) One of the grids is amplified showing five counting boxes. The outer four are further divided into 16 tertiary boxes, while the central one comprises 25 boxes. A central tertiary box is further enlarged showing two cells within it. The bent arrows on the top show where the cover slip is placed prior to loading the cell suspension. The straight arrow in the middle shows the grove in which the cell suspension is loaded.

The haemacytometer and the cover slip were wiped with 70% ethanol and the cover slip placed on the counting chambers. A 1:1 dilution for the cells with trypan blue (0.4%, Sigma, UK) was prepared and 10µl of the dilution was loaded into each counting chamber by a pipette. The suspension was drawn under the slip by capillary action, and cells, in the centre and in two of the corner squares, were counted under a light microscope (100x magnification, Nikon Eclipse TS1000, Japan) and with the aid of a hand-held counter. Whole blue-stained cells were excluded in the count, since only those staining yellow are viable. The following formula was used to calculate the cell number:

## "Cells/ml = average count of squares x dilution factor x $10^{4"}$

In some cases the Invitrogen automated cell counter, Countess<sup>®</sup>, was also used to count the cells. Here cells were again prepared at a 1:1 dilution factor with trypan blue (0.4%, Invitrogen, UK) and were loaded into both sides of the Countess<sup>TM</sup> call counting chamber slide. The manufacturer's protocol was followed in order to count the cells and predict the percentage of cell viability.

### 2.4.1.4 Freezing of cell lines

Cells for freezing were first transferred into 50ml test tubes, centrifuged for 5 minutes at 1000 rpm (GS-6R Centrifuge, Beckman, USA) at room temperature. The supernatant was discharged and the pellet was resuspended in 10ml of warm medium. Cells were counted, spun as stated previously and were resuspended in a volume of freezing medium, 90% FBS with 10% DSMO, (Sigma, UK) to obtain a concentration of 2 x  $10^6$  cells/ml. Cells were pipetted in 1ml aliquots into 2ml cyrovials. The vials were placed in a 1°C cryo freezing chamber (Nalgene<sup>TM</sup>, USA) to allow a gradual decrease in temperatue thus reducing the loss of cells. The camber containing the cells was left overnight in a -80°C freezer (Revco Scientific Inc, USA). The following day, cells were placed in a -150°C liquid nitrogen Dewar (HC series, Tayler-Wharton, UK).

## 2.4.2 Optimisation of Cell Line Transfection

The term transfection is usually used to indicate a process in which nucleic acids are intentionally introduced into cells. To date various methods for transfection are present and can be categorised as chemical and non-chemical based transfections. In this project, the K562 and KCL22 cell lines were transfected with the U6/shRNA or H1/shRNA pB12mcs-eGFP plasmids in order to observe and study the effects of the constructed shRNA cassettes. However, the K562 and KCL22 cell lines are cells in suspension and therefore are considered to be hard to transfect (Ovcharenko *et al.*, 2005, Oliveira and Goodell, 2003 and Merkerova *et al.*, 2007). Hence several transfection methods were tested to establish an acceptable transfection protocol giving a good efficiency.

## 2.4.2.1 Chemical based transfections

These transfections depend on the interactions of chemical materials with the DNA of interest in order to internalise the latter into the targeted cell lines. Several transfection reagents were tested, these included the followings:

- <u>SuperFect® Transfection Reagent (QIAGEN, UK)</u>: This transfecting reagent has a specifically designed activated dendrimer that possesses a defined spherical core with radiating branches that are positively charged at the ends. The reagent assembles the DNA particles into compact positively charged structures that in turn bind to the negatively charged receptors found on the surfaces of eukaryotic cells, hence internalising the target DNA molecules into the cells.
- FuGENE® HD Transfection Reagent (Roche, UK): This transfecting reagent consists of a proprietary blend of lipids and other components that are supplied in 80% ethanol, sterile-filtered, glass packaged vials. The reagent complexes with the target DNA molecules, thus transporting the DNA molecules into the cells during transfection.
- Lipofectamine<sup>™</sup> 2000 Transfection Reagent (Invitrogen, UK): This is a cationic lipid based reagent. The reagent is a mix of cationic and neutral lipids that form unilamellar positively charged liposome vesicles. When mixed with DNA, the cationic liposomes will absorb the DNA molecules then facilitate their

internalisation into the target cells by means of fusion to the cell's plasma membrane, thus forming an endocytic vesicle.

- <u>TransIT®-2020 Transfection Reagent (Mirus Bio, USA)</u>: According to the manufacturer, this transfecting reagent is lipid based, however it is not liposomal. Therefore, the reagent complexes with the DNA and causes it to be internalized into the cells during transfection.
- <u>HilyMax Transfection Reagent (Dojindo Moleculer Technologies, USA)</u>: is a highly purified lipid of a single molecule. The reagent forms cationic liposomes that complexes with the negatively charged DNA molecule. The DNA-HilyMax complexes electrostatically bind to the cell's membrane and the DNA molecules enter the cells by endocytosis.
- <u>NIMT®FeOfection Yellow Transfection Reagent (GENOVIS, SWEDEN)</u>: This transfection reagent is composed of superparamagnetic nanoparticles. The nanoparticles are composed of an iron oxide core and a positive surface. This composition facilitates the formation of non-covalent bonds between the nanoparticles and the DNA, and the resultant particle/DNA complexes are then internalized into the target cell lines.
- CalPhos Mammalian Transfection kit (Clontech): The action of this transfection reagent depends on the formation of calcium phosphate precipitate. In this method, a saline buffer 2x HEPES containing the phosphate ions, negatively charged, is combined with the DNA plasmid calcium solution, positively charged. This finally results in the production of a fine precipitate that is taken up by the target cell lines when added.
- <u>Nanofectin Kit (PAA)</u>: This transfection reagent is formed of two components, a positively charged polymer that has the capacity to bind to DNA and a porous nanoparticles. The unique size of the DNA-nanoparticle complex favours the

uptake of the complex by the target cells. In addition, the complex aids to protect the DNA from nuclease degradation.

To determine the amount of reagent needed for DNA transfection for each of the above transfection reagents, the pEGFPN-1 vector was initially used followed by the pB12mcs-U6-eGFP vector. The optimization reactions were conducted according to each manufacturer's protocol.

## 2.4.2.2 Non-chemical based transfections

Several non-chemical based transfection methods exist, however, only nucleofection, a subset of electroporation, was considered. The latter is a transfection method that depends on the discharge of electrical impulses that leads to the creation of transient micro-pores in the membrane of the target cell line. Nucleofection is a transfection method first developed by Amaxa to target cell lines that are difficult to transfect, such as primary cell lines, cells in suspension and non-dividing cells.

In nucleofection, the DNA molecules are directly delivered into the nucleus of the target cell. This requires both a nucleofector device and a cell-type specific nucleofector reagent. The Amaxa<sup>®</sup> Cell Line Nucleofector<sup>®</sup> Kit V and the Amaxa nucleofector device, Germany, were used. The control pmaxGFP<sup>®</sup> vector, supplied with the kit, was used to determine the appropriate nucleofection program to be used on the K562 and KCL22 cell lines. Furthermore, the pEGFPN-1 was initially used to determine the amount of plasmid DNA needed to optimize the nucleofection reaction. Once that was achieved, the constructed pB12mcs-U6-eGFP vector was used to optimize the amount of plasmid DNA needed to transfect the target cell lines. All nucleofection reactions were conducted according to the manufacture's protocol.

To assess the efficiency of nucleofection a 6-well plate with the transfected cell lines was observed under a fluorescence microscope (Nikon, UK). Once,

fluorescence was detected, the cells were counted and  $3 \times 10^5$  cells were fixed as described below for FACS analysis.  $1 \times 10^6$  cells were used for RNA extraction and  $1 \times 10^6$  cells were used for western blotting.

## 2.5 Fluorescence-Activated Cell Sorter (FACS) Analysis

FACS scan is a type of flow cytometry that involves the analysis and sorting of single cells or particles, in this case the K562 and KCL22 cell lines, depending of their physical and chemical properties. The cells of interest are suspended in a narrow precisely defined liquid stream also known as the sheath buffer. When the cell sample is passed through the sheath buffer, the cells will undergo hydrodynamic focusing. This type of pressure causes the cells of interest to be individually separated as single droplets that flow through the machine's cuvette. Furthermore, a laser beam of a single wavelength is directed on each cell in the sample. The laser beam passing through the flow stream causes the illuminating light to be scattered and fluorescence will be emitted. Two types of light scattering exist, forward scatter and side scatter. The first is linked with the cell's volume, whereas the second is more dependent on the internal properties of the cell. All light and fluorescent emissions are collected and directed to several special detectors. The detectors will proportionally convert these light pulses into electrical pulses, which can then be amplified by a linear or logarithmic amplifier. The electrical pulses can also be digitized for further computer processing and storage (Robinson, 2004). In the current work the cytometer used was a FACScan from Becton Dickinson, UK. Figure 2.7 illustrates the basic principles of the FACS scan.



Figure 2.7 The basic principle of the FACS scan. (Source <u>www.abcam.com</u>)

This figure illustrates the principles of the FACS scan. The cells of interest can be stained with a specific dye or express a fluorescent protein, for example eGFP. The sample cells are passed through a stream of liquid called the sheath fluid/buffer and through the process of hydrodynamic focusing, the cells will be separated into individual droplets that pass through a thin narrow glass cuvette. Once in the cuvette, a laser beam is directed in each individual cell, this causes the beam to separate into scattered light, in forward and side scattered, and emitted fluorescence. The emission is collected through a detector that will proportionally convert the collected light pluses into digitalized electrical pulses that can be further analysed by computer software.

As discussed previously, the K562 and KCL22 cell lines were transfected with the pB12mcs-U6-eGFP vector. The plasmid expresses eGFP that can be detected by FACS. The cell lines where centrifuged at 1,000 rpm (Beckman TJ-6 centrifuge, Ireland), the supernatant discharged and the pellet was washed with phosphate buffer saline (PBS) (Invitrogen, UK). The pellet was then resuspended in 500µl of PBS and the cells fixed by the addition of 500µl of 4% paraformaldehyde (4g PFA, (Sigma, UK) dissolved in 100ml of PBS, heated up to 60°C with moderate stirring, and the pH was adjusted between 7.2-8. Once the solution was clear, the heat was turned off, the solution was cooled down, filtered, and aliquoted into 15ml tubes. The solution was then stored at -20°C). The parameters were set for the type of cells used in this project, the K562 and KCL-22, and the cell suspensions were analysed using the FACS machine. The obtained data was analysed by the use of the computer software "CellQuest Pro" (Becton Dickinson, UK), and the collected data was plotted in a single dimension to produce a histogram. A twodimension plot in the form of a dot plot was also used, this was done when acquiring the data to ensure that the initial control sample fell in one group of cells. Figure 2.8 shows a graph generated from the CellQuest Pro software in the form of a histogram.



#### Figure 2.8 FACS Histogram Graph.

This figure shows a Histogram graph that was generated through the use of the "CellQuest Pro" software. The Histogram shows two samples, a control sample, in purple (this sample represents cells that have not been treated by any means), and a test sample overlaid in green. Both samples contain the same number of cells (10,000). The M1 Line corresponds to the gated cells. The histogram above shows that above 52% of cells have been transfected successfully.

#### 2.6 Real Time PCR

Real time PCR or quantitative PCR (qPCR) is a highly sensitive technique that allows simultaneous amplification and accurate quantification of a specific nucleic acid sequence from DNA or cDNA targets. PCR products in this technique are detected through using either fluorescent dyes, which bind to double stranded DNA, or by using fluorescently labelled probes that are sequence specific. Consequently, quantification is achieved by a determination of the cycle number at which the sequence of interest is first detectable.

In this study, real time PCR was used for gene expression analysis, the Roche Light Cycler 480 (Roche, UK) was used and PCR product quantification was done through the use of SYBR Green I and Taqman probes. SYBR Green I is a dye that binds to any double stranded DNA molecule, and emits a fluorescent signal of a particular wavelength. The detection of its fluorescence is achieved at the extension phase of the qPCR. Then with the accumulation of the double stranded PCR products, an increase of the fluorescent signal is also detected at the end of each cycle. This is due to the fact that the amount of fluorescence/signal is proportional to the amount of the accumulating PCR product. Figure 2.9 illustrates the basic principles of real time PCR when conducted using SYBR Green I.

Taqman probes also called hydrolysis probes are sequence specific oligonucleotides that carry a fluorophore moiety at the 5' end and a quencher moiety at the 3' end. Once the probe is cleaved by the 5'  $\rightarrow$  3' exonuclease activity of *Taq* polymerase (Holland P.M *et al.*, 1991), the fluorophore and the quencher moiety will be separated and a fluorescent signal will be detected. Again the signal is proportional to the amount of PCR product being accumulated. This process occurs during the annealing/extension phase of the qPCR. Figure 2.10 illustrates the basic principles of real time PCR using the Taqman probes. Taqman primers and probes were designed by the use of Roche Universal probe library (UPL) software and Primer3 software. Table 2.7 shows all Taqman primers and probes used in this project. The same primers were used without the probes when conducting real time PCR using SYBR Green I.



Figure 2.9 Real time PCR using SYBR Green I methodology.

SYBR Green dye binds to any double stranded DNA regardless its sequence. The dye will only fluoresce once bound to DNA and is excited via blue light. In (A) the DNA is denatured and the synthesis of double stranded DNA starts in (B). Once the double stranded DNA is being synthesised the SYBR Green dye starts to bind to the newly synthesised DNA molecules. The fluorescent signal increases to its maximum when the DNA synthesis is complete and the maximum amount of dye is bound (C). Hence, the fluorescent signals in this type of qPCR are measured at the end of each elongation phase.



Figure 2.10 Real time PCR in the presence of Taqman probes.

Taqman probes or hydrolysis probes are sequence specific that bind to DNA at a specific sequence, the probe has both a dye at its 5' end and a fluorescence quencher on its 3' end. In (A) the DNA is being denatured and the probe binds to its target site in (B) while DNA amplification begins. As the primer is being extended and the double stranded DNA is being synthesised to a point where it reaches the Taqman probe, the 5' nuclease activity of the polymerase enzyme will cleave the probe, this releases the dye and a fluorescent signal can be detected. This is demonstrated in part (C). Finally, in (D) the DNA synthesis is complete. The fluorescence in this type of qPCR is measured at the end of the elongation phase.

Taqman primer sets		Primer and probe sequences	
GAPDH	Forward	5' d-GAAGGTGAAGGTCGGAGTC'3 (NM_002046.3)	
(226bp)	Reverse	5' d-GAAGATGGTGATGGGATTTC'3 (NM_002046.3)	
	Probe	5' d-[6FAM]CAAGCTTCCCGTTCTCAGCC[BHQ1]'3	
ABL gene*	Forward	5' d-GGGCTGCAAATCCAAGAAG'3 (NM_005157.3)	
(77bp)	Reverse	5' d-ATGCTACTGGCCGCTGAA'3 (NM_005157.3)	
	Probe	UPL probe number 65	
BCR gene*	Forward	5' d-TCAGCCACTGGATTTAAGCAG'3 (NM_021574.2)	
(91bp)	Reverse	5' d-CGCGCGTCTTTGCTTTAT'3 (NM_021574.2)	
	Probe	UPL probe number 10	
$\beta 3\alpha 2$ fusion	Forward	5' d-GTCCACTCAGCCACTGGATT'3 (AJ131466)	
point	Reverse	5' d-TGTTATCTCCACTGGCCACA'3 (AJ131466)	
(190bp)	Probe	5' d-[TAM]AACTCCAAGGAAAACCTTCTCGCTGGACCC[BHQ2]'3	
$\beta 2\alpha 2$ fusion	Forward	5' d-GATGCTGACCAACTCGTGTG'3 (AJ131467)	
point	Reverse	5' d-GTTGGGGTCATTTTCACTGG'3 (AJ131467)	
(183bp)	Probe	5' d-[TxRd]TCGTTGGAACTCCAAGGAAAACCTTCTCGCT[BHQ2]'3	

**Table 2.7 Taqman primers and probe sets used in the project.** This table summarises all Taqman primer sets and their appropriate probes that have been used in this study. The sequence of each set is shown. Primer sets marked with (\*) were designed by the use of the UPL software (<u>http://www.roche-applied-science.com/sis/rtpcr/upl/ezhome.html</u>), while unmarked sets were designed using primer3 output software. The PCR product band sizes are shown in column 1 and the accession numbers of the transcripts are shown between brackets in column 3.

For qPCR reactions the Roche LightCycler<sup>®</sup> 480 SYBR Green I Master and the Roche LightCycler<sup>®</sup> 480 Probe Master mixes were used. Reactions were conducted on a 96 well plate (Roche, UK), with each qPCR reaction repeated in duplicate. The 96 well plate was covered with an adhesive seal and the reactions were conducted under the selection of the SYBR Green I or the multihydrolysis probe program with the SYBR Green or the Taqman template respectively. The parameters for the real time PCR are shown below in table 2.8 and the contents of each qPCR reaction can be found in table 2.9 and 2.10. Genes analysed by qPCR post transfection included the *GAPDH* gene, a house keeping gene, the normal *ABL* and *BCR* genes, to see whether or not the shRNA constructs have an effect on the normal genes, and the abnormal fusion genes. To achieve this, the first step was to generate standard curves for each gene. These curves are then used to determine the cross point value of the cDNA levels of the genes of interest in the transfected cell lines. The concentrations of each gene in ng were obtained from the Roche Lightcycler 480 software its self. The resultant concentrations where normalized with GAPDH. Statistical analysis was then conducted on the normalized data. The student t-test was used to calculate significance. Finally, the percentage of mRNA expression and depletion was calculated. "Microsoft Excel" was used to conduct all calculations.

### (A) Parameters for the Multi-hydrolysis probe program:

One cycle of initial denaturation at 95°C for 5 minutes		
	Denaturation at 95°C for 45 seconds	
40 cycles of amplification $\rightarrow$	Annealing temperature at 60°C for 1 minute	
(Quantification)	Extension at 72°C for 1 second	
One cycle of cooling at 40°C		

(B) Parameters for the SYBR Green I program:

One cycle of initial denaturation at 95°C for 5 minutes				
	Denaturation at 95°C for 15 seconds			
40 cycles of amplification $\rightarrow$	Annealing temperature at 60°C for 10 seconds			
(Quantification)	Extension at 72°C for 30 seconds			
	A single acquisition mode at 80°C for 1 second			
One cycle of Melting that goes up to 95°C for 1 second then down to 60°C for				
another 1 second				
One cycle of cooling at 40°C				

Table 2.8 Parameters used for real time PCR conducted in this project. (A) Shows theparameters used for real time PCR using the multi-hydrolysis probe program, while (B) showsthe real time PCR parameters used when the SYBR Green I program was conducted.

Component	Volume per 20µl reaction
cDNA	2-4µl
$10\mu M$ of Forward primer	1µl
10μM of Reverse primer	1µl
0.2μM of probe	1µl
Roche LightCycler <sup>®</sup> 480 Probe Master mix	10µl
Nuclease free H <sub>2</sub> O	xμl

**Table 2.9 Components of the qPCR Master Mix (20μl reaction mixture).** The preparation of master mix solution was done on ice.

Component	Volume per 20µl reaction
cDNA	2-4µl
$10\mu M$ of Forward primer	1µl
10μM of Reverse primer	1µl
Roche LightCycler <sup>®</sup> 480	10µl
SYBR Green I Master mix	
Nuclease free H <sub>2</sub> O	xμl

Table 2.10 Components of the qPCR Master Mix (20µl reaction mixture). The preparation of master mix solution was done on ice.

# 2.7 Western Blot Analysis

Western blots are an immunoanalytical method used for the purposes of protein analysis. The technique depends on antigen-antibody interactions in order to detect certain proteins of interest. The above technique allows a quantitative as well as a qualitative analysis of proteins.

## 2.7.1 Cell lysis and protein extraction

 $1 \times 10^{6}$  cells (K562 or KCL-22) were pipetted into a 1.5ml Eppendorf tube, washed once with PBS, then a pellet was obtained by centrifugation of the cells for 5 minutes at 14,000 g at 4°C. The supernatant was removed and 100µl of cell lysis buffer (Table 2.11) was added and the cells lysed by pipetting vigorously several times. The lysed samples were then kept on ice for 30 minutes, centrifuged for 5 minutes at 14,000 g at 4°C and the supernatant, containing the extracted proteins, was pipetted to a new Eppendorf. The lysate was then mixed with 1x sample Loading buffer (Table 2.11) in a 1:1 ratio and the samples were stored at -20°C.

## 2.7.2 SDS-Polyacrylamide Gel Electrophoresis

A ready prepared 12% polyacrylamide gel (Serva Electrophpresis, Germany) was placed in d was filled with 1 x Running buffer, Table 2.11 demonstrates the components of all buffers and solutions that are used during the running of the polyacrylamide gel and the Western blotting protocol. To denature the extracted proteins, previously described in section 2.7.1, the protein samples were placed in a heating block at 100°C and were allowed to boil for 3-4 minutes. The samples were left to cool down, and then 20-30µl was loaded onto the gel. 10µl of high range Rainbow marker (GE Healthcare, UK), see Appendix 3, was also loaded. The gel was then run at 100V for 90 minutes or when samples reached to the bottom of the gel.

Solution Name	Components	Storage Conditions
Lysis Buffer	50mM Tris-HCL (Sigma, UK)	Once the protease
	1% Triton	inhibitor tablet was
	150 NaCL (Sigma, UK)	added the solution was
	1mM EDTA (Sigma, UK)	stored at -20°C for 12
	- For every 10ml of buffer one	weeks and at 4°C for 2
	tablet of complete, mini, EDTA-free	weeks.
	protease inhibitor cocktail tablets	
	(Roche, UK) was added.	
Sample Loading	62.5mM Tris-HCL pH 6.8	4°C
buffer	25% Glycerol	
	2% SDS	
	0.01% Bromophenol Blue(Bio-RAD)	
	5% $\beta$ -mercaptoethanol	

Table 2.11 Components of the Lysis buffer and the Sample loading buffer.The Sample loadingbuffer is also known as the Laemmli loading buffer.

# 2.7.3 Western Blotting

The cassette containing the polyacrylamide gel was cracked open, the wells removed and the gel washed once with Milli Q water for 5 minutes. The gel was then covered with 1x Transfer buffer and placed on a rotary shaker for 20 minutes. The PVDF transfer membrane was treated according to the manufacturer's instructions (Thermo Scientific, UK), and the transfer cassettes, sponge pads, 6 pieces of Whatman paper were placed in a large container and equilibrated in 500ml of 1x transfer buffer for a few minutes.

The transfer sandwich was assembled, placed in the transfer tank with a block of ice and a magnetic stirring pin. The tank was placed on a magnetic stirrer, the whole apparatus was placed in a cold room at 4°C, and the transfer was conducted overnight at 20-30V.

Once the transfer was completed, the membrane was washed once with Milli Q water. The membrane was then dipped for a couple of seconds in 50ml of 1% (w/v) Ponceau S solution in 5% acetic acid (Sigma,UK). Once the proteins were observed, the membrane was scanned and the image was kept to assist the loading of the samples. The membrane was washed with TBS-Tween, washing buffer, for 10 minutes. Then the membrane was blocked with blocking buffer (see Table 2.12 for the composition of all solutions used in the processes of SDS-PAGE electrophoresis and Western Blotting) for 1 hour at room temperature.

The blocking buffer was then discharged and the membrane was incubated with the primary antibody, diluted in 15ml of blocking buffer, overnight at 4°C on a rotary shaker (Table 2.13 illustrates all antibodies used for the Western blots conducted in this study). The following day, the primary antibody was discharged and the transfer membrane was washed 3 times with washing buffer, 5 minutes each time at room temperature. The membrane was then incubated with the secondary antibody, again diluted in 15ml of blocking buffer, for 1 hour at room temperature. The secondary antibody was then removed and the membrane was washed for another 3 times in washing buffer, 5 minutes each time at room temperature. Finally, the blots were developed using the Pierce<sup>®</sup> ECL Western Blotting Substrate Kit (Thermo Scientific, UK). This was done according to the manufacturer's protocol. The membrane was then placed in a film cassette, in such a way that the treated protein side of the PVDF membrane can be exposed to the X-ray film (Fujifilm, UK). In the dark room and under a red safe light, the X-ray film was cut, to the appropriate size of the membrane, placed on the treated side of the membrane and the cassette was closed. The time of exposure varied from 1-60 minutes, depending on the intensity of the signal. Band intensities were analysed through the program "ImageJ", and all calculations were done by the use of "Microsoft Excel" software.
Solution name	Components	Additional
		Information
10x Running Buffer	25mM Tris-HCL	The buffer was
(Laemmli Buffer)	200mM Glycine	diluted to 1x before
	0.1% (wt/v) SDS	use
	- $dH_2O$ as required	
1x Transfer Buffer	100ml of:	
(1L)	1.92M Glycine	
	0.25M Tris-HCL	
	200ml of: MetOH	
	- 700ml of $dH_2O$	
TBS-Tween	1M Tris-HCL pH6.8	
(Wash Buffer)	5M NaCL	
	1% Tween	
	dH <sub>2</sub> O as required	
Blocking Buffer	5% (wt/v) Non Fat Milk (Marvel) in	
	1x TBS-Tween	
	5% (wt/v) BSA (Sigma) in 1x TBS-	
	Tween	

Table 2.12 Recipes of the solutions used in the process of SDS-PAGE electrophoresis andWestern Blotting. The above table shows the composition of all the buffers used for of SDS-PAGE electrophoresis and Western Blotting.

Primary	Dilution	Secondary Antibody		Dilution	Blocking	Buffer
Antibody	used			used	used	
c-ABL	1:1000	Anti-Rabbit	Secondary	1:1000	5% (wt/v)	BSA in
(Cell Signalling)		Antibody			1x TBS-Tw	een
$\beta 2\alpha 2$ junction	1:1000	Anti-Mouse	Secondary	1:1000	5%(wt/v)	Non
specific		Antibody			Fat Milk	in 1x
(Cell Signalling)					TBS-Twee	n
GAPDH	1:1000	Anti-Rabbit	Secondary	1:1000	5% (wt/v)	BSA in
(Cell Signalling)		Antibody			1x TBS-Tw	een

Table 2.13 A list of the primary and secondary antibodies used in the Western Blots in this

project. Antibodies used were all purchased from Cell signalling.

# Chapter 3 : Generation of a Reporter gene system for expression of shRNAi cassettes targeting the BCR-ABL fusion points

### 3.1 Introduction

The *BCR-ABL* fusion genes result from a reciprocal translocation between chromosomes 9 and 22. The two most common resultant fusion points associated with CML are the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  variants. These variants are characterised by an abnormally up-regulated Tyrosine kinase activity that causes the development of CML (Advani and Pendergast, 2002). Consequently, targeting of these fusion points as a therapeutic approach for CML could be rewarding.

In this project, short hairpin RNA molecules have been designed to target the two BCR-ABL variants. To achieve this, a reporter gene system was needed to monitor the synthesis and expression of the shRNA constructs. The shRNA oligonucleotides were designed and constructed in such a manner that they were under the regulation of either the human U6 or H1 promoters. The constructed cassettes were further incorporated into an expression vector that had the gene for the enhanced green fluorescent protein (eGFP). eGFP is a mutated form of GFP that has been optimised to have a brighter fluorescence and a higher expression in mammalian cells than GFP (Siemering K. R et al., 1996), and for the purposes of this project eGFP acted as a reporter gene for the shRNA constructs. Two plasmids were considered in order to generate the reporter gene system. The first was the pEGFP-N1 plasmid; however by using this plasmid it proved difficult to construct the system. The second plasmid was the pB12mcs-eGFP plasmid and constructing the reporter gene system in this plasmid was successful. This chapter shows the results obtained from attempts to construct such a system.

#### 3.2 Results

### **3.2.1** Sequencing of the $\beta$ 3 $\alpha$ 2 and the $\beta$ 2 $\alpha$ 2 Fusion Points

The  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  fusion genes present in the K562 and KCL-22 cell lines used in this study were sequenced. This step was done to verify that the sequences of the fusion points matched the sequences found in the EMBL website, accession numbers AJ131466 and AJ131467 respectively (Appendix 4).

# 3.2.1.1 PCR Amplification of the $\beta 3\alpha 2$ and the $\beta 2\alpha 2$ Fusion Points

Total RNA was extracted from K562 and KCL-22 cell lines. The RNA was quantified and a reverse transcriptase reaction was conducted. To determine whether the reaction worked, a PCR reaction using  $\beta$ -actin primers K, L and N, sequences found in Table 2.2, was conducted on the synthesised cDNA of both cell lines. When using the  $\beta$ -actin primers K and L as a pair, the  $\beta$ -actin gene PCR product is 635bp, whereas pairing primers N and L gives rise to a PCR product of 512bp. The last set of primers only works if genomic DNA is present in the sample; this is due to the fact that primer N is intronic. Figure 3.1 shows the results of the PCR reaction conducted on the synthesised cDNA from the K562 cell lines and shows that cDNA synthesis is successful with no genomic DNA contamination. Similar results were obtained with the KCL-22 cell line.



Figure 3.1 Electrophoretic Analysis of PCR reaction using  $\beta$ -actin primers on K562 cDNA.

Lane 1:  $\lambda$ /EcoRI/HindIII marker

- Lane 2: PCR products of the  $\beta$ -actin primers K and L
- Lane 3: PCR products of the β-actin primers N and L (negative control)

 $\beta$ -actin PCR products were analysed via electrophoresis using a 1% agarose gel .The 635bp band in lane 2 indicates that the RNA purification and K562 cDNA synthesis was successful. Lane 3 is the negative control and indicates the absence of any genomic DNA contamination.

Once the synthesis of cDNA was confirmed for the K562 and KCL-22 cell lines, a PCR reaction using primer set  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$ , sequences found in Table 2.2, was conducted. This PCR was done to amplify the region around the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  fusion points, and PCR products of 381bp and 522bp in size are expected to be produced, respectively. The PCR products are shown in Figures 3.2 and 3.3. The  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  PCR products were then purified, quantified, and 500ng was used for AT cloning. The remaining  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  PCR products were stored at -20°C.



#### Figure 3.2 Electrophoretic Analysis of the $\beta 2\alpha 2$ PCR product from KCL-22 cDNA.

Lane 1: pBluescript Hpa II marker Lane 2: PCR products Lane 3: repeat of lane 2

 $\beta 2\alpha 2$  PCR products were analysed via electrophoresis on a 2% agarose gel. The bands present in lanes 2 and 3 fall in the size range of interest, that is 522bp. This indicates that the KCL-22 cell line contains the  $\beta 2\alpha 2$  fusion point.





Lane 1: pBluescript Hpa II marker Lane 2: PCR products Lanes 3 and 4: repeat reactions

 $\beta 3\alpha 2$  PCR products were analysed via electrophoresis on a 2% agarose gel. The bands present in lanes 2-4 fall in the size range of interest, that is 381bp. This indicates that the K562 cell line contains the  $\beta 3\alpha 2$  fusion point.

# 3.2.1.2 AT Cloning of the $\beta$ 3 $\alpha$ 2 and the $\beta$ 2 $\alpha$ 2 PCR Products in the pGEM®T Easy Vector System

The amplified and purified  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  PCR products were cloned in the pGEM®T easy vector system. This was done through the process of AT cloning. The PCR products have an 'A' overhang, and the pGEM®T easy vector has a complimentary 'T' overhang in its multiple cloning region (MCR) for easy insertion of the PCR products.

# 3.2.1.3 Analysis of the $\beta 3\alpha 2$ and the $\beta 2\alpha 2$ Fusion Point Sequences

The sequences of the cloned  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  fusion points were analysed and compared to those sequences found in the EMBL website, accession numbers AJ131466 and AJ131467 respectively, Figure 3.4. The results confirmed that the sequences of the fusion points found in the K562 and KCL-22 cell lines used in this study matched their sequences found in the EMBL website. The full sequences of  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  fusion points are shown in Appendix 4, and the raw sequencing data are shown in Appendix 5.



#### Figure 3.4 The K562 and KCL-22 Fusion Point Sequencing Results

(a) Top: Boxed and underlined is the  $\beta 3\alpha 2$  fusion point sequence found in the EMBL website (EMBL/AJ131466)

Bottom: Boxed and underlined is the sequencing result of the  $\beta 3\alpha 2$  fusion point sequence found in the K562 cell line used in this study.

(b) Top: Boxed and underlined is the  $\beta 2\alpha 2$  fusion point sequence found in the EMBL website (EMBL/AJ131467)

Bottom: Boxed and underlined is the sequencing result of the  $\beta 2\alpha 2$  fusion point sequence found in the KCL-22 cell line used in this study.

# 3.2.2 Preparation of the pEGFP-N1 Vector for Cloning of the shRNA Cassettes

The first step was to remove the CMV promoter. Once the promoter was removed the prepared vector was self-ligated, creating a negative control. The vector was subsequently used for cloning the human U6 promoter (positive control), the human H1 promoter (positive control), and the shRNA cassettes.

# 3.2.2.1 Digestion and Removal of the CMV Promoter from the pEGFP-N1 Vector

For total removal of the CMV promoter, the restriction enzyme Asel could be used with any other restriction site found in the MCR. However the restriction site of the Asel enzyme is also found within the shRNA cassettes. So, instead the enzyme SnaBI, which has a site within the CMV promoter, and does not cut within the shRNA cassettes, was used.

2μg of the pEGFP-N1 vector was diluted and then digested with SnaBI enzyme for 3 hours at 37°C with the appropriate buffer, Figure 3.5 gel (a) lane 2. The vector was then precipitated, resuspended in Milli Q water and further digested with HindIII for another 3 hours at 37°C with the appropriate buffer. The digested vector was then precipitated, resuspended in Milli Q water and was run on a 1% agarose gel, Figure 3.5 gel (b) lane 3. The double digestion of the SnaBI and HindIII sites should give products of 283bp and ~4.4Kb. The presence of the 283bp band is clearly seen confirming that the double digestion has worked. The digested vector, band size of ~4.417Kb, was cut out of the gel, and was purified by the Wizard<sup>®</sup> SV kit. The vector was then quantified.





Gel (a) Lane 1: λ/EcoRI/HindIII marker Lane 2: pEGFP-N1 vector digested with SnaBI Lane 3: Wild type pEGFP-N1 vector Gel (b) Lane 1: λ/EcoRI/HindIII marker Lane 2: Wild type pEGFP-N1 vector Lane 3: pEGFP-N1 vector digested with SnaB1 and HindIII

To determine the success of the pEGFP-N1 digestion by the restriction enzyme SnaBI,  $1\mu$ l of the digested vector was run on a 1% agarose gel. Gel (a) lane 2. A single band, of a size approximately 4.7Kb is seen compared with the undigested wild type vector in lane 3. Gel (b), lane 3, two bands of approximately 283bp and 4.4KB are present, thus confirming that both digestions have worked. Lane 2 is the wild type vector that was run as a negative control.

## 3.2.2.2 Self Ligation and Cloning of the Modified pEGFP-N1 Vector

The ends generated from the SnaBI restriction enzyme are blunt ends, whereas the ends generated from the HindIII restriction enzyme are sticky with a couple of bases overhang, therefore the ends are incompatible for joining together. To resolve this issue the sticky ends were converted to blunt ones. A number of strategies were tried in order to achieve this self ligation.

The sticky ends of a vector can be either filled in with complementary bases or the overhang bases can be digested. To fill in the overhang created was considered including strategies such as filling the gap with *Taq* polymerase or treating the ends with the large Klenow fragment, and the Klenow fragment  $(3' \rightarrow 5' \text{ exo-})$ . *Taq* polymerase was first used to try to fill in the sticky ends. Several attempts were conducted but the vector failed to self ligate, indicating that the reaction was not successful. The large Klenow fragment retains its  $5' \rightarrow 3'$  polymerase activity, and its  $3' \rightarrow 5'$  exonuclease activity, but lacks its  $5' \rightarrow 3'$  exonuclease activity. The pEGFP-N1 plasmid was treated with this enzyme several times under different time and temperature conditions. Again the treated vector failed to self ligate. Finally, the Klenow fragment  $(3' \rightarrow 5' \text{ exo-})$  that retains its  $5' \rightarrow 3'$  polymerase activities was used and proved successful.

After treatment with the Klenow fragment  $(3' \rightarrow 5' \text{ exo-})$ , the treated pEGFP-N1 plasmid was purified and 500ng were used for self ligation. The self ligation reaction was done for an hour at room temperature, and overnight at 16°C and 4°C. The reaction was analysed by PCR using CMV-207 forward and EGFPN1 reverse primers. PCR of the wild type vector was included as a control. When using this set of primers a band of 530bp is expected in the wild type vector, whereas a band of 219bp is expected in the double digested, self-ligated vector. The results of this PCR can be seen below in Figure 3.6.





Lane 1: 100bp DNA Ladder from NEB

Lane 2: PCR of wild type pEGFP-N1 vector using CMV-207 forward and EGFPN-1 reverse Lane 3: PCR of self-ligated pEGFP-N1. The reaction was done at 16°C Lane 4: PCR of self-ligated pEGFP-N1. The reaction was done at 4°C

A 2% agarose gel was used to analyse the PCR products of the wild type and self-ligated pEGFP-N1 vectors. Lane 2 has shown the PCR product of the wild type vector, with a single band of ~530bp. Whereas, lane 4, shows a band ~219bp that corresponds to the double digested, self-ligated pEGFP-N1 vector.

Half the amount of the ligation mix was used to transform 50µl of JM109 bacterial cells. The cells were then plated on agar plates with the appropriate antibiotic and were kept overnight in an incubator at 37°C. The colonies on the plates were screened by means of PCR. The PCR was done using the set of primers CMV-207 forward and EGFPN1 reverse. However, all screened colonies were of wild type vector. Therefore, a higher concentration of ligation mix was used to transform bacterial cells. This also resulted in the growth of wild type colonies only, so the ligation reaction was digested with the enzyme Xhol. This was done to decrease the chances that a wild type vector would be taken into the bacterial cells, as an Xhol site is only present in the wild type vector and not in the self-ligated one. This experiment was repeated three times, and in all three no bacterial colonies grew on the agar plates. Consequently, work on

this vector was stopped and another vector was selected to generate a reporter system

#### 3.2.3 Preparation of the pB12mcs-eGFP Vector

This vector was selected to construct the reporter gene system for the shRNA cassettes after work on the pEGFP-N1 vector has been stopped. The vector has an eGFP gene and no promoter. Therefore, the wild type vector was used as the negative control.

### 3.2.3.1 Sequencing of the pB12mcs-eGFP Vector

The plasmid was sent for sequencing using the T7 forward primer and the T3 reverse primer and a new primer called GFPneo. The sequencing results confirmed that the plasmid matched its map shown in materials and methods chapter. However, some variation existed in the MCR; there was no EcoRI restriction site but an EcoRV instead. Based on the sequence analysis, the restriction sites for the enzymes HindIII and BamHI were selected, and the next step was to start preparing the pB12mcs-eGFP plasmid for insert intake.

# 3.2.3.2 Digestion and Purification of the pB12mcs-eGFP Vector

50µl of JM109 bacterial cells were transformed with 50ng pB12mcs-eGFP. The transformed bacterial cells were grown overnight in 500ml of LB with the appropriate antibiotic. 1ml of the grown bacteria was stored at -80°C as glycerol stock and then a maxi prep took place and the extracted plasmid was quantified. 2µg of the pB12mcs-eGFP vector was diluted and digested with BamHI enzyme. To confirm digestion, the digested plasmid was then analysed

through gel electrophoresis, Figure 3.7. The digested vector was then purified, further digested with the enzyme HindIII, purified and quantified.

The double digestion of the BamHI and HindIII sites should give products of 62bp and 3.887Kb. The presence of the 62bp band cannot be seen when the digestion reaction is analysed on the 1% agarose gel. Only the 3.887Kb band can be seen. Therefore, to ensure that the restriction enzyme HindIII digests the pB12mcs-eGFP plasmid efficiently, 2µg of the wild type pB12mcs-eGFP vector was freshly diluted, and was then digested separately with the HindIII enzyme. 5µl of this digestion was then analysed through electrophoresis. The digested pB12mcs-eGFP plasmid appeared as a single band, in comparison to the supercoiled wild type plasmid, hence confirming that the digestion conditions were correct (results are not shown).



Figure 3.7 Electrophoretic Analysis of the digested pB12mcs-eGFP vector

Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder

Lane 2: Wild type pB12mcs-eGFP vector

Lane 3: BamHI digested pB12mcs-eGFP vector

A clear single band of ~3.9Kb in size can be seen in lane 3, upon comparison with the multiband wild type vector observed in lane 2. The multiband appearance is the result of the wild type vector being supercoiled. Overall, the gel indicates digestion of the pB12mcs-eGFP vector with the enzyme BamHI.

#### 3.2.4 Amplification of the U6 and H1 Promoters

Once the pB12mcs-eGFP plasmid was digested and purified, the next step was to construct the positive controls by incorporating the U6 human promoter or the H1 human promoter in the MCR of the digested pB12mcseGFP plasmid. The sequences of both promoters were found through the EMBL website and the sequences had accession number M14486 for the U6 promoter, and accession number X16612 for the H1 promoter. The complete sequences of these promoters can be found in Appendix 6.

From the sequences found in the website, a U6 primer set and an H1 primer set were designed. The primer sequences were shown in Table 2.2. These primers were designed in such a way that a restriction site for the enzyme HindIII was incorporated in the forward primers, and a restriction site for the BamHI enzyme was incorporated in the reverse primers. PCR using these sets of primers was conducted on cDNA from K562 cell lines as a template. The size of the PCR products for these reactions was expected to be 292bp for the U6 promoter, and 219bp for the H1 promoter. Figure 3.8 demonstrates the electrophoretic analysis of the PCR products of the U6 and H1 human promoters respectively.

The PCR products of the U6 and H1 human promoters were purified, digested with the enzyme BamHI for 3 hours at 37°C with the appropriate buffer. The digested promoters were then precipitated and digested again for another 3 hours at 37°C with the HindIII enzyme and its appropriate buffer. Once, this digestion was achieved, the promoters were precipitated, resuspended in 10µl of Milli Q water, and were then quantified.



#### Figure 3.8 Electrophoretic Analysis of the PCR products of the U6 and H1 human promoters

(a) Top:

Lane 1: 100bp DNA Ladder from NEB Lanes 2-5: U6 promoter PCR products from four reactions (b) Bottom: Lane 1: 100bp DNA Ladder from NEB Lanes 2-5: H1 promoter PCR products from four reactions

The U6 promoter is 292bp in size, whereas the H1 promoter is 220bp. In lanes 2-5 in gel (a), bands of ~300bp are present, whereas in lanes 2-5 in gel (b), bands just above 200bp are present. These findings confirm that the PCR products are of the size expected.

## 3.2.4.1 Cloning of the U6 and H1 Promoters in the pB12mcseGFP Vector

A 1:10 vector to insert ratio was used to clone the prepared promoters into the double digested pB12mcs-eGFP vector. The ligation reaction was done overnight at 4°C in a total volume of 10µl. Half the volume of the ligation reaction was used to transform 35µl of JM109 bacterial cells. The cells were heat shocked and were left in a shaking incubator at 37°C for 2 hours before being plated on agar plates with the appropriate antibiotic.

The plate was examined for the growth of bacterial colonies and colonies were screened by means of PCR T7 forward and T3 reverse primers. The wild type pB12mcs-eGFP vector was also included as a negative control. PCR products were analysed by electrophoresis, and are illustrated in Figures 3.9 and 3.10. Positive colonies were grown overnight in 20ml of LB with the appropriate antibiotic in a shaker incubator.





Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder Lane 2: Wild type pB12mcs-eGFP vector PCR product Lanes 3-13: Colony screen PCR products Lane 11: Positive Colony

The primer set T7 forward and T3 reverse was used for the above colony screen and the expected PCR product for the wild type vector is ~1050bp, lane 2. The wild type vector was included as a negative control. Lanes 3-10 and 12-13 all have PCR products matching the size of the wild type vector, and were considered to be negative screens. However, in lane 11, highlighted in red, a band of size ~1310bp (~1050bp for the wild type and 292bp for the U6 promoter) is present. This colony was considered to be a positive colony, and was picked up from the agar plate and grown in 20ml of LB, with the appropriate antibiotic.



Figure 3.10 Electrophoretic Analysis of the PCR products of the Colony Screen of the pB12mcs-H1-eGFP vector

Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder Lane 2: Wild type pB12mcs-eGFP vector PCR product Lanes 3,4,6,8, and 9: PCR products of a positive colony screen Lanes 5 and 10-12: PCR products of a negative colony screen Lanes 7 and 13: PCR did not work

Lane 2 illustrates the PCR product of the wild type pB12mcs-eGFP vector. The band size of this PCR product is of around 1050bp when using the primer set T7 forward with T3 reverse. The band size for the positive colonies were around 1270bp, which is the standard size of the wild type PCR product (~1050) plus the size of the H1 human promoter (220bp). When comparing the PCR products of the colonies screened, the colonies in lanes 5 and 10-12 were negative, whereas colonies in all other lanes were positive (lanes 7 and 13 are excluded as the PCR reaction did not work here). The lanes highlighted in red are the colonies that were selected and grown in 20ml of LB with the appropriate antibiotic.

## 3.2.4.2 Sequence Analysis of the U6 and H1 Promoters

Plasmid extraction took place by means of mini preps, and plasmids were analysed by PCR first. Three sets of primers were used; for U6 analysis T7 forward with T3 reverse, T7 forward with U6 reverse, and U6 forward with T3 reverse. While the set of primers that were used for the plasmids positive for the H1 promoter were T7 forward with T3 reverse, T7 forward with H1 reverse, and H1 forward with T3 reverse. The PCR products of the reactions were analysed by gel electrophoresis, and are shown in Figures 3.11 and 3.12. Once, positive colonies were confirmed by PCR the plasmids were sent to Geneservice LTD for sequencing.





Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder

Lane 2: PCR product of the wild type pB12mcs-eGFP vector using primer set T7 forward and T3 reverse (negative control)

Lane 3: PCR product of the pB12mcs-U6-eGFP vector using primer set T7 forward with T3 reverse

Lane 4: PCR product of the pB12mcs-U6-eGFP vector using primer set T7 forward with U6 reverse

Lane 5: PCR product of the pB12mcs-U6-eGFP vector using primer set U6 forward with T3 reverse

Lane 6: GeneRuler<sup>™</sup> 100bp DNA Ladder

To confirm that the plasmid had the right insert in the right orientation, three PCR reactions were conducted on that plasmid. The three different sets of primers that were used were T7 forward with T3 reverse, T7 forward with U6 reverse and U6 forward and T3 reverse. If a U6 promoter has been inserted the sizes of the PCR products expected for the primer sets are 1320bp, 310bp, and 1350bp respectively. The negative control lane 2, was a PCR product of the wild type vector with the first set of primers, giving rise to a ~1050bp band. The electrophoretic analysis illustrates that the resultant PCR products are of the sizes expected, thus confirming that the plasmid in colony 11 had the correct insert.





Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder

Lane 2: PCR product of the wild type pB12mcs-eGFP vector, ~1050bp in size, primer set T7 forward with T3 reverse (negative control)

Lanes 3, 6, 9, and 12: PCR product of the pB12mcs-H1-eGFP vectors, primer set T7 forward with T3 reverse. Colonies 3, 4, 8 and 9 respectively

Lanes 4, 7, 10 and 13: PCR product of the pB12mcs-H1-eGFP vectors, primer set T7 forward with H1 reverse. Colonies 3, 4, 8 and 9 respectively

Lanes 5, 8, 11, and 14: PCR product of the pB12mcs-H1-eGFP vectors, primer set H1 forward with T3 reverse. Colonies 3, 4, 8 and 9 respectively Lane 15: GeneRuler<sup>™</sup> 100bp DNA Ladder

To confirm that the plasmids had the right insert in the right orientation, three PCR reactions were conducted on each plasmid. The three different sets of primers that were used in the reactions were T7 forward with T3 reverse, T7 forward with H1 reverse and H1 forward and T3 reverse. The sizes of the PCR products expected for the primer sets were 1270bp, 260bp, and 1300bp respectively. The analysis illustrates that the resultant PCR products are of the sizes expected, thus confirming that the extracted plasmids had the correct insert.

The sequencing results were analysed and the results for the pB12mcs-U6eGFP and pB12mcs-H1-eGFP used in this study are found in Appendix 8. The sequences of the U6 and H1 human promoters were compared to their sequences found in EMBL. Those plasmids with a 100% sequence match were grown in 500ml of LB medium, and a maxi prep was conducted. The extracted plasmids were purified, quantified and stored at -20°C.

### 3.2.5 Construction of the shRNA Molecules

Following preparation of the pB12mcs-eGFP vector for insert uptake and the construction of the positive control vectors. The next step was to design the shRNA molecules targeting the two different fusion genes, followed by construction of the shRNA cassettes.

### 3.2.5.1 Designing of the shRNA Molecules

The sequences of the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  fusion points found in EMBL website, accession numbers AJ131466 and AJ131467, were used as a basis for designing the shRNA molecules. The full sequences are shown in Appendix 4. For each fusion point three different shRNA molecules were designed, two molecules were 21nt long, whereas the third molecule was 22nt long.

The 21nt long shRNA molecule, shRNA1, had 11nt upstream of the fusion point and 10nt downstream, whereas shRNA2 had 10nt upstream of the fusion point and 11nt downstream. In the case of the 22nt long molecule, shRNA3, 11nt were on either side of the fusion points. Figure 3.13 demonstrates the predicted secondary structure of the shRNA1 molecule targeting the  $\beta 3\alpha 2$ fusion point. The predicted secondary structures for all the remaining shRNA molecules are given in Appendix 2.



Figure 3.13 The Secondary Structure of shRNA1 Antisense Oligonucleotide targeting the  $\beta 3\alpha 2$  fusion point

The figure shows the predicted secondary hairpin structure of the designed shRNA1 molecule targeting the  $\beta 3\alpha 2$  fusion point. The nucleotide sequence from 1-21, in green, represents the antisense strand, whereas the sense strand corresponds to the nucleotide sequence from 30-50, in red. The 8nt loop is in blue.

### 3.2.5.2 Synthesis of the U6 and H1 shRNA Cassettes

Once the secondary structures of the shRNA molecules were checked, U6 and H1 human promoter reverse primers, which contained the designed shRNA molecule sequences, were constructed and produced. The sequences of these primers are given in chapter 2 in Table 2.6.

To synthesise the shRNA cassettes, the U6 forward or the H1 forward primer was used together with the U6 or the H1 shRNA reverse primer to conduct a PCR reaction. K562 cDNA was used as a template, and the resultant PCR products were analysed through agarose gel electrophoresis. Figure 3.14 shows the analysis of these PCR products. The bands of the correct size were cut out of the gel, and the shRNA cassettes were purified. The shRNA cassettes were then quantified and digested with the restriction enzymes, BamHI and HindIII respectively. The shRNA cassettes were purified and quantified after each digestion. The final shRNA cassettes were then stored at -20°C.





Lane 1: PCR product of the U6  $\beta$ 3 $\alpha$ 2 shRNA1 cassette Lane 2: PCR product of the U6  $\beta$ 3 $\alpha$ 2 shRNA2 cassette Lane 3: PCR product of the U6  $\beta$ 3 $\alpha$ 2 shRNA3 cassette Lane 4: PCR product of the U6  $\beta$ 2 $\alpha$ 2 shRNA1 cassette Lane 5: PCR product of the U6  $\beta$ 2 $\alpha$ 2 shRNA2 cassette Lane 6: PCR product of the U6  $\beta$ 2 $\alpha$ 2 shRNA3 cassette Lane 7: GeneRuler<sup>TM</sup> 100bp DNA Ladder Lane 8: PCR product of the H1  $\beta$ 2 $\alpha$ 2 shRNA1 cassette Lane 9: PCR product of the H1  $\beta$ 2 $\alpha$ 2 shRNA2 cassette Lane 10: PCR product of the H1  $\beta$ 2 $\alpha$ 2 shRNA3 cassette

The PCR products of the shRNA cassettes were analysed by electrophoresis on a 2% agarose gel. Lanes 1-3 show the PCR products of the shRNA cassettes designed to target the  $\beta 3\alpha 2$  fusion point under the regulation of the U6 promoter. Lanes 4-6 show the PCR products of the shRNA cassettes designed to target the  $\beta 2\alpha 2$  fusion point under the regulation of the U6 promoter. The PCR products should be around 340bp in size, and when analysed above, the visualized bands are of the correct size. Lanes 8-10 show the PCR products of the shRNA cassettes designed to target the  $\beta 3\alpha 2$  fusion point that are under the regulation of the H1 promoter. The expected size of these PCR products is around 290bp.

# 3.2.5.3 Cloning of the shRNA Cassettes in the pB12mcs-eGFP Vector

The digested, purified shRNA cassettes were ligated into the pB12mcseGFP vector. The reaction was conducted overnight at 4°C with a 1:10 vector to insert ratio. Colony screening by means of PCR, using the T7 forward and the T3 reverse primer set, was conducted and the wild type pB12mcs-eGFP vector was included as a negative control. An example of the colony screen conducted on one of the U6 shRNA constructs is shown in Figure 3.15. Representative electrophoretic analysis for all the remaining shRNA cassettes can be found in Appendix 7.





Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder Lane 2: PCR product of the wild type pB12mcs-eGFP vector Lanes 3-15: PCR products of the colony screen Lanes 7 and 10: Positive colonies

This electrophoretic analysis was conducted on a 2% agarose gel. The above analysis is of the PCR products of the colony screen conducted on the U6 shRNA2 cassette that targets the  $\beta 2\alpha 2$  fusion point. The wild type vector gives a PCR product of ~1050bp. Any colony with the same band size indicates that the insert was not taken up by the vector. However, colonies 7 and 10, which are highlighted in red, show bands of ~1390bp, indicating that the vector has an insert of 340bp. These colonies were therefore picked up further analysis.

# 3.2.5.4 Confirmation by PCR of the shRNA Cassettes and the Sequencing of the Cassettes

The positive pB12mcs-eGFP-shRNA colonies were grown and purified plasmids were screened by three sets of primers prior to sequencing them. The sets of primers were the T7 forward with the T3 reserve primer, the T7 forward with the appropriate U6 or H1 shRNA reverse primer, and the U6 or H1 forward with the T3 reverse primer. PCR products were analysed by electrophoresis, and colonies positive in all three screens, with the correct band sizes, were sent for sequencing. Figure 3.16 shows an example of the analysis for one of the positive colonies. Representative analyses of the remaining colonies can be seen in Appendix 7.

The sequencing results were analysed and the sequences of both the U6 or H1 promoter and the shRNA molecules were checked. The raw sequencing data of all shRNA cassettes in Appendix 9 and Table 3.1 gives a brief summary and overview about all the shRNA cassettes that have been sequenced. A summary of the sequencing results of the shRNA constructs that were tested, and used for further expression studies can be found in Tables 3.2 - 3.5. Tables 3.2 and 3.4 show the sequencing results of the shRNA part of the constructs. Table 3.3 shows the sequencing results of the U6 part of the constructs and 3.5 the H1 part. The shRNA cassettes that were successfully synthesised included the U6 shRNA1; the U6 shRNA3 that targeted the  $\beta 2\alpha 2$  fusion point, the H1 shRNA2; and the H1 shRNA3 that targeted the  $\beta 2\alpha 2$  fusion point. Due to time limitations, the synthesis of further shRNA cassettes was stopped at this point.





Lane 1: GeneRuler<sup>™</sup> 1Kb DNA Ladder

Lane 2: PCR product of the wild type pB12mcs-eGFP vector with the primer set T7 forward and T3 reverse (negative control) Lane 3: PCR product of the primer set T7 forward with the T3 reverse Lane 4: PCR product of the primer set T7 forward with the U6 shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> reverse Lane 5: PCR product of the primer set U6 forward with the T3 reverse Lane 6: GeneRuler<sup>TM</sup> 100bp DNA Ladder

The electrophoretic analysis on a 2% agarose gel shows three PCR products from three different sets of primers. A negative control Lane 2 is included, which is the PCR product of the wild type plasmid with the primer set T7 forward and T3 reverse and a band size of ~1050bp is seen. In the case of a positive colony a band size of ~1390bp is expected, when using the same primer set. This is illustrated in lane 3. Lane 4 is the PCR product of the primer set T7 forward with the U6 shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> reverse. Here a band size of 380bp is expected and seen. For the last set of primers, the U6 forward and T3 reverse, a band size of ~1420bp is expected (lane 5).

shRNA	Promoter	Sequenced	Sequence	Used for	Number of
Construct			outcome *	expression	colonies
				studies	sequenced
β3α2 shRNA1	U6	Yes	2 mismatches	Yes	4
β3α2 shRNA2	U6	Yes	Inconclusive	No	3
β3α2 shRNA3	U6	Yes	3 mismatches	Yes	3
$\beta 2\alpha 2 \ sh RNA1$	U6	Yes	1 mismatch	Yes	4
$\beta 2\alpha 2 \ sh RNA2$	U6	No	-	-	-
$\beta 2\alpha 2 \ sh RNA3$	U6	No	-	-	-
β2α2 shRNA1	H1	Yes	ShRNA sequence	No	13
			was missing		
β2α2 shRNA2	H1	Yes	2 mismatches	No due to time	3
				limitations	
β2α2 shRNA3	H1	Yes	3 mismatches	No due to time	5
				limitations	

Table 3.1 A summary of the synthesis and sequencing of the shRNA constructs.

Nine constructs were designed in total, six regulated by the U6 promoter, and three by the H1 promoter. The U6-shRNA constructs targeted both fusion points, however, the H1-shRNA constructs only targeted the  $\beta 2\alpha 2$  fusion point. Several colonies for each construct were sequenced and only those with the correct shRNA sequence, and least number of promoter mismatches, were selected for further studies. \* All mismatches indicated are within the promoter sequences and not the shRNA molecules.

	DNA Fragment	Seque	ence							
1	β3α2 shRNA1 <sub>21nt</sub>	AAAAAAGCAG	AGTTCAAAAG	CCCTTCACAA	GCTTCTGAAG					
		GGC'I''I''I''I'GAA	CTCTGC							
2	β3α2 shRNA3 <sub>22nt</sub>	AAAAAAGCAG	AGTTCAAAAG	CCCTTCAGCA	AGCTTCCTGA					
	•	AGGGCTTTTG	AACTCTGC							
3	β2α2 shRNA1 <sub>21nt</sub>	ААААААСААТ	AAGGAAGAAG	CCCTTCACAA	GCTTCTGAAG					
	•	GGCTTCTTCC	TTATTG							
	Sequencing results									
4	U6 β3α2 shRNA1	AAAAAAGCAG	AGTTCAAAAG	CCCTTCACAA	GCTTCTGAAG					
		GGCTTTTGAA	CTCTGC							
5	U6 β3α2 shRNA3	AAAAAAGCAG	AGTTCAAAAG	CCCTTCAGCA	AGCTTCCTGA					
		AGGGCTTTTG	AACTCTGC							
6	U6 β2α2 shRNA1	ААААААСААТ	AAGGAAGAAG	CCCTTCACAA	GCTTCTGAAG					
	-	GGCTTCTTCC	TTATTG							
T	Table 3.2 The shRNA sequencing results of the pB12mcs-U6-shRNA-eGFP constructs.									

able 3.2 The shRNA sequencing results of the pB12mcs-U6-shRNA-eGFP const	ructs.
--------------------------------------------------------------------------	--------

The sequences of the originally designed shRNA molecules are shown in rows 1-3. Rows 4-6 show the consensus results of the shRNA molecules that were inserted into the pB12mcs-eGFP vector. The sequencing results confirm that the shRNA molecules found in the synthesised constructs completely match their original designed sequence. The sequences are given as the reverse sequence  $3' \rightarrow 5'$ .

<b>DNA Fragment</b>	Sequence				
U6 promoter	CAAGGTCGGG	CAGGAAGAGG	GCCTATTTCC	CATGATTCCT	TCATATTTGC
	ATATACGATA	CAAGGCTGTT	AGAGAGATAA	TTAGAATTAA	TTTGACTGTA
(Accession	AACACAAAGA	TATTAGTACA	AAATACGTGA	CGTAGAAAGT	AATAATTTCT
	TGGGTAGTTT	GCAGTTTTAA	AATTATGTTT	TAAAATGGAC	TATCATATGC
number M14486)	TTACCGTAAC	TTGAAAGTAT	TTCGATTTCT	TGGCTTTATA	TATCTTGTGG
	AAAGGACGAA	ACACCGTGCT	CGCTTCGGCA	GCACATATAC	ТА
		<u>Sequenc</u>	ing Results		
U6 B3a2 shRNA1	CAAGGTCGGG	CAGGAAGAGG	GCCTATTTCC	CATGATTCCT	TCATATTTGC
oo pour sinting	ATATACGATA	CAAGGCTGTT	AGAGAGA <mark>C</mark> AA	TTAGAATTAA	TTTGACTGTA
	AACACAAAGA	TATTAGTACA	AAATACGTGA	CGTAGAAAGT	AATAATTTCT
	TGGGTAGTTT	GCAGTTTTAA	AATTATGTTT	TAAAATGGAC	TATCATATGC
	TTACCGTAAC	TTGAAAGTAT	TTCGATTTCT	TGGCTTTATA	TATCTTGTGG
	AAAGGACGAA	GCACCGTGCT	CGCTTCGGCA	GCACATATAC	ТА
U6 β3α2 shRNA3	CAAGGTCGGG	CAGGAAGAGG	GCCTATTTCC	CATGATTCCT	TCATATTTGC
	ATATACGATA	CAAGGCTGTT	AGAGAGATAA	TTAGAATTAA	TTTGACTGTA
	A <mark>G</mark> CACAAAGA	TATTAGTACA	AAATACGTGA	CGTAGAAAGT	AATAATTTCT
	TGGGTAGTTT	GCAGTTTTAA	AATTATGTTT	TAAAATGGAC	TATCATATGC
	TTACCGTAAC	TTGAAAGTAT	TTCGATTTCT	TGGCTTTATA	TATCTTGTGG
	AAAGGACGAA	<b>G</b> CACCGTGCT	CGTTTCGGCA	GCACATATAC	ТА
U6 β2α2 shRNA1	CAAGGTCGGG	CAGGAAGAGG	GCCTATTTCC	CATGATTCCT	TCATATTTGC
	ATATACGATA	CAAGGCTGTT	AGAGAGATAA	TTAGAATTAA	TTTGACTGTA
	AACACAAAGA	TATTAGTACA	AAATACGTGA	CGTAGAAAGT	AATAATTTCT
	TGGGTAGTTT	GCAGTTTTAA	AATTATGTTT	TAAAATGGAC	TATCATATGC
	TTACCGTAAC	TTGAAAGTAT	TTCGATTTCT	TGGCTTTATA	TATCTTGTGG
	AAAGGACGAA	<b>G</b> CACCGTGCT	CGCTTCGGCA	GCACATATAC	ТА

Table 3.3 The U6 sequencing results of the pB12mcs-U6-shRNA-eGFP Constructs.

This table summarises the sequencing results from the pB12mcs-U6-eGFP constructs. The sequences shown above are of only the constructs that have been further used in expression studies described in the following chapter. The top of the table illustrates the human U6 promoter sequence found in the EMBL website. Below are the consensus results of the U6 promoter from each construct. Any mismatches in the sequenced data are highlighted in red. The sequences are given as the forward sequence 5'  $\rightarrow$  3'.

	DNA Fragment	Sequence	
1	β2α2 shRNA2 <sub>21nt</sub>	AAAAAAAATA AGGAAGAAGC CCTTCACCAA GCTTCGTGA GGGCTTCTTC CTTATTG	AA
2	β2α2 shRNA3 <sub>22nt</sub>	AAAAAACAAT AAGGAAGAAG CCCTTCACCA AGCTTCGTC AGGGCTTCTT CCTTATTGG	GΑ
		Sequencing results	
3	H1 β2α2 shRNA2	AAAAAAAATA AGGAAGAAGC CCTTCACCAA GCTTCGTGA GGGCTTCTTC CTTATTG	١A
4	H1 β2α2 shRNA3	AAAAAACAAT AAGGAAGAAG CCCTTCACCA AGCTTCGTG AGGGCTTCTT CCTTATTGG	JA

Table 5.4 The ShkinA sequencing results of the polyhilds-histikinA-edry constru	Table	3.4 T	Гhe	shRNA	sequencin	g results	of the	pB12mcs-	H1-shRNA-	-eGFP	construct	s.
---------------------------------------------------------------------------------	-------	-------	-----	-------	-----------	-----------	--------	----------	-----------	-------	-----------	----

The sequences of the originally designed shRNA molecules are shown in rows 1 and 2. Rows 3 and 4 show the consensus results of the shRNA molecules that were inserted into the pB12mcs-eGFP vector. The sequencing confirms that the shRNA molecules found in the synthesised constructs completely match their original designed sequence. The sequences are given as the reverse sequence  $3' \rightarrow 5'$ .

DNA Fragment	Sequence							
H1 promoter	GAACGCTGAC	GTCATCAACC	CGCTCCAAGG	AATCGCGGGC	CCAGTGTCAC			
	TAGGCGGGAA	CACCCAGCGC	GCGTGCGCCC	TGGCAGGAAG	ATGGCTGTGA			
(Accession	GGGACAGGGG	AGTGGCGCCC	TGCAATATTT	GCATGTCGCT	ATGTGTTCTG			
	GGAAATCACC	ATAAACGTGA	AATGTCTTTG	GATTTGGGAA	TCTTATAAGT			
number X16612)	TCTGTATGAG	ACCACTCTTT						
	Sequencing Results							
H1 β2α2 shRNA2	GAACGCTGAC	GTCATCAACC	CGCTCCAAGG	AATCGCGGGC	CCAGTGTCAC			
	TAGGCGGGAA	CACCCAGCGC	GCGTGCGCCC	TGGCAGGAAG	ATGGCTGTGA			
	GGGACAGGGG	AGTGGCGCCC	TGCAATAT <mark>C</mark> T	GCATGTCGCT	ATGTGTTCTG			
	GGAAATCACC	<b>G</b> TAAACGTGA	AATGTCTTTG	GATTTGGGAA	TCTTATAAGT			
	TCTGTATGAG	ACCACTCTTT						
H1 β2α2 shRNA3	GAACGCTGAC	GTCATCAACC	CGCTCCAAGG	AATCGCGGGC	CCAGTGTCAC			
	TAGGCGGGAA	CACCCAGCGC	GCGTGCGCCC	TGGCAGGAAG	ATGGCTGTGA			
	GGGACAGGGG	AGTGGCGCCC	TGCAATAT <mark>C</mark> T	GCATGTCGCT	ATGTGTTCTG			
	GGAAATCACC	<b>G</b> TAAACGTGA	AATGTCTTTG	GATTTGGG <mark>G</mark> A	TCTTATAAGT			
	TCTGTATGAG	ACCACTCTTT						

Table 3.5 The H1 sequencing results of the pB12mcs-H1-shRNA-eGFP Constructs.

This table summarises the sequencing results obtained from the pB12mcs-H1-eGFP constructs. The sequences shown above are of only the constructs that have been further used in expression studies described in the following chapter. The top of the table illustrates the human H1 promoter sequence found in the EMBL website. Below shows the consensus results of the H1 promoter from each construct. Any mismatches in the sequenced data are highlighted in red. The sequences are given as the forward sequence 5'  $\rightarrow$  3'.

### 3.3 Discussion

This chapter describes how a reporter gene system was generated. The pB12mcs-eGFP vector was used for this purpose. The vector had an enhanced GFP that is considered to be advantageous when using the system in mammalian cell lines. The plasmid was used to clone several synthesised shRNA cassettes. These shRNA molecules were designed to target the fusion points,  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$ , of the Philadelphia chromosome.

Cell lines chosen for use in this project were the human K562 cell line that bears the  $\beta 3\alpha 2$  fusion point, and the KCL-22 cell line that has the  $\beta 2\alpha 2$  fusion point. It was confirmed that both these cell lines contained the fusion point of interest by means of PCR and it was further confirmed that their sequences matched the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  sequences, that are found in EMBL website. Upon this confirmation, three short hairpin RNA molecules for each fusion point were designed. The secondary structures of the designed antisense oligonucleotides were predicted and upon confirmation that the designed shRNA molecules do have a hairpin like secondary structure, the molecules were synthesised. Synthesis was achieved by PCR, where the sequence of the H1 human promoters. The shRNA cassettes were therefore, synthesised in a manner where the molecules were downstream to the human U6 or H1 promoters.

The U6 and H1 promoters were selected due to the advantage of their natural presence in human cell lines and the ease of signalling their termination, where a stretch of 4 to 6 'T' nucleotides causes the promoter to terminate. The synthesised shRNA cassettes were then cloned into the double digested pB12mcs-eGFP vector. Positive colonies including both the vector and inserted shRNA cassettes were sequenced and those that have the U6 promoter and the correct shRNA sequence were selected for further expression studies.

The next step was to examine and test how efficient and specific the designed shRNA molecules were to their target sites. The following chapter deals with this issue, where expression studies were conducted on the generated pB12mcs-U6-shRNA-eGFP constructs.

# Chapter 4 : Expression of the shRNA Constructs in K562 and KCL-22 Cell Lines

## 4.1 Introduction

To examine the inhibitory effects that the synthesised shRNA constructs might have on their target sites, these constructs were expressed in two human cell lines K562 and KCL-22. The K562 cell line expresses the  $\beta$ 3 $\alpha$ 2 fusion protein, whereas the KCL-22 cell line expresses the  $\beta$ 2 $\alpha$ 2 fusion protein (Drexler *et al.*, 1999). The expression of the shRNA constructs was monitored through the reporter gene system, which has been described previously.

The effects of the synthesised constructs were then examined at both the RNA and protein levels, and the specificity of the shRNA molecules were also examined. In this part of the study, various chemical methods to transfect the synthesised shRNA constructs were undertaken; in addition, one type of physical transfection was also examined: nucleofection, which is a form of electroporation. In summary, this chapter deals with examining the expression of the generated shRNA constructs and their inhibitory effects on their target *BCR-ABL* fusion gene.

#### 4.2 Results

## 4.2.1 Transfection of the shRNA Constructs in K562 and KCL-22 Cell lines.

The K562 and KCL-22 cell lines are CML cell lines bearing the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$ BCR-ABL variants respectively. Leukaemic cell lines are characterised by having a growth advantage over normal cells, due to their rapid range of proliferation. These cells are unable to mature to functional non-dividing cells, hence leading to their accumulation (Koeffler and Golde, 1980). Lozzio and Lozzio, 1975, first reported the K562 cell line that was established from a pleural effusion of a terminal patient with CML in blast phase. The KCL-22 cell lines were established in the same manner as K562. Kubonishi and Miyoshi, 1983, first reported KCL-22 cell line in 1983.

These cell lines grow as single-cell suspension cultures (Lozzio and Lozzio, 1975 and Kubonishi and Miyoshi, 1983) and primary cells and cells in suspensions are hard to transfect (Ovcharenko *et al.*, 2005 and Oliveira and Goodell, 2003). Therefore, in order to select the best method of transfection, with the highest transfection efficiency, both chemical and physical methods were considered.

In this project, two of the three shRNA constructs designed to target the mRNA sequence of the  $\beta 3\alpha 2$  fusion variant were studied. While, one construct that targets the  $\beta 2\alpha 2$  fusion variant was examined.

### 4.2.2 Transfection by Chemical Methods and FACS Analysis

A range of chemical reagents was used to assess transfection efficiency into cell lines, and FACS analysis was performed after transfection. For FACS,  $3 \times 10^5$  cells were counted and the "Cell Quest" software used for analysis. The parameters were adjusted according to the cell line type when the FACS analysis was conducted. Examples of the generated data for the different transfection reagents tested are shown in Figures 4.1 - 4.4. Figures 4.1 and 4.2 demonstrate the highest transfection efficiencies achieved when using a chemical transfection reagent, while Figures 4.3 and 4.4 demonstrate transfection reactions that did not work. Table 4.1 summarises the transfection data obtained when using the different types of chemical transfection reagents. The highest levels of transfection in both K562 and KCL-22 cells were

seen when using feOfection/Yellow as a transfection reagent (K562 48% and KCL-22 38%).



Figure 4.1 Histogram generated by FACS demonstrating the transfection efficiency of the NIMT FeOfection/Yellow transfection reagent in K562 cell lines.

A histogram generated by the program CellQuest by the use of FACS for the K562 cell line transfected with the pEGFP-N1 vector by NIMT FeOfection/Yellow transfection reagent. The control, non-transfected cells, is shown in purple followed by the readings of the different dilutions of the transfection reagent, green, pink, blue, and yellow. These have the following dilutions of the transfection reagent 100, 120, 140, and 160µl, respectively, mixed with 5µl of DNA. The estimated transfection efficiencies are 43.4, 46.56, 43.5, and 48.7% respectively.



Figure 4.2 Histogram generated by FACS demonstrating the transfection efficiency of the NIMT FeOfection/Yellow transfection reagent in KCL-22 cell line.

A histogram generated by the program CellQuest by the use of FACS for the K562 cell line transfected with the pEGFP-N1 vector by NIMT FeOfection/Yellow transfection reagent. The control, non-transfected cells, is shown in purple followed by the readings of the different dilutions of the transfection reagent, green, pink, blue, and red. These have the following dilutions of the transfection reagent 100, 120, 140, and 160µl, respectively, mixed with 5µl of DNA. The estimated transfection efficiencies are 19.53, 29.01, 29.33, and 34.68% respectively.


Figure 4.3 Histogram generated by FACS demonstrating the transfection efficiency of the Nanofectin transfection reagent in K562 cell line.

A histogram generated by the program CellQuest by the use of FACS for the K562 cell line transfected with the pEGFP-N1 vector by Nanofectin transfection reagent. The control, non-transfected cells, is shown in purple followed by the readings of the different nanofectin ratios 2 (8µl), 3 (12.8µl), and 4 (16µl) that were mixed with 4µl of DNA. These are shown in green, pink, and red respectively.



Figure 4.4 Histogram generated by FACS demonstrating the transfection efficiency of the Nanofectin transfection reagent in KCL-22 cell line.

A histogram generated by the program CellQuest by the use of FACS for the KCL-22 cell line transfected with the pEGFP-N1 vector by Nanofectin transfection reagent. The control, non-transfected cells, is shown in purple followed by the readings of the different nanofectin ratios 2 (8µl), 3 (12.8µl), and 4 (16µl) that were mixed with 4µl of DNA. These are shown in green, pink, and blue respectively.

Name of Kit	Supplier	Transfection efficiency	
		K562	KCL-22
Superfect	Qiagen	21-31%	Didn't work
Lipofectamin 2000	Invitrogen	17.7-27%	14.8-18.5%
Hilymax	Dojindo	10-15%	Didn't work
TransIT-2020	Mirus	21.8-40.5%	25.5-26.5%
NIMT FeOfection/Yellow*	Genovis	43-48.7%	29-38%
FuGENE HD	Roche	16-32%	25-30%
Nanofectin	РАА	Didn't work	Didn't work
Lentiphos HT	Clontech	Didn't work	Didn't work

Table 4.1 A summary of different transfection methods on the transfection efficiency of K562and KCL-22 cell lines.

Several transfection kits were tested in order to transfect a control vector, the pEGFP-N1 vector, into the K562 and KCL-22 cell lines. The above table shows a list with all the transfection reagents that have been tested and the range of resultant transfection efficiency obtained from each kit. Some kits didn't work at all, where no fluorescent cells were visible under the fluorescent microscope nor any difference between the control and transfected sample using FACS. \*The highest transfection efficiency was by using the NIMT FeOfection/Yellow kit that gave an efficiency of up to 48.7% in the K562 cell lines.

### 4.2.3 Transfection Through Nucleofection

Chemical methods of transfecting K562 and KCL-22 cells did not give high efficiencies so nucleofection was tried. Nucleofection is a form of electroporation, where specific electrical impulses are used to weaken the target cell's membrane. Pores in the cell's membrane are created, through which DNA fragments can be internalized. In nucleofection, the DNA of interest is mixed with a special reagent (its composition is unknown as the manufacturer does not supply this information), which is then added to the cell line, and a wave of electrical impulses is introduced. The Amaxa nucleofector system was used in this study. It should be noted that the company supplies different transfecting reagent kits and nucleofector programs depending on the cell line type.

For the K562 cell line a commercially available reagent and two nucleofection programs were available. Both programs were used with the K562 cells and control vector supplied with the nucleofection Kit to determine which program was best for this cell line. However, in the case of the KCL-22 cells, neither reagent nor programs were available. Therefore, the same reagent and programs, previously tested for the K562 cells, were used for the KCL-22 cell line. The program T-016 was found to give best transfection efficiencies and cell viability for both cell lines. The transfection efficiency and cell viability results obtained from both nucleofection programs are summarized in Table 4.2 below.

	K562 cell line	T-003	T-016
1	Transfection efficiency	40.4%	54.7%
2	Viability	76%	90%
	KCL-22 cell line	T-003	T-016
3	Transfection efficiency	35%	40%
4	Viability	78%	90%

 Table 4.2 A Summary of the transfection efficiency obtained from FACS analysis and cell

 viability of K562 and KCL-22 cell lines by Nucleofection with the pB12mcs-U6-eGFP vector.

K562 and KCL-22 cell lines were transfected using the Amaxa cell line nucleofector<sup>®</sup> kit V and two nucleofection programs, T-003 and T-016. Each program was for nucleofecting a different K562 clone. The two programs were tested because the clone of the given K562 cell line was unknown. Transfection and viability results for both cell lines showed that the T-016 program had higher transfection efficiency and viability than the T-003. This was observed in both cell lines, consequently, the T-016 program was used for all following nucleofections.

The T-016 program was used in all subsequent studies. To investigate the transfection efficiency with pB12mcs-eGFP vectors, groups of 2.5 x10<sup>6</sup> cells were transfected with the following: negative control (pB12mcs-eGFP

wildtype), positive control (pB12mcs-U6-eGFP), shRNA1<sup> $\beta$ 3a2</sup>, and shRNA3<sup> $\beta$ 3a2</sup> constructs for the K562 cell line, and negative control (pB12mcs-eGFP wildtype), positive control (pB12mcs-U6-eGFP), and shRNA1<sup> $\beta$ 2a2</sup> construct, for the KCL-22 cell line. Successful transfection was confirmed by detection of GFP fluorescence of the target cell lines under the fluorescent microscope, see Figures 4.5 and 4.6. Once the fluorescence of the eGFP was detected, the percentage of cell viability was calculated, through the use of an automated cell counter.



Figure 4.5 Nucleofection of the K562 cell line with the pB12mcs plasmids.

(A) Shows the K562 cell line under the fluorescent microscope 24 hours after nucleofection with the pB12mcs-U6-eGFP plasmid (positive control, U6 promoter), in bright field. (B) Shows the cells when excited with blue light, fluorescing in green. (C) Shows the K562 cells transfected with wild type pB12mcs-eGFP vector (negative control, no promoter) in bright field. The same cells are shown in (D) after excitation with blue light.



Figure 4.6 Nucleofection of the KCL-22 cell line with the pB12mcs plasmids.

(A) Shows the KCL-22 cell line under the fluorescent microscope 24 hours after nucleofection with the pB12mcs-U6-eGFP plasmid (positive control, U6 promoter), in bright field. (B) Shows the cells when excited with blue light, fluorescing in green. (C) Shows the KCL-22 cells transfected with wild type pB12mcs-eGFP vector (negative control, no promoter) in bright field. The same cells are shown in (D) after excitation with blue light.

Once successful transformation was achieved, cell viability was determined and 3 x 10<sup>5</sup> cells from an original number of 2.5 x 10<sup>6</sup> transfected cells were counted and fixed and the transfection efficiency was determined by FACS analysis. Five groups of K562 cell were tested: non-transfected cells, cells transfected with the negative control vector (wild type pB12mcs-eGFP no promoter), cells transfected with the positive control vector (pB12mcs-U6eGFP), cells transfected with pB12mcs-U6-shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup>-eGFP, and cells transfected with pB12mcs-U6-shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>-eGFP. Four groups of KCL-22 cell lines were tested: non-transfected cells, cells transfected with the negative control vector (wild type pB12mcs-eGFP), cells transfected with the negative control vector (pB12mcs-U6-eGFP), and cells transfected with the positive control vector (pB12mcs-U6-eGFP), and cells transfected with pB12mcs-U6shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup>-eGFP. Figures 4.7 and 4.8 are examples of the FACS results generated from the nucleofections of K562 and KCL-22 groups, and Table 4.3 summarises the transfection efficiency results of three successful nucleofections conducted on each of the cell lines. All the other data is given in Appendix 10.

It was interesting to observe that the shRNA constructs also generated fluorescent signals indicating that the stop signal did not cause termination of the U6 promoter. Advantage of the observed GFP fluorescence generated was taken in measuring efficiency of transfection.



Figure 4.7 Histogram generated by FACS demonstrating the transfection efficiency of nucleofection in the K562 cell line.

The histogram is a representation of one of three nucleofections conducted on the K562 cell lines, transfection number 2 in Table 4.3. In purple are the non-transfected K562 cells. In pink are the K562 cells transfected with the positive control vector (pB12mcs-U6-eGFP). In green are the K562 cells transfected with the negative control, pB12mcs-eGFP wild type vector. In blue are the cells transfected with the shRNA1<sup> $\beta$ 3\alpha2</sup> construct, and in orange are the cells transfected with the shRNA3<sup> $\beta$ 3\alpha2</sup> construct. The transfection efficiencies for each with regards to the non-transfected cells are 57%, 11.63%, 35%, and 23% respectively. The transfection efficiencies with regards to the cells transfected with the negative control are 45.08% for the positive control and 23.29% for shRNA1<sup> $\beta$ 3\alpha2</sup> construct and 11.39% for shRNA3<sup> $\beta$ 3\alpha2</sup> construct.



Figure 4.8 Histogram generated by FACS demonstrating the transfection efficiency of nucleofection in the KCL-22 cell line.

The histogram is a representation of one of three nucleofections conducted on the KCL-22 cells, transfection number 2 in Table 4.3. In purple are the non-transfected KCL-22 cells; in pink are the cells transfected with the positive control vector (pB12mcs-U6-eGFP); in green are cells transfected with the negative control (pB12mcs-eGFP wild type vector), and in yellow are the cells transfected with the shRNA1<sup> $\beta$ 2\alpha2</sup> construct. The transfection efficiencies for each with regards to the non-transfected cells are 64%, 19.08%, and 29.1% respectively. The transfection efficiencies with regards to the cells transfected with the negative control are 44.9% for the positive control and 10.02% for shRNA1<sup> $\beta$ 2\alpha2</sup> construct.

Clearly the negative control in this transfection exhibiting GFP fluorescence and will be discussed latter.

K562 Cell Line	(A) Transfection	(B) Cell viability of cells transfected with:				
efficienc positive plass (pB12m eGI	efficiency of the positive control plasmid (pB12mcs-U6- eGFP)	Non transfected cells	Negative control	Positive control	shRNA1 β3α2	shRNA3 β3α2
Transfection 1	28%	99%	89%	90%	86%	80%
Transfection 2	57%	98%	90%	92%	84%	82%
Transfection 3	72%	99%	90%	90%	87%	84%
KCL-22 cell line	(C) Transfection	(D) Cell viability of cells transfected with:				with:
	efficiency of the					
	positive control	Non	Negative		Positive	shRNA1
	plasmid	transfected	contro	bl	control	β2α2
	(pB12mcs-U6-	cells				-
	eGFP)					
Transfection 1	53%	98%	90%		89%	87%
Transfection 2	64%	98%	88%		91%	86%
Transfection 3	67%	99%	90% 88%		87%	

 Table 4.3 A Summary of the three nucleofections conducted on each of the human leukaemic

 cell lines, K562 and KCL-22, and the cell viability in each of the transfected cell groups.

Each K562 cell transfection consisted of 5 groups; non-transfected cells, cells transfected with the wild type pB12mcs-eGFP vector (negative control), cells transfected with the pB12mcs-U6-eGFP vector (positive control), cells transfected with the shRNA1<sup> $\beta$ 3\alpha2</sup> construct, and cells transfected with the shRNA3<sup> $\beta$ 3\alpha2</sup> construct. The transfection efficiency (positive control plasmid) obtained from FACS analysis for each transfection is shown in part (A) of the table. The cell viability for each group of transfected cells is indicated in part (B). The bottom parts of the table (C) and (D) illustrate a similar table but for the KCL-22 cell line with cells transfected with the wild type pB12mcs-eGFP vector (negative control), cells transfected with the shRNA1<sup> $\beta$ 2\alpha2</sup> construct.

## 4.2.4 Real Time PCR Analysis of the Transfected Cell Lines

qPCR was used to study the effects of the generated shRNA constructs at the mRNA levels of the fusion gene of interest. To do so, several qPCR primers were designed. Primers that spanned the fusion points,  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$ , and their appropriate probes were designed by the program `primer3'. Primers that amplified the normal *BCR* and *ABL* genes and the *GAPDH* reference gene were generated through the Roche UPL library software, which has a set of 250 probes. All primers gave rise to PCR products that were no more than 230bp in size. Primers were tested for their specificity to their target sequences. This was achieved through conducting a PCR reaction on cDNA extracted from K562 and KCL-22 cell lines. The electrophoretic analysis of the products of these PCR reactions can be seen in figures 4.9 - 4.11. Figure 4.9 shows an electrophoretic analysis of PCR products with GAPDH primers using K562 cDNA as a template. Figure 4.10 shows the electrophoretic analysis of PCR products with ABL and BCR primers. The electrophoretic analysis with  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  primers are shown in Figure 4.11.



#### Figure 4.9 Electrophoretic analysis of PCR products of GAPDH primers using K562 cDNA

Lane 1: pBluescript Hpa II marker Lane 2: PCR product of GAPDH primers using K562 cDNA Lane 3: PCR product of GAPDH primers using MilliQ H<sub>2</sub>O

GAPDH PCR products were analysed by the use of a 2% agarose gel. Lane 2 shows a 226bp band, which is the size of the GAPDH PCR product. PCR in lane 3 is a negative control.



Figure 4.10 Electrophoretic analysis of PCR reactions using the ABL and BCR primers on K562 cDNA

Lane 1: GeneRuler<sup>™</sup> 100bp DNA ladder

Lane 2: PCR product of MilliQ H<sub>2</sub>O using the ABL primers (negative control)

Lane 3: PCR product of K562 cDNA using the ABL primers

Lane 4: PCR product of K562 cDNA using the BCR primers

Lane 5: PCR product of MilliQ H<sub>2</sub>O using the BCR primers (negative control)

The PCR products of ABL and BCR primers were analysed through electrophoresis using a 2% agarose gel. In Lane 3, the ABL PCR product of 77bp is present, in lane 4, a BCR PCR product of 91bp is shown. PCR in lanes 2 and 5 didn't give a product.



Figure 4.11 Electrophoretic analysis of PCR reaction using  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  primers with K562 and KCL-22 cDNA, respectively.

Lane 1: GeneRuler<sup>™</sup> 100bp DNA ladder

Lane 2: PCR product of MilliQ  $H_2O$  using the  $\beta 3\alpha 2$  primers (negative control)

Lane 3: PCR product of K562 cDNA using the  $\beta 3\alpha 2$  primers

Lane 4: PCR product of MilliQ  $H_2O$  using the  $\beta 2\alpha 2$  primers (negative control)

Lane 5: PCR product of KCL-22 cDNA using the  $\beta 2\alpha 2$  primers

The PCR products with  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  primers were analysed through electrophoresis using a 2% agarose gel. In Lane 3, a  $\beta 3\alpha 2$  PCR product of 190bp is present, in lane 5, a  $\beta 2\alpha 2$  PCR product of 183bp is shown. PCR in lanes 2 and 4 didn't give a product.

Electrophoretic analysis of all hydrolysis primers confirmed the specificity of the primers, as only one band of the right size is present in each PCR reaction. In addition, no bands were observed in the negative controls when MilliQ  $H_2O$  was used as a template.

A reference gene, GAPDH, is included in this analysis as a control. The normal BCR and ABL levels were also examined; this was done to investigate whether the shRNA constructs had an inhibitory effect on mRNA production of the normal genes as well as the fusion genes.

## 4.2.4.1 The Generation of Standard Curves

Preliminary experiments used SYBR green I methodology to validate the primers. Once validated standard curves for each gene product were generated by the use of Taqman and UPL probes and primers. This was achieved by conducting a qPCR on several dilutions of plasmids in the case of the  $\beta 3\alpha 2$  and the  $\beta 2\alpha 2$  fusion points or on several dilutions of a PCR product, in case of the *GAPDH*, normal *BCR*, and normal *ABL* genes. The starting point for the concentration of each gene ranged from 10-30ng (per 100µl), then each sample was diluted 100x to give rise to a total of 5 samples. The qPCR results were analysed through the Roche LightCycler 480 software. Figures 4.12-4.16 show the generated standard curves.





Figure 4.12 The qPCR of GAPDH standards and the generated Standard Curve.

The qPCR results of the GAPDH standards are shown in (A). The qPCR was conducted on five dilutions of a GAPDH PCR product. The starting concentration, of 0.15ng/µl of a purified GAPDH PCR product, and the dilutions used are also shown here. The qPCR reactions were performed in duplicate, and the results analysed by the Light Cycler 480 software to determine the values of the GAPDH standard curve in (B). The standard curve is a least square fit line drawn through all dilutions, which is the mean as the reactions were done in duplicate. The calculated amplification efficiency was 91.3%. This curve was further used to calculate the concentration of the GAPDH gene product in the cell lines of interest.



Figure 4.13 The qPCR of BCR standards and the generated Standard Curve.

The qPCR results of the BCR standards are shown in (A). The qPCR was conducted on five dilutions of a BCR PCR product. The starting concentration, of  $0.12 ng/\mu l$  of a purified BCR PCR product, and the dilutions used are also shown here. The qPCR reactions were performed in duplicate, and the results analysed by the Light Cycler 480 software to determine the values of the BCR standard curve in (B). The standard curve is a least square fit line drawn through all dilutions, which is the mean as the reactions were done in duplicate. The calculated amplification efficiency was 96.4%. This curve was further used to calculate the concentration of the *BCR* gene product in the cell lines of interest.





Figure 4.14 The qPCR of ABL standards and the generated Standard Curve.

The qPCR results of the ABL standards are shown in (A). The qPCR was conducted on five dilutions of a ABL PCR product. The starting concentration, of 0.195ng/µl of a purified ABL PCR product, and the dilutions used are also shown here. The qPCR reactions were performed in duplicate, and the results analysed by the Light Cycler 480 software to determine the values of the ABL standard curve in (B). The standard curve is a least square fit line drawn through all dilutions, which is the mean as the reaction was done in duplicate. The calculated amplification efficiency was 96.6%. This curve was further used to calculate the concentration of the ABL gene product in the cell lines of interest.





Figure 4.15 The qPCR of  $\beta 3\alpha 2$  standards and the generated Standard Curve.

The qPCR results of the  $\beta 3\alpha 2$  standards are shown in (A). The qPCR was conducted on five dilutions of a  $\beta 3\alpha 2$  containing plasmids. The starting concentration, of  $0.2 \text{ng/}\mu \text{l}$  of a  $\beta 3\alpha 2$  construct, and the dilutions used are also shown here. The qPCR reactions were performed in duplicate, and the results analysed by the Light Cycler 480 software to determine the values of the  $\beta 3\alpha 2$  standard curve in (B). The standard curve is a least square fit line drawn through all dilutions, which is the mean as the reaction was done in duplicate. The calculated amplification efficiency was 95%. This curve was further used to calculate the concentration of the  $\beta 3\alpha 2$  fusion gene product in the cell lines of interest.





Figure 4.16 The qPCR of  $\beta 2\alpha 2$  standards and the generated Standard Curve.

The qPCR results of the  $\beta 2\alpha 2$  standards are shown in (A). The qPCR was conducted on five dilutions of a  $\beta 2\alpha 2$  containing plasmid. The starting concentration, of 0.3ng/µl of a purified  $\beta 2\alpha 2$  construct, and the dilutions used are also shown here. The qPCR reactions were performed in duplicate, and the results analysed by the Light Cycler 480 software to determine the values of the  $\beta 2\alpha 2$  standard curve in (B). The standard curve is a least square fit line drawn through all dilutions, which is the mean as the reaction was done in duplicate. The calculated amplification efficiency was 89.7%. This curve was further used to calculate the concentration of the  $\beta 2\alpha 2$  fusion gene product in the cell lines of interest.

Once the standard curves were established,  $2.5 \times 10^6$  cells, K562 or KCL-22, were nucleofected, and 3 x  $10^5$  cells were used for FACS analysis, and 1 x  $10^6$ cells were used for total RNA extraction, followed by cDNA synthesis. Another sample of 1 x 10<sup>6</sup> of transfected cells was used for protein extraction. A PCR reaction on the newly synthesised cDNA, to confirm that the RT-PCR had worked, was routinely done. Primers for the RPL13 gene (ribosomal protein L13 gene), a housekeeping gene, were also used in this reaction, and a band size of 96bp was expected; Figure 4.17 shows an example of the electrophoretic analysis. Normal ABL and BCR levels were also examined; this was done to investigate whether the shRNA constructs reduced the mRNA levels of the normal genes. Five qPCR reactions (for GAPDH, ABL, BCR,  $\beta 3\alpha 2$ , and  $\beta 2\alpha 2$ ) were done in duplicate for each sample. The standard curves were used for each transfection to determine the concentrations of genes of interest, the results were analysed by the Roche LightCycler 480 software. The software calculates the mRNA concentration levels. The levels were analysed and the percentages of depletion were calculated.



Figure 4.17 An example of an electrophoretic Analysis of the PCR reaction using RPL13 primers on transfected K562 cells cDNA.

Lane 1: GeneRuler<sup>™</sup> 100bp DNA Ladder

Lane 2: PCR product of the non-transfected K562 cDNA Lane 3: PCR product of the K562 transfected cDNA wild type pB12mcs-eGFP Lane 4: PCR product of the K562 transfected cDNA wild type pB12mcs-U6-eGFP Lane 5: PCR product of the K562 transfected cDNA wild type pB12mcs-U6-shRNA1<sup>β3α2</sup>-eGFP Lane 6: PCR product of the K562 transfected cDNA wild type pB12mcs-U6-shRNA3<sup>β3α2</sup>-eGFP

RPL13 PCR products were analysed via electrophoresis using a 2% agarose gel .The 96bp band in lanes 2-6 indicates that the RNA purification and K562 cDNA synthesis were successful in all the transfections including the non-transfected K562 control.

GAPDH was used as a reference gene, and the qPCR data generated in this reaction were used to normalize the data obtained from the other genes.

# 4.2.4.2 Effects of shRNA1 $^{\beta 3\alpha 2}$ and shRNA3 $^{\beta 3\alpha 2}$ in K562 and KCL-22 cells

To study the effects of the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> constructs, the RNAi molecules were transfected into K562 cells that express the target site, the  $\beta$ 3 $\alpha$ 2 BCR-ABL variant. To examine the specificity of the constructs KCL-22 cells expressing the  $\beta$ 2 $\alpha$ 2 variant that has the same *ABL* sequence as  $\beta$ 3 $\alpha$ 2 were used. To measure the  $\beta$ 3 $\alpha$ 2 mRNA levels, qPCR was conducted using primers specific only to the  $\beta$ 3 $\alpha$ 2 fusion gene. The forward primer binds to exon 14 of the *BCR* part of the gene, while the reverse primer binds to the *ABL* part. Similarly, the  $\beta$ 2 $\alpha$ 2 variant was measured using primers specific only for this fusion gene. The forward primer binds to exon 13 of the *BCR* part, while the reverse primer binds to the *ABL* part of the fusion gene.

Table 4.4 demonstrates the normalized qPCR data generated from the K562 cell lines transfected with the shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> constructs. The data were then converted to percentage of expression with regards to the positive control and a bar chart is shown in Figure 4.18. Note, the `negative' control are the cells transfected with the wildtype vector pB12mcs-eGFP, whereas the `positive' control are cells transfected with the pB12mcs-U6-eGFP vector.

Evp Efficiency		`Negative'	shRNA1 <sup>β3α2</sup>	shRNA3 <sup>β3α2</sup>	`Positive'
exp enciency	Control (pg*)	(pg*)	(pg*)	Control (pg*)	
1	28%	0.0043	0.0030	0.0036	0.0045
2	57%	5.1176	2.4459	3.746	5.0037
3	72%	7.7561	2.5397	5.1694	7.5672

Table 4.4 The qPCR data of the  $\beta 3\alpha 2$  mRNA levels of the transfected K562 cells.

The RNA extraction was conducted 24hr after each transfection. The generated data from the qPCR reactions of the  $\beta 3\alpha 2$  gene is presented as concentrations in pg. The results from three transfections (Exp. 1-3) with their corresponding transfection efficiencies are shown. The data are normalized with regards to the reference gene, *GAPDH*. The `negative' controls are cells transfected with pB12-eGFP and the `positive' controls are cells transfected with pB12mcs-U6-eGFP. Similar values are seen with the `negative' control and `positive' control because qPCR was measuring only the  $\beta 3\alpha 2$  fusion point. \* Per pg of *GAPDH*.



Figure 4.18 The percentage of expression of  $\beta 3\alpha 2$  mRNA levels in K562 cell line transfected with shRNA molecules targeting the  $\beta 3\alpha 2$  fusion point.

The figure shows the results for three transfection experiments. The `positive' controls are K562 cells transfected with the pB12mcs-U6-eGFP vector and are arbitrarily chosen to be 100%. The percentages of expression of all other samples were calculated with regards to the positive control. The `negative' controls are the K562 cells treated with the wild type vector (pB12mcs-eGFP). K562 cells transfected with shRNA1<sup> $\beta$ 3\alpha2</sup> and shRNA3<sup> $\beta$ 3\alpha2</sup> showed  $\beta$ 3\alpha2 mRNA expression levels of 33.6-68% (p<0.034) and 68-81.5% (p<0.023), respectively.

The data show a reduction in the  $\beta 3\alpha 2$  mRNA levels when using both constructs. However, shRNA1<sup> $\beta 3\alpha 2$ </sup> constantly showed a higher level of depletion in comparison with shRNA3<sup> $\beta 3\alpha 2$ </sup>. The third transfection reaction had the highest transfection efficiency of 72%, and shRNA1<sup> $\beta 3\alpha 2$ </sup> decreased the  $\beta 3\alpha 2$  mRNA levels to 33.6% in comparison with shRNA3<sup> $\beta 3\alpha 2$ </sup> that had 68% expression.

To test the specificity of the  $\beta 3\alpha 2$  shRNA molecules, the KCL-22 cell line was treated with the constructs and the normalized data were converted into percentage of expression with regards to the positive control, the KCL-22 cells transfected with the pB12mcs-U6-eGFP. This is shown in Figure 4.19. The levels

of the  $\beta 2\alpha 2$  mRNA expression were on average 99.2% of the controls, which indicates that the shRNA1<sup> $\beta 3\alpha 2$ </sup> and shRNA3<sup> $\beta 3\alpha 2$ </sup> do not target the  $\beta 2\alpha 2$  gene.



Figure 4.19 Percentage of expression of  $\beta 2\alpha 2$  mRNA levels in KCL-22 cells after treatment with  $\beta 3\alpha 2$  shRNA constructs.

The figure shows the results of three transfection experiments. The controls are the KCL-22 cells treated with the pB12mcs-U6-eGFP vector and are arbitrarily chosen to be 100%. The data obtained from the cells treated with  $shRNA1^{\beta3\alpha2}$  and  $shRNA3^{\beta3\alpha2}$  were converted to percentages in regards to the `positive' control. The expression levels of the  $\beta2\alpha2$  mRNA was of 99-99.6% (±2 SD) for  $shRNA1^{\beta3\alpha2}$  and of 98.4-99.6 (± 0.5 SD) for  $shRNA3^{\beta3\alpha2}$ .

## 4.2.4.3 Effects of shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> in KCL-22 and K562 cell lines

shRNA1<sup> $\beta^{2\alpha^2}$ </sup> constructs were transfected into KCL-22 cells in order to study their effects on their target site; KCL-22 cells express the  $\beta^{2\alpha^2}$  gene. To study the specificity of the shRNA molecule, K562 cells, which express the other BCR-ABL variant ( $\beta^{3\alpha^2}$ ), were used for transfection. The  $\beta^{2\alpha^2}$  mRNA levels were measured by qPCR with the use of primers specific to only the  $\beta^{2\alpha^2}$  gene. Table 4.5 demonstrates the normalized qPCR data of the effects of shRNA1<sup> $\beta^{2\alpha^2}$ </sup> on  $\beta 2\alpha 2$  fusion point in the KCL-22 cell line. These data were calculated as percentages of expression with regards to the `positive' control. The `positive' control was transfected with the pB12mcs-U6-eGFP vector and the `negative' control was transfected with the pB12mcs-eGFP vector. These data were used to generate the bar chart shown in Figure 4.20. The data showed a reduction of the  $\beta 2\alpha 2$  mRNA expression. The expression levels of the  $\beta 2\alpha 2$  mRNA treated with shRNA1<sup> $\beta 2\alpha 2$ </sup> reduced to a maximum of 66% when the transfection efficiency of nucleofection was 67%.

		`Negative'	shRNA1 <sup>β2α2</sup>	`Positive'
Ехр	Effeciency	control (pg*)	(pg*)	control (pg*)
1	53%	5.072	2.3719	4.8399
2	64%	17.7292	6.6635	16.6261
3	67%	14.6786	4.816	14.2657

#### Table 4.5 The qPCR data of the $\beta 2\alpha 2$ mRNA levels of the transfected KCL-22 cells.

The RNA extraction was conducted 24hr after each transfection. The generated data from the qPCR reactions of the  $\beta 2\alpha 2$  gene is presented as concentrations in pg. The results from three transfections (Exp. 1-3) with their corresponding transfection efficiencies are shown. The data are normalized in regards to the reference gene, *GAPDH*. The `negative' controls are cells transfected with pB12-eGFP and the `positive' controls are transfected with pB12mcs-U6-eGFP. Similar values are seen with the `negative' control and `positive' control because qPCR was measuring only the  $\beta 2\alpha 2$  fusion point. \* Per pg of *GAPDH*.





The figure shows the results of three transfection experiments on KCL-22 cell line. The `positive' and `negative' controls were transfected with pB12mcs-U6-eGFP and pB12mcs-eGFP vectors respectively. The positive control was chosen arbitrarily to be 100%. The normalized data were then calculated into percentages of expression in regards to the positive control. KCL-22 cells treated with shRNA<sup> $\beta 2\alpha 2$ </sup> showed expression levels of 34-49%. (p<0.005)

The specificity of the shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup> construct was also tested in the K562 cell line, Figure 4.21, where the  $\beta^{3\alpha^{2}}$  mRNA levels of expression were on average 98.9% of the controls. The results demonstrate that the shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup> is specific for the  $\beta^{2\alpha^{2}}$  target site.



# Figure 4.21 Percentage of $\beta 3\alpha 2$ mRNA levels in K562 cells after treatment with $\beta 2\alpha 2$ shRNA constructs.

The figure shows the results of three transfection experiments. The control is the K562 cells treated with the pB12mcs-U6-eGFP vector and are arbitrarily chosen to be 100%. The data obtained from the cells treated with  $shRNA1^{\beta 2\alpha 2}$  were converted to percentages in regards to the `positive' control. The expression levels of the  $\beta 3\alpha 2$  mRNA were of 98.2-99.5% (± 0.66 SD).

# 4.2.4.4 Effects of the shRNA constructs on normal BCR and ABL mRNA levels

To further confirm the specificity of the shRNA constructs to their target sites, the levels of the normal *ABL* and *BCR* genes were examined in K562 cells when transfected with shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>, and in KCL-22 cells when transfected with shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup>. This was done by qPCR with primers that are specific only for the normal *BCR* and *ABL* genes. In each primer set, at least one primer was binding to an exon not present in the *BCR-ABL* fusion genes.

GAPDH was used as a reference gene, and the data generated were normalized. The mRNA levels of the normal ABL and BCR genes were not

affected by any of the shRNA constructs. This is shown in Figures 4.22 and 4.23. Figure 4.22 shows the levels of the *BCR* and *ABL* genes in relation to the positive control in transfected K562 cells. Figure 4.23 shows results for KCL-22 cells. The `negative' controls are the cells transfected with the wildtype vector pB12mcs-eGFP, whereas the `positive' controls are cells transfected with the pB12mcs-U6-eGFP vector.



Figure 4.22 The percentage of expression of the normal *BCR* and *ABL* genes in K562 cell line transfected with shRNA constructs targeting the  $\beta 3\alpha 2$  fusion gene.

This figure shows the results of three transfection reaction experiments (Exp 1-3). The `positive' and `negative' controls are K562 cells transfected with pB12mcs-U6-eGFP and pB12mcs-eGFP vectors, respectively. The `positive' control was selected arbitrarily to be 100%; the remaining data generated for all others samples were converted to percentages with regards to the positive control. *BCR* and *ABL* mRNA expression levels for shRNA1<sup> $\beta$ 3\alpha2</sup></sup> ranged from 103.3-106% (± 1.5 SD), and 98.5-102.8% (± 1.88 SD), respectively. For shRNA3<sup> $\beta$ 3\alpha2</sup> BCR and ABL mRNA expressions ranged from 102.5-106.2% (± 2.4 SD), and for 98.7-102.9% (± 2.1 SD), respectively. The percentage of expression is shown in the bar chart, red for shRNA1<sup> $\beta$ 3\alpha2</sup> and green for shRNA3<sup> $\beta$ 3\alpha2</sup>.



Figure 4.23 The percentage of expression of the normal *BCR* and *ABL* genes in KCL-22 cell line transfected with the shRNA1<sup> $\beta 2\alpha^2$ </sup> construct.

This figure shows the results of three transfection reaction experiments (Exp 1-3). The `positive' and `negative' controls are KCL-22 cells transfected with pB12mcs-U6-eGFP and pB12mcs-eGFP vectors, respectively. The `positive' control was selected arbitrarily to be 100%; the remaining data generated for all others samples were converted to percentages with regards to the `positive' control. BCR and ABL mRNA expression levels for shRNA1<sup> $\beta$ 2\alpha2</sup> ranged from 99.4-103.8% (± 2.37 SD), and 96.4-105.3% (± 4.46 SD), respectively.

# 4.2.5 Protein analysis of the K562 transfected Cell Line by Western Blotting

To observe the effects of the constructed shRNA molecules on their target genes at the protein level, Western blots were conducted. Proteins extracted from the K562 cell line transfected with pB12mcs-U6-eGFP and all pB12mcs-U6-shRNA-eGFP vectors, targeting both *BCR-ABL* genes, were examined. Figure 4.24 shows the Western blot analysis conducted on the K562 cell lines.



Figure 4.24 Western blot analysis of the  $\beta 3\alpha 2$  fusion protein from K562 transfected cell line.

The figure shows a representative Western blot analysis of the  $\beta 3\alpha 2$  protein from the transfected K562 cell line. The first band in Row 1 is the  $\beta 3\alpha 2$  protein from K562 cells transfected with pB12mcs-U6-shRNA1<sup> $\beta 2\alpha 2$ </sup>-eGFP plasmid. The control band row 1 refers to the  $\beta 3\alpha 2$  protein from cells that were transfected with the `positive' control plasmid, pB12mcs-U6-eGFP. The last two bands in Row 1 show the  $\beta 3\alpha 2$  proteins from the K562 cell line treated with shRNA1<sup> $\beta 3\alpha 2$ </sup> and shRNA3<sup> $\beta 3\alpha 2$ </sup> constructs. Row 2 shows the GAPDH housekeeping gene that was included as a control for loading and was further used to normalise the results. The blot was probed with an anti *c-ABL* antibody (row 1) and anti *GAPDH* antibody (row2).

The band intensities for all samples were measured and normalized with regards to the *GAPDH* housekeeping gene. The data generated were converted to percentage of expression with regards to the `positive' control, which was chosen to be 100%. Upon analysis of the band intensities a reduction in the  $\beta 3\alpha 2$  fusion protein was observed. Each western blot was repeated three times. Table 4.6 shows the statistical analysis done on the results obtained from this Western blot.

	Measurement	of band	Percentage		Student
Transfected	intensities		of	SD (±)	t-test
shRNA molecule	β3α2 fusion		reduction		P value
	protein	Control			
shRNA1 β3α2	1.90	3.16	40%	0.105	<0.002
shRNA3 β3α2	2.10	3.16	33.5%	0.232	<0.014
shRNA1 β2α2	3.00	3.16	5.1%	0.148	<0.163

Table 4.6 Statistically Analysed data from the Western blot conducted on the  $\beta 3\alpha 2$  fusion protein extracted from K562 cells that were transfected with shRNA1<sup> $\beta 3\alpha 2$ </sup>, shRNA3<sup> $\beta 3\alpha 2$ </sup>, shRNA1<sup> $\beta 2\alpha 2$ </sup>, and the blot probed with *c-ABL* Ab, and band intensities measured.

The arithmetic means for the results obtained from three Western blots were first calculated. A reduction in the  $\beta 3\alpha 2$  protein treated with shRNA molecules specific for the gene showed a reduction in comparison with the control. The percentage of reduction was then calculated, column 4. The results showed 40% reduction in protein levels when treated with shRNA1<sup> $\beta 3\alpha 2$ </sup> and 33.5% with shRNA3<sup> $\beta 3\alpha 2$ </sup>. No significant reduction is observed when treated with shRNA1<sup> $\beta 2\alpha 2$ </sup>, column 4 row 3. The SD and the student t-test are shown in columns 5 and 6.

Both the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> molecules reduced the levels of  $\beta$ 3 $\alpha$ 2 protein. It was found that a higher reduction in the protein level was observed with shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> construct in comparison with shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>. In addition, the shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> molecule targeting the  $\beta$ 2 $\alpha$ 2 fusion gene showed no effects on the levels of the  $\beta$ 3 $\alpha$ 2 fusion protein. These results support the previous qPCR findings (previous section) and demonstrate that shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> are specific for their target sites. Figure 4.25 demonstrates the generated Western blot data in the form of a bar chart.



Figure 4.25 Western blot analysis of the  $\beta 3\alpha 2$  fusion protein from K562 transfected cell line.

The figure shows Western blot results from three different transfection experiments. Each bar is generated from the average of three Western blot repeats. The `positive' control, cells transfected with the pB12mcs-U6-eGFP vector, was selected to be 1 and the data from other transfected cells were calculated with regards to the `positive' control. The chart demonstrates that shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> constructs reduced  $\beta$ 3 $\alpha$ 2 protein levels by 40 and by 33.5% respectively. The data also show that the shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> molecule has little effect on the  $\beta$ 3 $\alpha$ 2 fusion gene product. \*\*p < 0.002 and \*p < 0.01

# 4.2.6 Protein analysis of the KCL-22 transfected Cell Line by Western Blotting

To observe the effects of shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup> on its target gene at the protein level, three Western blots were conducted. Proteins extracted from the KCL-22 cell line transfected with pB12mcs-U6-eGFP and pB12mcs-U6-shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup>-eGFP vectors, were examined. Figure 4.26 is a representation of a Western blot analysis done on the KCL-22 cell lines.



# Figure 4.26 Western blot analysis of the $\beta 2\alpha 2$ fusion protein from the KCL-22 transfected cell line.

The figure shows a representative Western blot analysis of the  $\beta 2\alpha 2$  protein from the transfected KCL-22 cell line. The first band in Row 1 refers to the control band that is the  $\beta 2\alpha 2$  protein from cells transfected with the `positive' control plasmid, pB12mcs-U6-eGFP. The remaining bands in row 1 show the fusion protein from KCL-22 cells transfected with pB12mcs-U6-shRNA-eGFP plasmids. Row 2 shows the GAPDH housekeeping gene that was included as a control for loading and was further used to normalise the results. The blot was probed with anti  $\beta 2\alpha 2$  junction specific antibody (row 1) and anti GAPDH antibody (row 2).

Band intensities for all samples were measured and normalized with regards to the *GAPDH* housekeeping gene. The data generated were converted to percentages of expression with regards to the `positive' control, which was chosen to be 100%. Table 4.7 shows the statistical analysis done on the results obtained from this Western blot.

	Measurement of band				
Transfected	intensities		Percentage		Student
shRNA	β2α2 fusion		of	SD (±)	t-test
molecules	protein	Control	Reduction		p value
shRNA1 β2α2	1.7	3.1	45%	0.0248	0.0001
shRNA1 β3α2	3.0	3.1	4.5%	0.0984	0.1335
shRNA3 β3α2	2.9	3.1	6.2%	0.1299	0.1240

Table 4.7 Statistically analysed data from Western blot conducted on the  $\beta 2\alpha 2$  fusion protein from KCL-22 cells transfected with shRNA1<sup> $\beta 2\alpha 2$ </sup>, shRNA1<sup> $\beta 3\alpha 2$ </sup>, shRNA3<sup> $\beta 3\alpha 2$ </sup>, and the blot probed with  $\beta 2\alpha 2$  Ab, and band intensities measured.

The arithmetic means for the results obtained from the Western blots were first calculated. The percentage of reduction was then calculated, column 4. The results showed a 45% reduction in protein levels when treated with  $shRNA1^{\beta 2\alpha 2}$ . No significant reduction is observed with cells treated with  $shRNA1^{\beta 3\alpha 2}$  or  $shRNA3^{\beta 3\alpha 2}$ , column 4 rows 3 and 4. The SD and the student t-test are shown in columns 5 and 6.

The data were used to generate the bar chart shown in figure 4.27.



Figure 4.27 Western blot analysis of the  $\beta 2\alpha 2$  fusion protein from KCL-22 transfected cells.

The bar chart shows Western blot results from three different transfection experiments. Each bar is generated from the average of three Western blot repeats. The `positive' control, cells transfected with the pB12mcs-U6-eGFP vector, was selected to be 1 and the data from other transfected cells were calculated with regards to the positive control. The chart demonstrates that the shRNA1<sup> $\beta$ 2\alpha2</sup> construct is specific for its fusion point and has reduced its target's protein levels to 55%. In the case of the shRNA1<sup> $\beta$ 3\alpha2</sup> and shRNA3<sup> $\beta$ 3\alpha2</sup> constructs, little effect on the  $\beta$ 2\alpha2 protein levels was seen. \*\*\*p < 0.0001.

The data generated from measuring band intensities of the Western blots, show a reduction in  $\beta 2\alpha 2$  BCR-ABL fusion protein only when cells are treated with shRNA1<sup> $\beta 2\alpha 2$ </sup>, and not with shRNA1<sup> $\beta 3\alpha 2$ </sup> and shRNA3<sup> $\beta 3\alpha 2$ </sup>. Hence, confirming the specificity of the constructs to their target gene.

### 4.3 Discussion

In this chapter the effects of some of the constructed shRNA molecules were tested and studied. This was conducted through *in vitro* studies. Three of the constructed shRNA molecules were transfected by nucleofection into K562 and KCL-22 cell lines. Following confirmation of successful transfection by fluorescence microscopy, FACS analysis was performed to quantify transfection and the transfection efficiencies were calculated. mRNA and protein extractions were conducted and the effects of the shRNA constructs were examined at the mRNA levels using qPCR and protein levels using Western blotting.

FACS results have indicated that the negative control, pB12mcs-eGFP with no promoter, gave GFP fluorescent signals. This background fluorescence maybe due to cell damage by the physical process of nucleofection, or due to intrinsic fluorescence that are present in all cells due to mitochondria, lysosomes, aromatic amino acids, and other endogenous fluorophores such as NADPH (Srivastava *et al.*, 2011 and Mosiman *et al.*, 1997).

The qPCR results showed that the mRNA levels of the targeted genes were reduced. This was true for both cell lines. In the K562 cell line the shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> construct was more efficient in reducing its target's mRNA levels than the shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> construct. The first exhibited a 66.4% depletion in the concentration of the  $\beta$ 3 $\alpha$ 2 mRNA levels, whereas the second only exhibited a 32% depletion. The specificity of both constructs was also tested, and it was found that both construct were specific to their target gene. This was concluded because the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> constructs did not reduce the mRNA levels of the  $\beta$ 2 $\alpha$ 2 variant when transfected into KCL-22 cell line. In addition, the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> construct did not affect the normal levels of the *BCR* and *ABL* genes.

In the case of the second BCR-ABL variant  $\beta 2\alpha 2$ , the KCL-22 cell line showed a reduction in the mRNA levels of the  $\beta 2\alpha 2$  fusion point, when transfected with the shRNA1<sup> $\beta 2\alpha 2$ </sup> construct. The construct caused a maximum
of 66% depletion in the mRNA levels. The specificity of this construct was also tested, where the construct was used to transfect the K562 cell line. qPCR results of this transfection demonstrated that there was no significant reduction in the mRNA levels of the  $\beta 3\alpha 2$  fusion point. In addition, the shRNA1<sup> $\beta 2\alpha 2$ </sup> construct did not affect the normal expression levels of the *BCR* and *ABL* genes when transfected into the KCL-22 cells, only reductions in the  $\beta 2\alpha 2$  gene were observed. Hence, confirming the specificity of the synthesised shRNA1<sup> $\beta 2\alpha 2$ </sup> molecule to its target site.

Studies conducted at the protein level were performed by Western blotting. The results obtained from the Western blots with shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA<sup> $\beta$ 2 $\alpha$ 2</sup> constructs confirmed those from the qPCR, where reductions of both BCR-ABL fusion proteins were observed. In the case of the  $\beta$ 3 $\alpha$ 2 fusion protein, shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> showed a higher reduction in the protein levels in comparison with shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>. Statistical analysis showed that the reduction levels were significant. The Western blots also confirmed the qPCR findings that the synthesised shRNA molecules were specific to their target sequence.

In conclusion, the results shown in this chapter give evidence that the constructed shRNA molecules do indeed specifically target their mRNA sequences causing a reduction in both mRNA and protein levels.

#### Chapter 5 : Discussion

#### 5.1 Discussion

Chronic myeloid leukaemia (CML) was one of the first cancers to be linked to a chromosomal abnormality (Sawyers, 1999), the Philadelphia chromosome. This chromosome results from a translocation between chromosomes 9 and 22, where the *ABL* gene on chromosome 9, a tyrosine kinase, is translocated to the *BCR* gene region on chromosome 22 giving rise to an abnormal *BCR-ABL* fusion gene (Rowley, 1973). The resultant fusion gene has an abnormally upregulated tyrosine kinase activity (Weisberg and Griffin, 2000) that results in an increase in the proliferation of immature white blood cells, thus leading to the development of CML. There are several breakpoints that can occur in the *BCR* gene two of which give rise to 95% of CML cases (Evans and Sillibourne, 1996). These fusion points are called the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  depending on where the chromosomal breakages occur in the *BCR* gene (Van Der Burg *et al.*, 2004).

Like all cancers, the treatment approach in CML is multiple sessions of chemotherapy via the usage of drug cocktails (Hoffbrand *et al.*, 2006). Drugs for the disease have been developed over the years, differing in their actions and target sites. With further understanding of the molecular pathology of CML, new drugs have been designed and developed. These drugs have aimed at inhibiting the *ABL* tyrosine Kinase activity (Milojkovic and Apperiey, 2008, Puttini *et al.*, 2008) and, are named tyrosine kinase inhibitors (TKI). However, these drugs are only treatments, not cures, for CML; patients tend to relapse, and the issue of drug toxicity still remains. The only cure available for this disease is allogeneic bone marrow (BM) transplants (Fausel, 2007),

Various drugs exist for CML and the TKI Imatinib is considered to be the first line of treatment for patients with the disorder. Imatinib (Gleevec<sup>®</sup>), a TKI that was developed by Novartis, USA, has shown promising outcomes with

patients establishing a 90% complete hematologic response upon treatment with the drug. However, cases of drug resistance have occurred leading to patient relapse, which led to the development of other tyrosine kinase inhibitors such as Nilotinib (Tasigna®), produced by Novaris, USA (Shieh *et al.*, 2011), and Dasatinib (Sprycel®) (Steinberg, 2007, Talpaz *et al.*, 2006, and Shieh *et al.*, 2011), produced by Bristol Pharmaceuticals, Switzerland. Unfortunately, both drugs display toxic side effects and the issue of drug resistance still remains, favouring the need for development of further drugs and other therapeutic approaches, and making the modulation of gene expression a promising strategy for potential CML treatment.

This project aims to develop an alternative therapeutic approach for CML by the use of shRNA that target and down regulate the expression of the two BCR-ABL variants accounting for 95% of CML cases.

## 5.1.1 Development and Expression of shRNA constructs targeting the BCR-ABL variants

K562 and KCL-22 cell lines were selected for the purpose of this study. The K562 cell line is a CML cell line with the  $\beta 3\alpha 2$  variant (Lozzio and Lozzio, 1975), whereas the  $\beta 2\alpha 2$  variant is present in the KCL-22 cell line (Kubinishi and Miyoshi, 1983). shRNA molecules were designed based on the sequences of these fusion points, and delivered endogenously along with a reporter gene system. The first attempt to construct a reporter gene system used the pEGFP-N1 vector. This vector has an eGFP and a CMV promoter. The eGFP is a mutated form of GFP that is found to be thermostable at 37°C in mammalian cultured cells. Hence, it has greater and improved levels of fluorescence when compared the wild type protein (Siemering *et al.*, 1996), making its use advantageous in the current study. Also, the CMV promoter was to be removed from the vector and the shRNA molecules placed under the regulation of the human U6 and H1 promoters. This was because these

promoters are members of the RNA Polymerase III (Pol III) promoter family, and transcribe successfully any sequence no more than 400 bases (Tuschl, 2002) without addition of extra bases flanking sequences of interest. Previous studies have shown that chemically and T7-synthesized shRNAs exhibit 3' or 5' overhang extensions, which tend to reduce the efficacy of the synthesized shRNA (Paddison *et al.*, 2002). In addition, transcription with U6 and H1 promoters is easily terminated by a stretch of 6 'T' residues, making them ideal for the expression of siRNAs and shRNAs (Tuschl, 2002). Furthermore, these promoters can transcribe small RNA transcripts that lack the 5' cap and 3'polyadenosine (Wu *et al.*, 2005).

Attempts to remove part of the CMV promoter and re-circularize the double digested pEGFP-N1 vector were undertaken and were successful (previously shown in chapter 3, sections 3.2.2.1 and 3.2.2.2). However, it proved difficult to transform E.coli with the altered pEGFP-N1 plasmid. In every attempt, colony screening showed that E.coli were transformed with only the wild type vector. This may be due to the fact that not all the CMV promoter was removed and the part still intact within the vector was toxic for the growth of E.coli bacterial cells. This explanation was considered due to the fact that the wild type vector always got cloned, rather then the altered one. In cases where attempts to eliminate the presence of the wild type vector in the sample was undertaken, none of the vectors were cloned. In future studies, it would be interesting to see which part of the CMV promoter is toxic to the cells. A previous study done by Trudel et al. in 1996 found that a part of the GATA-1 eukaryotic transcription factor was toxic to bacterial cells. This toxic peptide was then used to develop a positive selection vector called pGATA (Clonesure<sup>™</sup>), where inserts caused a disturbance to the gene encoding the peptide. Something similar may be done after identifying the toxic part of the CMV promoter.

Because of the above difficulties, another plasmid was considered in order to construct the reporter gene system. This was the pB12mcs-eGFP plasmid, which has an eGFP as a reporter gene, and lacks a promoter. Therefore, it was a suitable choice because the human U6 or the H1 promoter could be inserted into it.

The shRNA expression cassettes, containing both the promoter and shRNA molecules, were synthesised by PCR, and inserted into the pB12mcs-eGFP plasmid. The shRNA molecules designed in this study were designed to be as symmetrical as possible to aid the binding to their target site and cleavage of the correct gene. All constructs were sequenced before testing in expression studies. To conduct expression studies, the pB12mcs-U6-shRNA-eGFP constructs were delivered into K562 and KCL-22, suspension cell lines. Because primary cells and cells in suspension are hard to transfect (Ovcharenko et al., 2005, Oliveira and Goodell, 2003 and Merkerova et al., 2007), several chemical transfection agents were tested, none was found to give good transfection efficiency (45-50% for both cell lines). Consequently, other methods of transfection were used. It was found that nucleofection, a form of electroporation, where the DNA molecules of interest are directly delivered into the target cell's nucleus gave acceptable transfection efficiencies, a maximum of 72% for K562 and a maximum of 67% for KCL-22. In addition nucleofection gave a good percentage of cell viability, an average of 89.3% for K562 and 90.8% for KCL-22. The `negative' control (pB12mcs-eGFP with no promoter) showed auto-fluorescence; therefore, the transfection efficiencies with regards to the `negative' control, as well as the non-transfected cells were calculated. However, the transfection efficiencies shown in the current study are those with regards to the non-transfected cell lines. The auto-fluorescence observed on the `negative' control maybe due to cell damage or due to intrinsic levels of fluorescence due to mitochondria, lysosomes, aromatic amino acids, and other endogenous fluorophores such as NADPH (Srivastava et al., 2011 and Mosiman et al., 1997). Cell damage might have occurred due to the physical nature of nucleofection. Thus, suggesting that other methods of transfection avoiding cell damage and at the same time giving high transfection efficiencies are required for CML cell lines. In future developments, attempts to quench auto-fluorescence through staining with trypan blue before FACS analysis can be applied (Srivastava *et al.,* 2011 and Mosiman *et al.,* 1997).

Merkerova *et al.*, 2007, examined two physical transfection methods including nucleofection on CML primary cells and three chemical methods to transfect leukemic cell lines and primary CML cells. Their study reported that two chemical methods gave 35-40% efficiency while the third one gave no satisfactory transfection in leukaemic cell lines; the efficiency for the last was not reported nor the type of the cell lines used in the study. The efficiency of nucleofection using the program T-16 on primary CML cells was 25-29%. Another study done by Mahmodabady *et al.*, 2010, also compared nucleofection to other chemical methods in K562 cells. The study found that nucleofection gave higher transfection efficiencies, 50%, when compared to lipofectamine and CaCl<sub>2</sub> methods, <50% and <20% respectively; cell viability was not mentioned. Nevertheless, both studies give evidence supporting the results in the current study on the difficulty of transfection in regards to CML cell lines.

The reason behind the K562 and KCL-22 cell lines being difficult to transfect maybe due to the fact that these cells are morphologically spherical cells (Bártová *et al.*, 2005) that float within the media, in contrast to adherent cells that become immobilized and adhere to the bottom of the culture flask. Adherent cells become flatted and spread, gaining more surface area in comparison to the spherical K562 and KCL-22 cell lines. This would result in easier, more abundant take up of the chemically manipulated DNA complexes used in transfection. No program, or a transfection kit, was available for the KCL-22 cell line, therefore the same program and Kit used for the K562 cell line were tested and were found to give satisfactory results, average of 61.3% for transfection efficiency and 90.8% for viability.

# 5.1.2 Efficiency of shRNA constructs targeting the β3α2 fusion point

Two shRNA constructs, shRNA1 $^{\beta 3\alpha 2}$  and shRNA3 $^{\beta 3\alpha 2}$ , under the regulation of the U6 promoter were tested against the  $\beta 3\alpha 2$  and  $\beta 2\alpha 2$  variants. The  $\beta 3\alpha 2$ and  $\beta 2\alpha 2$  BCR-ABL mRNA expression levels were examined 24 hr after transfection, and were quantified by the use of real time PCR. The results of this reaction demonstrated a decrease in the levels of only  $\beta 3\alpha 2$ . The results show that the mRNA expression levels of  $\beta 3\alpha 2$  fusion product were depleted by 66.4% with shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and 38% with shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>. Further studies showed that the constructs were specific for the  $\beta 3\alpha 2$  fusion point and that they did not target the  $\beta 2\alpha 2$  fusion point. The different efficiencies of shRNA1<sup> $\beta 3\alpha 2$ </sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> could be due to the fact that the U6 promoter expressing the shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> molecule had 3 mismatches compared to 2 mismatches in the U6 promoter regulating the expression of the shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> molecule. Mismatches occurred in the U6 promoter even though a high fidelity Taq polymerase was used (Fast Start High Fidelity PCR system, Roche, UK). For future studies, it would be interesting to examine what effect the mismatches in U6 may have on the effectiveness of the promoter. Another reason could be due to the difference in sequences of the shRNA molecules, or how the molecules bind to their targets in the cells in regards to the target's secondary structure; the shRNA1 $^{\beta 3 \alpha 2}$  sequence may give the molecule a better efficiency to induce RNAi based silencing than shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>.

Western blot results supported the real time PCR results and demonstrated a reduction in the protein levels and confirmed the specificity of the shRNAs. This reduction was 40% and 33.5% for shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup> and shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup>, respectively, in relation to the positive control. These results demonstrate that the shRNA molecules tested had an effect on both mRNA and protein levels of the *BCR-ABL* fusion gene.

### 5.1.2.1 Delivery and production of the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> Constructs

Mahmodabady et al., 2010, used shRNA targeting the  $\beta 3\alpha 2$  variant in K562 cell lines. Their publication reported that they have established 50% transfection efficiency with the use of electroporation, however, Lipofectamine<sup>TM</sup> 2000 that gave lower transfection efficacy was used to carry out their transfection experiments (the actual percentage of transfection efficiency was not reported but only a bar chart represented the results). Arthanari et al., 2010, also transfected K562 cells with shRNA constructs; transfection was done through chemical methods that involved lipofectamine showing transfection efficiencies between 20-50%. In the current study, K562 cells were transfected by nucleofection and a maximum transfection efficiency of 72% was established when using non-viral vectors for shRNA delivery. Other studies have achieved higher transfection efficiencies for shRNA delivery including those using a lentiviral system. Examples include studies done by Myssina et al., 2008 and Li et al., 2003. In general lentiviral vectors, have higher transfection efficiencies than other non-viral systems, and the advantage of long stable expression. However, these vectors are highly toxic (Wang *et al.*, 2008).

Different promoters can be used to produce shRNA molecules, however, as discussed earlier, the small nuclear RNA Pol III promoters have been used to develop transcripts to induce RNAi, especially the U6 and H1 promoters (Wu *et al.*, 2005). Mahmodabady *et al.*, 2010, used the H1 promoter to express shRNA molecules used in their study. Myssina *et al.*, 2009, used a CMV promoter to produce their shRNA molecules. The CMV promoter is a Pol II promoter; these promoters have also been shown to produce shRNA molecules (Rossi, 2008). A study done by Li and Mahato, 2009, to see the effects of H1, U6, and CMV promoters on shRNA mediated gene silencing, found that when the same shRNA molecule was produced from the three different promoters, there was no significant difference in gene silencing between the U6 and CMV promoters. However, H1 had less effect than CMV and U6. Other studies have

also reported that the U6 promoter is more potent than H1 (An *et al.*, 2006 and Snyder *et al.*, 2009). In the present study, the U6 promoter was used to produce shRNA molecules.

# 5.1.2.2 Effects of the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> Constructs on $\beta$ 3 $\alpha$ 2 mRNA and protein levels

Mahmodabady et al., 2010, showed significant reduction in the BCR-ABL mRNA levels, up to 42%, 24hr after shRNA transfection. However, to measure the mRNA levels, RT-PCR was performed and PCR was used to amplify the RT-PCR products. A more accurate way of measuring  $\beta 3\alpha 2$  mRNA levels is through qPCR as attempted in this study. Myssina et al., 2009, showed that their designed shRNA against the  $\beta 3\alpha 2$  variant had an effect on mRNA and protein levels, where a 75% and 95% reduction, respectively were reported. However, this was measured 48hr after transfection on a 100% transfected cell population (FACS was used to select GFP+ cells) that was transfected with high concentrations of shRNA ( $5\mu g/10^6$  cells), which was expressed from a lentiviral system. These can be the reasons for achieving the high levels of reduction. This was also the case in the study conducted by Li et al., in 2003 that used a lentiviral vector to express shRNAs, under the regulation of the U6 promoter. The latter, as discussed previously, also yields high levels of gene expression, hence the observation of >90% depletion in both mRNA and protein levels. Nevertheless, in the current study a non-viral system has demonstrated successful depletion of both mRNA (66.4%) and protein levels (40%) when under regulation of the U6 promoter, and a shRNA concentration of  $2\mu g/2.5 x$ 10<sup>6</sup> cells. The effects of the shRNA constructs used in the current work might have greater effects if a lentiviral system were used.

Arthanari *et al.*, 2010, have shown that an shRNA expressed in the K562 cell line leads to a decrease in  $\beta 3\alpha 2$  mRNA levels of 70% and protein of 75%. mRNA levels were measured after 48hr and protein levels after 96hr, when cells were

treated with 10µg of shRNA. A further decease of mRNA to 85% and protein to 90% was observed at 30µg of shRNA. The depletion levels were measured after 48 and 96hr in comparison to only 24hr in the present work. These high levels of depletion might be due to this reason. Another explanation for achieving such high depletion levels could be due to the high shRNA vector concentrations used by Arthanari *et al.*, 2010. As they have used 10-30µg of shRNA vector per 1.5 x  $10^6$  cells, which is 5-15x more the concentration of shRNA, used in the present study (2µg/2.5 x  $10^6$ ). Greater levels of depletion might have been accomplished in the current project, if higher shRNA vector concentrations had been used.

In the current work, the shRNA<sup> $\beta$ 3 $\alpha$ 2</sup> constructs demonstrated high specificity to their target site because these molecules induced RNAi mediated gene silencing only to their target sites and not to other genes such as *BCR-ABL* fusion point ( $\beta$ 2 $\alpha$ 2), the normal *ABL*, and *BCR* genes, in which 48-50% similarity in sequence exist. Other studies testing shRNA molecules against the  $\beta$ 3 $\alpha$ 2 either considered their molecules specific on the basis that the levels of *GAPDH* housekeeping were not affected, or by having a scramble sequence of the same shRNA used that did not target any human gene (Mahmodabady *et al.*, 2010). Myssina *et al.*, 2009 and Li *et al.*, 2003, demonstrated specificity of their shRNA molecules in regards to the  $\beta$ 2 $\alpha$ 2 variant and did not consider the normal *ABL* and *BCR* genes.

#### 5.1.2.3 Conclusion

In conclusion, the shRNA molecules targeting the  $\beta 3\alpha 2$  fusion gene showed specific reduction of their target site mRNA (66.4%) and protein levels (40%), which were achieved 24hr after transfection with the shRNA constructs and with a relatively low concentration of shRNA (2µg/2.5 x 10<sup>6</sup> cells). The BCR-ABL protein has been reported to have a long half-life (Myssina *et al.*, 2008 and Wilda *et al.*, 2002) that is estimated to be >48hr (Spiller *et al.*, 1998).

Therefore, higher levels of reduction at protein levels can be expected for the shRNAs used in the current study over a longer period of time.

### 5.1.3 Efficiency of shRNA constructs targeting the β2α2 fusion point

The  $\beta 2\alpha 2$  BCR-ABL mRNA expression levels were examined 24 hours after transfection, and were quantified by the use of real time PCR. The results demonstrated a decrease in the level of the  $\beta 2\alpha 2$  BCR-ABL variant. One shRNA constructs, shRNA1<sup> $\beta 2\alpha 2$ </sup>, under the regulation of the U6 promoter was tested. The results show that the mRNA expression levels of the  $\beta 2\alpha 2$  fusion product, and not  $\beta 3\alpha 2$  were depleted by 66% with shRNA1<sup> $\beta 3\alpha 2$ </sup>. Western blot results demonstrated a reduction in the protein levels of 45%, in relation to the `positive' control. These results show that the shRNA molecules tested had an effect on both mRNA and protein levels of the *BCR-ABL* fusion gene.

### 5.1.3.1 Delivery and production of the shRNA<sup> $\beta 2\alpha 2$ </sup> Construct

The current study demonstrated that KCL-22 cell lines can be transfected by means of nucleofection, and the shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup> construct achieved a relatively high transfection efficiency of 67% by this method. shRNA1<sup> $\beta^{2\alpha^{2}}$ </sup> was cloned in a non-viral vector, pB12mcs-eGFP plasmid, and was under the regulation of the snRNA Pol III U6 promoter. Li *et al.*, 2003, constructed a  $\beta^{2\alpha^{2}}$  shRNA molecule under the regulation of the U6 promoter, however, the construct was inserted into a lentiviral system, and the transduction efficiency reached ~100% in Meg01 leukaemic cell line. Meg01 is a CML cell line that expresses the  $\beta^{2\alpha^{2}}$  variant, and was developed by Ogura *et al.*, in 1985. This is a differentiated cell line where mostly large immature megakaryocytes predominate, unlike the

KCL-22 cell line, used in the current study, that is an undifferentiated haematopoietic cell line (Kamamoto *et al*, 1986). In summary, Li *et al.*, 2003, used a different cell line and a viral system to express their shRNA constructs that can both explain the high transduction efficiency achieved in their study, in comparison to the non-viral vectors used in the current study.

# 5.1.3.2 Effects of the shRNA<sup> $\beta^{2\alpha^{2}}$ </sup> Constructs on $\beta^{2\alpha^{2}}$ mRNA and protein levels

In the present study a reduction in the  $\beta 2\alpha 2$  mRNA levels of 66% and protein levels of 45% was observed 24hr after transfection with shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup>. In contrast Li *et al.*, 2003, showed no significant reduction in the  $\beta 2\alpha 2$  variant. This maybe the result of differences in the shRNA structure and sequence used in their study. The current study used an shRNA molecule of 21 nt long and an open stem loop of 8 nt. Li et al., used the same shRNA sequence but with a 9 nt loop and a G overhang at the 5' end on the antisense strand. It has been reported that the loop size, structure, and its sequence, affects the knockdown efficiencies of shRNA molecules (Brummelkamp et al., 2002 and Schopman et al., 2010). Schopman and colleges, 2010, have found that improvements on the 9 nt long stem loop used in Brummelkamp et al., 2002, studies are achievable by using different structures of the stem loop. Schopman et al., 2010, demonstrated that even weak shRNA inhibitors could be turned into strong gene silencers by simply differing the loop length, sequence, or structure. Consequently, these could be the reasons why the shRNA molecules used in Li et al., 2003, study demonstrated lack of  $\beta 2\alpha 2$  gene silencing in comparison to current work.

In the current study, qPCR results also demonstrate that the shRNA1<sup> $\beta$ 2 $\alpha$ 2</sup> construct was specific to its target sequence and no significant reductions in the mRNA expression levels were observed in related and non-related genes, including normal *ABL*, normal *BCR*, *GAPDH* (when transfected into KCL-22 cells)

and the  $\beta 3\alpha 2$  fusion gene. The protein analysis confirmed that the shRNA1<sup> $\beta 2\alpha 2$ </sup> construct was specific to its target site, supporting the qPCR results.

#### 5.1.3.3 Conclusion

In conclusion, the current study demonstrates successful and specific targeting of the  $\beta 2\alpha 2$  variant by a non-viral system expressing shRNA molecules under the regulation of the U6 promoter. The effects of the shRNA used in the present work might have greater effects if a lentiviral system was used. Lentiviral systems have higher transfection efficiencies than non-viral systems; this is mainly because lentiviruses have the ability to integrate into the host's genome, establishing long-term stable expression of the integrated DNA sequence. However, drawbacks exist for these types of system, which include toxicity and oncogenesis (Hacein-Bey-Abina *et al.*, 2008 and Davidson and McCary, 2011).

#### 5.1.4 siRNA against the BCR-ABL Variants

Other studies have targeted the *BCR-ABL* fusion genes through siRNA technology have for example Scherr *et al.*, 2003, Koldehoff *et al.*, 2010, Wohlbold *et al.*, 2003, Bártová *et al.*, 2005, and Withey *et al.*, 2005 using siRNA targeting the  $\beta$ 3 $\alpha$ 2 variant, and Rapozzi and Xodo, 2004 for targeting the  $\beta$ 2 $\alpha$ 2 variant. Findings from Scherr *et al.*, 2003, demonstrated that the siRNA tested had reduced the mRNA levels up to 87% after 24hr. However, this efficiency was obtained from a co-transfection of both a BCR-ABL expressing vector, which was fluorescently labelled, and chemically synthesised siRNA molecules in Hela cells. It was further reported, in the same publication, that when the same siRNA was transfected into K562 cell lines, the mRNA levels of the *BCR-ABL* fusion gene were reduced up to 24.8% after 24hr. Withey *et al.*, 2005, reported a 84% reduction of mRNA levels in K562 cell line. However, these

results were obtained from PCR amplification of the cDNA of the transfected cell lines. A more precise method to measure the mRNA levels would be to use qPCR, which was used in the current study. The shRNA molecules used in the current study have higher potency, sustainable effects using low copy number, and less off-target effects in comparison with siRNAs (Rao *et al.*, 2009).

Studies have also been conducted where antisense technology and drug treatment have been combined to try and reduce CML effect. For example Koldehoff *et al.*, 2010 and Wohlbold *et al.*, 2003 showed that siRNA targeting the BCR-ABL fusion point had additive effects with Nilotinib and Imatinib. In addition, the study done by Wohlbold *et al.*, 2003, suggested that targeting the mRNA by the process of RNAi sensitizes the drug resistant leukaemic cells to TKI. The findings of these studies suggest future strategies combining both molecular targeting of the *BCR-ABL* fusion gene at the mRNA level by the use of RNAi and using TKI on the protein level.

Bártová *et al.*, 2005, demonstrated the use of siRNA against the *BCR-ABL* fusion gene. However, the siRNA designed in this study targeted the *BCR-ABL* downstream of the fusion point, thus also targeting the normal *c-ABL*. In addition, the transfection levels of the siRNA only ranged between 25-30% which is considered to be quite low.

A study done by Rapozzi and Xodo, in 2004, showed that significant reductions in mRNA and protein levels of the  $\beta 2\alpha 2$  BCR-ABL variant could be achieved with single stranded (40% for mRNA and 28% for protein) and double stranded siRNA (20-40% for mRNA and 83% for protein) within 24hr. Analysis was carried out 24 and 48hr after two electroporation treatments. Thus, high doses of siRNA were required to achieve these results. In addition, the effects of the siRNA molecules only lasted for 72hr after treatment, indicating the transient effect of siRNA molecules.

In conclusion, RNAi technology is a powerful tool for down regulation of genes of interest. In the current project, this technology was used to down regulate the expression of the abnormal *BCR-ABL* gene, more precisely the upregulated tyrosine kinase activity of the *ABL* part of the fusion gene. The constructed shRNA molecules, targeting both BCR-ABL variants, show

specificity to their target sites and caused depletion in both mRNA and protein levels. The U6 promoter showed ease of production and construction of shRNA expression cassettes. The promoter also displayed efficient synthesis of shRNA molecules. In this context, shRNA constructs demonstrated in the current work can be promising tools for CML therapy when used alone or coupled with TKI's.

#### 5.1.5 Future developments

Future developments to be considered in this project include the testing of the other constructs that have been synthesised and sequenced; transfection for longer periods of time, such as 48 and 72 hours; and a comparison the effects of different concentration of the shRNA molecules on their target sites. Once an appropriate time period and concentration has been achieved, it would also be appealing to compare the effects of the same shRNA molecules but under the regulation of the two different human promoters, the U6 and H1, as only the U6 promoter was tested in the current project. The effects of the shRNA on growth inhibition, cell proliferation, and differentiation could be also investigated.

It would also be intriguing to examine the tested shRNA constructs in an animal model. To do so, first, a suitable delivery system needs to be chosen. These systems can be either viral or non-viral systems, with each option having its own pro and cons. Non-viral vectors avoid issues such as immunogenicity and insertional mutagenesis; however, these vectors suffer from low transduction efficiency, and non-permanent expression. Viral vectors, on the other hand, overcome the disadvantages of non-viral vectors, in terms of efficiency and provide stable expression, but, give rise to issues relating to toxicity, safety, immunogenicity, and specificity of delivery. Other vectors to consider are the new hybrid vectors, engineered to have qualities of more than one vector (Wang *et al.*, 2008). Rumpold *et al.*, 2005, constructed a non-viral transposon-based vector that demonstrated stable expression of siRNAs, that

caused knockdown to their target genes. However, the system was described to have a laborious production, systematic instability and had a pathogenicity issue. Therefore, great care should be taken when deciding what type of vector system is chosen to express the shRNA molecules *in vivo*.

Secondly, once the appropriate vector system has been chosen, another concern is the method and specificity of delivery to the target cells. The target cells in CML will not only be the cancerous cells but also the abnormal hematopoietic stem cells that are hard to identify. Recently, researchers at Lund University have shown that CML stem cells might be isolated through a cell surface marker called IL-1 receptor accessory protein (Luo, 2010). This can be advantageous, as it can help the identification of stem cells with the Philadelphia chromosome. These stem cells can be therefore isolated and used for the conduction of *ex vivo* studies, providing a way to achieve specific targeting of the abnormal cells of interest. Another aspect to consider is cellpenetrating peptides. This technology has been successfully considered as an application for in vivo delivery of various molecules including shRNAs (Wang et al., 2008). A study done by Arthanari el al., 2010, showed that by combining the cell penetrating peptide HIV-Tat with the membrane lytic peptide LK-15, higher transfections efficiencies could be achieved for delivery of shRNA targeting the BCR-ABL fusion gene in K562 cell lines. These finding make the technology of cell-penetrating peptides an appealing area to explore when considering in vivo studies.

The next stage would be to choose the appropriate animal model. This is of great importance. It would be highly interesting to see how the shRNA molecules react *in vivo* and how the animal reacts to such treatment. The Severe Combined Immunodeficiency (SCID) mouse or murine CML-like models could be considered. The SCID mouse results from a mutation that impairs the cell development in the lymphoid lineage making the animal severely immune-deficient. This leads to the animal accepting human exogenic engraftments without initiating an immune response. A study done by Wang *et al.*, in 1998 showed the establishment of high levels of engraftments by SCID mice with

haematopoietic cells from patients with CML. Thus, making this model highly considerable for future studies.

Other murine CML-like models have also been developed to study CML through *in vivo* studies. These include mice where both transgenic and retroviral transduction approaches have been employed, the latter seems to be more promising in regards to establishing a realistic murine CML-like model (Van Etten, 2002). A study done by Wertheim *et al.*, in 2002 has shown a successful attempt to generate a CML-like murine model. Scientists in this study developed a protocol to transfuse high titre retroviral vectors into murine bone marrow causing the haematopoietic stem cells to have an increased expression of the BCR-ABL protein. Furthermore, lethally irradiated mice were transplanted with the transfused bone marrow cells leading to the development of fatal myeloproliferative disorders in 100% of the mice. Therefore, this model can also be considered. However, more research into this area should be conducted in order to select the best CML model before clinical trials are attempted.

# Appendix 1: The Molecular Weight Markers Used in The Current Project

Five types of molecular weight markers were used to determine band sizes. The  $\lambda$ /EcoRI/HindIII marker, and the GeneRuler<sup>TM</sup> 1Kb marker were used for determination of large band sizes and on a 1% agarose gel. The pBluescript Hpa II marker, the 100bp DNA Ladder, and the GeneRuler<sup>TM</sup> 100bp marker were used for smaller size bands ranging from 1000bp to as small as 100bp in size. These markers were usually used on 2% agarose gels.



Figure 67 The  $\lambda$ /EcoRI/HindIII Marker. Taken from the Promega website

### GeneRuler<sup>™</sup> 1 kb DNA Ladder O'GeneRuler<sup>™</sup> 1 kb DNA Ladder, ready-to-use

	bp ng/	0.5 µg	%
#R0491)	10000 8000 5000 4000 3500 3500 2500 2000 1500	30.0 30.0 <b>70.0</b> 30.0 30.0 30.0 <b>70.0</b> 25.0 25.0 25.0	6.0 6.0 6.0 6.0 6.0 <b>14.0</b> 5.0 5.0 5.0
arose	1000 750	<b>60.0</b> 25.0	<b>12.0</b> 5.0
- egg 40	500	25.0	5.0
ÿ <b>—</b> —	250	25.0	5.0
1% TopVisio			
0.5 μg/lane, 8 α 1X TAE, 7 V/cm	em lengt , 45 min	h gel,	

Figure 68 The GeneRuler<sup>™</sup> 1Kb marker. Taken from the Thermo Scientific website



Figure 69 The pBluescript Hpall Marker.



Figure 70 The 100bp DNA Ladder. Taken from the NEB website



Figure 71 The GeneRuler<sup>™</sup> 100bp Marker. Taken from the Thermo Scientific website

### **Appendix 2: The Secondary shRNA Structures**

The following figures are the secondary structures predictions of the designed shRNA molecules. The program "mfold" was used to predict these secondary structures. The program is found in the following website: (http://mfold.rna.albany.edu/?q=mfold/RNA-Folding-Form)



Figure 72 The Secondary structure prediction of shRNA2  $^{\beta3\alpha2}$  by "mfold"



Figure 73 The Secondary structure prediction of shRNA3  $^{\beta3\alpha2}$  by "mfold"



Figure 74 The Secondary structure prediction of shRNA1  $^{\beta 2\alpha 2}$  by "mfold"



Figure 75 The Secondary structure prediction of shRNA2  $^{\beta 2\alpha 2}$  by "mfold"



Figure 76 The Secondary structure prediction of shRNA3  $^{\beta2\alpha2}$  by "mfold"

Appendix 3: The Protein Molecular Weight Marker used in the Current Project



Figure 77 The High-Range Rainbow protein molecular weight marker.

### Appendix 4: The $\beta 3\alpha 2$ and $\beta 2\alpha 2$ Sequences

• The β3α2 fusion point sequence from the EMBL website (Accession number AJ131466).

1	gcgaacaagg	gcagcaaagc	tacggagagg	ctgaagaaga	agctgtcgga	gcaggagtca
61	ctgctgctgc	ttatgtctcc	cagcatggcc	ttcagggtgc	acagccgcaa	cggcaagagt
121	tacacgttcc	tgatctcctc	tgactatgag	cgtgcagagt	ggagggagaa	catccgggag
181	cagcagaaga	agtgtttcag	a <mark>agcttctcc</mark>	ctgacatccg	<mark>tgga</mark> gctgca	gatgctgacc
241	aactcgtgtg	tgaaactcca	gactgtccac	agcattccgc	tgaccatcaa	taaggaagat
301	gatgagtctc	cggggctcta	tgggtttctg	aatgtcatcg	tccactcagc	cactggattt
361	aagcagagtt	c <mark>aaaa</mark> gccct	tcagcggcca	gtagcatctg	actttgagcc	tcagggtctg
421	agtgaagccg	ctcgttggaa	ctccaaggaa	aaccttctcg	ctggacccag	tgaaaatgac
481	cccaaccttt	tcgttgcact	gtatgatttt	gtggccagtg	gagataacac	tctaagcata
541	actaaaggtg	aaaagctccg	gg <mark>tcttaggc</mark>	tataatcaca	<mark>atggg</mark> gaatg	gtgtgaagcc
601	caaaccaaaa	atggccaagg	ctgggtccca	agcaactaca	tcacgccagt	caacagtctg
661	gagaaacact	cctggtacca	tgggcctgtg	tcccgcaatg	ccgctgagta	tctgctgagc
721	agcgggatca	atggcagctt	cttggtgcgt	gagagtgaga	gcagtcctgg	ccagaggtcc
781	atctcgctga	gatacgaagg	gagggtgtac	cattacagga	tcaacactgc	ttctgatggc
841	aagctctacg	tctcctccga	gagccgcttc	aacaccctgg	ccgagttggt	tcatcatcat
901	tcaacggtgg	ccgacgggct	catcaccacg	ctccattatc	cagccccaaa	gcgcaacaag
961	cccactgtct	atggtgtgtc	ccctaactac	gacaagt		

The fusion point is underlined and highlighted in yellow. The forward (in green) and reverse (in blue) primers to amplify the fusion point are highlighted.

• The  $\beta 2\alpha 2$  fusion point sequence from the EMBL website (Accession number AJ131467).

1	gcgaacaagg	gcagcaag <mark>gc</mark>	tacggagagg	<mark>ctgaagaa</mark> ga	agctgtcgga	gcaggagtca
61	ctgctgctgc	ttatgtctcc	cagcatggcc	ttcagggtgc	acagccgcaa	cggcaagagt
121	tacacgttcc	tgatctcctc	tgactatgag	cgtgcagagt	ggagggagaa	catccgggag
181	cagcagaaga	agtgtttcag	aagcttctcc	ctgacatccg	tggagctgca	gatgctgacc
241	aactcgtgtg	tgaaactcca	gactgtccac	agcattccgc	tgaccatcaa	taagga <mark>agaa</mark>
301	gcccttcagc	ggccagtagc	atctgacttt	gagcctcagg	gtctgagtga	agccgctcgt
361	tggaactcca	aggaaaacct	tctcgctgga	cccagtgaaa	atgaccccaa	ccttttcgtt
421	gcactgtatg	attttgtggc	cagtggagat	aacactctaa	gcataactaa	aggtgaaaag
481	ctccgggtct	taggctataa	tcacaatggg	gaatggtgtg	aagcccaaac	caaaaatggc
541	caaggctggg	tcccaagcaa	ctacatcacg	ccagtcaaca	gtctggagaa	acactcctgg
601	taccatgggc	ctgtgtcccg	caatgccgct	gagtatctgc	tgagcagcgg	gatcaatggc
661	agcttcttgg	tgcgtgagag	tgagagcagt	cctggccaga	ggtccatctc	gctgagatac
721	gaagggaggg	tgtaccatta	caggatcaac	actgcttctg	atggcaagct	ctacgtctcc
781	tccgagagcc	gcttcaacac	cctggccgag	ttggttcatc	atcattcaac	ggtggccgac
841	gggctcatca	ccacgctcca	ttatccagcc	ccaaagcgca	acaagcccac	tgtctatggt
901	gtgtccccca	actacgacaa g	ŋt			

The fusion point is underlined and highlighted in yellow. The forward and reverse primers to amplify the fusion point are highlighted in green and blue respectively.

# Appendix 5: The Raw Sequencing Data of the $\beta 3\alpha 2$ and $\beta 2\alpha 2$ Fusion Point Sequences in the pGEM<sup>®</sup>-T Easy Vector

 3 colonies were sequenced, and all showed similar results. The β3α2 fusion point sequence for a representative colony is shown bellow, colony A5. The fusion point is marked by "]":

NNNNNNNNNNNNNNATGNNNNNNGCCGCCATGGCGGCCGCGGGAATTCGATTAGCTTCTCCCTGGCAT CCGTGGNAGCTTCTCCCTGGCATCCGTGTCTTAGGCTATAATCACAATGGGGCTCCCTGGCATCCGTGGAG GATCTTAGGCTATAATCACAATGGGACTAGCTTCTCCCTGGCATCCGTGGCTCGTAGTGTTCTGGCGAGCA GTTTTGATTGATTTAACTGTTGAGGCTTAGGCTATAATCACAATGGGACTAGCTTCCCCCTGGCATCCGTGG AGCCGCAGATGCTGACCAACTCGTGTGTGAAACTCCAGACTGTCCACAGCATTCCGCTGACCATCAATAAG GAAGATGATGAGTCTCCGGGGGCTCTATGGGTTTCTGAATGTCATCGTCCACTCAGCCACTGGATTTAAGCA GAGTTCAA AAGCCCTTCAGCGGCCAGTAGCATCTGACTTTGAGCCTCAGGGTCTGAGTGAAGCCGCTCGT TGGAACTCCAAGGAAAACCTTCTCGCTGGACCCAGTGAAAATGACCCCAACCTTTTCGTTGCACTGTATGA TTTTGTGGCCAGTGGAGATAACACTCTAAGCATAACTAAAGGTGAAAAGCTCCGGGTCTTAGGCTATAATC ACAATGGGAATCACTAGTGAATTCGCGGCCGCCTGCAGGTCGACCATATGGGAGAGCTCCCCAACGCGTTGG **ATGCATAGCTTGAGTATTCTATAGTGTCACCTAAATAGCTTGGCGTAATCATGGTCATAGCTGTTTCCTGT GTGAAATTGTTATCCGCTCACAATTCCACAACATACGAGCCGGAAGCATAAAGTGTAAAGCCTGGGGTG** NCTAATGAGTGAGCTAACTCACATTAATTNGCGTTGCGCTCACTGCCCGCTTTCCAGTCGGGAAACCTGTC TCCTCGCTCACTGACTCGCTGCGCTCGGNCGTTCGGCTGCGGCGAGCGNNNNCAGCTCACTCNAAGGCGGN ANNNNNGTNTCCNCAGAATCAGNNNNNACGCAGGAAANAACATGNTGANCNANAAGNCNAGCNAANNCNNN AANCNNNAAAANGCNGNNNNNNNNNNNTTTTNNNNNNCGCCCNNCNGNNNNNNTNNCNAANNGNNNNTC NNNNN

3 colonies were sequenced, and all showed similar results. The β2α2 fusion point sequence for a representative colony is shown bellow, colony KCL-22 `1'. The fusion point is marked by "]":

### Appendix 6: The Human U6 and H1 Promoter Sequences

• The Human snRNA U6 Pol III promoter sequence from the EMBL Website (Accession number M14486).

1	ccccagtgga	aagacgcgca	ggcaaaacgc	accacgtgac	ggagcgtgac	cgcgcgccga
61	gcc <mark>caaggtc</mark>	gggcaggaag	<mark>agggccta</mark> tt	tcccatgatt	ccttcatatt	tgcatatacg
121	atacaaggct	gttagagaga	taattagaat	taatttgact	gtaaacacaa	agatattagt
181	acaaaatacg	tgacgtagaa	agtaataatt	tcttgggtag	tttgcagttt	taaaattatg
241	ttttaaaatg	gactatcata	tgcttaccgt	aacttgaaag	tatttcgatt	tcttggcttt
301	atatatcttg	tggaaaggac	gaaacacc <mark>gt</mark>	gctcgcttcg	gcagcacata	tactaaaatt
361	ggaacgatac	agagaagatt	agcatggccc	ctgcgcaagg	atgacacgca	aattcgtgaa
421	gcgttccata	tttttacatc	aggttgttt	tctgttttta	catcaggttg	tttttctgtt
481	tggtttttt	tttacaccac g	gtttatacgc c	ggtgcacgg t	ctacca	

The forward and reverse primers to amplify the U6 promoter are highlighted in yellow and green respectively.

• The Human RNA H1 Pol III promoter sequence from the EMBL Website (Accession number X16612).

1	ttatagggag	ctgaagggaa	gggggtcaca	gtaggtggca	tcgttccttt	ctgactgccc
61	gccccccgca	tgccgtcccg	cgatattgag	ctccgaacct	ctcgccctgc	cgccgccggt
121	gctccgtcgc	cgccgcgccg	ccatggaatt	<mark>cgaacgctga</mark>	cgtcatcaac	ccgctccaag
181	gaatcgcggg	cccagtgtca	ctaggcggga	acacccagcg	cgcgtgcgcc	ctggcaggaa
241	gatggctgtg	agggacaggg	gagtggcgcc	ctgcaatatt	tgcatgtcgc	tatgtgttct
301	gggaaatcac	cataaacgtg	aaatgtcttt	ggatttggga	atcttat <mark>aag</mark>	ttctgtatga
361	gaccactctt	<mark>tc</mark> ccataggg	cggagggaag	ctcatcagtg	gggccacgag	ctgagtgcgt
421	cctgtcactc	cactcccatg	tcccttggga	aggtctgaga	ctagggccag	aggcggccct
481	aacagggctc	tccctgagct	tcggggaggt	gagttcccag	agaacggggc	tccgcgcgag
541	gtcagactgg	gcaggagatg	ccgtggaccc	cgcccttcgg	ggaggggccc	ggcggatgcc
601	tcctttgccg	gagcttggaa	cagactcacg	gccagcgaag	tgagttcaat	ggctgaggtg
661	aggtaccccg	caggggacct	cataacccaa	ttcagactac	tctcctccgc	ccatttttgg
721	aaaaaaaaaa	aaaaaaaaaa	aacaaaacga	aaccgggccg	ggcgcggtgg	ttcacgccta
781	taatcccagc	actttgggag	gccgaggcgg	gcggatcaca	aggtcaggag	gtcgagacca
841	tccaggctaa	cacggtgaaa	cccccccca	tctctactaa	aaaaaaaaaa	tacaaaaaat
901	tagccattag	ccgggcgtgg	tggcgggcgc	ctataatccc	agctacttgg	gaggctgaag
961	cagaatggcg	tgaacccggg	aggcggacgt	tgcagtgagc	cgagatcgcg	ccgactgcat
1021	tccagcctgg	gcgacagagc	gagtctcaaa	aaaaaaa		

The forward (in yellow) and reverse (in green) primers to amplify the H1 promoter are highlighted.

Appendix 7: Electrophoretic Analysis of Representative Colony Screens PCR Products of the U6 shRNA and H1 shRNA Construct



Figure 78 Electrophoretic analysis of colony screen PCR products of the U6 shRNA1  $^{\beta 3\alpha 2}$ 

#### constrcut

Gel (A): PCR products of the U6 shRNA1  $^{\beta 3\alpha 2}$  constrcut using the T7 forward and T3 reverse primers

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3-7: PCR products of the colony screen

Lanes 3, 6, and 7: positive colonies (~1390bp)

Gel (B): PCR products of pB12mcs-U6-shRNA1<sup> $\beta$ 3 $\alpha$ 2</sup>-eGFP extracted plasmids

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3 and 6: PCR product of the primer set T7 forward and T3 Reverse (~1390bp)

Lanes 4 and 7: PCR products of the primer set T7 forward with U6 shRNA1 $^{\beta 3\alpha 2}$  reverse (~380bp)

Lanes 5 and 8: PCR products of the primer set U6 forward and T3 reverse (~1420bp)



Figure 79 Electrophoretic analysis of colony screen PCR products of the U6 shRNA2 $^{\beta 3 \alpha 2}$  construct

Gel (A): PCR products of the U6 shRNA2  $^{\beta 3\alpha 2}$  construct using the T7 forward and T3 reverse primers

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3-12: PCR products of the colony screen

Lanes 11 and 12: positive colonies (~1390bp)

Gel (B): PCR products of pB12mcs-U6-shRNA2  $^{\beta 3 \alpha 2}$ -eGFP extracted plasmids

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3 and 6: PCR product of the primer set T7 forward and T3 Reverse (~1390bp)

Lanes 4 and 7: PCR products of the primer set T7 forward with U6 shRNA2 $^{\beta 3 \alpha 2}$  reverse (~380bp)

Lanes 5 and 8: PCR products of the primer set U6 forward and T3 reverse (~1420bp) Lane 9: Gene Ruler<sup>™</sup> 100bp DNA Ladder



### Figure 80 Electrophoretic analysis of colony screen PCR products of the U6 shRNA3 $^{\beta 3\alpha 2}$ construct

Gel (A): PCR products of the U6 shRNA3  $^{\beta 3\alpha 2}$  constrcut using the T7 forward and T3 reverse primers

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3-9: PCR products of the colony screen

Lanes 3, 6, and 9: positive colonies (~1390bp)

Gel (B): PCR products of pB12mcs-U6-shRNA3  $^{\beta 3 \alpha 2}$ -eGFP extracted plasmids

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3, 6, and 9: PCR product of the primer set T7 forward and T3 Reverse (~1390bp)

Lanes 4, 7, and 10: PCR products of the primer set T7 forward with U6 shRNA3<sup> $\beta$ 3 $\alpha$ 2</sup> reverse (~380bp)

Lanes 5, 8, and 11: PCR products of the primer set U6 forward and T3 reverse (~1420bp) Lane 12: Gene Ruler<sup>™</sup> 100bp DNA Ladder


Figure 81 Electrophoretic analysis of colony screen PCR products of the H1 shRNA1  $^{\beta 2\alpha 2}$  construct

Gel (A): PCR products of the H1 shRNA1 $^{\beta 2\alpha 2}$  construct using the T7 forward and T3 reverse primers

Lanes 1 and 15: Gene Ruler<sup>™</sup> 1KB DNA Ladder
Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)
Lanes 3-14: PCR products of the colony screen
Lanes 3-14: positive colonies (~1340bp)
Gel (B): PCR products of pB12mcs-H1-shRNA1 <sup>β2α2</sup>-eGFP extracted plasmid
Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder
Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)
Lane 3: PCR product of the primer set T7 forward and T3 Reverse (~1340bp)

Lane 4: PCR products of the primer set T7 forward with U6 shRNA1 $^{\beta 3 \alpha 2}$  reverse (~330bp)

Lane 5: PCR products of the primer set H1 forward and T3 reverse (~1370bp)

Lane 6: Gene Ruler<sup>™</sup> 100bp DNA Ladder





Gel (A): PCR products of the H1 shRNA2  $^{\beta 2\alpha 2}$  construct using the T7 forward and T3 reverse primers

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp) Lanes 3-15: PCR products of the colony screen Lanes 9-10, and 15: positive colonies (~1340bp)

Gel (B): PCR products of pB12mcs-H1-shRNA2  $\beta^{2\alpha^2}$ -eGFP extracted plasmid

Lane 1: Gene Ruler<sup>TM</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lane 3: PCR product of the primer set T7 forward and T3 Reverse (~1340bp)

Lane 4: PCR products of the primer set T7 forward with H1 shRNA2<sup> $\beta$ 3 $\alpha$ 2</sup> reverse (~330bp)

Lane 5: PCR products of the primer set H1 forward and T3 reverse (~1370bp)

Lane 6: Gene Ruler<sup>™</sup> 100bp DNA Ladder





Gel (A): PCR products of the H1 shRNA3  $^{\beta 2\alpha 2}$  construct using the T7 forward and T3 reverse primers

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp) Lanes 3-8: PCR products of the colony screen Lanes 3, 5, and 6: positive colonies (~1340bp)

Gel (B): PCR products of pB12mcs-H1-shRNA3<sup> $\beta$ 2\alpha2</sup>-eGFP extracted plasmid

Lane 1: Gene Ruler<sup>™</sup> 1KB DNA Ladder

Lane 2: PCR pruduct of wild type pB12mcs-eGFP vector (~1050bp)

Lanes 3 and 6: PCR product of the primer set T7 forward and T3 Reverse (~1340bp)

Lanes 4 and 7: PCR products of the primer set T7 forward with H1 shRNA2  $^{\beta 3\alpha 2}$  reverse

(~330bp)

Lanes 5 and 8: PCR products of the primer set H1 forward and T3 reverse (~1370bp)

# Appendix 8: The Raw Sequencing Data of the U6 and H1 Promoters Inserted into the pB12mcs-eGFP Plasmid

- Sequencing of the U6 promoter inserted into the pB12mcs-eGFP
- 1. Colony number 5. The following U6 sequence is given as the forward sequence  $5' \rightarrow 3'$  and is highlighted in yellow. The U6 sequence shows no mismatches to the U6 sequence found in the EMBL website (accession number M14486).

NNNNNNNNNNNNGNNNATCTCGAGAGCTTCAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATTCCTT TTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAAATTATGTTTTAA AATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAG GACGAAACACCGTGCTCGCTTCGGCAGCACATATACTAGGATCCACAGCCACCATGAGTAAAGGAGAAGAA CTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAG CTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCAT ATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGGAGGACCATCTTCTTCAA GGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGC TTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACNACTCCCAC AACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCGA AGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTTAC CAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATGGNC CTTCTTGAGTTTGTACGGCTGCTGGGNNTACNNATGGNATGNNGANTATACAAANNANTCGANCNCTANNA **CTNNANNGNGTCGTANTACGTANATCNNANNTGATAANNANNNATTGATNANTTNGNNNNNNNNANTANNN** NTNNNNNNNCNNNNNTNNNNNTNNGNNNNNNNNNNNGNNNNNNANTTTTTNTNNNN

The following is the same U6 sequence, however, it is given as the reverse sequence  $5' \rightarrow 3'$ .

Colony number 11. The following U6 sequence is given as the forward sequence 5' → 3' and is highlighted in yellow. The U6 sequence shows no mismatches to the U6 sequence found in the EMBL website (accession number M14486).

NNNNNNNNNNNNNNNNNCTCGAGAAGCTT<mark>CAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATTCCT</mark> ATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAAATTATGTTTTA AAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAA GGACGAAACACCGTGCTCGGCTCGGCAGCACATATACTAGGATCCACAGCCACCATGAGTAAAGGAGAAGA ACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCA CCTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCA TATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTTCTTCA AGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAG CTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCA CAACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCG AAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCCAATTGGCGATGACCCTGTCCTTTTA CCAGACAACCATTACCTGTCCACACAAACTGCCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATGGN **CCTTCTTGAGTTTGTAACGGCTGCTGGGATTACNCATGGNATNNNGANTATACNAATAANTCNANNNCTAN ANTATAGTGAGTCGTANTACGTANATCNNANNGANNAGNNNNNNATNNNTTTNGNNNNNCCNNNNNANAN** 

The following is the same U6 sequence, however, it is given as the reverse sequence  $5' \rightarrow 3'$ .

NNNNNNNNNNNNNNANNGGNNAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCCACACCTCCCCCTG AACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATA AAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAAC

Another 6 colonies were also sequenced and were found to have mismatches (raw sequences are not shown).

- Sequencing of the H1 promoter inserted into the pB12mcs-eGFP
- 1. Colony number 8. The following U6 sequence is given as the forward sequence  $5' \rightarrow 3'$  and is highlighted in yellow. The U6 sequence shows no mismatches to the U6 sequence found in the EMBL website (accession number X16612).

NNNNNNNNNNNGNNATCTCGAGAAGCTT<mark>GAACGCTGACGTCATCAACCCGCTCCAAGGAATCGCGGGCC</mark> CANTGTCACTAGGCGGGAACACCCAGCGCGCGCGCGCCCTGGCAGGAAGATGGCTGTGAGGGACAGGGGAG TGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCATAAACGTGAAATGTCTTTGGAT TTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACAGCCACCATGAGTAAAGGAGAAGAA CTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAG CTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCAT ATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTTCTTCAA GGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGC TTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCAC AACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCCAACTTCAAGACCCGCCACAACATCGA AGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTTAC CAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATGGTC CTTCTTGAGTTTGTAACGGCTGCTGGGATTACACATGGCATGGNTGAACTATACAAATAACTCGAGCCTCT AGAACTATAGTGAGTCGTANTACGTANATCCNGACATGATAGATACATTGATGAGTTTGGNNAANNACACT AGAATGCAGTGAAAAAATGNTTNATTTGNGAAATTNGNGATGCTNTGCTTNNNNGNNNNTNNNAGCTGCAN CNGNNCNNNNTNNNANNNNNCNNNNNNNNNNNNNNNNNCNNTNGNTNCNNNNNNNGNNGGGN

The following is the same H1 sequence, however, it is given as the reverse sequence  $5' \rightarrow 3'$ .

NNNNNNNNNNNNGNNANNGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCC TGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAA TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCCAATTGGACTATTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT TGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTG ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA GTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTT GTACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTAC TCATGGTGGNTGTGNNCCGAAAGAGTGNNNCATACNGAACTTATANATTCCCAAAATCCAANACNTTTCACG TTTNATGGTGATTTNCCANANNCNTANCGNCNTGCANNNNNTGCNNNCACTNCCCNNTCCNNNNNAN 

Another 4 colonies were also sequenced and were found to have mismatches (raw sequences are not shown).

# Appendix 9: Raw Data of Sequencing of the shRNA Constructs

- <u>The followings are the raw sequence data of all colonies sequenced for the</u> <u>shRNA constructs targeting the β3α2 fusion point and under the regulation of</u> <u>the U6 promoter.</u>
- 1. The following are sequences from colonies with the pB12mcs-U6-shRNA1<sup> $\beta$ 3\alpha2</sup>- eGFP plasmid
- Colony 1, the forward sequence is given from  $5' \rightarrow 3'$

NNNNNNNNNNNNNNNNNNATCTCGAGAAGCTT<mark>CAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATT</mark> GATATTAGTACAAAATACGTGACGTANAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAATTATGTT **TTAAAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGG** AAAGGACGAAGCACCGTGCTCGGCTCGGCAGCACATATACTAGCAGAGTTCAAAAGCCCTTCAGAAGCTTG TGAAGGGCTTTTGAACTCTGCTTTTTTGGATCCACCACCACCATGAGTAAAGGAGAAGAACTTTTCACTGG AGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGGAGAGGGTG CCAACACTTGTCACTACTCACTTATGGNGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGGCA CGACTTCTTCNAGAGCGCCATGCCTGAGGGATACNTGCAGGAGGACCATCTTCTTCAAGGACGACGGGA ACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAATC GATTTCAAGGANGACGGANACATCCTCGGCCACAAGTTGNAATACAACTACAACTCCCACAACGTATACAT CATGGNNGACNAGCAANGAACGGCATCNAAGCCAACTTCAAGACCCGCCACAACATCGAAGACNGNCGGCG TGCAGCTCGCTNATCATTATCAACAAAATACTCCNAATTGGCGATGANCNCTGTCCTTTTACCAGACNNNN NTNACCTGNCCACACATCTGCCCTTCCNAAGATCCCNANCGANANNNGNGACCGNNTGNTCCNTNCTNGNG TATGNNNNNCTGCNGGNATACNNNTGNANGNNNNNANNNTCCNNNNACTCNNNCCTNNNNAACTATNNG NNNNNAAANNNNNTNNNTNNNNNNNTNGNNNNNNNGNNNTNNNNTNNNNNANCNNNNNTANNNNN NN

- Colony 1, the reverse sequence is given from 3'  $\rightarrow$  5'

NNNNNNNNNNNNNNNNNGGNNNNCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCC GANNNGAAACATAAAATGAATGCAATTGTTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATA AAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAAC TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT

## - Colony 4, the forward sequence is given from 5' $\rightarrow$ 3'

NNNNNNNNNNNNNNGNTATCTCGAGAAGCTT<mark>CAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATTCCT</mark> ATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAATTATGTTTTA AAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTGGCTTTATATATCTTGTGGAAA GAACGAAACACTGTGCTCGCTTCGCAAGAAATTATCCAAGAAAAGTTCAAAAGCCCTTTCAAAACCTGGGAA <mark>AGGGTTTTTAAACTCGGTTTTTTT</mark>GGATCCCCACCCCCAGGAGTAAGGAAAAAAATTTTTCATTGAATT TGCCCCATTTCTGGTTAAATTAAAGGGGGAGGTTAAGGGGAACAAATTTTCGGCCANGGAAAAGGGTGAAG **GNGATGCACCTTACGAAAAATTNACCCTTAAATTTATTTGCACTATGGAAAAACTACCGGTCCNTTGNCCA** ACNNTGGTCATTACTTCNATTTAGGNGGTCNATTGGTTTTCAAAATTTCCGAATANNATGAANCGGNCCAC CTTCTTNNNAANNCCCTTGCNNAAGGAANNGNTGANGAANAGGACNTCCTCCTTCANGACNAANGGAANTT NCNNGANNCTNCCGGAGTCCAGGTTGAAGGNAGANNNCCTCCNNNNNNGAATCAANCTAANGGAANTCAAT TTCAAGGAGNANNNAATCNNCNTCGNCNNNANNTGGNAANANNNTACAACTCCCCCACCGANACTTCNGNN CNNANAGGCAAAGGAANGNNTCNAGNCAACTTNNGNNCCNNCCCCNACATCCAAAACGTCNGNNTGCNNCN NNTNNTCTTNNCANCAATGACTCAANTNNNNATGNCCNNNCNTTTTACNNNNNACNNTNNNTNNCCNNCNN NNCNGCCNNTNNANNNNCCCNNNNAANANNNNNNCNNNNGNTCNNNTTNNTTNNTNNGNCNNCTGNNNT NCNNGNNNNNNNNNNNNNNN

# - Colony 4, the reverse sequence is given from $3' \rightarrow 5'$

- Colony 6, the forward sequence is given from 5'  $\rightarrow$  3'

- Colony 6, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNGCGTTGGGAGCTCTCCCATATGGTCGACCTGCAGGCGGCCGCGAATTCACTAGTGATT ATGGATCCAAAAAACAATAAGGAAGAAGCCCTTCACAAGCTTCTGAAGGGCTTCTTCCTTATTGTAGTATA TGTGCTGCCGAAGCGAGCACGGTGCTTCGTCCTTTCCACAAGATATATAAAGCCAAGAAATCGAAATACTT TCAAGTTACGGTAAGCATATGATAGTCCATTTTAAAACATAATTTTAAAACTGCAAACTACCCAAGAAATT ATTACTTTCTACGTCACGTATTTTGTACTAATATCTTTGTGTTTACAGTCAAATTAAATTCTAATTATATCTCT CTAACAGCCTTGTATCGTATATGCAGATATGAAGGAATCATGGGAAATAGGCCCTCTTCCTGCCCGACCTT GAATTCCAAATCGAATTCCCGCGGCCGCCATGGCGGCCGGGAGCATGCGACGTCGGGCCCAATTCGCCCTA TAGTGGAGTCGTATNAA

- Colony 7, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNGNNNNNNNNNNTNGNNNAGNNNNNCCATATGGTCGACCTGCAGGCGGCCGCNAATTCACTAG TGATTTGGAATT<mark>CAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATTCCTTCATATTTGCATATACGAT ACAAAGCTGTTANAGAGATAATTAGAATTAATTTGACTGTAAACACAAAGATATTAGTACAAAATACGTGA</mark> CGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAATTATGTTTTAAAATGGACTATCATATGCC TACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACCGTGCTCG CTTCGNCAGCACANATACTAGCAGAGTTCAAAAAGCCCTTCAGAAGCTTGTGAAGGGCTTTTGAACTCTGCT TTTTT GGATCCATAATCGAATTCCCGCGGCCGCCATGGCGNCCGGNAGCATGCGACGTCGGGCCCAATTCG CCCTATAGTGAGTCGNATTAANNN

- Colony 7, the reverse sequence is given from  $3' \rightarrow 5'$ 

1. The following are sequences from colonies with the pB12mcs-U6-shRNA2<sup> $\beta$ 3 $\alpha$ 2</sup>eGFP plasmid. The sequencing results were inconclusive.

- Colony 11, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNGNNNGNNATCTCGAGNGCTTCAGGTCGGGCAGGAAGAGGGCCTATGTGTACAGACATT TGTTCCAGGGGGCTTTAAATAGCTGGTGGTGGAACCCAATATTCGTGCTCGGCTCGGCAGCACATATACTAG CAGAGTTCAAAAGCCCTTCAGGAAGCTTGCTGAAGGGCTTTTGAACTCTGTTTTTTGGATCCACAGCCACC ATGAGTAAAGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGG GCACTACTGGAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTT TCAAGATATCCAGATCATATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGA GAGGACCATCTTCTTCAAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCC TCGTCAACAGGATCGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAA TACAACTACAACTCCCACAACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAA GACCCGCCACAACATCGAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCG ATGACCCTGTCCTTTTACCAGACAACCATTACCTGTCCACAATCTGCCCTTTCNAAAGATCCCAACGAA AAGAGAGACCACATGGTCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACACATGGCATGNNTGANCTATA CAAATAACTCGAGCCTCTAGAACTATAGTGAGTCGTATTACGTANATCCAGACNTGATAAGATACATTGAT GANNTTGGNCAANNANANTANAATGCAGTGAANNAAAATGCTTTANTTGNGANNNTTNNGATGCTANTGNT TTANTNGNNNNNTANNAGCTGCAATANANCNAGNTNNNANNNATGCATTCNNTTATGNTTCANNNCAGN NANGNNANTNNNNNNTTNNNNNANNNNN

## - Colony 11, the reverse sequence is given from $3' \rightarrow 5'$

NNNNNNNNNNNNNNGNAANNGGNNATCTTNNTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCT GAANNNNNNNNAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGNAGCTTATAATGGTTACAAAT AAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCAAA CTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTAT TTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCT CTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGT CATCGCCNATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGG GTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTT GTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTTAAGCTCGATCCTGTTGA CGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTC CTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGNATATCT TGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGNACNGGTAGTTTTCCAGTAG TGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCCTTCACCCTCTCCACTGACAGAAATTTGT ACCCATTAACATCACCATCTAATTCAACAAGAANTNNNNANTNNNNGAAAAGTNNTTCNNCTTTACTCATG **GTGGNTGTGNNCAAAAAAACAGANTTNNANNCNNTNNNANCTNCNNANGGNTTTTGAANTCTGTANNNNAT** GNGCTGNCNAAGNNNNNCNANNNNGGNNNCNCNCANNNANTNANNCCNGANNNANGNCTGTANNNNNNNCN NNNCNGNNCNNNNNNNN

## - Colony 12, the forward sequence is given from 5' $\rightarrow$ 3'

NNNNNNNNNNNNNNNNNNGNTATCTCGAGNGCTTCAAGGTCGGGCAGGAAGAGGGCCTATGTGTACAGAC ATTTNNTNNNGGGCTTTAAATAGCTGGTGGTGGAACCCAATATTCGTGCTCGGCTTCGGCAGCACATATAC TAGCAGAGTTCAAAAAGCCCTTCAGGAAGCTTGCTGAAGGGCTTTTGAACTCTGTTTTTTGGATCCACAGCC ACCATGAGTAAAGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAA TGGGTACAAATTTTCTGTCAGTGGAGAGGGGTGAAGGTGATGCAACATACGGAAAACTTACCCTTAAATTTA TTTGCACTACTGGAAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTCACTTATGGTGTTCAATGC TTTTCAAGATATCCAGATCATATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCA GGAGAGGACCATCTTCTTCAAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACA CCCTCGTCAACAGGATCGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTG GAATACAACTACAACTCCCACAACGTATACATCATGGCCGACAAGCAAAGAACGGCATCAAAGCCAACTT CAAGACCCGCCACAACATCGAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTG GCGATGACCCTGTCCTTTTACCAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCAAC GAAAAGAGAGACCACATGGTCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACACATGGCATGGATGAACT ATACAAATAACTCNAGCCTCTAGAACTATANTGAGTCGTATTACGTANATCCAGACATGATAAGATACNTT GATGANTTTGGACAAACNCACTNNNANGCAGTGNAANAAAATGCCTTNNTTTGNNAANNNTNGNNGATGCT NTNNNTTNANNNGNANCATNNAAGCNNCANTNAAANNNANNTTNNNANNANNNNGNANTNCNNTTNNNNNT 

## - Colony 12, the reverse sequence is given from $3' \rightarrow 5'$

**CCTGANNNNNNNNNATGAATGCAATTGTTGTTGTTAACTTGTTTNNNNCAGCTNATAATGGTTACAAA** TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCNAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT **TGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTG** ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCNTGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGNCATGGAACAGGTAGTTTTCCAGTA **GTGCAAATAAATTTAAGGGGTAAGTTTTCCGTATGTTGCATCNACCTTCACCCTCTCCACTGACAGAAAAT** TTGTACCCATTAACATCACCATCTAATTCAACAAGAAATTGGGACAACTCCNGTGAAAAGTTCTTCTCCNT TNACTCATGNNNNCTGTGNATCNNAAAAAACAGAGNTCAAANGCCCCTTCNNNNAAGNNTTCCTGNNNGGN 

- Colony 15, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNATCTCGAGNGCTTCAAGGTCGGCAGGAAGAGGGCCTATGTGTACAGACATTTGT TCCAGGGGCTTTAAATAGCTGGTGGTGGAACTCAATATTCGTGCTCGCTTCGGTAGCACATATACTACAGA GTTCAAAAGCCCTTCAGGAAGCTTGCTGAAGGGCTTTTGAACTCTGTTTTTGTATCCACAGCCACCATGA TTAAAGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTAC AAATTTTCTGTCAGTGGAGAGGNTGAANNTGATGCAACATACGGAAAACTTACCCTTAAATTTATTTGCAC TACTGGAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTCACTTATGGAGTTCAATGCTGTGCAA GATATCCNGATCATATGAANCGGCNGNCTTCTTCANNAGCGCCATGCCTGAGGGATACGTGCGNGAAAGGA CCATCTTCTTCAAGGACGACNGGAACTACAAGACACGTGCTGNGNCAAGTTTGAGGGANACTCCCTCGTCA ACCGGAACGAGCTTAAGGNAATCNATTTCAATGAAGACCGAANCGTCCTCGACCACGANTTGGAATACNNC TACNNCTCCCACANCNNANACNTCNNGNNNNACNNNNNAGNAACGGTCTCACNNCCNACTTCNAGACCCN **CNNNNNCATCCAANACNGCNNNNTGAAANTNGCTGATCATTGTCCACAAAATACTCGAATTGGNNNTGACC** CTGANNTTTTACCANANCACCATAACCGGACCACACGANCTGCNCTTTCNAANNANCCCAGACGAAAAGNN ANNNNAGNTGGTCNTTCNNTGAGNTNGTNACCNCTGCTGNCATNNCACNTGNCGNGGANGAACTATANAAG TANCTCNAGNCNCTANAACNATNCTGAGTGGTAATNCGNANATNCNNNACATGCANNAGATACATTGATGA GNNTNGGNNAANNCNNCNNCTNNAAANNNNNNNNNNNCAANTGCTNNANTTTGGGCAGNNNTNNNNNAN NNNANTAGNNTTNNNNNNNNNNCCNNNNNNANNNNTNNCCATNNANGNNCNNNNNANNNNCNGNTNNGNNN 

- Colony 15, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNNGGGATAATCTTAAGTCGCGNNNGCGAATTNNNNCCTCCNANACCTCCCCC **TGANNNNNNAGAAAATGAATGCAAGTGGTGGTGTTAACTTGTTTNTTGCAGCTTATAATGGTTACAAAT** AAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCAAA CTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTAT TTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCT CTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGT CATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGG GTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTT GTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGA CGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTC CTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCT TGAAAAGCATTGAACACCATAAGTGAGAGAGTGGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAG TGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCCTTCACCCTCTCCACTGACAGAAAATTTG TACCCATTAACATCACCATCTAATTCAACAANAATTGGGACAACTCCNGTGAAAAGTTCTTCTCCTTTACT CATGGNGGCTGTGGNTCCAAAAAAAAAAAGTTCAAAAGCCCTTCAGCAAGCTTCNTGAANGGNNTTTTGAA NNGNNNNNNNNNNNNNNNNNNNNNNNN

2. The following are sequences from colonies with the pB12mcs-U6-shRNA3<sup> $\beta$ 3\alpha2</sup>-eGFP plasmid.

- Colony 3, the forward sequence is given from 5'  $\rightarrow$  3' (shRNA sequence was inconclusive)

- Colony 3, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNAGGNNAATCTTAAGTCGCGGCCGCGAATTAAAAAAACCTCCCACACCTCCCCCT GAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAAT AAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGGTTTGTCCAAA CTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTAT TTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCT CTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGT CATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGG GTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTT **GTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGA** CGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTC CTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCT **TGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAG** TGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCCTTCACCCTCTCCACTGACAGAAAATTTG TACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACT CATGGNGGCTGTGGATCNAAAAAGCANNNAGTTCAAAAGCCCNTTCAGCANGCTTNNTNNNGGCTTTGAAN CNNNNNNNNGNNANNAANNNNNTTNNNNNNNNN

#### - Colony 6, the forward sequence is given from 5' $\rightarrow$ 3'

NNNNNNNNNNNNNNNNTATCTCGAGANGCTTCAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATT GATATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAATTATGTT **TTAAAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGG** <mark>AAAGGACGAAGCACCGTGCTCGTTTCGGCAGCACATATACTA</mark>GCAGAGTTCAAAAGCCCTTCAAGAACCTT GCTGAAGGGCTTTTGAACTCTGGATTTNTGGATCCCCCTTTTCCNTGAGATCTTTCGAAAAGCTTTACTGT **GTGNACGGCCCATGTCTGGCTGGNNNAGGACAGGGTCATCGCNAATTGGATNTNTTGTTGATAATGATCAG** CGGCCATGATGTANNTCTTGCGTGAGGTGTAGTTGGATTCCNACTAGTGGCCGATCATGTTACCGGCCTCC NTGACTTCGATACCNTCANGCTCGATCCAGTNGANNAGGNNGNCACCNTCNNNCTTGACNTCNACACGNGN NNTGNAATTCCCGCTGANNTTGANNANGATGNANNNCNCCNGNTCGTATCCNTCAAGCNTGGNGCTNNNGA NNAANTCNNGACGCNNNNNATNANCTNNATATCTTGAAAAGCATTGAACACTCNAANTGAGANTAGTGANA NNNGNNNNNCATGNNNAACGGNATNTNTCCNGANTTGNAATANATNNNNAGNGTAAANTATCNNNATGTGC NANNCNCTNNNCCCTCTCCACCTGACAGACNANTTGNCNNNNTANCAGTCCNCTTTANNATTCANCNNNAA **TTGGNACNAATTCNNTGACNATTTCTNCNANCTTNACTCAAGGNGGANNGNGNNTCCANNNANNTNNANTN** NNAANNCCNTNNNNAAGTCTTCNNTGNANGNNATTNANNNNTTCNGCTNANNNNNTCNTNNNGAAANNNN

- Colony 6, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNGANNNGGNNNTCTTAAGTCGCGGCCGCGAATTAAAAAAACCTCCCACACCTCCCCCT GANNNNNNNANAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGNAGCTTATAATGGTTACAAATA TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT TGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCTCTC TTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTC ATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGG **TCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTG** TATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGAC GAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCC TCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTT GAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGT GCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGT **ACCCATTAACATCACCATCTAATTCAACAAGAATTGGGANNACTCCNGTGAAAAGTTCTTCTCCTTTACTC** ATGGNGGCTGTGGATCCAAAAAAGCAGAGTTCAAAANNCNTTCANCAAGCTTNNGAAGGGCTTTTNNACTC TGCTANNATATGTGCNNCCNAAACGNNNNNNGGNGNNTNCNTCTTNNNCNAGAATATNNNAAGNCNAANAA NTATNNNNNCNNCNNGNANTTNNNNNAAAAANNNNNN

- Colony 9, the forward sequence is given from 5'  $\rightarrow$  3'

(shRNA sequence was inconclusive)

- Colony 9, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNGNNANNGGNTANTCTTAAGTCGCCGCCGCGAATTAAAAAACCTCCCACACCTCCCCCT CTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTAT TTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCT CTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGT CATCGCCAATTGGAGTATTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGG GTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTT **GTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGA** CGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTC CTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCT TGAAAAGCATTGAACACCATAAGTGAGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAG TGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTG TACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCNGTGAAAAGTTCTTCTCCTTTACT CATGGTGGCTGTGGATCCAAAAAGCANNANNTTCAAAAAGCCCTTCAGCAAGCTTNCNGAANGGCTTTTGA 

- <u>The followings are the raw sequence data of all colonies sequenced for the</u> <u>shRNA constructs targeting the β2α2 fusion point and under the regulation of</u> <u>the U6 promoter.</u>
- 1. The following are sequences from colonies with the pB12mcs-U6-shRNA1  $^{\beta 2\alpha 2}$  eGFP plasmid

- Colony 7, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNATGCNCNGGCCGCCATGGCGGCCGCGGGAATTCGATTTGGAATT<mark>CAAGGTCGGGCA GGAAGAGGGCCTATTTCCCATGATTCCTTCATATTTGCATATACGATACAAGGCTGTTAGAGAGATAATTA GAATTAATTTGACTGTAAACACAAAGATATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGG</mark> TAGTTTGCAGTTTTAAAATTATGTTTTAAAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGA TTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAGCACCGTGCTCGCTTCGGCAGCACATATACTAGCAA TAAGGAAGAAGCCCTTCAGAAGCTTGTGAAGGGCTTCTTCCTTATTGTTTTTT GGATCCATAATCACTAGT GAATTCGCGGCCGCCTGCAGGTCGACCATATGGGAGGAGCTCCCAACGCGTTGGATGCATAGCTTGAGTATT CTATANNGNNCCTAANN

- Colony 7, the reverse sequence is given from  $3' \rightarrow 5'$ 

- Colony 10, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNATGNNNCGGCCGCCATGGCGCGCGCGGGAATTCGATTTGGAATT<mark>CAAGGTCGGGCAG</mark> GAAGANNNNCTATTTCCCATGATTCCTTCATATTTGCATATACGATACAAGGCTGTTAGAGAGATAATTAG AATTAATTTGACTGTAAACACAAAGATATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGT AGTTTGCAGTTTTAAAATTATGTTTTAAAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGAT TTCTTGGCTTTATATATCTTGTGGGAAAGAACGAAACACTGTGCCTCGCTTCGGCAGCACATATACTAGCAAT AAGGAAGAAGCCCTTCAGAAGCTTGTGAAGGGCTTCTTCCTTATTGTTTTTT GGATCCATAAACCAAAGCTTGTGAAGGGCTTCTTCCTTATTGTTTTTTG GATCCATGACCTTGCAGGTCGACCATATGGGAGAGCTCCCAACGCGTTGGATGCATAGCTTGAGTATTC TTCTTGCCGCCGCCTGCAGGTCGACCATATGGGAGAGCTCCCAACGCGTTGGATGCATAGCTTGAGTATTC TATANNGNNNCCTANNN

- Colony 10, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNANGCGTTGGGAGCTCTCCCATATGGTCGACCTGCAGGCGGCCGCGAATTCACTAGTGATT ATGGATCCAAAAAACAATAAGGAAGAAGCCCTTCACAAGCTTCTGAAGGGCTTCTTCCTTATTGTAGTATA TGTGCTGCCGAAGCGAGCACAGTGTTTCGTTCTTTCCACAAGATATATAAAGCCAAGAAATCGAAATACTT TCAAGTTACGGTAAGCATATGATAGTCCATTTTAAAACATAATTTTAAAACTGCAAACTACCCAAGAAATT ATTACTTTCTACGTCACGTATTTTGTACTAATATCTTTGTGTTTACAGTCAAATTAAATTCTAATTATATCTCT CTAACAGCCTTGTATCGTATATGCAAATATGAAGGAATCATGGGAAATAGGCCCTCTTCCTGCCCGACCTT GAATTCCAAATCGAATTCCGCGCGCCGCCATGGCGGCCGGGAGCATGCGACGTCGGGCCCAATTCGCCCTA TAGTGAGNCGTATNAN - Colony U61A, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNCNNATGCTNNGGCCGCCATGGCGGCGCGCGGGAATTCGATTTGGAATT<mark>CAAGGTCGGGCAGG</mark> AAGAGGGCCTATTTCCCATGATTCCTTCATATCTGCATATACGATACAAGGCTGTTAGAGAGATAATTAGA ATTAATTTGACTGTAAACACAAAGATATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTA GTTTGCAGTTTTAAAATTATGTTTTAAAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATT TCTTGGCTTTATATATCTTGTGGAAAGGACGAAGCACCGTGCTCGGCTCGGCAGCACATATACTAGCAATA AGGAAGAAGCCCTTCAGAAGCTTGTGAAGGGCTTCTTCCTTATTGTTTTTT GGATCCATAATCACTAGTGTGAAGGGCTTCTTCCCAACGCGTTGGATGCATAGCTTGAGTATTCT ATTCGCGGCCGCCTGCAGGTCGACCATATGGGAGGGCTCCCCAACGCGTTGGATGCATAGCTTGAGTATTCT ATANTGNNNCCTAANN

#### - Colony U61A, the reverse sequence is given from $3' \rightarrow 5'$

- Colony 3, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNATCTCGANAGCTT<mark>CAAGGTCGGGCAGGAAGAGGGCCTATTTCCCATGATTCCT</mark> **ATTAGTACAAAATACGTGACGTAGAAAGTAATAATTTCTTGGGTAGTTTGCAGTTTTAAAATTATGTTTTA** AAATGGACTATCATATGCTTACCGTAACTTGAAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAA GAACGAAACACTGTGCTCGCCTCGGCAGCACATATACTAGAATAAGGAAGAAGCCCTTCACGAAGCTTGGT <mark>GAAGGGCTTCCTTACTTTTTTTTTT</mark>GGATCCACAGCCACCATGAGTAAAGGAGAAGAACTTTTCACTGGAG **NGTGATGCAACATACGGAAAACTTACCCTTAAATTTATTTGCACTACTGGAAAACTACCTGTTCCATGGNC** AACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGGCACG ACTTCTTCAAGAGCGCCATGCCTGAGGGGATACNTGCAGGAGGAGCATCTTCTTCAAGGACGACGGGGAAC TACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAATCGA TTTCAAGGAGGACGGANACATCCTCAGCCACAAGTTGNNNNACANCTACAACTCCCCACAACGNATACATCA TGNNGACTAGCAAAAGAACGGCGTCAAAGTCAACTTCAANANCCNNCACNACATCGAAGACNGCGGCGTGC AACTCGCTGATCATTATCANCAAATACNNNATTGGNNATGANNCTGNCCTTTTACCANACACNNNTNACNT AANNNNNNTTTNATNNNGNNNNNNN

- Colony 3, the reverse sequence is given from  $3' \rightarrow 5'$ 

- <u>The followings are the raw sequence data of all colonies sequenced for the</u> <u>shRNA constructs targeting the β2α2 fusion point and under the regulation of</u> <u>the H1 promoter.</u>
- 1. The following are sequences from colonies with the pB12mcs-H1-shRNA1<sup> $\beta$ 2\alpha2</sup>eGFP plasmid. All sequences were missing the shRNA sequence.
  - Colony 3, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNNNNNNNTCTCGANNGCTTGAACGCTGACGTCATCAACCCGGCTCCAAGGAATCGCGGM NNNANNGNNNCTAGGCGGGAACACCCAGCGCGCGCGCGCGCCCTGNNAGGAAGATGGCTGTGAGGGGACAGGGG AGTGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGGAAATCACCGTAAACGTGAAATGTCTTTGG ATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTT CGGATCCACAGCCACACATGTCTGTGAATGAGACCACCCTTTACAGGGTACAAATTTCTGTC AGTGGAGAGGGTGAAGGTGATGCAACATACGGAAAACTTAGATGGTGTACAATGTCTTTCAAGGATAAAGGAGAAACT ACCTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATC ATATGAAGCGGCACGACTTCTTCAAGAGCGCCCATGCCTGAGGGATACGTGCAGGAGAGGACCACCTTCTTCT AAGGACGGCACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAAGGGAGACACCCTCGTCAACAGGATCGA GCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTACAACTCC ACAACGTATACATCATGGCCGACAAGCAAAGAACAACATCCTCGGCCACAAGTTGGAATACAACTACAACTAC GAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTT ACCAGACAACCATTACCATGCCCACACACTGCCTGTCAACAGGACCACCTGTCTTTT ACCAGACAACCATTACCACGCTGCTGGGATTACACAAGACTCCAACAGAAACAACAACTACAACAAGG TCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACACAAGGCAATGGNTGAACTATACAAAAAAACTCCGAGCCT

Colony 3, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNAGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCCACACCTCCCCCCT GAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAAT AAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAA CTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTAT TTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCT CTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGT CATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGG GTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTAGTGTGGGGAGTTGTAGTT GTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTTAAGCTCGATCCTGTTGA CGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTC CTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCT TGAAAAGCATTGAACACCATAAGTGAGAGAGTGGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAG TGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTG TACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACT CATGGNGGCTGTGGATCCGAAAGANTGGNNTCATACAGAACTTANANANNCCCCAAATNCNAAAGACNTTTN ACGTTACGGNGATTTNCCAGAACNCNTANCGACATGCAAATNTGNNGGNNCCNNNTNCCNNNNCCNNNNAN NACNNCNNNNTNCANNNNNNN

- Colony 4, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNGGNNATCTCGA<mark>GAAGCTTGAACGCTGACGTCAACCCGCTCCAAGGAATCGC</mark> GGGAGTGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTT TGGATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCCGGATCCACCACCATGAGTAAAGGAG **Δ**ΔGΔΔCͲͲͲͲϹΔCͲGGΔGͲͲGͲCCCCΔΔͲͲCͲͲGΔΤͲGΔΔͲͲΔGΔͲGGͲGΔͲGCͲΔΔΔΠGGGTΔCΔΔΔͲͲͲͲCͲ ACTACCTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAG <u>ΑΨĊΑΨΑΨĠΑΑĠĊĠĠĊAĊĠAĊŦΨĊŢΨĊAAĠAĠĊĠĊĊAΨĠĊĊĊŦĠAĠĠĠAŦĂĊĠŦĠĊĊAĠĠAĠAĠĊĊAŦĊŦŦĊ</u> TTCAAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGAT CGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACT CCCACAACGTATACATCATGGCCGACAAGCAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAAC ATCGAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCT TGGNCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACACATGGCATGGNTGAACTATACAAATAACTCGAG **CCTCTAGAACTATANNGAGTCGTANTACGTANATCCNGACATGANNNNATACATTGATGAGTTTNGNNNN** CCNCNNNANNNGCNNNGAAANAAATGCTTTNNNNNNAANNNNTGCNNNNNTTTNNNNNNNACNTNNAN

- Colony 4, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNAGGNTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCTG AACCTGAAACATAAAATGAATGCAATTGTTGTTGTTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATA AAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAAC TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT **TGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCTC** TTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTC ATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGG TCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTG TATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGAC GAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCC TCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTT GAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGT GCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGT **ACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTC** ATGGNNGCTGTGGNTCCGAAAGANNNGNCTCATACAGAACTTATANATTCCCAAATCCNAAGACATTTNNC NNNNNNNAANNN

- Colony 5, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNGGNNATCTCGAGANGCTTGAACGCTGACGTCATCAACCCGCTCCAAGGAATCGCGGG **CCCAGTGTCACTAGGCGGGAACACCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTGAGGGACAGGGG** AGTGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTTTGG ATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACAGCCACCATGAGTAAAGGAGAAG AACTTTTCACTGGAGTTGTCCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTC ACCTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATC ATATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGGATACGTGCAGGAGAGGACCATCTTCTTC AAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGA GCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCC ACAACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATC GAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTT ACCAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCCAACGAAAAGAGAGAACAACATGG TCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACACATGGCATGNTGAACTATACANTAACTCGAGCCTCT AGAACTATAGTGAGTCGTATTACGTAGATCCAGANATGANANNTACATTNGATGANTNNGNNNNNACANT ANNNGCANNNAAAAAAATGNNNNATNNNAAANNNNNTGCTNTGNTTNNTTNNACNTNNANCTGCANNAA NNNTANNNATNNN

- Colony 5, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNACNGGNNNNCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCTG AACCTGAAAACATAAAATGAATGCAATTGTTGTTGTTGATGTTTATTGCAGCTTATAATGGTTACAAATA TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT TGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCTCTC TTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTC ATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGG TCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTG TATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGAC GAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCC TCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTT GAAAAGCATTGAACACCATTAAGTGAGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGT GCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGT ACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTC ATGGNGGCTGNNNNCCGAAAGNNNGGNCTCATACAGAACTTATANANTCCCNAANNCNAAAGACATTTCAC **GTTACGNNGATTTNCCANANNNATANCGACATGCAANTATGCNNNNCCNNNTCCCNNNTCCNNNNNNNNTN** NNNNCANNNNTTNNNNN

- Colony 6, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNGNNNNGGNNATCTCGANNGCTTGAACGCTGACGTCATCAACCCGCTCCAAGGAATCGCGG GCCCAGTGTCACTAGGCGGGAACACCCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTGAGGGACAGGG GAGTGGCCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTTTG GATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACCACCATGAGTAAAGGAGAA GAACTTTTTCACTGGAGTTGTCCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGT TACCTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGAT CATATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTTCTT CAAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCG AGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCC CACAACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACAT CGAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTT TACCAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATG **TCTAGAACTATANNGAGTCGTANTACGTAGATCCNGACNTGATANNNNCNTNGATGANTTTNGNNNNNNN** NNAGANGCNNNAAAAAAATGNTTNANTNNNAAATTNNNNTGCNNTGNTTTNNNTNGNANNTNNNNCTGCAN TANNNNNNN

- Colony 6, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNNNNGGGNNNCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCTG AACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATA AAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAAC TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT TGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCTC TTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTC <u>ΑΨĊĠĊĊ</u>ΑΑΨΨĠĠAĞΨΑΨΨΨΨĠΨΨĠAΨŎAΨĠAΨĊAĠĊĠAĠΨΨĠĊAĊĠĊĊĠĊĊĠŦĊΨΨĊĠAΨĠŦΨĠŦĠĠĊĠĠĠ TCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTG TATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGAC GAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCC TCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTT GAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGT **GCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGT** ACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTC **ATGGNGGNTGTGGATCCGAAAGAGTNNNTCATACAGAACTTATANANTCCCAAANNCNAANNANATTTNAC GTTACGNNGATTTCCCAGAACACNTANCGACATGCAANNNTGNNNNNNCCNNNNCCTNNNNCCTCNNANNNT** GNNNNNTCAANANNNNNN

- Colony 7, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNGGNNATCTCGANNGCTT<mark>GANGCTGACGTCATCAACCCGCTCCAAGGAATCGCGGGCC</mark> CAGTGTCACTAGGCGGGAACACCCAGCGCGCGCGCGCCCTGGCAGGAAGATGGCTGTGAGGGACAGGGGAG TGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTTTGGAT TTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACAGCCACCATGAGTAAAGGAGAAGAA CTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAG CTGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCAT ATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGGACCATCTTCTTCAA GGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGC TTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCAC AACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCGA AGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTTAC CAGACAACCATTACCTGTCCACACAATCTGCCCCTTTCGAAAGATCCCCAACGAAAAGAGAGACCACATGGNC NNNCANNNNAAAANANGNNTTNATTNNGAAATNGNGNTGCNNTGCTTNATTNNNNNNTNNNANCTGCAATA NNNNNNNNNN

- Colony 7, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNGGNNACNGGNTATCTTAAGTCGCGCCGCGAATTAAAAAAACCTCCCACACCTCCCCC **TGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAA** TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT TGTATTCCAACTTGTGGCCCGAGGATGTTTCCCGTCCTTCGAAATCGATTCCCTTAAGCTCCGATCCTGTTG ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA **GTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTT GTACCCATTAACATCACCATCTAATTCAACAAGAATTGGGANANTCCAGTGAAAAGTTCTTCTCCTTTACT** CATGGNNGCTGTGGNTCCGAAAGNANTGGNCTCATACAGAACTTANANANTCCNNAATCCNNGACATTTCA CGNNNCGGNGATTNCCANANNNNNNNNANNGACNTGCAANNNTGCNNNNGNCNNNTCCNNNNCCNNNNNNTN NNTNANNNNNNNNNN

- Colony 8, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNNNNNNNCTCGAGNAGCTT<mark>GAACGCTGACGTCNTCNACACCATAAGTTCTGTA</mark> TGAGACCACTCTTTCCAATAAGGAAGAAGCCCTTCAGAAGCTTGAACGCTGACGTCATCAACCCGCTCCAA GGAATCGCGGGCCCAGTGTCACTAGGCGGGAACACCCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTG AGGGACAGGGGGAGTGGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGA **AATGTCTTT**GGATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACAGCCACCATGAG TAAAGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACA **ACTGGAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTTTCAAG** CCATCTTCTTCAAGGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTC AACAGGATCGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAA CTACNACTCCCACAACGTATACATCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCC GCCACAACATCGAAGACGGCGGCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGAC CCTGTCCTTTTACCAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCCNACGAAAAGAG AGACCACATGGNNCTTNNGAGTTTGNAACGGCNGCTGNNANNCNCATGNNTGNNNANNATACNAANNANTC NNTANNNGCNNNANNANGNNTNNTNNNNNNNNNGNNTGNNNATNNNNCNTNANCNNNNNNAANNANNN

- Colony 8, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNNNANNGGGTANTCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCC TGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAA TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA **ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA** TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT **TGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTG** ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA **GTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTT** GTACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTAC TCATGGNGGCTGTGGATCCGAAAGANNNGNCTCATACAGAACTNNNAGANTCCCAAATCCNAAGACATTTN NCGTTTANGNNATTTCCANANNNNNAGCNACNTGCAANTNTGCNGGNNCNCTCCCNNTCCTCNNANNNTN NNNNCAGNNNNNCANNNNN

The following sequences were inconclusive:

- Colony 9, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNGNNTAGGCGCGCCATGAGCTCGGGGCTCGAGTTATTCCAGTGCCCANAATTCTTCTCA TCCNTGGNTTTATTATCTTTGCCTTTCAGGGCATCATCATTAAACATAACTTCAAAAATTTTCTTCTTGTC CTGGCCTCATGGGCCTTCCGCTCACTGCCCGCTTTCCAGTCGGGAAACCTGTCGTGCCAGCTGCATTAACA TGGTCATAGCTGTTTCCTTGCGTATTGGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTCG TTCGGGTAAAGCCTGGGGTGCCTAATGAGCAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGT TGCTGGCGTTTTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTGG CGAAACCCGACAGGACTATAAAGATACCAGGCGTTTCCCCCTGGAAGCTCCCTCGTGCGCTCTCCTGTTCC GACCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCCCCTTCGGGAAGCGTGGCGCTTTCTCATAGCTCAC GCTGTAGGTATCTCAGTTCGGTGTAGGTCGTTCGCTCCAAGCTGGGCTGTGTGCACGAACCCCCCGTTCAG CCCGACCGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTTATCGCCACT GGCAGCAGCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTCTTGAAGTGGT GGCCTAACTACNGCTACACTAGAAGAACAGTATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGGA AAAAGAGTTGGTAGCTCTTGATCCGGCAAACAAACCACCGCTGGTAGCGGTGGTTTTTTTGTTGCAAGCA GCANATTACGCGCAGAAAAAAAGGATCTCAAGAAGATCCTTTGATCTTTCTACGGGGTCTGACGCTCAGNG TAAAATGANNTTNATCNNTCTANNNTATATANGANTAANNNNNTGACNNTNNNNGCTNNTCNNNNGCAC NNNNNCAGCNATNNNNNATTCGNTCATCNTANNNNCNGNNNNCCCNNNCNGNNNANACTACGANNNGGNNN

# - Colony 9, the reverse sequence is given from $3' \rightarrow 5'$

NNNNNNNNNNNNNNGGNTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCTG AACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATA TCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATT TGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCTCTC TTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTC ATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGG TCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTG TATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGAC GAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCC TCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTT GAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGT **GCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGT** ACCCATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTC ATGGNGGCTGTGGATCCGAAAGAGTGGTCTCATACAGAACTTATAAGATTCCCAAATCCAAAGACATTTCA CGTTTANNGTGATTTCCCAGAACACATAGCGACATGCAATATTGCAGGGCGCCACTCCCTGTCCNTCACAG CCATCTTCCTGCCNGGNGCANGCGCGCTGGGNNNTCCCCGNCTANTGACACTGGNNNCGCGATNNNGNANCG **GNNNATGACGTCANNNNNAGCTNNNNANATACCNNNNATCCNANNACCNNNNNCCNANNGNNNNNCNNNN** ACNCNNNCNNNNNNNNNCNTTTNNNANNNNNNCNNNNACCNNNNNNNCNNNNNANCNCNNNNANNN 

# - Colony 10, the forward sequence is given from 5' $\rightarrow$ 3'

NNNNNNNNNNNGNNTAGGCGCGCCATGAGCTCGGGGCTCGAGTTATTCCAGTGCCCANAATTCTTCTTCA TCCNTGGNTTTATTATCTTTGCCTTTCAGGGCATCATCATTAAACATAACTTCAAAAATTTTCTTCTTGTC CTGGCCTCATGGGCCTTCCGCTCACTGCCCGCTTTCCAGTCGGGAAACCTGTCGTGCCAGCTGCATTAACA **TGGTCATAGCTGTTTCCTTGCGTATTGGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTCG** TTCGGGTAAAGCCTGGGGTGCCTAATGAGCAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGT TGCTGGCGTTTTTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAATCGACGCTCAAGTCAGAGGTGG CGAAACCCGACAGGACTATAAAGATACCAGGCGTTTTCCCCCCTGGAAGCTCCCTCGTGCGCTCCTCCTGTTCC GACCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCCCTTCGGGAAGCGTGGCGCTTTCTCATAGCTCAC GCTGTAGGTATCTCAGTTCGGTGTAGGTCGTTCGCTCCAAGCTGGGCTGTGTGCACGAACCCCCCGTTCAG CCCGACCGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTTATCGCCACT GGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTCTTGAAGTGGT GGCCTAACTACNGCTACACTAGAAGAACAGTATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGGA GCANATTACGCGCAGAAAAAAAGGATCTCAAGAAGATCCTTTGATCTTTCTACGGGGTCTGACGCTCAGNG GANCGAAACTCACGTTNNGGGATTTTGGNCATGAGATTATCAAAANGATCTTCAC

- Colony 10, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNNTCTANATNCNANCTGCNGGCATGCGNNCTTGGCNTAATCNGGCCCATAGCTGT TTCCTGTGNNAGATTGTTATCCGCTCANATTCCACACAACATACGAGCCGGAATCNTAAAGTGGAAAGCCN GGAGTGCCTAATGAATGACTAACTCACATTAATTGCGTTGCGCTCACTGCCCGCTTTCCAGTCNGCAAACC TGTCGNGCCANCTGCATTAATGAATCNGCCAAGGCGGGGGAGAGGCGGTTTGAGTATTGNGCNCTCATCC GCTTCCTCNCTNACTGACTCNCTGCGCTCGNGCGTTGNGCTGCGNAGAGCGGTATCGGGTCACTCANAGGC AGNAATACGGTTATCNAGAGAATCAGGGGATAACGCNGGAAANAACATGTGAACAAAAGGCCGNCNAAAGG CCANGACCCGTAAAAAAGGCCGCGTTGCTGGCAGTTTGCCATAGGCTCCNNCCCTGTTACNAGCATCANAGA AATCGACGTTNNAGTCNAGGTGGTGAAAGGGATTGGACTATAANNATACTAGGGGTGTCNCCTGGAAGCAC TCTCGCGCGCTCACCTGTTCCCACTCT

- Colony 11, the forward sequence is given from  $5' \rightarrow 3'$ 

NNNNNNNNNNNGNTATCTCGANNGCTTGNNNGCTGACGTCATCAACCCGNTCCAAGGAATCGCGGGCCC AGTGTCACTAGGCGGGAACACCCCAGCGCGCGCGCGCCCCTGGCAGGAAGATGGCTGTGAGGGACAGGGGAGT GGCGCCCTGCAATATTTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTTTGGATT TGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCGGATCCACAGCCGCCATGAGTAAAGGAGAAGAAC TTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGT TGTTCCATGGCCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAAAATCATA TGAAGCGGCACGACTTCTTCAAGAGCGCCNTGCCTGANGGATACGTGCAGGAGAGGACCATCTTCTTCAAG GACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCT TAANGGAATCGATTTCAAGGAGGACGGANACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCACA ACGTATACNTCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCNAA GACGGCGGCGTGCAACTCGCTGATCATTATCNACANNATACTCCAATTGGCGATGACCCTGTCCTTTTACC AGACAACCATTACCTGTCCACAACATCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATGGTCC **TTCTTGAGTTTGTAACNNNTGCTGGGATTACACATGNNNTGNNTGAACTATACANATAACTCGAGCCTCTA** GAACTATAGTGAGTCGNATTACGTANATCCAGANNNGANANANACATTGATGAGTTTGGACNNNCNCAACT AGAATGCANNGAANAAAATGCTTTATTTGNGAAATTTGTGANGCTANTGCTCTATTTGTANNNNTTANANC TGCAANNANNCANNTTAANANNNNANTGCNTTCNTTTNANNNTTNNAGGTNNGNGGNNNNGNGGNNNNTTT NNNNNNNNNNNCNNGNNNGNNNCTANNGN

- Colony 11, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNGNNANNGGNTATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCCTGAACCT **GNNNCNNNAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCANCTTATAATGGTTACAAATAAAGCA** ATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAAACTCATC <u>Ϸ</u> ϷͲϾͲϷͲϹͲͳϷͲϹͲϹͲϾϾϷͲϹͲϷϹϾͲϷϷͲϷϹϾϷϹͲϹϷϹͲϷͲϷϾͲϷϹͲϷϾͳϷϾ AGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTCTCTTTTC GTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTCATCGC CAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGGGTCTTG AAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTGTATTC CAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAAATCGATTCCCTTAAGCTCGATCCTGTTGACGAGGG TGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCCTCTCC TGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTTGAAAA GCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGTGCAAA TAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGTACCCA TTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTCATGGT GGCTGTGGATCCGAAAGAGTGGTCTCATACAGAACTTATAAGATTCCCAAAATCCAAAGACATTTCACGTTT ACGGTGATTTCCCAGAACACATAGCGACATGCAATATTGCNGGNNCCACTCCCCTGTCCNTCACAGCCATC TTNCTGNCAGGGNGCACGCNNGCTGGGNGTTCCCGCCTANNGNNNCTGGNNNGNGATTCCTNNNNCGGNNN GATGACGTCAGCNTTCAGCTNNTCGANATTNNCCNNNNATCCCTANNACCCNNNNCCCNNNAGNGANNCNN NANTACNNNNCGNCNNNNNNNNNNNNNNNNGNCNNGNANNGNNNNACCNGNNNNTNCCNANTNANCN NNNNNNGNNNNNTCCCNNNNNNNNNNNNNNGNANNNNCNNAAANNNCNNNNCNNNAATNNNCCCNTT CCNANN

- Colony 12, the forward sequence is given from  $5' \rightarrow 3'$ 

NNNNNNNNNNNNCTAGGCGCGCCNTGAGCTCGGGGCTCGAGTTATTCCAGTGCCCAAAATTCTTCT **TCCNNGGNTNTATTATCTTTGCCTTTCAGGGCATCATCATTAAACATAACTTCAAAAATTTTCTTCTTGTC** CTGGCCTCATGGGCCTTCCGCTCACTGCCCGCTTTCCAGTCGGGAAACCTGTCGTGCCAGCTGCATTAACA TGGTCATAGCTGTTTCCTTGCGTATTGGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTCG TTCGGGTAAAGCCTGGGGTGCCTAATGAGCAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGT TGCTGGCGTTTTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTGG CGAAACCCGACAGGACTATAAAGATACCAGGCGTTTCCCCCTGGAAGCTCCCTCGTGCGCTCTCCTGTTCC GACCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCCCTTCGGGAAGCGTGGCGCCTTTCTCATAGCTCAC GCTGTAGGTATCTCAGTTCGGTGTAGGTCGTTCGCTCCAAGCTGGGGCTGTGTGCACGAACCCCCCCGTTCAG CCCGACCGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTTATCGCCACT GGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTCTTGAAGTGGT GGCCTAACTACGGCTACACTAGAAGAACAGTATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGGA **GCANATTACGCGCAGAAAAAAAGGATCTCAAGAAGATCCTTTGATCTTTCTACGGGGTCTGACGCTCANN GGAACGAAAACTCACGTTANGGGATTTTGGNCATGAGATTATCAAAAAGGATCTTCNCNANATCCTTTTAA ATTAAAANNANNTTTAAATCAATCTAAAGTANATATGAGTAACTNGNCTGACNNTTACCNANGNTNNNCAN** NGANGCACCTANNNANNNATNTGNCTANTTCNNTCATCNNANNNCNNTNANNNCCCNNTNNNNANANNN

# - Colony 12, the reverse sequence is given from $3' \rightarrow 5'$

- Colony 13, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNGNNTAGGCGCGCCATGAGCTCGGGGCTCGAGTTATTCCAGTGCCCAAAATTCTTCTTCAT CCATGGCTTTATTATCTTTGCCTTTCAGGGCATCATCATTAAACATAACTTCAAAATTTTCTTCTTTGTCT GGCCTCATGGGCCTTCCGCTCACTGCCCGCTTTCCAGTCGGGAAACCTGTCGTGCCAGCTGCATTAACATG GTCATAGCTGTTTCCTTGCGTATTGGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTCGTT CGGGTAAAGCCTGGGGTGCCTAATGAGCAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGTTG CTGGCGTTTTTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTGGCG AAACCCGACAGGACTATAAAGATACCAGGCGTTTCCCCCTGGAAGCTCCCTCGTGCGCTCTCCTGTTCCGA CCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCCCCTTCGGGAAGCGTGGCGCTTTCTCATAGCTCACGC TGTANGTATCTCAGTTCGGTGTAGGTCGTTCGCTCCAAGCTGGGGCTGTGTGCACGAACCCCCCCGTTCAGCC CGACCGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTTATCGCCACTGG CAGCAGCCACTGGTAACAGGATTAGCANAGCGAGGTATGTANGCGGTGCTACAGAGTTCTTGAAGTGGTGG CCTAACTACNGCTACNCTAGAAGAACAGTATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCNGAAA AAGAGTTGGTAGCTCTTGATCCGGCAAACAAANNACCGCTGGTAGCGNNNNNNTTTTTTGTTTGCAAGNNN CANANTANNCGNNNNAANAAAANNGNTNNNANAANANCNTTTGANCTTTTCNANGGGNTCNGANGNNNNNN NANTTNNNCNNNNNNGNATNNNANNANNGNNNNNGNTNCNNNNNNNNCNGNGNGNNNNNNANNNNNNNNN CNNNNNNNNNGNNNTNCNNNNTN

## - Colony 13, the reverse sequence is given from $3' \rightarrow 5'$

NNNNNTNNNGNNNNNNNNNNNNNNNAAGNCGCGGCCGANCANNNNGNNNNTNNNCACCTCGGCA TGAAGCTGTNNNAAAGNNNGAANGNANTGCTGTTGTTANTTGTTTATTGCNGNNTGTNNAGGTTACAAATG AAGAAATAGAATCTNNAATTTCAGAAAAAGTGCTTTTTTTTCNNTGTANNCTNNTTNNGGNNTGTCCNNNC TTNTCAATGGGNTTAANTGNCTGGAGCTACGTAGTACNACTCACGATNGGCCTANAGGCTNCAGATATTTG TAATATTCATCCATGNTATGGGGGCCTCCCNNCAGCCGTTACAAACTCCAGAANGACCGTGANNACTCTCTT GTCNTTGGGATCAATCGACATGGCACATTGTGCGGACACGTCAGGGTTGTCAGGNAAAAGGAGAGGTTCAC CGCCNATTGGAGNATTTTGATGACAATGCCCGNCGGGTGGCAGGCCGCCGCCAGGAATGTTGTGTCAACCC NNGAAGTTGGCTNTGATGNNNTTCTTAAGCTNNTCGCCNATGATGCNNACNNGNGAAAGAAGGACCTGCAT TCCNAAATGTGNNNGAGGATGTTTCANNCCTCTTGANACTAATTCCTTAAGCTCACTTCTGTGGACAAGGG **GGGCCCAATCCCACTTGACTTCNNNNTGGANGTNGTCCCTCCNGCCGTCCATGTAGAANATGNTCCCCCCT** CGCACTATCCCTCANGNNTGNCGAATCTNGAANAANTAAGTGCCNCNTCNTNTGATCTGGACCTCCTGNAA ANNTTTTGNACNNGCTANCGNNANNGNNNNCNNNAGTNNNNANGNNANAGNANNNCTACAGNACNGNNAT AANTGAAGGACNAGNTNTCNGATATGNNAATTGATCTTCACCCTNTGAACTNTCCGGAANANNNGNACCCC **CTGACCNTCTCCATTNNATTGCANNNNGGANTNGNGACNACNCNNGGANNNNTGATCTCNTNNANNNNNGN** ANNCCGANNANANNNANNNNGCNTCTCATNNANANNTTNCAGAANTGCCCTNNNTCNNNACNTNNNACGN TNTCNNNNANNNCCTNNAGCCTGNNNNNNNNNNNNATCATTNNNGGTNGCCNNCCNNNNNGNTCCNTCNA NTNNNATCCATCTNTCANANGNNNNNAACGNACACNCTANGCT

#### - Colony 14, the forward sequence is given from 5' $\rightarrow$ 3'

NNNNNNNNNNNNNTAGGCGCGCCNTGAGCTCGGGGCTCGAGTTATTCCAGTGCCCAAAATTCTTCTTC ATCCNNNNNNTATTATCTTTGCCTTTCAGGGCATCATCATTAAACATAACTTCAAAAATTTTCTTCTTTGT CTTCTTCCTGCACCACTTTTTCCGGTTTTTTTCTTTGCGCAGAAAATAGGATCCGGGGGGGTACCTCTTAATTA ATGGTCATAGCTGTTTCCTTGCGTATTGGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTC GTTCGGGTAAAGCCTGGGGTGCCTAATGAGCAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCG TTGCTGGCGTTTTTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTG GCGAAACCCGACAGGACTATAAAGATACCAGGCGTTTCCCCCTGGAAGCTCCCTCGTGCGCTCTCCTGTTC CGACCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCCCTTCGGGAAGCGTGGCGCTTTCTCATAGCTCA CGCTGTAGGTATCTCAGTTCGGTGTAGGTCGTTCGCTCCAAGCTGGGCTGTGTGCACGAACCCCCCGTTCA GCCCGACCGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTTATCGCCAC TGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTCTTGAAGTGG TGGCCTAACTACGGCTACACTAGAAGAACAGTATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGG AGCANATTACGCGCAGAAAAAAAAAGATCTCAAGAAGATCCTTTGATCTTTTCTACGGGTCTGACGCTCAN TGGAACGAAACTCACGTTAANGGNATTTTGGNCATGAGATTATCAAAAAGGATCTTCACCTAGATCNNTTT NAATTAAANANGANTTTTNANTCANTCNNAAGTATATNNGANTAAANTNNCTGACNNTTACCNATGCNTAN NNNNAGCACNATNNNAGNNATNNNNNATTNNNNATNNNNNNNGNCTGANNCCCCGNNNNNANNNANNTA NNANNANNCCNNNNNNNANN

## - Colony 14, the reverse sequence is given from $3' \rightarrow 5'$

NNNNNNNNACNNNGNTCCNNNNNNNAAAGNNNGNGCGGAGCNNNANGNNNTGAGNCCNGGTAANNTGA GGGANNNNNGNNNNNTNNCNGGGNNAGNNGNNGGNANGNNGNNTNNAGAAGACAAAGAAGAAGAAGATTGTG AAGTTNAGGTTAATGAGGATGCCCTGGAAGGCGAAGATAATAAANNCATGNANGACAANAATTTTGGGCAC TGGAATAACTCNAGCCCCNAGCTCATGGCGCGCCTAGGCCTTGACGNNCTTCNGCNNATTCGCCCTATNNT GAGACGGATTACGTCGCGCTCACTGGCCGTCGTTNTACANCGCCGTGACNGGNAAAACCCTGGCGTTACCN AACTTANNCGACTTGAGCNNATCCCCCCTTTCGCCANCTGGCNTAGAGCGAAGAGGCCCGCCCGAAACNNNN NTCNCAACANNGCGCNGCCTGATGGGAATGGGAGCGCCCTGTAGCGGCCACTCGCCCNGTCTCGGNCTACN TTCGATTTNTAANGGATTTTGCCGATTTCGGCATATTGGNNACANAATGATCTGATTTANCNNAAANATAA CGCGAATTTTAACGNANNNTANCGCTTACATTTANGTGGNACTCCACGGGAAAATGTGTGCGGNGANCCCTA TCTGTCTATGATTGTCANTCCCCTCCAAAAATGTATACGCTCATGAAAACACTAANCCTGATAACTGTTTC GGATGATATTGAAAAAGGAAGAGTAGGACGCACACAACATATCNGTGTCNCCTGTANNNCTTTCNTCCCAGC ANGTTTGCGTTCCTGNTTTNGCNNNNNNTAAANGNNNNGNAAGTAGTANNGGTGANNATNANNGNGTGC ACGAGANTGGNNNATGNNANCTGATNCTANCNCCGNNNCNNANNNNNNNTATNNCCCCNANNACNNNNTGN

- Colony 15, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNANGNNNATCTCGAGAGCTTCTGAGGGCTTCTTCCTTATTGGAAAGAGTGGTCTCATACAGAA CTTCAACCTCGTTGATGACGTCAGCGTTCACGCTTGAACGCTGACGTTCTCAATTTTCTTCTTGGAATCCT GGAGTGGCGCCCTGCAATATTTGCCTGTCTCTATGTGGTCTGGGAAGTCGTGCCAAACGTGAAATGTCTTT GGATTTGGGAATCTTATAAGTTCTGTATGAGACACTCTTTCGGATCCACCACCATGACTAAAGGAGAA GAACTTTTCCCGGGANGTGTCCCCGATTCTTGTTGAATTAGATGGTGATGTTAATGGNNACNAATTTTCTGT CGGNGTTNNNNNGANGNNNATGCCNCNTNCNGAAAACTTACCCTTAAATTTATTTGCACTACTGGAAAAC TACCTGTTCCATGGTCAACACTTGTCGCTACTCTCACTGATGGCGTTCAATGCTTTTCTNGAGATCCNAAN CATATGAAGCGNNNCGACNTCTTCNANANCGCCATGCCTGANGNATACGTGCATGANAGGACCNTCTTCTT NNAGGACTACNNTAACTACAAGACACGTGCTGAAGTCNAGNNTGAGGGAGACACCCTCGTCAACAGGATCN AGCTTACGGGAATCGATTTCAAGGAAGACNGAAACNTCCTCCGCCACAAGTTGGAATACCNCTACTGNTCC CACNNCGTANACATCATGNCCGACAAAACAAAAGAACGGCATCNGAGCCACCTACNTGACCCGCCGCAACAT CNAATACGGCGGCNCGCAANTCGCTGATCNTTATCACNNAATNCTCCAGATGNCGATGACCCTGTCCNNTT ACCTGACAACCTNTACCTGNCCACANAATCTGCCCTTNNAAAGATCCCATCTAATAGATAGACACATGGNC NNNNNGAGNTNGAAACGGATCGCNGGGATACNCNTGGNNTGNNTNACCTATACAGANNCTCGANNCACTAN 

- Colony 15, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNANNNNNNNNGGNATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCTGAACC TGANACNTAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAATAAAGC **AATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCAAACTCAT** CAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTATTTGTA TAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCTCTCTTTT CGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGGTCATCG CCAATTGGAGTATTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCGGGGTCTT GAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGTTGTATT CCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTGACGAGG GTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGTCCTCTC CTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATCTTGAAA AGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTAGTGCAA ATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTTGTACCC ATTAACATCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTACTCATGG TGGCTGTGGATCCGAAAGANTGTCTCATACAGAACTTATAAGATTCCCAAATCCAAAGACATTTCACGTTT ACGGTGATTTCCCAGAACNCNTAGCGACATGCAAATATTGCAGGGCGCNNCTCCCCTGTCCNTCACANCCA TCTTNCTGNCAGGNNNACGCNCGCTGGGNGNTCCCGNNANNGANNCTGGNNNGCGATCNNAGNNNNTNGA 

- 2. The following are sequences from colonies with the pB12mcs-H1-shRNA2<sup> $\beta$ 2\alpha2</sup>eGFP plasmid. All sequences were missing the shRNA sequence.
- Colony 9, the forward sequence is given from 5'  $\rightarrow$  3'

- Colony 9, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNAGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAAACCTCCCACACCTCCCC **CTGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAA** ATAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCA AACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTT ATTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCT CTCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGG GTCATCGCCAATTGGAGTATTTGTTGATGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGC GGGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAG **TTGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTCAAATCGATTCCCTTAAGCTCGATCCTGTT** GACGAGGGTGTCTCCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGG TCCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATAT AGTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCCCCCTGACAGAAAATT TGTACCCATTAACATNNNNNTCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTTA CTCATGNNGCTGTGGATCCAAAAAANNNAAGGNAAGAAGCCNTCACCNAGCTTCGTGAANGGCTNCTTNCT TANTGAAANANNGNNTCATACNGAACTTNTANANNNNNANTCNNNGANNNTNACNNNNNNNATTNCNNAN NNNANCNNNNTGCANAANNTGNNGGNNNCNNNNNCNTCNNNNNNCTTNNNNNNNCANNNGNCNNNT NNNNNCNNCNNNGNANNNGNNNNNNNANNNCNNNNNNGNNNN

- Colony 10, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNTATCTCGAGAAGCTTCGTGAAGGGCTTCTTCCTTATTAAAAGAGTGGTCTCA TACAGAACTTATAAGATTCCCCAAAATCCCAAAGACATTTCACGGTTACGGTGATTTCCCCAGAACACATAGCGA CATGCAGATATTGCAGGGCGCCACTCCCCTGTCCCTCACAGCCATCTTCCTGCCAGGGCGCACGCGCGCTG GGTGTTCCCGCCTAGTGACACTGGGCCCGCGATTCCTTGGAGCGGGTTGATGACGTCAGCGTTCAAGCTTG GTGAAGGGCTTCTTCCTTATTTTTTTTGGATCCACAGCCACCATGAGTAAAGGAGAAGAACTTTTCACTGG AGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGGAGAGGGTG CCAACACTTGTCACTACTCTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGGCA CGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGCACCATCTTCTTCAAGGACGACGGGA ACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAATC GATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCACAACGTATACAT CATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCGAANACGGCGGCG TGCAACTCGCTGATCATTATCACAAAATACTCCCAATTGGCGATGACCCTGTNCNTTTACCANANANCATTA CCTGTCCNCACATCTGCCCTTTCGAAGATCCCAACGAAAAGAGAGACCACATNNNNNNNAGTTNNACGGC TGCTGGNANNNNATGNNNNNGANTATNCAANANTCNANCNNNANACTNNNNTGNNTCNANTNCGTNNAT CNNANNGATANANNCNTNNTNANNNNNNNNNNACTNNNANGNNNNAAAANGNTNANTNNGANNNNNTG NTTNNNTNNGNCNACTNNNNTNCCCNNNNNN

- Colony 10, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNGNANNAGGGTAATCTTAAGTCGCGGCGCGAATTAAAAAAACCTCCCACACCTCCCC CTGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTATTGCAGCTTATAATGGTTACAA ATAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCA AACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTT ATTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCT CTCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGG GTCATCGCCAATTGGAGTATTTGTTGATGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGC GGGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAG **TTGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTCAAATCGATTCCCTTAAGCTCGATCCTGTT** GACGAGGGTGTCTCCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGG TCCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATAT CTTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGT AGTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCCCCCTGACAGAAAATT TGTACCCATTAACATCACCATNNNNNTCAACAAGAATTGGNACACTCCAGTGAAAAGTNNTCTNCTTTACT CATGGNGGNTGTGNTCCAAAAAAAAAAAAANNNANAAGCCNTTCNCNAGCTTGAACGCTGACGTCATCANNCN GNTNAGNNCNCGNNCCANNNTNNCTNGNNNNNNCCNNNNNGNNNNNNCNNNNGANANGNNTGNNNNNN TNNANNNNNNNNNNNNNNNNNNNNNNNN

The following sequence was inconclusive:

- Colony 15, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNGGGNATCTCGAGNGCTTCGTGAAGGGCTTCCTTATTAAAAGAGTGGTCTC ATACAGAACTTATAAGATTCCCAAATCCAAAGACATTTCACGTTTACGGTGATTTCCCAGAACACATAGCG ACATGCAGATATTGCAGGGCGCCACTCCCCTGTCCCTCACAGCCATCTTCCTGCCAGGGCGCACGCGCGCC GGGTGTTCCCGCCTAGTGACACTGGGCCCGCGATTCCTTGGAGCGGGTTGATGACGTCAGCGTTCAAGCTT GGTGAAGGGCTTCTTCCTTATTTTTTGGATCCACAGCCACCATGAGTAAAGGAGAAGAACTTTTCACTG GAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGGAGAGGGT GCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGGC ACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTTCTTCAAGGACGACGGG AACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAAT CGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACAACTCCCACAACGTATACA TCATGGCCGACAAGCAAAAGAACGGCATCAAAGCCAACTTCAAGACCCGCCACAACATCGAAGACGGCGGC **GTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTTTTACCAGACAACCA** TTANNNGNNCACACAATCTGCCCTTTCGAAAGATCCCCAACGAAAGAGAGACCACATGNNCCTTCTTGAGTT TGNAACGGCTGCTGGGATTACNCATGGCATGNTGNANTATNCNAANNNNNGAGNCTCTANANTATNNNGAG TCGTATNCGTANATCNGNNNTGATAGANNNNTGNTNANNTNNNNNNNNNANCTNNNANGNNNNNAAAAANG TNNNNNNNGGNANNNNNNNNNNNNNTTTNNNN

- Colony 15, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNANGNANNAGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAAACCTCCCACACCTCCCCC TGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAA TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA **ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA** TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT **TGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTG** ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA **GTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATTT** GTACCCATTAACNNCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTNCTTNTCCTTTAC NNCGCTCNNNNTCNNGGGNCCNNNNNCNNTAGNNGGGNNNCCNAGCGNGNNNGNGCCCNNNNGNNNANGNTN 

- 3. The following are sequences from colonies with the pB12mcs-H1-shRNA3<sup> $\beta$ 2\alpha^2</sup>eGFP plasmid. All sequences were missing the shRNA sequence.
- Colony 3, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNNGNTATCTCGAGAAGCTTGAACGCTGACGTCATCAACCCCGCTCCAAGGAATCGC **GGGAGTGGCGCCCTGCAATATCTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTT** TGGATTTGGGGATCTTATAAGTTCTGTATGAGACCACTCTTTCCAATAAGGAAGAAGCCCTTCACGAAGCT **TGGTGAAGGGCTTCTTCCTTATTGTTTTT**GGATCCACAGCCACCATGAGTAAAGGAGAAGAACTTTTCAC TGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGGAAAGG TGGCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCG GCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTTCTTCAAGGACGACG GGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGA **ATCGATTTCNAGGAGGACGGANACATCCTCGGNCACNAGTTGGAATACNACTACNACTCCCACAACGTATA** CATCATGGNCGACAAGCAAAAGAACGGCATCANAGCCAACTTCAAGACCCGCCACAACATCGAAGACGGCG **GCGTGCAACTCGCTGATCATTATCAACAAAATACTCCAATTGGCGATGACCCTGTCCTNTTACCANNNNN** ATTACCTGTCCNCACAATCTGCNCTTTCGAAAGANCNNNACGANANANAGACNNNNTGNNCNTTCNTGAGT TGTACGGCTGCTGNNATTANNCATGCNNGNATGANNTNNACNNNNNCTCGNGNCTCNAGACNNTAGNNAGN NNNNNACNNCNANCNNANNNNANNNGANNNNTNTNNTNNNNNNANNNNNNNNNANNNNNNAANNNNNAANNNAA NNNTNATNNNANNNGGNNNGNNNNGNNNNTNNANNNTNNANCTGCANNAANCAGNNNNNNN
- Colony 3, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNNNNNGGNTATCTCGAGACCCAAGCTTGAACGCTGACGTCATCAACCCGCTCCAAGG AATCGCGGGCCCAGTGTCACTAGGCGGGGAACACCCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTGAG GGACAGGGGGAGTGGCGCCCTGCAATATCTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAA TGTCTTTGGATTTGGGAATCTTATAAGTTCTGTATGAGACACTCTTTCCAATAAGGAAGAAGCCTTTCACG AAGCTTGGTGAAGGGCTTCTTCCTTATTGTTTTTGGATCCACAGCCACCATGAGTAAAGGAGAAGAACTT **TTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGG** TTCCATGGCCAACACTTGTCACTACTCCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATG AAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGGAGCCATCTTCTTCAAGGA CGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTA ANGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACACTCCCACAACG TATACNTCATGNCGACAGCAAAGANGGCATCAAAGCCAACTTCAGACCGCACACATCGANANGCGCGTGCA NTCGCTGANCATATCACAATACTCCATTGCNATGACCNGNNNTTNACAGANACNTNACTGNCCACACATCT **GNCCTTTCNAANATCCNANNAAAANANAGACCNNATGNNCNNTNNNNNACGNCTGCTGGNATNNNNTG** NNNTNNNNNNNNNNGNNNNNNNNNNNNNNNN

- Colony 5, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNNNGGNNNNNCTCGAGACCCAAGCTT<mark>GAACGCTGACGTCATCAACCCGCTCCA</mark> GAGGGACAGGGGAGTGGCGCCCTGCAATATCTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTG AAATGTCTTTGGATTTTGGGAATCTTATAAGTTCTGTATGAGACACTCTTTCCAATAAGGAAGAAGCCTTTC **ACGAAGCTTGGTGAAGGGCTTCTTCCTTATTGTTTTT**GGATCCACCACCATGAGTAAAGGAGAAGAA **CTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAG** CTGTTCCATGGCCAACACTTGTCACTACTCCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCAT ATGAAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGGAGCATCTTCTTCAA GGACGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGC TTAAGGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACNAGTTGGAATACAACTACACTCCCACA ACGTATACATCATGGCCGACAGCAAAGANNGCATCAAAGCCAACTTCNANACCCGCACACATCGANACGNG CGTGNACTCGCTNANCATATCACAAATACTCAATTGCGATGACCTGTCTTTTACNNACACCATACCNGTCC NNCATCTGNCTNNNNNNNCCACNAAANANAACNNNNGGNCNTNCTGANTTNACGCTGCTGGNANNNNNNGC ATGNNGACNNNCNATACTCNAGCCTCNNANNNANNNNCNTNNNCNNNNTCCGANTGNNNNNNNTGTGNN NNGNNNNNNNANATGNNNNANAANNNNTATTNNATNNNNTTGNTTNNNNNANNNNNNAAANNNAN NTNNNNNANNNNNNNNNNNNNNNNNNNN

- Colony 5, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNGGNANNAGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCCCC TGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAAA TAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCCAA **ACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTTA** TTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTCTC TCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGGACAGGTAATGGTTGTCTGGTAAAAGGACAGGG TCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGCG GGTCTTGAAGTTGGCCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAGT **TGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTTG** ACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATGGT CCTCTCCTGCACGTATCCCTCAGGCATGGCGCCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATATC TTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA **GTGCAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCTCCACTGACAGAAAATNN** GTACCCATTAACATCACCATCTAATTCAACAAGAATTNNNNNCTCCAGTGAAAAGTTCTTCTCCTTTACTC **ATGGNGGCTGTGGATCCAAAAAANANAAGNAAGAGCCCNTCNCNNCTTCNTGAAAGNTTCTTCNNTNNNN** NNNNNNN

- Colony 6, the forward sequence is given from 5'  $\rightarrow$  3'

NNNNNNNNNNNNNNNNNGGNTATCTCGAGACCCAAGCTTGAACGTCATCAACCCGCTCCAAGG AATCGCGGGCCCAGTGTCACTAGGCGGGAACACCCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTGAG GGACAGGGGAGTGGCGCCCTGCAATATCTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAA TGTCTTTGGATTTGGGAATCTTATAAGTTCTGTATGAGACACTCTTTCCAATAAGGAAGAAGCCTTTCACG AAGCTTGGTGAAGGGCTTCTTCCTTATTGTTTTTTGGATCCACAGCCACCATGAGTAAAGGAGAAAACTT **TTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGG** TTCCATGGCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATG AAGCGGCACGACTTCTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGGAGCCATCTTCTTCAAGGA CGACGGGAACTACAAGACACGTGCTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTA ANGGAATCGATTTCAAGGAGGACGGAAACATCCTCGGCCACAAGTTGGAATACAACTACACTCCCACAACG TATACNTCATGNCGACAGCAAAGANGGCATCAAAGCCAACTTCAGACCGCACACATCGANANGCGCGTGCA NTCGCTGANCATATCACAATACTCCATTGCNATGACCNGNNNTTNACAGANACNTNACTGNCCACACATCT **GNCCTTTCNAANATCCNANNAAAANANAGACCNNATGNNCNNTNNNNNACGNCTGCTGGNATNNNNTG** NNNTNNNNNNNNNNGNNNNNNNNNNNNNNNN

- Colony 6, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNTNTANGNNNNAGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCC CCTGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTATTGCAGCTTATAATGGTTACA **AATAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTGTCC** AAACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGT TATTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTC TCTCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAG GGTCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGG CGGGTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTA GTTGTATTCCAACTTGTGGCCCGAGGATGTTTCCCGTCCTTGAAATCCATTCCCTTAAGCTCCATCCTGT TGACGAGGGTGTCTCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATG GTCCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATA TCTTGAAAAGCATTGAACACCATAAGTGAGAGAGTAGTGACAAGTGTTGGNCATGGAACNGTAGTTTTCCAGT ANTGCAAATAAATTTNNGNNAAGTTTTNCGTATGNTGCATCACCTCCCCTCTCNCTGACAGNNNNTTNGT ACCCATTAANATCANCATCTAATNAACAAGANTNNNCNACTCCAGTGAAAAGTNTCTCNTTACTCATGNNG CTGNGNNCNAAAANAANNANNAGAAGCCNTNNCANNTNNNAANNTCTCNNTNAANNNNNNNNNANGAC CNGCNGNNNNNNNNNNNNNNNNNNNNATCTNANNNNNANNANNNN

The following sequences was inconclusive:

- Colony 4B, the forward sequence is given from 5'  $\rightarrow$  3'

TTTTTGGATCCACCACCATGAGTAAAGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAA TTAGATGGTGATGTTAATGGGTACAAATTTTCTGTCAGTGGAGAGGGTGAAGGTGATGCAACATACGGAAA **ACTTACCCTTAAATTTATTTGCACTACTGGAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTCTCA** CTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGGCACGACTTCTTCAAGAGCGCCATG CCTGAGGGATACGTGCAGGAGGAGGACCATCTTCTTCAAGGACGACGGGAACTACAAGACACGTGCTGAAGT CAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACA TCCTCGGCCACAAGTTGGAATACAACTACAACTCCCACAACGTATACATCATGGCCGACAAGCAAAAGAAC GGCATCAAAGCCAACTTCAAGACCCGCCACAACATCGAAGACGGCGGCGTGCAACTCGCTGATCATTATCA ACAAAATACTCCAATTGGCGATGACCCTGTCCTTTTACCAGACAACCATTACCTGTCCACACAATCTGCCC TTTCGAAAGATCCCAACGAAAAGAGAGAGACCACATGGTCCTTCTTGAGTTTGTAACGGCTGCTGGGATTACA CATGGNATGGNTGAACTATACAAATAACTCGAGCCTCTAGAACTATAGTGAGTCGTATTANNNNNTCCAGA CATGATAAGATACATTGATGAGTTTGGACNAACCACANTAGANGNAGTGAAAAAATGCTTANTTGNGAAAN TTNNNTGCNNTNNTTNANTNGTANCATNNNGCTGCANNAANNNNNACACANNNNNNTNCATTNNNTNNTT NNNNNGNGNNGGNNNGGNAGNNTNNNNNTNNNGNNNNNTNNNNTACCCNGNTANNCNNNNNCNGNTT NNNNNNNANAN

- Colony 4B, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNTNTNNGNAANNGGNTANNCTTAAGTCGCGCCGCGAATTAAAAAACCTCCCACACCTCCCC **CTGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACAA** ATAAAGCAATAGCATCACAAATTTCACAAATAAAGCATTTTTTTCACTGCATTCTAGTTGTGGTTTTGTCCA AACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGTT ATTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAAACTCAAGAAGGACCATGTGGTCT CTCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAGG GTCATCGCCAATTGGAGTATTTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGGC GGGTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTAG TTGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGTT GACGAGGGTGTCTCCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCCGTCGTCCTTGAAGAANATGG TCCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATAT CTTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAGTGTTGGCCATGGAACAGGTAGTTTTCCAGTA NTGCAAATAAATTTANGGTAAGTTTTCCGTATGNTGCATCACTTCACCTCNNCACTGACNGAAAATTTGTA CCCATTANATCACCATCTANTCAANANAATTGGGACANTCNGTGAAAAGTCTTTCNTNCNGGNGNTNGGNN 

## - Colony 5B, the forward sequence is given from 5' $\rightarrow$ 3'

GGGCCCAGTGTCACTAGGCGGGAACACCCCAGCGCGCGTGCGCCCTGGCAGGAAGATGGCTGTGAGGGACAG GGGAGTGGCGCCCTGCAATATCTGCATGTCGCTATGTGTTCTGGGAAATCACCGTAAACGTGAAATGTCTT TGGATTTGGGAATCTTATAAGTTCTGTATGAGACCACTCTTTCAATAAGGAAGAAGCCCTTCACGAAGCTT GGTGAAGGGCTTCTTCCTTATTTTTTTGGATCCACAGCCACCATGAGTAAAGGAGAANAACTTTTCACTG GAGTTGTCCCAATTCTTGTTGAATTAGATGNNGATGTTAATGGGTACAAATTTTCTGTCAGTGGAGANGGT NCCAACACTTGTCACTACTCACTTATGGTGTTCAATGCTTTTCAAGATATCCAGATCATATGAAGCGNC ACGACTTCTTCAAGAGCGCCATGCCTGANGGATACGTGCAGGAGGAGCATCTTCTTCANGGACNACGGN AACTACNAGACACGTGCTGAANTCAAGTTTGAGGGAGACACCCTCGTCACAGGATCGAGNTAANGGAATCG AGCAAAGACGNATCAAGCAACTCANACCCGCNCAACATCGANACGNGCNNNNNCTCGNTGATCATTATCAN NNAATACNNCANGGNNATNACCNTGTCNTTTACAGANNCNTTACNNNCACNCATCNGCCCTTNCGAGANCC NGNNNNNNNNNNNANNATCNNNNNNNNNNNNTNNNTGCNNNNNNNCNNNNNNGNNNN

- Colony 5B, the reverse sequence is given from  $3' \rightarrow 5'$ 

NNNNNNNNNNNNGGNANNNGGGTAATCTTAAGTCGCGGCCGCGAATTAAAAAACCTCCCACACCTCCC CCTGNACCTGAAACATAAAATGAATGCAATTGTTGTTGTTGTTAACTTGTTTATTGCAGCTTATAATGGTTACA AAACTCATCAATGTATCTTATCATGTCTGGATCTACGTAATACGACTCACTATAGTTCTAGAGGCTCGAGT TATTTGTATAGTTCATCCATGCCATGTGTAATCCCAGCAGCCGTTACAAACTCAAGAAGGACCATGTGGTC TCTCTTTTCGTTGGGATCTTTCGAAAGGGCAGATTGTGTGGACAGGTAATGGTTGTCTGGTAAAAGGACAG GGTCATCGCCAATTGGAGTATTTGTTGATAATGATCAGCGAGTTGCACGCCGCCGTCTTCGATGTTGTGG CGGGTCTTGAAGTTGGCTTTGATGCCGTTCTTTTGCTTGTCGGCCATGATGTATACGTTGTGGGAGTTGTA GTTGTATTCCAACTTGTGGCCGAGGATGTTTCCGTCCTCCTTGAAATCGATTCCCTTAAGCTCGATCCTGT TGACGAGGGTGTCTCCCCTCAAACTTGACTTCAGCACGTGTCTTGTAGTTCCCGTCGTCCTTGAAGAAGATG GTCCTCTCCTGCACGTATCCCTCAGGCATGGCGCTCTTGAAGAAGTCGTGCCGCTTCATATGATCTGGATA TCTTGAAAAGCATTGAACACCATAAGTGAGAGTAGTGACAAGTGTTGGCCATGGAACAGGTAGTTTTCCAG TAGTGCAAAATAAATTTAAGGGTAAGTTTTCCGTATGTTGCATCACCTTCACCCTCCACTGACAGAAAAT TTGTACCCATTNNNNTCACCATCTAATTCAACAAGAATTGGGACAACTCCAGTGAAAAGTTCTTCTCCTTT ACTCATGGNGGCTGTGGATCCAAAAAAAANNAAGGGAAGAAGCCNNTCACCNNCTTCGTGAAGGGCNTTCT **TCTTANTGAAANAGTGNNTCATNNNNANTTATANANNCCCAAANNNNNNGANNTTNACGTTANGNGATTNC** 

## Appendix 10: FACS Generated Data of Transfection of K562 and KCL-22 cell lines

- $\mathsf{Supp}_{\mathsf{res}}$
- FACS data generated from nucleofection of K562 cells



Sample		Transfection efficiency with		Transfection efficiency with	
		regards to non-transfected		regards to negative control	
		cells			
		(35.95%)	-Purple	(47.40%)	-Green
Positive control -Blue		28.06%		28%	
shRNA1 <sup>β3α2</sup>	- Pink	26.04%		2	26.07%
shRNA3 <sup>β3α2</sup>	-Orange	14.01%		2	2.53%



Figure 85 FACS histogram showing the transfection efficiency in K562 cells, transfection number 2. The results as shown below:

Sample		Transfection	efficiency	Transfection	efficiency
		regards to non-transfected		regards to negative control	
		cells			
		(23.05%)	-Purple	(68.33%)	-Green
Positive control -Pink		72.15%		27	<b>.</b> 17%
shRNA1 <sup>β3α2</sup>	- Blue	54	.18%	8.9	9%
shRNA3 <sup>β3α2</sup>	-Orange	47.72%		2.4	44%



Figure 86 FACS histogram showing the transfection efficiency in K562 cells, using  $shRNA1^{\beta 2\alpha 2}$ . The results as shown below:

Sample		Transfection	efficiency
		regards to nor	n-transfected
		cells	
		(21.70%) –Pur	ple
Positive contro	ol -Green	43.18%	
$shRNA1^{\beta 2 \alpha 2}$	- Pink	5.77%	
$shRNA1^{\beta 2 \alpha 2}$	- Blue	9.86%	
$shRNA1^{\beta 2 \alpha 2}$	- Orange	6.77%	

• FACS data generated from nucleofection of KCL-22 cells



Figure 87 FACS histogram showing the transfection efficiency in KCL-22 cells, transfection number 1. The results as shown below:

Sample	Transfection	efficiency	Transfection	efficiency
	regards to nor cells	n-transfected	regards to neg	ative control
	(32.29%)	-Purple	(58.69%)	-Green
Positive control -Pink	53	.14%	26	.74%
$shRNA1^{\beta 2 \alpha 2}$ - Bue	28.33%		1.9	93%



Figure 88 FACS histogram showing the transfection efficiency in KCL-22 cells, transfection number 3. The results as shown below:

Sample	Transfection regards to nor cells	efficiency n-transfected	Transfection regards to neg	efficiency ative control
	(26.06%)	-Purple	(61.66%)	-Green
Positive control -Pink	67	%	31	.35%
$shRNA1^{\beta 3 \alpha 2}$ - Blue	36.94%		1.3	34%



Figure 89 FACS histogram showing the transfection efficiency in KCL-22 cells, using shRNA1 $^{\beta 3 \alpha 2}$ . The results as shown below:

Sample		Transfection	efficiency
		regards to no	on-transfected
		cells	
		(18.87%)	– Purple
Positive control -Green		79.27%	
shRNA1 <sup>β3α2</sup>	- Pink	50.36%	
shRNA1 <sup>β3α2</sup>	- Blue	51.35%	
shRNA1 <sup>β3α2</sup>	- Yellow	44.29%	



Figure 90 FACS histogram showing the transfection efficiency in KCL-22 cells, using shRNA3<sup> $\beta$ 3\alpha<sup>2</sup></sup>. The results as shown below:

Sample		Transfection efficiency with
		regards to non-transfected cells
		(15.44%) – Purple
Positive control	-Green	77.49%
shRNA3 <sup>β3α2</sup>	- Pink	53.67%
shRNA3 <sup>β3α2</sup>	- Yellow	51.81%
shRNA3 <sup><math>\beta</math>3<math>\alpha</math>2</sup>	- Blue	54.88%

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