

Using a multi-omics approach to identify novel therapeutics in multiple myeloma capable of overcoming drug resistance



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Abstract

Multiple myeloma (MM) is an incurable cancer of plasma cells, with an average five-year survival rate of approximately 50%. Newer therapeutics, namely proteasome inhibitors (PI) and immunomodulatory imide drugs, have almost doubled median survival time of MM patients. However, most patients relapse and become resistant to drugs they previously have been treated with. Acquired anti-cancer drug resistance remains one of the biggest barriers in the treatment of myeloma. Therefore, identifying novel therapeutics effective against MM, which are capable of overcoming drug resistance is of the utmost importance. Recently the prolyl-tRNA synthetase inhibitor, Halofuginone, has been shown to be effective against cancer, including one study demonstrating effectiveness against MM.

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List of Abbreviations

MM	Multiple Myeloma
BM	Bone marrow
MGUS	Monoclonal gammopathy of unknown significance
SMM	Smoldering multiple myeloma
PI	Proteasome inhibitor
IMiDs	Immunomodulatory imide drugs
ER	Endoplasmic reticulum
UPS	Ubiquitin proteasome system
UPR	Unfolded protein response
DNA	Deoxyribonucleic acid
RNA	Ribonucleic acid
tRNA	Transfer RNA
NGS	Next generation sequencing
WGS	Whole genome sequencing
RNA-Seq	Ribonucleic acid sequencing
scRNA-Seq	Single cell RNA-Seq
dscRNA-Seq	Droplet-based scRNA-Seq
CB	Cellular barcode
UMI	Unique molecular identifier
LC-MS/MS	Liquid chromatography with tandem mass spectrometry
PCA	Principle component analysis
DMSO	Dimethyl sulfoxide
UMAP	Uniform Manifold Approximation and Projection
tSNE	t-distributed Stochastic Neighbor Embedding
NK	Natural killer

aaRS	Aminoacyl tRNA synthetase
EPRS	Glutamyl-prolyl-tRNA synthetase
ProRS	Prolyl-tRNA synthetase
MSC	Multi-tRNA synthetase
AIMP	aaRS interacting multifunctional proteins
FF	Febrifugine
HF	Halofuginone

1

Introduction

1.1 Overview

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1.2 The adaptive immune system

Humans are exposed to millions of potential pathogens every day and therefore require defences to be able to protect themselves against infection. These defences can be innate or adaptive. An example of an innate defence is the skin acting as a physical barrier between the outside world and the body. Another example of an innate defence is non-specific engulfing (phagocytosis) of foreign pathogens by macrophages (a type of white blood cell). Innate responses are relied upon as the first line of defence, however sometimes a more sophisticated, specialised response is required- called the adaptive immune response. (REF-mol biology of the cell).

Adaptive immune responses are specific to the pathogen that induced the response and are dependent on B cells and T cells, two major classes of lymphocytes (a class of white blood cell). Two classes of adaptive immune responses exist: antibody responses, co-ordinated by B cells, and cell mediated immune responses, co-ordinated by T cells. T-cell-mediated immune responses recognise foreign antigens (antibody generators; substances capable of eliciting an immune response by stimulating B or T cell activation) on the surface of cells and can either kill the pathogen-infected cells or stimulate B cells or phagocytes to help eliminate the pathogen.

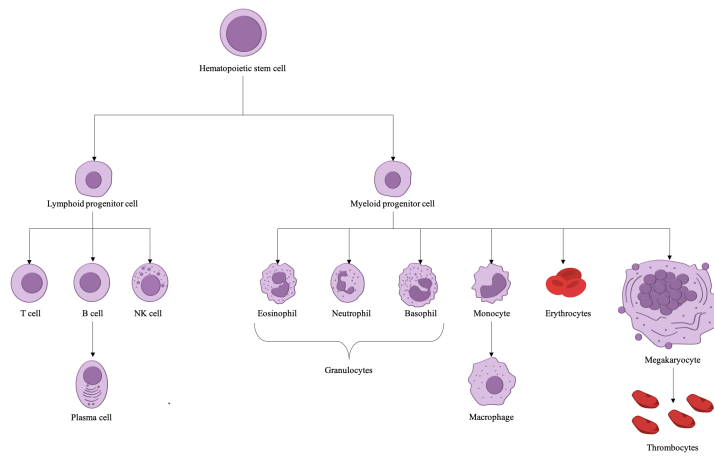


Figure 1.1: Hematopoietic stem cell (HSC) cell differentiation. HSCs divide into myeloid or lymphoid progenitor cells. Dendritic cells and a number of precursor states have been ommitted.

In antibody responses, B cells and plasma cells secrete antibodies, also known as immunoglobulins. Immunoglobulins are large Y-shaped proteins, which recognise and bind to the specific foreign antigen on the pathogen which stimulated their production. Binding of immunoglobulins to antigens renders the virus or microbial toxin inactive as it blocks their ability to bind to host cells. Additionally, antibody binding makes it easier for phagocytic cells to ingest the pathogen.

1.2.1 Plasma cells

Plasma cell development

Stem cells are precursor cells which can give rise to at least one type of differentiated (mature) cell, with the capability of indefinite self-renewal. Hematopoietic stem cells (HSC) are stem cells that give rise to all the cells of the hematopoietic system. Two predominant cell populations are produced by HSCs: the common myeloid progenitor (CMP) and the common lymphocyte progenitor (CLP). CMP differentiation produces erythrocytes (red blood cells), mast cells, monocytes, macrophages, neutrophils, eosinophils, basophils and myeloid dendritic cells. CLP differentiation results in B cells, T cells, natural killer (NK) cells and lymphoid dendritic cells.

Most B cells die in the bone marrow soon after developing, however some will develop in the bone marrow, where initial stages of maturation occur and then migrate to secondary lymphoid organs, such as the spleen. Within secondary lymphoid organs, numerous critical decisions on B cell fate are made, involving complex transcriptional networks, cell interactions, gene rearrangements, and mutations[1, 2]. Upon antigenic-stimulation, naive B cells differentiate into memory B cells or plasma cells. Terminally differentiated plasma cells are the final effectors of the B cell lineage, each dedicated to secreting large amounts of a single type of antibody. Plasma cells have an extensive rough endoplasmic reticulum (ER), and have numerous genes involved in immunoglobulin secretion upregulated, including *XBP-1* and *CHOP*[3], to enable the production of copious amounts of antibody. Plasma cells appear to consist of two distinct categories: short-lived plasma cells, which have life-spans of several months and are located in extrafollicular locales such as in medullary chords of lymph nodes or the red pulp of the spleen, and long-lived plasma cells, which have life-spans of decades and are mainly found in the bone marrow[4, 5].

1.3 Multiple myeloma

1.3.1 Multiple myeloma cells

Multiple myeloma is a malignancy of terminally differentiated plasma cells. It is characterised by aberrant proliferation of clonal, long-lived plasma cells in the bone marrow[6].

1.3.2 Epidemiology

Multiple myeloma accounts for 1-2% of all cancers and has the second highest incidence of hematological malignancies, after non-Hodgkin's lymphoma[7]. MM is rare in individuals under the age of 40, with the average age at time of diagnosis centering around 70[8, 9]. MM is more prevalent in males than females and is around twice as common in black populations than in Caucasian or Asian populations[10]. The average incidence rate is approximately 1-6 cases per 100000 individuals[8, 9,

11], with the highest age-standardised incidence rates in the regions of Australasia, North America, and Western Europe[12]. Five-year survival rate of MM patients is approximately 49%, whilst approximately a third of MM patients survive ten years or greater[13, 14].

1.3.3 Presentation

Precursor states

All cases of MM are preceded by asymptomatic precursor states, monoclonal gammopathy of unknown significance (MGUS) and smoldering multiple myeloma (SMM). However, only some patients with SMM or MGUS progress to active MM.

MGUS is a pre-malignant condition where patients have the presence of monoclonal immunoglobulins in their blood or urine, <10% clonal plasma cells in their bone marrow, but lack any myeloma-related end-organ damage[15]. Patients with SMM have between 10 and 60% clonal plasma cells in their bone marrow, serum monoclonal immunoglobulin of ≥ 3 g/dL, and like MGUS, have no signs of end-organ damage[16]. Progression risk of MGUS into symptomatic MM is about 1% per year, whilst progression risk of SMM to MM is higher, at around 10% per year for the first 5 years, after which it decreases[17, 18].

Active MM

There are multiple classifications of active MM. The International Myeloma Working Group's definition[19] is as follows: Greater than 10% clonal plasma cells located in the bone marrow and one or more myeloma-defining event or biomarker of malignancy. Myeloma defining events consist of evidence of end-organ damage that can be attributed to the surplus of M protein and clonal plasma cells, namely the CRAB features:

- Hypercalcemia
 - Serum calcium > 1 mg/dL higher than the upper limit of normal, or
 - Serum calcium > 11 mg/dL

- Renal insufficiency
 - Creatinine clearance < 40 mL per min, or
 - Serum creatine > 2 mg/dL
- Anemia
 - Hemoglobin value of > 20 g/L below the lower limit of normal, or
 - Hemoglobin value < 100 g/L
- Bone lesions
 - One or more osteolytic lesions on skeletal radiography, CT or PET-CT

Biomarkers of malignancy include greater than or equal to 60% clonal plasma cells in the bone marrow, an involved:uninvolved serum free light chain ratio greater than or equal to 100, and more than one focal lesion on an MRI study[19].

It is currently unclear what causes the malignant transformation between precursor states and active MM. However certain factors have been identified as risk factors, including point mutations, a large array of up-regulated transcription factors, and numerous immune events.

1.3.4 Treatment of multiple myeloma

Multiple myeloma may be an incurable disease, however it is treatable. In fact, in the last decade median survival time for newly diagnosed MM patients has almost doubled[20]. Novel therapeutic advances have contributed to this improvement (Table 1.1). Myeloma is usually treated with a combination of drugs, often comprising a corticosteroid, a proteasome inhibitor, and an immunomodulatory drug (IMiD). A common regimen, approved in the USA, European Union and UK for untreated myeloma is the triplet VRd regimen. This consists of the proteasome inhibitor bortezomib (brand name Velcade), the IMiD lenalidomide (brand name Revlimid), and the corticosteroid dexamethasone.

Year	Treatment	Usage	Ref
1958	Melphalan	The alkylating agent melphalan was first used in plasma cell myeloma in 1958.	[21]
1960s	Corticosteroids	Placebo-controlled double-blind trial of prednisone in multiple myeloma. Combinations of prednisone and melphalan showed an increased survival over melphalan alone. Dexamethasone and prednisone have become a cornerstone in the treatment of multiple myeloma.	[22, 23]
1980s	Stem-cell transplantations	Numerous successful allogenic and autologous bone marrow transplantations in patients with multiple myeloma	[24–27]
2003	Proteasome inhibitors	Bortezomib, a first-in-class proteasome inhibitor, was first approved by the FDA for use in relapsed and refractory multiple myeloma. In 2008 it was approved for patients with no prior treatment. Carfilzomib was approved in 2012 for advanced MM and later in 2015 for treatment of relapsed MM. The oral proteasome inhibitor, ixazomib, was approved as a combination treatment with lenalidomide and dexamethasone in 2016 for people who have received at least one previous treatment.	[28–30]
2006	IMiDs	The antitumour activity of thalidomide was demonstrated in 1999, this led to the development of lenalidomide, the first approved immunomodulatory imide drug (IMiD) for use in multiple myeloma. Currently, thalidomide, lenalidomide and pomalidomide are approved for use in multiple myeloma	[31–33]
2015	Monoclonal antibodies	In 2015, daratumumab, an anti-CD38 monoclonal antibody and elotuzumab, an anti-SLAMF7 monoclonal antibody, were approved for MM treatment.	[34, 35]

Table 1.1: Timeline of treatment options for multiple myeloma. Listed by first usage or FDA approval for MM.

1.3.5 Proteasome inhibitors

Proteasome inhibitors have contributed greatly to the improved prognosis of MM since their introduction into treatment regimes. The first-in-class proteasome inhibitor bortezomib (Velcade®) was approved by the FDA in 2003 as a single-

agent for injection of relapsed MM[28]. Since then it has been approved for use in combination therapies. Bortezomib in combination with melphalan-prednisone proved to be superior to the previous standard of care for patients ineligible for HDT-ASCT of melphalan-prednisone alone, increasing time until tumour progression[36]. The combination of bortezomib, dexamethasone and thalidomide was also shown to be superior to previous standard of care for patients prior to ASCT[37]. In 2010, bortezomib was approved as a frontline therapy for treatment-naive MM patients. Since then, two more proteasome inhibitors have been approved, carfilzomib and ixazomib. Carfilzomib is structurally and mechanistically different to bortezomib and shows activity on bortezomib resistant primary MM cells[37]; it is approved for relapsed or refractory MM.

The ubiquitin-proteasome system

Proteasome inhibitors work by blocking the action of the proteasome in the cell. Misfolded proteins can be harmful to a cell, so the combined activity of molecular chaperones, which aid in protein folding, and the ubiquitin-proteasome system (UPS), which acts to digest misfolded proteins, is needed to prevent massive protein aggregation. Unneeded, misfolded or damaged proteins are tagged with lysine-48-linked poly-ubiquitin chains, marking them for degradation by the proteasome (Figure 1.2a). The proteasome is sometimes described as a complex ‘protein destruction machine’. The proteasome consists of the 20S core particle, a central hollow cylinder, and the 19S regulatory caps associated with each end of the cylinder. The 19S regulatory caps perform substrate recognition, deubiquitination, unfolding and threading of the protein substrate into the 20S core. The core is made up of four stacked heptameric ring structures. The outer rings are responsible for docking to the 19S cap and for acting as a gate to the inner rings. The inner rings consist of seven β subunits, containing inward-facing protease active sites for degrading proteins[38, 39] (Figures 1.2a and 1.2b).



Figure 1.2: Structure of the proteasome. 1.2a shows the structure of the 26S proteasome, comprised of the 19S regulatory caps and 20S core particle. A misfolded protein tagged with a poly-ubiquitin chain is recognised by the 19s regulatory cap, which cleaves the ubiquitins from the protein and threads the protein through to the core, where it is degraded into small peptides. The 20S core particle is made up of two outer rings of α -subunits and two inner rings of β -subunits. 1.2b shows the β -subunit arrangement in one of the inner rings of the 20s particle. β_1 (caspase-like), β_2 (trypsin-like) and β_5 (chymotrypsin-like) are the proteolytically active subunits. Proteasome inhibitors are designed to primarily inhibit β_5 .

Mechanism of action

Of the seven proteasome β subunits, only β_1 , β_3 and β_5 are proteolytically active (Figure 1.2b). Proteasome inhibitors are designed to target β_5 as it has been shown as the rate limiting protease for proteasomal protein turnover [40]. Bortezomib reversibly co-inhibits β_5 and β_1 subunits, whilst carfilzomib irreversibly binds to β_5 , with greater selectivity than bortezomib, and at higher doses binds to β_2 as well [40].

The precise downstream effects of β subunit proteasome inhibition are not fully understood, however the unfolded protein response (UPR), NF- κ B signalling, JNK signalling, apoptotic factors and p53 are thought to be involved in the anti-MM effects [41]. Specifically, the action of the UPR has been demonstrated as an important mechanism in the anti-MM effect of PIs. MM cells secrete large amounts of monoclonal protein, leading to the rapid accumulation of misfolded proteins within the endoplasmic-reticulum (ER) lumen. This results in heightened ER stress,

which is compensated by the UPR by reducing global protein translation and up-regulating UPS machinery [42]. Therefore, by inhibiting the proteasome, fewer ubiquitin tagged proteins are degraded and more misfolded proteins accumulate in the ER lumen. ER stress is then further increased, causing the UPR to switch from a homeostatic, pro-survival system to a pro-apoptotic pathway [41, 42].

Another important mechanism for PI is the attenuation of NF- κ B signalling. I κ B α , a specific endogenous inhibitor of the transcription factor NF κ B, is a protein degraded by the proteasome. Inhibition of the proteasome increases levels of I κ B α , thereby abolishing NF κ B signalling. NF κ B is a key transcription factor in many cancers, contributing to overall tumour growth and chemoresistance. NF κ B has been shown to promote tumour cell proliferation, anti-apoptotic and angiogenic factors [43].

1.4 Drug resistance in multiple myeloma

Although PIs are extremely effective at killing MM cells initially, long-term treatment inevitably results in a drug-resistant relapse. Drug resistance is one of the biggest barriers in the treatment of MM. Patients follow a pattern of peaks and troughs of treatment cycles, remission and relapse, until all therapies have little effect. In order to overcome resistance and increase overall survival of MM patients, the molecular mechanisms of resistance to proteasome inhibitors needs to be understood. This will aid in the design of novel therapies and inform better use of existing therapies. Previous studies on proteasome drug resistance have been performed and certain mechanisms and genes have been identified. For example, point mutations have been noted in the *PSMB5* gene (coding for the β 5 subunit of the proteasome), as well as and over-expression of the β 5 subunit [44]. Other upregulated genes have been identified, for example *ABCB1*, coding for P-glycoprotein, responsible for pumping various substrates out of the cell, often referred to as the multidrug resistant protein 1. *XBP1*, involved in the UPR, has been seen to be downregulated in PI resistance [44]. Although many genes have been identified to be differentially expressed in drug resistant MM, the mechanism is not fully elucidated and further research is

imperative in the progression of treatment for multiple myeloma. Identifying novel therapeutics effective against MM is of critical importance.

1.5 Transcriptomics, proteomics and epigenomics

It has been shown that changes in the genome, transcriptome, epigenome and proteome all contribute to acquired-drug resistance in myeloma. Therefore, to sufficiently investigate the multiple layers driving this development of resistance, a multi-omics approach must be employed.

1.5.1 DNA and the genome

The genome is the genetic material of an organism, it consists of deoxyribonucleic acid (DNA). DNA consists of two polynucleotide chains (or strands), running anti-parallel to each other, held together in a double helix structure by hydrogen bonds. Nucleotides are composed of a five-carbon sugar (deoxyribose for DNA), attached to one or more phosphate group (a single phosphate group in the case of DNA) and a nitrogenous base. Nucleotides are covalently linked to form an alternating sugar-phosphate backbone, with bases extending from each sugar towards the inside of the double helix. Nucleotides contain four different types of bases: adenine (A), cytosine (C), guanine (G) and thymine (T). The two DNA chains are held together by hydrogen bonds via complementary base pairing between the bases of the strands, A pairing with T and G pairing with C. Often sections of DNA are denoted as their sequence of A, C, T and Gs (in order reading from the 5' to 3' direction).

Every individual has approximately 6 billion base pairs of DNA per cell, which would amount to about 2 metres of DNA if laid end-to-end. The nucleus of a human cell is approximately 6 μm in diameter, therefore chromosomal DNA must be folded tightly to fit. DNA packaging is a complex task involving numerous specialised proteins. Negatively charged DNA is complexed with an octomer of positively charged proteins called histones to form nucleosomes. The histone core is made up of eight subunits, two copies of H2A, H2B, H3 and H4 subunits. DNA wraps tightly around the histone core 1.65 times. Linker DNA connects adjacent nucleosomes, to resemble ‘beads on a string’. Nucleosomes fold tightly to form 30nm chromatin fibre, which in turns forms loops averaging 300nm in length. This fibre is folded and compressed again to form fiber 250nm in width with loops of 700nm

in length. Tight coiling of this fiber forms the single chromatids of chromosomes [45, 46]. Human cells contain 23 pairs of chromosomes.

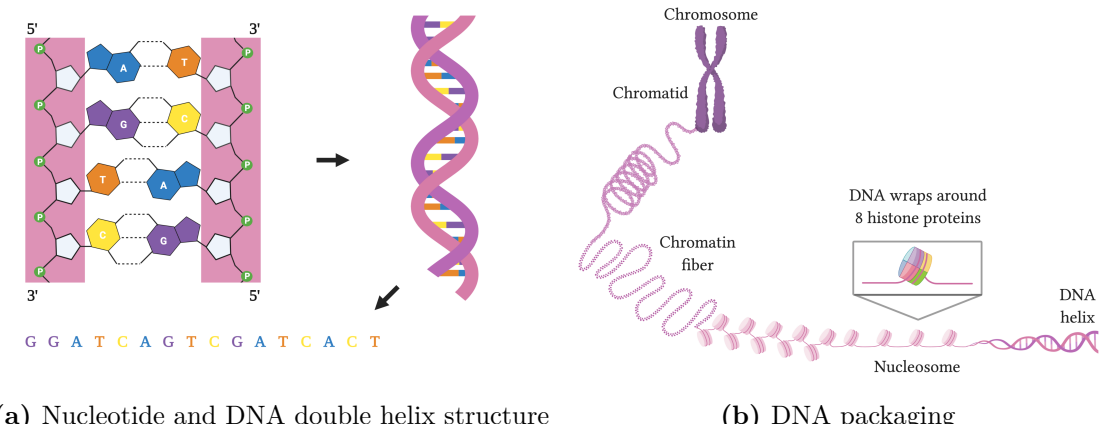


Figure 1.3: 1.3a shows the DNA nucleotides and the DNA double helix structure. DNA consists of two polynucleotide chains. Nucleotides are covalently linked to one another, forming a sugar-phosphate backbone. They contain one of four bases adenine (A), cytosine (C), guanine (G) and thymine (T). DNA strands are held together by hydrogen bonds between complementary base pairs, A pairing with T and G pairing with C. Sections of DNA are often read by their sequence of bases from the 5' direction to the 3' direction. 1.3b shows how chromosomal DNA is packaged in the cell. DNA wraps 1.65 times around an octomer of histone proteins, to form a structure called a nucleosome. Nucleosomes are linked by linker DNA to form a structure that resembles ‘beads on a string’. Nucleosomes fold to create chromatin fiber. This in turn forms loops and coils tighter and tighter until it makes up the single chromatids of chromosomes.

Created with BioRender.com.

The complete genome is made up of coding DNA (genes), non-coding DNA, as well as mitochondrial DNA and ribosomal DNA. An alteration in the nucleotide sequence of the genome is called a mutation. There are a number of types of mutations, including insertions, deletions, inversions, substitutions and duplications. A technique called whole genome sequencing (WGS) can be used to determine the sequence of nucleotides in an individual’s DNA and therefore it can be used to determine any variations in the genome.

1.5.2 The epigenome

Epigenetics is the study of any heritable phenotypic changes that do not involve alterations of the DNA sequence itself. These changes occur at the chromatin

level. Epigenetic changes include histone modifications, DNA methylation and chromatin remodelling.

DNA methylation is the addition of methyl groups to the C5 position of cytosines in DNA. This happens extensively at CpG sites (cytosine followed by a guanine). Stretches of DNA with a high CpG ratio (CpG islands) are often found in the promoter region of genes. Increased DNA methylation at CpG islands results in transcriptional silencing of those genes. Genome wide DNA methylation is often examined using DNA-methylation-seq or DNA methylation microarrays.

DNA wraps tightly around histones (section 1.5.1), they contribute to the tight packaging of DNA. Histone modifications are post-translational modifications. They include methylation, acetylation, phosphorylation, ubiquitination and sumoylation. Histone modifications affect transcriptional activity either by directly influencing the structure of chromatin and DNA accessibility or by regulating binding of effector molecules to ‘read’ histone marks to mediate downstream biological effects. Histone modifications also regulate DNA processes, such as repair, replication and recombination [47]. ChIP-seq can be used to investigate and measure various post-translational histone modifications.

Chromatin remodelling is the process of modifying chromatin architecture to regulate the accessibility of DNA. Gene expression is regulated by allowing certain gene regions better access to transcription machinery. This is achieved by ATP-dependent chromatin-remodelling complexes moving, ejecting or restructuring nucleosomes. ATAC-seq can be used to identify accessible DNA regions.

1.5.3 The transcriptome

Transcription is the first of many steps in gene expression. During transcription, the enzyme RNA polymerase reads a DNA sequence and produces an anti-parallel, complementary ribonucleic acid (RNA) strand. The transcriptome is the set of all RNA transcripts of an individual. RNA is a nucleic acid similar to DNA. Like DNA it has a sugar-phosphate backbone and 4 different types of bases attached to each

sugar. However, unlike DNA, RNA is single-stranded, it contains the sugar ribose in place of deoxyribose, and the nucleotide uracil (U) in place of thymine (T).

Despite the chemical differences between DNA and RNA, they are essentially written in the ‘same language’ and one-to-one mapping of nucleotides can be performed. Transcription begins with the unwinding and opening of a small part of the DNA double helix, so bases are exposed. One strand of DNA acts as a template and the RNA chain is formed by complementary base pairing with the template. RNA polymerases catalyse the reaction of forming phosphodiester bonds between nucleotides, forming the RNA chain. The RNA polymerase moves stepwise along the DNA chain, unwinding the chain just ahead exposing a new region of the template strand. Just behind the region where ribonucleotides are being added, the DNA helix reforms.

The genes in a cell’s DNA that specify the amino acid sequence and result in protein synthesis are called messenger RNA (mRNA) molecules. Genes that produce the RNA molecule itself are called non-coding RNAs, because they do not code for proteins. There are many other types of RNA, such as transfer RNA (tRNA), ribosomal RNA (rRNA) and micro RNA (miRNA).

Traditionally microarrays were used to measure gene expression. Now RNA-seq (outlined in section 1.5.5) is more commonly used study to gene expression and the transcriptome. Depending on the library preparation, different types of RNAs can be selected for or excluded, to study different RNA molecules.

1.5.4 The proteome

The proteome is the entire set of proteins that is or can be expressed by an organism. mRNAs are translated into protein molecules. mRNA is made up of only four different nucleotides, but proteins are made up of 20 amino acids, therefore a direct one-to-one function matching nucleotides to amino acids is impossible. Instead, the sequence of mRNA is read in groups of three consecutive nucleotides, called a codon. 4^3 gives a possible 64 combinations, therefore some combinations map to the same amino acid (many-to-one function), or signal to terminate translation of

the current protein, named a stop codon. This genetic code directs the translation from mRNA to protein. Translation takes place in the ribosome.

Codons on mRNA do not directly recognize their given amino acid, they require tRNA molecules that bind to both the codon on mRNA and the correct amino acid. tRNAs possess an anticodon, a set of three nucleotides complementary to a given codon. Firstly, tRNAs are coupled to their cognate amino acid. This reaction is catalysed by aminoacyl-tRNA synthetase (aaRS) enzymes. aaRSs attach amino acids to the 3' end of tRNA. Most cells have a specific aaRS for each amino acid. Once the tRNA is charged with the correct amino acid, the tRNA molecule binds to its complementary codon on mRNA. Subsequent aminoacylated tRNA molecules bind to mRNA codons. A polypeptide chain grows by stepwise addition of amino acid to the C-terminal end. The formation of the new peptide bond between amino acids releases the tRNA molecule. The peptide chain grows until a stop codon is reached and synthesis of the current protein is complete.

Protein translation is partly regulated by availability of mRNAs, but it also depends on other factors such as RNA silencing and post-transcriptional modifications. Proteins have a large array of functions, such as transporting small molecules, catalysing reactions, cell-cell signalling and providing structural support.

Proteomics is the study of the proteome. CyTOF and LC-MS/MS are techniques often employed to examine the proteome.

1.5.5 RNA-seq

Modern RNA sequencing (RNA-seq) implements next generation sequencing (NGS) technology to analyse RNA across the transcriptome of a biological sample and allows for the quantification of gene expression.

Bulk RNA-seq

Bulk RNA-seq measures the average expression across a sample. Creating a bulk RNA-seq library involves isolating RNA from a biological sample, filtering for a specific type of RNA (most commonly mRNA), fragmentation of RNA into



Figure 1.4: Outline of bulk RNA-seq library prep. Cells are lysed and RNA is extracted. The specific RNA of interest is selected and enriched, for example selecting for mRNA using polyA selection or ribo-depletion. The mRNA is fragmented into smaller pieces of RNA. First and second stranded cDNA are reverse transcribed from the RNA fragments using random primers. The ends of the cDNA are repaired and dAMP (dA) tails are added to the 3' end of the DNA. Adaptors are ligated to the 3' and 5' end of the cDNA. These adaptors contain complementary sequences that allow the fragments to hybridize to the flow cell during sequencing. Universal (P5/i5) and index (P7/i7) primers are added to the adaptor ligated DNA. The libraries are then amplified using PCR and cleaned-up, ready for sequencing.

fragments, reverse transcription of the fragments to generate a complementary DNA (cDNA) library, end repair and adaptor ligation of the cDNA library, followed by PCR amplification ready for sequencing.

Single-cell RNA-seq

Single-cell RNA-seq (scRNA-seq) measures gene expression for each individual cell across a population of cells and therefore provides information on clonal diversity that may be lost when pooling cells into bulk samples. Since its inception in 2009[48], there have been numerous scRNA-seq techniques, such as SMART-seq2[49], Drop-seq[50], STRT[51] and inDrops[52]. scRNA-seq library preparation shares many steps with bulk RNA-seq workflow, however preliminary steps are required to isolate single cells and barcode reads that originated from the cell.

For droplet-based scRNA-seq (dscRNA-seq) methods, single cells are isolated using microfluidic devices by individually encapsulating them in aqueous droplets contained in oil. Below, a dscRNA-seq method, Drop-seq, is outlined (Figure 1.5).

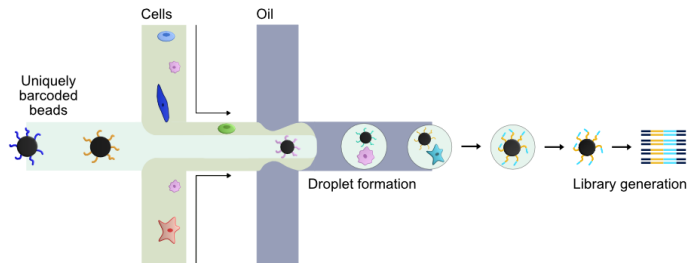


Figure 1.5: Outline of Drop-seq, a droplet-based scRNA-seq method. A microfluidic device combines two aqueous flows, one containing cells and the other containing barcoded primer beads suspended in lysis buffer. The two aqueous channels flow across an oil channel to form aqueous droplets surrounded by oil. Relatively few droplets contain both a cell and a bead. Following droplet formation, the cell is lysed and its mRNAs are released, which then hybridise to the primers on the bead surface. A reagent is added to break up the droplets and the beads are collected and washed. The mRNAs are reverse-transcribed into cDNAs, generating a set of “STAMPS” (single-cell transcriptomes attached to microparticles) and template switching is used to introduce a PCR handle. The barcoded STAMPS can then be amplified using PCR.

1.5.6 ATAC-seq

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1.5.7 Next generation sequencing

Next generation sequencing (NGS), differs from its predecessors in that it is highly scalable and massively parallel. With NGS you can rapidly sequence the entire genome if desired. It is quicker and cheaper than traditional Sanger sequencing, and progressed data output from the kilobase range up to potentially multiple terabases per run.

1.5.8 CyTOF

...

1.5.9 Liquid chromatography with tandem mass spectrometry

Liquid chromatography with tandem mass spectrometry (LC-MS/MS) based proteomics is a popular analytical technique to measure the protein abundance of

a sample. The general steps for LC-MS/MS-based proteomics include: cell lysis, protein extraction, protein digestion using an enzyme to cleave proteins into peptides, peptide purification, and analysis by mass spectrometry. The resultant data includes mass and charge (m/z) information and peak intensities. Software is then employed which performs database searches and calculates the most likely peptide for each peak. From this data, protein abundance can then be calculated and normalised.

LC-MS/MS-based proteomics can also be used to search for specific proteins within the proteome. For example, immobilized metal affinity chromatography (IMAC) can be used to enrich for phosphorylated peptides (phosphoproteomics), and anti-ubiquitin antibodies can be used to enrich for ubiquitinated peptides (ubiquitinomics).

1.6 Summary

1.7 Aims

This thesis aims to identify novel therapeutics with anti-MM properties, capable of overcoming acquired drug resistance in MM. Chapter 2 introduces the structure and function of aminoacyl-tRNA synthetases (aaRS). It then gives examples of aaRSs in disease and therapeutics that target aaRSs, focusing on the prolyl-tRNA synthetase inhibitor, Halofuginone (HF), and reviewing literature of HF's application, particularly in multiple myeloma. Chapter 3 outlines the methodology used in this work. Chapter 4 outlines the computational workflows generated to support this work and gives the results of a benchmark conducted to analyse the effectiveness of a computational pipeline developed. Chapter 5 investigates the use of ProRS inhibitors in PI-sensitive and PI-resistant MM cell lines. Bulk-RNA seq is employed and the transcriptional landscape following ProRS treatment is characterised. Chapter 6 investigates ProRS inhibitor treatment of primary BM samples from MM patients at the single cell level. Newly-diagnosed and relapsed MM patient tissue is assessed. Chapter 7 describes the changes seen in tRNA ... following ProRS inhibition.

2

Aminoacyl tRNA synthetases and Halofuginone

2.1 Introduction

Aminoacyl tRNA synthetases (aaRS) are a highly-conserved family of enzymes, responsible for “charging” tRNAs with their cognate amino acid. Human cytoplasmic aaRSs are either “free” as individual species or bound in a macromolecular complex, comprised of eight aaRSs and three auxiliary proteins, known as the multi-tRNA synthetase complex (MSC). On top of their canonical enzymatic role, aaRSs also engage in non-enzymatic functions in numerous pathways, including angiogenesis, inflammation and metabolism. Often species are released from the MSC to regulate these non-canonical activities. aaRSs have been shown to be involved in numerous diseases, including cancer. Initially, due to their high fidelity and complex evolution over millennia, aaRSs were seen as an attractive drug target for antimicrobials, to enable specifically targeting microbial aaRSs with minimal effects on human cells. The mechanism of action of Febrifugine (FF), a quinazoline alkaloid that has long been used as an antimalarial remedy, has recently been revealed; it acts as a competitive inhibitor of ProRS (part of the bifunctional glutamyl-prolyl-tRNA synthetase enzyme; EPRS), responsible for charging tRNA^{Pro} with proline. Although it has potent antimalarial effects, Febrifugine exhibits high liver and gastrointestinal (GI) toxicity, so cannot be used as a widespread drug, therefore several analogues of Febrifugine were developed in the hope of minimizing toxicity to the host’s cells.

One such analogue, Halofuginone (HF) was synthesized and was shown to have the most potent antimalarial properties of all the derivatives, with lower toxicity to the host than Febrifugine, but still some liver toxicity and GI side effects remain. Halofuginone has been applied to and showed promise in many other non-parasitic diseases too. It has received orphan drug status for scleroderma and HIV-Related Kaposi's Sarcoma. Recently, Halofuginone's application in various cancers has become of great interest, including but not limited to: metastatic brain tumours, bladder carcinomas, prostate cancer, renal carcinomas, hepatocellular carcinomas, lung cancer, breast cancer and multiple myeloma.

This review will introduce the structure and function of aminoacyl tRNA synthetases, provide an insight into their role in pathology and potential as therapeutic targets. EPRS1 and its inhibitors will be the primary focus, and exploring...

2.2 Function and structure of aminoacyl tRNA synthetases

Aminoacyl tRNA synthetases (aaRS) are an ancient family of ubiquitous enzymes, conserved across three major domains of life (but not present in viruses). They can be traced back prior to the “Last Universal Common Ancestor” (LUCA) [53]. aaRSs are essential for protein biosynthesis, and catalyse the first step in translation (see section 1.5.4). aaRSs catalyse the charging of tRNAs with their cognate amino acid. This is a two-step process. Firstly, aaRSs catalyse the formation of an aminoacyl-adenylate (activated amino acid) from their corresponding amino acid and an ATP molecule, releasing an inorganic pyrophosphate. Next, aaRSs catalyse the reaction between the aminoacyl-adenylate and their cognate tRNA to release an AMP molecule and generate an aminoacyl (charged)-tRNA, ready to be used by the ribosome to decode mRNA (see equation 2.1). An example of this process would be prolyl-tRNA synthetase (abbreviated to ProRS) charging tRNA^{pro} with proline.



Eukaryotes have 20 cytoplasmic aaRS and 20 nuclear-encoded mitochondrial aaRS. These are localised in distinct cellular compartments. aaRSs are often denoted by their one letter amino acid symbol, followed by ARS and either 1 (indicating they are cytoplasmic) or 2 (indicating they are mitochondrial), for example PARS1 for cytoplasmic ProRS. This review will focus on cytoplasmic aaRS enzymes. aaRS can be divided into two distinct classes based on the structure of the fold of their catalytic domains. Class I aaRS enzymes are functional monomers that contain a dinucleotide or Rossman fold (RF) of alternating alpha-helices and parallel beta-sheets. This fold is where ATP and amino-acid binding takes place and therefore facilitates the aminoacylation reaction. The active site of class I aaRS is marked by the signature motifs “HIGH” (His-Ile-Gly-His) and “KMSKS” (Lys-Met-Ser-Lys-Ser). Within the first half of the RF the HIGH motif helps to correctly position the adenine base of ATP and interacts with the phosphates. The second K of the KMSKS motif is thought to be involved in stabilising the transition state for the primary step of aminoacylation [54]. Amino acid recognition and binding takes place in the catalytic site when the KMSKS motif is open. The KMSKS loop closes after the aaRS binds ATP and the aminoacyl-adenylate is formed [55].

Class II aaRS enzymes are functional dimers or tetramers with an uncommon catalytic core, comprising seven anti-parallel beta-sheets, flanked by alpha-helices. Class II aaRS enzymes are defined by three conserved sequence motifs. Motif 1 is located at the interface of the dimer and enables oligomerization. Motifs 2 and 3 comprise part of the aminoacylation active site and facilitate amino acid/ ATP binding and adenylate formation. Motif 3 binds ATP, and motif 2 is involved in coupling ATP and the amino acid and then transferring the amino acid to the 3'-tRNA [55]. The distinct active-site structures of class I and II enzymes confer markedly different binding mechanisms for the aminoacylation reaction. For example, class I aaRSs bind the tRNA acceptor stem via the minor groove side and bind ATP in an extended conformation, whilst class II aaRSs bind the tRNA acceptor stem from the major groove side and bind ATP in a bent conformation. The two classes of aaRSs split the twenty amino acids into two groups. Val, Leu, Ile, Met,

Glu, Gln, Trp, Tyr, Arg and Cys are activated by their cognate class I aaRS; and Gly, Pro, Ala, Thr, Ser, Hist, Asp, Asm, Lys and Phe are activated by their cognate class II aaRS. Class I and class II can be further divided into different sub-groups, however that is beyond the scope of this review. The structural diversity of aaRSs is likely attributable to the need to exclude similar non-cognate amino acids and to discriminate the correct tRNA isoacceptor.

Both class I and class II aaRSs are multi-domain proteins- in addition to their catalytic domains, they include other domains such as their anti-codon recognition domain or an editing domain. The editing domain found in some aaRSs is to ensure that the essential step of aminoacylation in protein biosynthesis is as accurate as possible, so incorrect amino acids can be removed from aminoacyl-adenylates or mischarged tRNAs [55]. Theoretically, it was estimated that mistranslation rate should be approximately 1 in 200 for amino acids differing by just a methyl group (such as valine and isoleucine) [56], however in-vivo work demonstrated that the error frequency is closer to 3 in 10,000 (approximately 1 in 3000) [57]. This suggested the existence of proof-reading capabilities of aaRSs, to account for the difference between observed and predicted error rates. Editing capability has since been shown to be of high functional importance to some aaRSs. For example, a study in mice in which there was a missense in the editing domain of AlaRS. The impaired proof reading activity of the enzyme lead to an accumulation of misfolded proteins, resulting in the activation of the unfolded protein response and substantial neurodegeneration [58]. Not all aaRSs possess editing activity, only about half do, however the high specificity of the active site of those aaRSs is enough to alleviate proofreading need.

2.2.1 Multi-tRNA synthetase complexes

Higher eukaryotes contain macromolecular complexes, which consist of nine enzymes and three auxiliary proteins, known as multi-tRNA synthetase complexes (MSC). The 11 cytoplasmic aaRSs not located in the MSC remain free as individual species. The nine cytoplasmic aaRS enzymes of the MSC are GluRS (EARS1), ProRS (PARS1), IsoRS (IARS1), MetRS (MARS1), GlnRS (QARS1), LysRS

(KARS1), ArgRS (RARS1), AspRS (DARS1), LeuRS (LARS1). GluRS and ProRS are covalently fused via triple repeats of WHEP domains to form a bifunctional enzyme, EPRS1. The non-enzyme component of the MSC consists of three aminoacyl-tRNA synthetase-interacting multifunctional proteins (AIMP), AIMP1, AIMP2 and AIMP3. Human MSCs contain more class II aaRS enzymes than other species, namely DARS1, KARS1, and PARS1, they also contain more auxiliary proteins. Human MSC components have several additional domains or motifs (Figure 2.1A), for instance GST-homology domains in EPRS, MetRS, AIMP1 and 2, and WHEP domains in EPRS and MetRS [59–61].

The structure of human MSC has not been fully elucidated, however some sub MSC-complex structures have been revealed. LysRS forms a homodimer and is anchored to the N-terminal peptide region of AIMP2 within the main body of the MSC. MetRS, AIMP3, EPRS1 and AIMP2 are compactly linked through their GST-homology domains. ArgRS, GlnRS and AIMP1 assemble into a heterotrimeric complex [59–61]. A proposed bisymmetrical model of the human MSC, via homodimerization of AspRS and ProRS, is shown in Figure 2.1B, based on subcomplex and interaction data [62–64]. This hypothesis proposes that the MSC is a super-complex of two identical, symmetrically arranged subunits (symmetrical along the y-axis in Figure 2.1B), each containing one copy of the constituent elements, except for LysRS which is present as a dimer in each subunit.

The function of the MSC was originally thought to be to increase efficiency of protein biosynthesis by localising aaRSs. Another proposed function of the MSC was to increase stability of its components. It has been shown using systematic depletion analysis that some of the components are in fact intrinsically less stable in isolation and dependent on their neighbours for stability [65]. More recently, examples have emerged where the MSC seems to work as a ‘molecular reservoir’ which can control the release of its components. The release of components from the MSC has been linked to numerous non-canonical pathways, including cell signalling, metabolism, inflammation and angiogenesis.

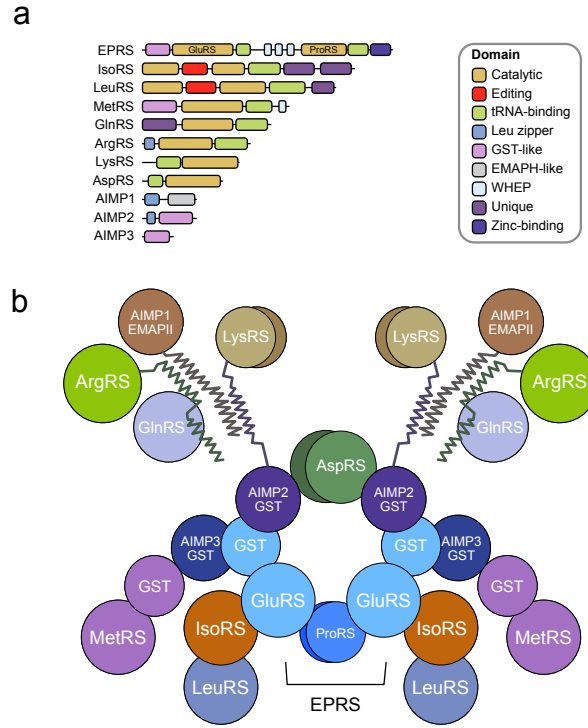


Figure 2.1: The human multi-tRNA synthetase (MSC) and its components. **A)** The domains of the aminoacyl tRNA synthetases and auxiliary proteins (AIMP1, 2 and 3) making up the human multi-tRNA synthetase (MSC). The bifunctional enzyme EPRS1 is made up of the class 1 enzyme GluRS and class II enzyme ProRS (dimer) covalently linked by three WHEP domains. **B)** Cartoon representation of a proposed bisymmetrical model structure of the human multi-tRNA synthetase complex (MSC). An adaption of a figure created by Myung Hee Kim and Sunghoon Kim [61].

Higher eukaryotes usually have extra-domains at the N- or C- terminus of aaRS enzymes compared with lower eukaryotes and prokaryotes, which may partly contribute to MSC assembly. Most human cytoplasmic aaRS enzymes have at least one new sequence extension or domain, most of which are dispensable for enzymatic activity, suggesting they may contribute to the non-canonical roles of aaRS. Additionally, aaRSs are often found in the nucleus of cells, where protein biosynthesis does not occur. The additional evolutionary complexity in human aaRSs and MSC seems to explain the increased physiological complexity and their functionality in non-enzymatic processes.

Examples of non-canonical MSC functionality include—LARS1 translocating from the MSC to lysosomes, facilitating mTORC1 activation [66]; KARS1 translo-

cating to the nucleus upon immune activation and activating MITF-dependent gene expression in mast cells [67]. Another example is EPRS1 release from the MSC in myeloid cells upon IFN- γ stimulation [68]. IFN- γ induces a network of kinase events (Cdk5, mTORC1 and S6K1 activation) which causes a two-step phosphorylation of two serines in the linker region of human EPRS, and causes its release from the MSC. EPRS1 combines with other proteins (namely NSAP1, L13a and GAPDH) to form the cytosolic IFN- γ activated inhibitor of translation (GAIT) complex, which represses translation of numerous inflammatory-related transcripts, including VEGFA and ceruloplasmin [69].

In addition to the enzymatic components of the MSC, the auxiliary proteins AIMP1, 2 and 3 are also involved in fundamental biological processes. AIMPs exhibit non-canonical functions aside from their roles as scaffolds in the MSC. AIMPs have been linked to numerous biological processes, including involvement in immune regulation, nervous system functions, viral replication, genome stability, angiogenesis, and cancer. AIMP1 interacts with RARS1 and facilitates incoming tRNA substrates to its catalytic site to enhance its enzymatic activity [70]. In addition to improving amino-acyl synthetase activity, secreted AIMP1 has also been shown to be involved in angiogenesis, inflammation induction, wound closure, and maintaining glucose homeostasis [71]. TGF- β and the DNA damage response have both been shown to cause phosphorylation of AIMP2 and disassociation from the MSC. Released AIMP2 has shown to act as a pro-apoptotic mediator and tumorigenesis suppressor via various pathways [72]. AIMP3 largely interacts with MARS1, and under conditions initiating the DNA damage response, MARS1 undergoes a conformational change that releases AIMP3 from the MSC [73]. Released AIMP3 acts as a tumour suppressor, translocating to the nucleus and upregulating expression of the tumour suppressor gene p53.

The functional and structural complexity of the MSC is still being revealed. The canonical and non-canonical functionality of MSC components promises an unexplored rich source of potential therapeutic targets, but also lends itself to associated pathology.

2.3 aaRSs in disease

With the diversity of functionality in human aaRSs comes an increase in functionality that can be associated pathologically with human disease. Structural and functional variations in aaRSSs' enzymatic and non-enzymatic activities have been linked to various human diseases. Changes in gene expression, copy number, mutations and genetic variations of aaRSs have been documented in relation to disease [55].

Charcot Marie Tooth (CMT) is a genetically and clinically-presenting heterogeneous group of hereditary peripheral neuropathies. CMT is characterised by progressive degeneration of distal sensory and motor neuron function [74]. Six aaRSs have been linked to CMT through dominant mono-allelic mutations, including GARS1 and YARS1, which are among numerous genetic-loci to have been linked causally to CMT. Drosophila models of CMT have demonstrated that CMT-causing YARS1 mutations lead to a conformational change in YARS1, leading to aberrant interactions with transcriptional regulators in the cell nucleus and aberrant expression of certain transcription factors [75].

aaRSs have also been implicated in autoimmune diseases. “Anti-synthetase syndrome” (ASS) is a heterogeneous group of autoimmune diseases, including interstitial lung disease (ILD), arthritis, idiopathic inflammatory myopathies, myositis and Reynaud’s phenomenon. Autoimmune antibodies against histidyl-, threonyl-, alanyl-, isoleucyl-, phenylalanyl-, glycyl-, tyrosyl-, asparaginyl-tRNA synthetase have been found in approximately 30% of all autoimmune patients. Dysregulation of aaRS has also been noted in other autoimmune diseases, for example multiple sclerosis and immune thrombocytopenia [76].

aaRSs have been linked to viral and bacterial infection. For example, it has been shown that viral infection leads to the phosphorylation of EPRS and dissociation from the MSC, ultimately blocking PCBP2-mediated mitochondrial antiviral signalling (MAVS) ubiquitination and inhibiting viral replication [77]. Additionally, HIV-1 infection leads to KRS release from the MSC, which is partially transported to the nucleus. Blocking this release reduced the infectivity of progeny virions, implying that HIV-1 utilizes a dynamic MSC for enhanced viral replication [78].

WARS1 was shown to be increased approximately 27-fold in sepsis patients with a bacterial infection compared with healthy controls. Following a range of infections by various pathogens, host monocytes were shown to rapidly secret WRS. The secreted WRS increased cell surface levels of CD40, CD80 and CD86, markers of macrophage activation [79].

2.3.1 aaRSs in cancer

A growing number of studies have implicated aaRSs and MSC components in tumorigenesis. Firstly, aaRS enzymatic activity is essential to sustain tumour growth. In cancer metabolism, biosynthesis of aminoacyl-tRNAs has been shown to be highly up-regulated [hu2013heterogeneity]. In cancer, we see often see dramatic rapid cell growth, this demands an intense increase in overall protein synthesis. To keep up with this demand, the canonical aminoacylation role of aaRSs is crucial as the first step in protein synthesis.

On top of the enzymatic role of aaRSs, their non-canonical functionality has also been associated with both promoting and inhibiting cancer. The hallmarks of cancer- enhanced growth signalling and proliferation, vascularization, metastasis, altered metabolism, and immune/tumour microenvironment invasion, all have links to tRNA synthetase function. Cancer cells require enhanced growth signalling and proliferation to maintain their rapid growth beyond the capacity of normal cells, several aaRSs have been linked to this aberrant growth signalling. GlyRS has been shown to be integral for cancer-promoting neddylation to occur, and reduced MetRS expression resulted in reduced tumorigenicity in p16INK4a-negative breast cancer cells *in vivo* [80–82]. For tumours to grow and metastasize they need to hijack existing vasculature to get blood flow to growing area, or make new vessels by promoting angiogenesis. Endothelial cells (EC) exposed to TNF- α or VEGF secrete ThrRS. ThrRS promotes EC migration and angiogenesis. Inhibition of ThrRS was shown to inhibit angiogenesis, with and without inducing the uncharged tRNA response [83, 84]. LysRS has been shown to support metastasis by increasing migration. Following phosphorylation by the MAPK pathway, LysRS binds to the 67kDa

membrane bound laminin receptor protein (67LR), preventing its degradation and sustaining laminin-dependent migration. Once bound to LysRS, 67LR also binds integrin $\alpha 6\beta 1$, which initiates ERK and paxillin signalling, increasing migration by altering cell-cell and cell-ECM adhesion.

aaRSs have also been linked to altering metabolism in cancer. To make rampant growth feasible, cancer cells adjust metabolism to meet energy demands and provide building blocks for biosynthesis. LeuRS activates the mTORC1 pathway, which controls translation and autophagy. Cancer cells utilize the mTORC1 pathway to proliferate more efficiently. The mTORC1 pathway also causes phosphorylation of EPRS and the release of it from the MSC. In adipocytes, released EPRS interacts with FATP1 and directs it to the plasma membrane. Inhibition of FATP1 leads to increased cell viability in breast cancer cell lines, and its expression correlates with decreased patient survival in triple negative breast cancer [85].

2.3.2 AIMP_s in cancer

As well as the association between aaRSs and cancer, AIMP_s have also been shown to play a role in signalling pathways relevant to numerous cancers. The MSC-bound aaRSs seem to predominantly promote tumorigenic functions when released from the MSC. In contrast, the AIMP_s bound with them seem to have more tumour-suppressive effects. AIMP2 has been shown to be a potent tumour suppressor, working via key regulators in the p53, c-Myc, Wnt, TGF- β and TNF- α signalling pathways. Loss of a single allele of AIMP2 in mice resulted in a far higher susceptibility to tumour formation [86]. AIMP1 has also demonstrated tumour-suppressive effects. In mouse xenograft models, administered AIMP1 was found to reduce tumour volume [87, 88]. AIMP1 has been shown to induce apoptosis of endothelial cells, such that it suppresses tumour vascularization [89]; it also stimulates anti-tumour immune responses, for example activating natural killer (NK) cells via macrophages, dramatically reducing lung metastasis of melanoma cells [90]. AIMP3 activates the tumour-suppressor gene p53 following DNA damage

or oncogenic stress. Loss of an AIMP3 allele results in higher susceptibility to spontaneous tumour formation [91].

2.4 aaRSs as therapeutic targets

aaRSs are considered very attractive drug-targets. Initially the interest in aaRSs as therapeutic targets arose with the detection of differences between prokaryotic and eukaryotic aaRSs. Thus, enabling specific targeting of microbial aaRSs with minimal effect on the homologous human aaRSs, making aaRS inhibitors attractive anti-microbial candidates.

2.4.1 Antibacterials and antifungals

In the 1990s, Mupirocin (brand name Bactroban) was approved as an antibiotic for the topical treatment of bacterial skin infections. Mupirocin selectively inhibits bacterial IleRS, by simultaneously occupying the isoleucine and AMP binding sites and inhibiting aminoacylation [92]. Mupirocin has shown high selectivity for bacterial IleRS over mammalian IleRS (greater than 8000 fold) [93]. This conferred selectivity seems to be due to only a two-amino acid residue difference in the active site of eukaryotic and prokaryotic IleRS [94]. Another example is Kerydin (Tavaborole or AN2690), an anti-fungal used to treat onychomycosis, which targets fungal LeuRS. Kerydin is a boron-containing compound (Benzoxaborole) and the first drug to target an aaRS editing site. The boron atom of Kerydin binds to the terminal tRNA_{Leu} ribose, trapping tRNA_{Leu} in the editing site and causing a non-productive enzyme conformation, which inhibits protein biosynthesis. [[REF]]

2.4.2 Anti-parasitics

On top of the success of the druggability of aaRS enzymes for bacterial and fungal infections, aaRSs have also showed promise as an anti-parasitic target. Much like cancer, parasites are extremely reliant on protein synthesis to keep up with rapid cell growth and continuous proliferation, so are likely to be more sensitive to disruptions to aminoacylation. Additionally, the evolutionary distance between parasitic aaRSs

and human aaRSs is quite large, in fact several parasites have bacterial-like protein translation pathways, not shared by humans [95]. Numerous aaRSs have previously shown promise as targets for anti-parasitic agents. Several naturally occurring compounds target the AsnRS site of parasites, such as *Brugia malayia*, a nematode which causes Lympathic Filariasis. *Trypanosoma brucei* has also been shown to be susceptible to aaRS inhibition, for example by Benzoxaboroles targeting LeuRS, or by Aminoquinoles and Benzimidazoles targeting MetRS. The parasite *Plasmodium falciparum* has been shown to be affected by numerous aaRS inhibitors, including Mupirocin, Cladosporin and Febrifugine derivatives.

2.4.3 Febrifugine and its derivatives

Dichroa febrifuga has been used for centuries in Chinese medicine as an antimalarial remedy, it is considered one of the 50 fundamental herbs. In 1948, two quinazoline alkaloids, named Febrifugine (FF) and Isofebrifugine, were first isolated from the plant Dichroa febrifuga (Figure 2.2) [96], as part of a directive to find new anti-malarials from plant sources. Although Febrifugine has excellent anti-parasitic activity, it also has strong liver and gastrointestinal toxicity, limiting its use as a widespread therapeutic. This motivated the generation of Febrifugine derivatives with the hope of reducing off-target toxicity. The medical applications of the long-used traditional anti-parasitic agent Febrifugine and its derivatives have recently attracted much attention. Febrifugine derivatives have been used to treat malaria, fibrosis, inflammatory disease and cancer.

2.4.4 Halofuginone

One such analogue, a synthetic racemic halogenated derivative of Febrifugine, Halofuginone (HF; Figure 2.2), was synthesized in 1967 by American Cyanamid Company [97]. Halofuginone was found to have the most potent anti-malarial activity of the FF analogues in vitro and affected all three stages of *P. falciparum* (ring stages, trophozoites and schizonts) with equal speed, unlike many other chemicals with antimalarial effects. The addition of bromine on the quinazoline ring in HF was

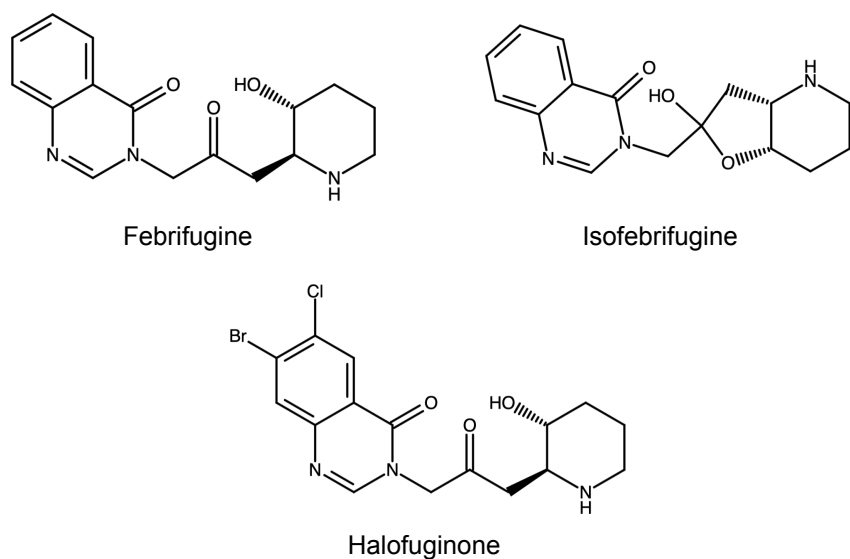


Figure 2.2: Chemical structures of prolyl-tRNA synthetase (ProRS/PARS1) inhibitors. Febrifugine and isofebrifugine were first isolated from *Dichroa febrifuga* in 1948. Halofuginone is a derivative of Febrifugine, first synthesized in 1967.

found not affect the antimalarial properties of FF, whilst lowering the cytotoxicity for host cells compared to FF. However, HF does still demonstrate some toxicity to the liver, among other side effects, including diarrhoea and vomiting [98]. In an attempt to reduce the side effects of HF and increase the therapeutic window, trans-enantiomers (2R,3S / +) and (2S,3R / -) of HF have been prepared. Although (-)-HF was found to have lower toxicity than its optical antipode, it was also found to be less efficacious than (+)-HF [99, 100]. This suggests that the biological activity and mammalian toxicity of HF reside with the same enantiomer, therefore there is no advantage to using a specific enantiomer over the racemic mixture.

Recently Halofuginone has been researched extensively in association with its applications to non-parasitic diseases. HF is FDA-approved as a feed additive for poultry to prevent coccidiosis from the protozoa *coccidian*. HF has also received orphan drug status for scleroderma and Duchenne muscular dystrophy. HF has undergone clinical trials as a potential therapeutic in a number of conditions, including cancer [101, 102].

Halofuginone's mechanism of action

Until the last decade, the mechanism of action of Halofuginone was unclear, until two papers from 2009 [103] and 2012 [104] elucidated HF's target and downstream effects. Mouse T_H17 cells were treated with HF or an inactive derivative, MAZ1310, for 3 or 6 hours and microarray analyses were performed. ATF4 target genes were found to be activated by HF expression, including Asns, Chop, eIF4Ebp, Gpt2, as well as amino acid transport genes, such as Slc6a9 and Slc7a3, both patterns that correspond with activation of the amino acid starvation response (AAR). Using western blots, the group also showed that GCN2 autophosphorylation was activated by HF treatment, further indicating HF activates the AAR pathway. This effect was not limited to T_H17 cells, the AAR pathway was also activated by HF treatment in fibroblasts and epithelial cells [103]. However, this paper did not reveal how HF activated the AAR.

In 2012, the group demonstrated that HF and FF activate the AAR by competing with proline as potent inhibitors of tRNA^{Pro} charging activity of EPRS1. Rabbit reticulocyte lysate (RRL) was used as an in-vitro translation system. Following supplementation with excess amino acids, only proline was shown to restore translation inhibited by HF in the RRL system. Moreover, HF-derivatives that were shown to be inactive in functional cell-based assays, such as MAZ1320, also lacked activity in the RRL assay. Together, this suggests that HF functionality is linked to blocking proline utilization. To further demonstrate that HF and FF affect proline utilization, the group synthesized DNAs encoding two epitope-tagged polypeptides, one encoding a proline-dipeptide (ProPep), the second encoding a proline-free peptide (NoProPep). HF and FF treatment prevented translation of ProPep, but had no effect on NoProPep translation [104].

Next the group investigated the effect of HF on prolyl-tRNA charging and the bifunctional enzyme EPRS1 (comprised of GluRS and ProRS fused together). The addition of EPRS from purified-rat-liver reduced the sensitivity of RRL to HF. They then investigated the inverse using siRNA-mediated knockdown to reduce EPRS levels in lung fibroblasts. Lung fibroblasts have high levels of EPRS endogenously,

so are quite resistant to HF treatment. The reduction of EPRS levels sensitized the cells to HF treatment and AAR pathway activation—GCN2 autophosphorylation was induced as well as ATF4 response genes, such as CHOP and ASNS. Together this established for the first time that EPRS is a critical target of inhibition for HF and FF, through which the compounds elicit AAR activation. The group demonstrated that HF inhibits EPRS in a competitive fashion with proline at the prolyl-tRNA synthetase active site. HF binding is an ATP-dependent process. ATP directly locks onto and positions HF onto human ProRS so that one part of HF mimics bound proline and the other mimics the 3' end of bound tRNA^{Pro} [105]. Excess proline addition was shown to abrogate AAR activation and reversed the biological effects of HF [104].

By binding the active site of ProRS, HF blocks proline from binding and inhibits ProRS enzymatic activity. This results in an intracellular build-up of unaminoacylated (uncharged) tRNA^{Pro}s, mimicking the cellular state of proline deficiency, thus triggering the amino acid starvation response. Uncharged tRNAs bind to the protein kinase GCN2 and stimulates its dimerization and autophosphorylation. Activated GCN2 phosphorylates eukaryotic translation initiation factor 2A (eIF2 α), this leads to a reduction in most protein synthesis, whilst increasing translation of ATF4. ATF4 is a transcription factor of the cAMP response element binding protein (CREB) and induces the expression of many genes involved in the integrated stress response (for example DDIT3/CHOP), amino acid synthetases and transporters, aminoacyl tRNA synthetases, and autophagy regulators (figure 2.3) [103, 106].

Halofuginone's downstream signalling pathways

— TO FILL in — To complete...

2.5 Halofuginone and cancer

HF has exhibited anti-cancer effects in numerous studies and different cancers, including metastatic brain tumours, bladder carcinomas, prostate cancer, renal carcinomas, pheochromocytomas, hepatocellular carcinomas, esophageal squamous



Figure 2.3: A diagram of Halofuginone’s (HF) mechanism of action and relationship with the amino acid starvation response (AAR). HF binding with the catalytic site of prolyl-tRNA synthetase (ProRS) of the bifunctional aminoacyl-tRNA synthetase, EPRS, causes an accumulation of uncharged tRNAs, mimicking the same cellular environment as if the cell were amino acid deprived. Uncharged tRNAs bind to the cellular sensor GCN2 and cause it to autophosphorylate and activate. Activated GCN2 then phosphorylates eIF2- α . eIF2- α -p reduces global protein synthesis, except for mRNAs containing an upstream ORF cluster in their 5’ untranslated region (UTR) which are efficiently translated upon eIF2-alpha phosphorylation [106], including the transcription factor ATF4. Upregulated ATF4 results in increased expression of many genes involved in stress responses (e.g. CHOP/DDIT3), amino acid metabolism, amino acid synthetases (e.g. ASNS) and aminoacyl tRNA synthetases. Figure created with Biorender.com

carcinomas, lung cancer and breast cancer [107–115]. HF has been shown to exert anti-cancer effects in numerous manners, including reducing tumour growth, reducing angiogenesis, activating autophagy and apoptosis, and disrupting the collagen network of tumours, among other mechanisms.

2.5.1 Halofuginone and multiple myeloma

As mentioned in section 1.3, multiple myeloma (MM) is an incurable cancer of plasma cells. Drug resistance is a massive problem in MM, with patients becoming resistant to drugs they’ve previously been treated with, cycling through treatment and relapse cycles. Therefore, identifying novel therapeutics for the treatment of MM is of critical importance.

Following the success of HF treatment in numerous preclinical cancer studies and the phase II study of HIV-related Kaposi's sarcoma [116], Leiba et al. (2012) investigated the treatment of HF in multiple myeloma, both in-vitro and in-vivo [117]. 17 MM cell lines were treated for 48 hours with a range of HF concentrations. HF was shown to induce a reduction in cell viability in a dose-dependent manner across all 17 MM cell lines, with an IC₅₀ of approximately 100nM in most cell lines. The effect of HF on primary cells was then investigated. CD138+ cells from BM samples from five MM patients and PBMCs from two healthy donors were treated with a range of HF concentrations. A greater dose-dependent reduction in cell viability was seen in the primary MM cells compared to the healthy PBMCs, with an IC₅₀ ranging from 101-253nM for the MM cells. Demonstrating that, at this concentration range, HF specifically inhibits the viability of MM cells while having no significant effect on normal cells; this also gave a therapeutic window for HF in MM. Next, the group demonstrated that HF induces apoptosis in MM—HF treatment triggered caspase 3, 8 and 9 activities in MM cell lines in a dose dependent manner; it increased the quantity of apoptotic cells (Annexin V-FITC apoptosis assay); it caused an accumulation of cells in sub G1 phase of the cell cycle, associated with DNA fragmentation; and it elevated expression of the heat shock protein Hsp-90. They also showed that exogenous IL-6 and IGF-1, which are central for MM growth and survival, did not rescue HF-induced cytotoxic effects on MM cell lines, indicating that paracrine MM cell growth and the BM environment are unlikely to reverse the biological effects of HF.

The group also exhibited the anti-MM effects of HF in-vivo, using in a xenograft model of SCID mice injected with MM.1S cells. Once tumours reached sufficient size, mice were treated with either PBS or HF for five days a week for the duration of the experiment. HF treatment was found to inhibit tumour growth and increase overall survival compared to the control mice.

Synergy of HF with existing MM drugs was investigated [117]. Cells were cultured for 48 hours with HF (25, 50 and 100nM) in combination with 5nM Bortezomib, 25uM lenalidomide, 500nM dexamethasone, or 500nM doxorubicin. Cells were

cultured for 24 and 72 hours with HF (25, 50 and 100nM) in combination with 10uM melphalan. CalcuSyn software (Biosoft, Ferguson, MO, USA) was used to evaluate synergism. Lenalidomide, dexamethasone and doxorubicin were found to be synergistic or additive with HF in all MM cell lines. HF showed moderate antagonism in combination with Bortezomib. However, only one concentration value was used for each of the established MM agents, and a small range of concentrations were used for HF treatment. A larger range of concentrations of both HF and the other agents would be required to gain a greater insight into the drugs interactions with one another.

From this study, it is clear HF is effective against MM, and could show promise as a potential line of therapy. However, Leiba et al. (2012) did not show how HF was exerting its effect, or if the AAR was activated. AAR activation results in upregulated levels of the transcription factor ATF4. It would be interesting to explore the transcriptional landscape of MM cells and the tumour microenvironment following HF treatment, to see how AAR activation affects this. The group used MM cell lines, mouse models and primary BM samples from MM patients. The primary BM samples were compared against healthy donors' PBMCs and not MM patients' own non-myeloma cells, so only limited conclusions can be drawn about HF's specificity for MM cells over normal cells. Moreover, MM cells are known to interact substantially with their microenvironment. In this study, the cells were studied in isolation, the effect of HF on the immune microenvironment was not investigated. The paper states that BM samples were taken from five MM patients, however it is not stated what stage of disease progression the patients were in, or if they were a mixed group of patients in various disease stages. Therefore, conclusions cannot be drawn whether HF is equally or preferentially effective against newly-diagnosed and relapsed MM patients.

2.6 Discussion and future directions

Considering aaRSs are such a highly conserved and ancient family of enzymes, it is surprising how much about their structure and function is still unknown. Concerted

efforts are being made to elucidate the full structure of the MSC, with this knowledge, the full functionality of the MSC might be more clearly understood too. More non-canonical functions of aaRSs are being revealed, and with it, associated pathologies. This also presents unexplored potential for aaRS therapeutics.

Drugs targeting aaRSs have showed effectiveness in a wide range of clinical settings including numerous cancers. aaRS inhibitors are an exciting drug in the application of cancer, as tumours often have such a high protein biosynthesis burden, especially MM with the overproduction of large quantities of paraprotein. Therefore, targeting the first step in protein biosynthesis is highly attractive. Halofuginone, an inhibitor of the ProRS active site of EPRS, has been researched extensively in recent years. HF has shown anti-MM effects against MM cell lines, in-vivo mouse models and primary patient BM samples.

HF has been shown to activate the amino acid response in rabbit reticulocyte lysate and lung fibroblasts. HF's effects are abrogated by excess proline supplementation. HF also demonstrates substantial liver and GI toxicity, so may not be able to be used as a widespread drug in cases without orphan drug status. A ProRS inhibitor which was less toxic, with a wider therapeutic range and whose effects could not be overcome by excess proline would likely be a much more effective anti-cancer agent therapeutically.

The mechanism of action of HF in MM has not been clarified and the transcriptional changes of HF treatment in MM have not yet been described. MM patient cells have only been studied in isolation—MM patients' immune microenvironment following HF treatment has not been investigated. The specificity of HF for MM cells has also not been demonstrated fully. MM patients' transcriptome, epigenome and genome evolves greatly during disease progression, and the changes are not limited to MM cells and plasma cells. Therefore, the effect of HF treatment on MM cells must be compared to the patient's own non-myeloma cells to be able to assess specificity, rather than healthy donor's PBMCs, as they are so different from the PBMCs of MM patients. It would be hugely pertinent to employ single-cell sequencing of BM samples following HF treatment. scRNA-seq could capture transcriptional changes

of the MM cells and their surrounding immune microenvironment, and would also allow composition analysis, so that proportional changes of each cell type could be quantified, and HF's specificity for MM cells evaluated.

So far, only BM samples from MM patients of unknown disease progression have been treated with HF. Therefore, it cannot be commented on whether HF works on BM samples from relapsed patients. Various MM cell lines have been treated with HF, however the resistant variants were resistant to traditional chemotherapy agents (Doxorubicin, Mitoxantrone and Melphalan) or Dexamethasone. MM is conventionally treated with a three-drug regimen (as discussed in section 1.3.4), comprising a corticosteroid (e.g. Dexamethasone), a proteasome inhibitor (PI; e.g. Bortezomib or Carfilzomib), and an immunomodulatory drug (IMiD; e.g. Lenalidomide). MM patients eventually accrue resistance to all three drugs in the regimen. HF's anti-MM effects were not demonstrated against either PI-resistant or IMiD-resistant cell lines or relapsed patients. It would be interesting to see if HF's anti-MM effects are maintained in PI-resistant MM cells. Proteasome inhibition leads to an accumulation of misfolded, damaged or unneeded proteins, which activates the unfolded protein response (UPR), which in part contributes to the anti-MM effects of PIs. The UPR and AAR share many joint effectors, such as ATF4 and CHOP, both contributing to ER stress and leading to apoptosis. As there is some overlap with HF's and PIs' mechanisms of action, and that Leiba et al. (2012) reported mild antagonism of HF in combination with the PI bortezomib [117], you may not expect HF to be effective against PI-resistant MM. Many new MM agents are approved for relapsed MM initially, rather than as first-line treatments; therefore, it would be critical for ProRS inhibitors' success in myeloma that their anti-MM effects extend to PI-resistant and relapsed MM. This is crucial question that must be answered.

Another thing that would be of interest to investigate, is the interaction of HF treatment and EPRS's non-canonical functionality. Following IFN- γ stimulation, the EPRS dissociates from the human MSC to participate in the GAIT complex in myeloid cells. The GAIT complex represses translation of inflammatory-related genes, including VEGFA. It would be interesting to see if HF has any effect on

this non-canonical function of EPRS, or in fact, if the GAIT complex impacts HF treatment in myeloid cells.

Summing up paragraph... aaRSs are a very exciting area of study, particularly as drug targets. So far, the application of aaRS inhibitors in disease has only scratched the surface of their potential as therapeutics. Much more research is required to fully explore the...

3

Methods

3.1 Cell culture

3.1.1 AMO-1 cells

AMO-1 cells, plasma cells from a 64-year old female myeloma patient, were used as a model cell-line for multiple myeloma. Proteasome inhibitor-sensitive AMO-1 cells are referred to as WT cells. Bortezomib resistant (aBTZ) and carfilzomib resistant cells (aCFZ), believed to be AMO-1 cells were generated and gifted by the Driessen lab[118]. After typing these cells, they were found to be a mix of AMO-1 cells and L363 cells. AMO-1 cells were cultivated in RPMI-1640 medium (Thermofisher, UK), supplemented with 10% fetal bovine serum (FBS), 100 μ g ml⁻¹ streptomycin and 100 U/ml penicillin (P/S) and 2mM L-glutamine (Invitrogen, UK). Cells were passaged when they reached approximately 1.5-2 million cells per ml. AMO-1 cells are suspension cells and were split twice a week to approximately 0.5 million cells per ml.

3.1.2 L363 cells

After typing the cells gifted by the Driessen lab, they were found to be a mix of AMO-1 MM cells and L363 MM cells. In-house PI-resistant cell lines were by Dr James Dunford by continual and escalating drug exposure of drug-sensitive (WT) AMO-1 cells. However after these cells were typed, they were found to be L363 cells. This was due to the drug exposure selecting the L363 contaminant population over the AMO-1 cells, due to their natural increased resistance to PI, compared to

AMO-1 cells. Once this mistake made by our collaborators was noticed, WT L363 cells were purchased from (<INSERT HERE>). WT, aCFZ and aBTZ cells were cultivated in RPMI-1640 medium (Thermofisher, UK), supplemented with 10% fetal bovine serum (FBS), 100 μ g ml⁻¹ streptomycin and 100 U/ml penicillin (P/S) and 2mM L-glutamine (Invitrogen, UK), and kept in 100nM of their respective proteasome inhibitor. Cells were passaged when they reached approximately 1.5-2 million cells per ml. L363 cells are suspension cells and were split twice a week to approximately 0.5 million cells per ml.

3.2 Compounds

3.2.1 Proteasome inhibitors

<WHERE were they obtained> etc etc.

3.2.2 PRS inhibitors

Halofuginone (MAZ1392)

MAZ1805

NCP22

NCP26

ProSA

3.2.3 Epigenetic inhibitors

The Oppermann group has an epigenetic compound screening library, consisting of 144 compounds. The compounds were obtained XYZ <where did Jim get compounds> SGC???? A dual TRIM24/BRPF inhibitor was identified as a possible candidate to reverse drug-resistance in AMO-1 cells. The structure of the inhibitor is shown below in figure 3.1.



Figure 3.1: TRIM24 inhibitor chemical structure

3.3 Assays

3.3.1 Cell viability assays

10X presto blue (alamar??) was added in a 1:10 ratio to cells in suspension and incubated at 37°C for two to three hours. Plates were read [DETAILS OF MACHINE AND PROTOCOL, e.g. wavelength]

3.3.2 Dose response curves

90 μ l of cells in fresh media were seeded into 96-well plates a day prior to treatment with compound. A total of 20,000 cells were seeded into each well. No cells were placed in edge wells, to avoid edge effects. The following day, media 0% viability controls were placed in the first and last row. Drug concentrations were made up 1000x the desired final concentration. Drugs were diluted once in media (usually 1 in 100), then into the final plate with seeded cells (usually in 10), depending on the experiment. All drug concentrations/combinations were performed in triplicate. Cells were treated with DMSO in triplicate as 100% viability controls.

3.4 Bulk RNA-seq

3.4.1 RNA extraction

RNA was extracted and purified using the Direct-Zol RNA MiniPrep kit (Zymo, USA), following the manufacturer's protocol. In brief, for each sample, approximately 100,000 cells were lysed in 300 μ l of TRIzol and the lysate was transferred to a microcentrifuge tube. 300 μ l of ethanol was added to the lysed samples and vortexed. The mixture was transferred to miniPrep columns and centrifuged at 10,000-16,000g for 30 seconds. The column was washed twice with 400 μ l of Direct-Zol pre-wash and once with 700 μ l of RNA wash buffer. The column was transferred to an RNase-free tube and eluted with 50 μ l of nuclease-free water and centrifuged.

The RNA concentration was quantified using a NanoDrop ND-1000 Spectrophotometer (Thermo Fisher Scientific, USA), and samples were stored at -80°C. Samples were normalised to 100ng with nuclease-free water.

3.4.2 RNA library preparation

NEBNext® Ultra II directional RNA library prep kit for Illumina® with TruSeq indexes was used to prepare RNA libraries, following the manufacturer's protocol. RNA concentration was normalised to 100ng with nuclease-free water, made up to 50 μ l. The NEBNext Poly(A) mRNA Magnetic Isolation Module (NEB, USA) was used to enrich polyadenylated RNA. READ booklet in lab

3.4.3 Pre-sequencing preparation

The molarities of the libraries were determined by electrophoresis on a TapeStation (Agilent, USA). The samples were then pooled according to their peak molarity. The pooled library was then denatured and diluted ready for sequencing. In short, 10 μ l of the approximately 2nM library was denatured with 0.2N 10 μ l NaOH. The mixture was vortexed briefly, centrifuged at 300g for 1 minute, and then incubated at room temperature for 5 minutes. 10 μ l 200mM Tris-HCl (pH 7.0) was then added, vortexed and spun as above. The denatured library was then diluted to 20pM. 970 μ l chilled HT1 buffer was added, vortexed and spun. 117 μ l of the 20pM library

was then mixed with 1183 μ l chilled HT1 buffer to give a 1.8pM library ready for sequencing on the Illumina NextSeq platform.

3.5 Single-cell RNA-seq

3.5.1 Drop-Seq

Cell encapsulation

The Drop-Seq protocol[50] was followed for single-cell RNA-seq sample preparation. Cells were loaded into a microfluidics cartridge. Nadia, an automated microfluidics device (Dolomite Bio, UK), performed cell capture, cell lysis and reverse transcription. Reverse transcription reactions were performed using ChemGene beads.

Library preparation

Beads were collected from the device and cDNA amplification was performed. The beads were treated with Exo-I prior to PCR. The amplified, purified cDNA then underwent fragmentation reactions. A TapeStation (Agilent, USA) was used to assess library quality. The samples were pooled together and split across multiple sequencing runs.

3.5.2 10X Chromium V3

Bone marrow samples were collected from two newly diagnosed multiple myeloma patients and two relapsed multiple myeloma patients; anonymised human tissue samples used in this project were obtained with informed consent by the Haem-Bio Tissue Bank (REC reference: 17/SC/0572). After Ficoll gradient separation, mononuclear bone marrow cells were diluted to 500,000 cells/ml in RPMI media supplemented with 2mM L-glutamine and 10% FBS and 1ml was added to 15ml polypropylene tubes. Compounds were dissolved in DMSO, and 1 μ l of compound solution was added to achieve a final concentration of 1 μ M and incubated for 24 hours. Cells were counted and single-cell RNA-seq library preparation was performed using the Chromium Next GEM Single Cell 3' GEM, Library and Gel Bead Kit v3.1

according to the manufacturer's instructions. Indexed libraries were quantitated by TapeStation, pooled and sequenced on an Illumina NovaSeq 6000 (Novogene, UK).

3.6 QuantM tRNA-seq

Approximately 1 million cells were collected per sample and seeded overnight in six-well plates. Cells were treated with 700nM NCP26, 300nM Halofuginone or 2 μ M ProSA. Controls were treated with equal volumes of DMSO. Cells were collected at time =0, time = 3 hours or time = 6 hours. Samples were centrifuged at 300g for 5 minutes, the supernatant was discarded and the pellets were resuspended in 300 μ l of Trizol. RNA was extracted as above (section 3.4.1) and quantified using a nanodrop. Concentrations ranged from 189.7ng/ μ l to 398.85ng/ μ l.

QuantM tRNA-seq as outlined in [119] was used for library preparation with significant adaptations made to the protocol. Each sample was normalised to 500ng total RNA in 2.65 μ l of nuclease-free water. Samples were deacylated with deacylation buffer (Tris-HCl pH9.0; final concentration 20mM) and incubated at 37°C for 45 minutes. Samples were not demethylated.

3.6.1 Annealing and ligating adapters

The samples were transferred to LoBind PCR plates, and 10pM of 3' adaptor and 2.5pM of each 5' adaptor (A, U, C and G) were added (1 μ l of mix that is 10 μ M for 3' and 2.5 μ M for 5'). The plate was incubated at 95°C for 2 minutes in a thermocycler. 1 μ l of 5x annealing buffer (table 3.1) was added to each sample and the plate was incubated at 37°C for 15 minutes. Ligataion of the adapters to the tRNA was

Annealing buffer	Volume (μ l)
1M Tris-HCl (pH 8.0)	250
0.5M EDTA (pH 8.0)	50
1M MgCl ₂	400
Nuclease-free water	9200
Total	10,000 (10ml)

Table 3.1: Annealing buffer recipe

performed. 0.5 μ l T4 RNA ligase 2 (NEB), 1 μ l 10x reaction buffer and 3.2 μ l nuclease-free water was added to each 5.3 μ l of sample, to total a final volume of 10 μ l (0.5U/ μ l). The plate was placed in a thermocycler and incubated at 37°C for an hour, and then 4°C for an hour. The ligated samples were then transferred to 1.5ml eppendorfs.

3.6.2 RNA precipitation

1.5 μ l GlycoBlue was added to each tube. Each sample was made up to 100 μ l with nuclease-free water. 10 μ l of 3M sodium acetate (pH 5.2) and 250 μ l 100% ethanol was added to each tube and vortexed. Samples were precipitated overnight at -80°C. The following morning, tubes were centrifuged at >12,000g at 4°C for 30 minutes to form a pellet. 2 washes were performed with ice-cold, freshly prepared 75% ethanol, spinning for 10 minutes at 12,000g. All ethanol was removed with an extra 10 second top speed spin, and 10 minutes of air drying with the tube cap off.

3.6.3 Hybridization of RT primer

Samples were resuspended in 10 μ l nuclease-free water and transferred to a PCR plate. 1 μ l 10 μ M RT primer <PRIMER TABLE of sequences ref AT BOTTOM>, 1 μ l 10 μ M dNTP mix, and 1 μ l nuclease free water was added to each sample. The PCR plate was placed in a thermocycler and incubated at 70°C for 2 minutes.

3.6.4 cDNA synthesis

cDNA was synthesised using SuperScript IV Reverse Transcriptase (Invitrogen), following the manufacturer's instructions. 4 μ l 5x SuperScript IV buffer, 1 μ l 100mM DTT, 1 μ l SuperScript IV Reverse Transcriptase and 0.25 μ l RNase Nxgen inhibitor (Lucigen) was added to each sample (totalling 19.25 μ l). The plate was heated in a thermocycler at 55°C for an hour. 19.25 μ l 0.2N NaOH (final concentration 0.1N) was added to each sample, and heated in a thermocycler at 98°C for 20 minutes to hydrolyze RNA. The samples were then transferred to 1.5ml eppendorfs. Ethanol precipitation was performed overnight (as in section 3.6.2), and nucleic acids were resuspended in 12 μ l.

3.6.5 Separating cDNA libraries

Two 18-well 10% Criterion TBE-Urea Polyacrylamide Gels (Bio-Rad) were used to separate cDNA libraries, following the manufacturer's instructions. 1X TBE (89mM Tris, 89mM boric acid, 2mM EDTA) was used as running buffer. 5x sample buffer (89mM Tris, 89mM boric acid, 2mM, 12% Ficoll 400, 0.01% bromophenol blue, 0.02% xylene cyanole FF, 7M urea) was made up, and 3 μ l was added to each sample. 20 μ l 1xTBE, 2 μ l PCR marker (N3234; NEB) and 5.5 μ l 5x sample buffer was mixed and used as a ladder for each gel. cDNA libraries and ladders were pipetted into their corresponding well and the gels were ran for approximately an hour at 90V. Gel were removed and placed back into their plastic tray for staining. Gels were covered in excess 1xTBE and 3 μ l SYBR gold (Invitrogen) was added to each tray and stained for 15 minutes on a mixing tray. Gels were excised on 300 (????)nM UV light. A clean scalpel was used to cut out gel between 75nt and 300nt, the region representing tRNAs, and to place in labelled eppendorfs. Eppendorfs were placed in the fridge overnight.

A 25-gauge needle was used to pierce holes at the bottom of 500 μ l eppendorfs. The pierced 500 μ l eppendorfs were nested inside 1.5ml eppendorfs, and gel pieces were transferred into their corresponding, labelled nested tube. The nested eppendorfs were centrifuged at 18,500g for 5 minutes, until the gel was completely sheared into the bottom tube. The 500 μ l eppendorfs were discarded. 400 μ l of DNA extraction buffer (table 3.2) was added to each 1.5ml eppendorf and the tubes were frozen on dry ice for 30 minutes. The tubes were placed on a rotator overnight at room temperature.

DNA extraction buffer	Volume (μ l)
4M NaCl	1500
1M Tris-HCl (pH 8.0)	200
0.5M EDTA	40
Deionised water	18260
Total	20000 (20ml)

Table 3.2: Extraction buffer recipe

Stage	Temperature	Time	Cycles
Initial Denaturation	98°C	30 seconds	1
Denaturation	98°C	10 seconds	12
Annealing/extension	65°C	75 seconds	
Final extension	65°C	5 minutes	1
Hold	4°C	Inf	-

Table 3.3: tRNA libraries PCR amplification thermocycling conditions

UltraFree MC-VV centrifugal filters with a 0.1µM pore were used to remove small gel pieces. The filters were pre-wet with 7µl DNA extraction buffer, and the gel/ extraction buffer slurry was transferred to them. Tubes were spun at 20,000g for 3 minutes. The filter columns were discarded. 1.5µl GlycoBlue and 500µl propan-2-ol was added to each tube and precipitated on dry ice for 30 minutes. The samples were centrifuged at 18,500g at 4°C for 30 minutes, and washed once with 70% ethanol for 5 minutes. Ethanol was removed completely and air dried for 10 minutes. Samples were resuspended in 12µl nuclease-free water and transferred to a PCR plate.

3.6.6 Circularization

The cDNA libraries were circularized using CircLigase II (Lucigen). 1.5µl 10X reaction buffer, 0.75µl 50mM MnCl₂ and 0.75µl CircLigase II ssDNA Ligase was added to each sample. The plate was placed in a thermocycler and incubated at 60°C for an hour, followed by 80°C for 20 minutes. The libraries were transferred to eppendorfs and ethanol precipitation was performed (section 3.6.2). Samples were resuspended in 15µl nuclease-free water and transferred to a LoBind PCR plate.

3.6.7 Amplification

The circularized cDNA libaries were amplified using NEBnext Ultra II Q5 master mix and custom i5 and i7 primers. 5µl 10µM PCR primer, 5µl 10µM specific itRNA primer, and 25µl Q5 master mix was added to each sample (a master mix was made up of PCR primer and Q5 master mix). The PCR plate was placed in a thermocycler and amplified following the thermocycling conditions in table 3.3.

3.6.8 Library purification

The amplified libraries were purified using 2% agarose gels stained with SYBR gold. 20 μ l 10,000x SYBR gold was added to 200ml 2% agarose gel. 5 μ l 10x bluejuice gel loading buffer (Invitrogen) was added to each sample. 4 μ l PCR marker, 36 μ l nuclease-free water and 4 μ l loading buffer were combined to form the ladders of the gel. The gel was ran at 120V for an hour. Bands corresponding to cDNA libraries (100bp-250bp) were excised on a UV light box with a clean scalpel and transferred to 2ml eppendorfs. Gel extraction was performed using the GeneJET gel extraction kit, following the manufacturer's protocol. In short, approximately 700 μ l binding buffer was added to each tube and the gel mixtures were incubated at 55°C until dissolved. The solubilized gel solutions were added to purification columns and centrifuged at 12,000g for 1 min and the through-flow discarded. 100 μ l of binding buffer was added to each column and centrifuged again. Two washes with 700 μ l of wash buffer were performed, followed by an additional spin and air dry to remove any residual ethanol from the columns. The collection tubes were discarded of and the columns were nested in 1.5ml eppendorfs. 50 μ l of elution buffer was added to each column and centrifuged for 2 minutes. The columns were discarded and the eppendorfs stored at -20°C.

The libraries were quantified on a tape station (Agilent) and pooled according to their peak molarity. The pooled library was then denatured and diluted ready for sequencing (as in section 3.4.3).

3.7 ATAC-seq

3.7.1 Cell lysis

Approximately 2 million cells were collected in 15ml falcon tubes for each condition. The cells were centrifuged at 300g for 5 minutes at 4°C and the supernatant was discarded. The cell pellets were resuspended in 1ml of cold PBS and centrifuged at 300g for 5 minutes at 4°C, the supernatant was then discarded. Fresh lysis buffer was prepared (see tables ??) with occasional gentle flicking. The falcons

were then centrifuged at 500g for 10 minutes at 4°C. The supernatant (cytoplasm) was discarded, leaving the nuclei pellet.

Resuspension buffer	Volume (μl)
1M Tris-HCl (pH 7.5)	500
5M NaCl	100
1M MgCl ₂	150
Nuclease-free water	49,250
Total	50000 (50ml)

Table 3.4: Resuspension buffer recipe

Lysis buffer	Volume (μl)
Resuspension buffer	940
10% non-iodet P40	50
10% tween 20	10
Total	1000 (1ml)

Table 3.5: Lysis buffer recipe

3.7.2 Transposition

Pellets were resuspended in 890μl transposition mix (500μl 2X TD buffer, 330μl 1X PBS, 10μl 10% Tween-20, 10μl 5% Digitonin, 40μl nuclease-free water). For each condition, 176μl was taken in triplicate and transferred to LoBind 1.5ml eppendorfs (Eppendorf, UK). 4μl Tn5 enzyme was added to each eppendorf. The samples were then incubated at 37°C for an hour at 500rpm.

3.7.3 DNA purification

Magic bead clean-ups were performed to purify the DNA. 220μl of magic beads was added to each tube (1.2X), vortexed, centrifuged for 1-2 seconds and incubated at room temperature for 5 minutes. Tubes were placed on a magnetic rack for 2 minutes, until the solution was clear. The liquid from the tubes was aspirated away, leaving about 10μl of liquid remaining. 200μl of 80% ethanol was dispensed over the beads, the tubes were vortexed, spun and placed back on the magnetic rack until the solution was clear and then the ethanol was aspirated away. This wash was

repeated for a total of two ethanol washes. Following aspiration on the 2nd wash, an additional spin was performed and the tubes were placed back on the magnetic rack and any remaining liquid was aspirated away, to ensure all ethanol was removed. The beads were left to air dry for 3-5 minutes on the magnetic rack with the lids of the tubes open. The tubes were removed from the magnetic rack and eluted with 26µl 0.1X TE buffer (Zymo Research, UK). The tubes were vortexed, spun and left to incubate for 5 minutes at room temperature, before being placed back on the magnetic rack. The eluant was transferred to fresh LoBind tubes. The purified DNA was then stored at -20°C until PCR amplification was ready to be performed.

3.7.4 PCR amplification

20µl of purified DNA from each sample was mixed with 20µl nuclease-free water, 5µl ATAC-seq universal primer, 50µl Nebnext high fidelity 2X master mix and 5µl unique ATAC-seq index primer, and split across two PCR tubes. The PCR tubes were put in a thermocycler with a lid temperature of 103.5°C, they were heated to 72°C for 5 minutes, 98°C for 30 seconds, and then cycled at 98°C for 10 seconds, 63°C for 30 seconds and 72°C for 1 minute, 13 times. Samples were then held at 4°C. The paired PCR tubes for each sample were then combined into single 1.5ml LoBind eppendorfs. Magic bead clean-up (as above) was performed, with 110µl magic beads (1.1X). The purified amplified DNA was eluted in 20µl 0.1X TE buffer and transferred to new LoBind tubes. D1000 high sensitivity screen tapes and 2200 TapeStation (Agilent, USA) were used to quantify libraries.

3.8 Pooling, denaturing and diluting libraries

Libraries were then denatured and diluted, following the NextSeq denature and dilute libraries guide, ready for sequencing.

3.9 Sequencing

Sequencing of the resultant libraries was performed on the NextSeq 500 (Illumina, USA) platform using a paired-end run, according to the manufacturer's instructions.

3.10 Phosphoproteomics

3.10.1 Collecting cell pellets

Greater than 20 million cells for each condition (in triplicate) was taken. The cell suspension was centrifuged at 1500g for five minutes. The supernatant was removed, the pellet was re-suspended in 500 μ l of ice-cold PBS, transferred to a 1.5ml eppendorf and centrifuged for a further five minutes. The supernatant was removed using a pipette and the pellet was stored at -80°C.

3.10.2 Cell lysis

300 μ l of fresh lysis buffer (10ml RIPA buffer, 3 μ l benzonase, 1 tablet phos stop) was added to each pellet, vortexed and left for 10 minutes on ice and then sonicated. The supernatant was transferred to a fresh tube.

3.10.3 Protein quantification

Protein concentrations were determined by BCA protein assay (Thermofisher, UK). 400 μ g of protein was taken from each sample. Samples were made up to a volume of 200 μ l with MilliQ-H₂O.

3.10.4 Protein Digestion

Kessler lab protocols were followed (<https://www.tdi.ox.ac.uk/research/research/tdi-mass-spectrometry-laboratory/mass-spectrometry/protocols-and-tools>). The lysed samples were reduced with 5 μ l of 200mM DTT in 0.1 M Tris buffer and incubated for 40 minutes at room temperature. The reduced samples were alkylated with 20 μ l of 200mM iodoacetamide in 0.1M Tris buffer, vortexed and then incubated for 45 minutes in the dark at room temperature. The protein was precipitated using methanol/chloroform extraction. The alkylated samples were transferred to 2ml eppendorfs. 600 μ l of methanol was added to each sample, followed by 150 μ l of chloroform and then vortexed gently. 450 μ l of MilliQ-H₂O was then added and vortexed gently. The samples were centrifuged at maximum speed on a table top centrifuge for one minute. The upper aqueous phase was removed, without disturbing

the precipitate at the interface. 450 μ l of methanol was added to each sample, without disturbing the disc and centrifuged for two minutes. Protein pellets were resuspended, one sample at a time: the supernatant was removed and 100 μ l of 6M urea in 0.1M Tris buffer was added. The samples were vortexed and then sonicated (???). Samples were diluted with 500 μ l MilliQ-H₂O, to ensure the final urea concentration was below 1M. Porcine trypsin (Sequencing Grade Modified Trypsin; Promega, USA) was added in a 1:50 ratio of enzyme:total protein content of sample, such that 40 μ l of trypsin solution containing 8 μ g trypsin in 0.1M Tris buffer was added to each sample. Samples were left to digest overnight at 37°C in an incubator shaker.

3.10.5 Peptide purification

The following day, the reaction was stopped, acidifying samples to 1% Trifluoroacetic acid (TFA). Samples were desalted and concentrated using 1ml C-18 Sep-Pak (Waters) cartridges. Two reagents were used: solution A (98% MilliQ-H₂O, 2% Acetonitrile (CH₃CN) and 0.1% TFA) for washing and solution B (65% Acetonitrile, 35% MilliQ-H₂O and 0.1% TFA) for activation and elution. The columns were flushed with 1ml of solution B and then washed with 1ml of solution A. The digested samples were added to the columns and vacuumed through slowly. Two 1ml washes with solution A were performed. Fresh, labelled eppendorfs were placed beneath the columns and peptides were eluted with 500 μ l of solution B. For phosphopeptide-enrichment, 90% of the peptides were removed for Immobilized Metal Affinity Chromatography (IMAC) on a Bravo Automated Liquid Handling Platform (Agilent). 10% of the peptides were used for total proteome analysis. Eluted peptides were dried using a vacuum concentrator (Speedvac, Eppendorf) and stored at -20°C until analysis by mass spectrometry (MS). Prior to MS analysis, dried peptides were resuspended in solution A.

3.11 Ubiquitinomics

3.11.1 Collecting cell pellets

100 million cells were taken for each condition in triplicate. The cell suspension was centrifuged at 1500g for five minutes. The supernatant was removed, the pellet was re-suspended in 500 μ l of ice-cold PBS and centrifuged for a further five minutes. The supernatant was removed and the pellet was stored at -80°C.

3.11.2 Cell lysis

PMTScan Ubiquitin Remnant Motif Kit (K-ε-GG; Cell signalling, USA) was used, following the manufacturer's protocol (REF). Pellets were solubilized and denatured in 4ml urea lysis buffer (20mM HEPES, pH 8.0, 9M urea, 1mM sodium orthovanadate, 2.5mM sodium pyrophosphate, 1mM β -glycerophosphate). The lysates were sonicated on ice, with two bursts of 15 seconds with a one minute break in-between.

3.11.3 Protein quantification

Protein concentrations were determined by BCA protein assay (Thermofisher, UK). All samples were found to contain between 10mg and 20mg of protein, so all of the available protein was used, with no normalisation.

3.11.4 Protein digestion

Lysates were reduced using dithiothreitol (DTT) at a final concentration of 4.5 mM for 30 minutes at room temperature. The reduced samples were alkylated using iodoacetamide (100mM final) for 15 minutes in the dark at room temperature. The alkylated samples were diluted four-fold with 20mM HEPES (pH 8.0) and digested with 400 μ l trypsin solution, containing 1mg ml⁻¹ trypsin-TPCK (Worthington, LS003744) in 1mM HCl. Samples were left to digest overnight at room temperature on a rotator.

3.11.5 Peptide purification

The following day, the reaction was stopped, acidifying samples to 1% Trifluoroacetic acid (TFA). Samples were desalted and concentrated using 10ml C-18 Sep-Pak (Waters) cartridges. The columns were activated using 5ml of solution B, washed with 10ml of solution A. The samples were added to the columns and ran through slowly. The peptides were washed with 10ml of solution A. The cartridges were then removed from the vacuum and the peptides were eluted into fresh falcon tubes with 6ml of solution B, using the plunger of the syringes. 20 μ g of digested protein was removed from each sample for matching total proteome analysis. The eluate was kept at -80°C overnight. The frozen peptide solutions were lyophilized for two days and then stored at -80°C.

3.11.6 Immunoaffinity purification

10x immunoaffinity purification (IAP) buffer provided with PTMScan Kit was diluted to 1x concentration with MilliQ-H₂O. Purified peptides pellets were resuspended in 1.4ml of IAP buffer by pipetting up and down and transferred to 1.7ml eppendorfs. The samples were centrifuged at 4°C for 5 minutes at 10000xg and kept on ice whilst preparing antibody beads. The anti-body bead slurry was centrifuged (30 seconds at 2000 g) and 1ml of PBS was added and then centrifuged. The supernatant was removed and the antibody beads were washed a further four times with PBS and resuspended in 40 μ l of PBS. The peptide solution was transferred to the antibody vial and the solution was incubated on a rotator for two hours at 4°C. The samples were centrifuged, put on ice and the supernatant was removed. The beads were washed twice with 1ml IAP, followed by three washes with 1ml chilled HPLC water. Immunoprecipitated material was eluted at room temperature in 55 μ l and 50 μ l 0.15% TFA in water, letting the sample stand for 10 minutes after each elution, with gentle mixing every two-three minutes. The eluates were centrifuged and the supernatant was transferred to new tubes. Peptide material was desalted and concentrated using 1ml C-18 Sep-Pak cartridges as above. Prior to mass spectrometry

analysis, purified GlyGly-modified peptide eluates and matching proteome material were dried by vacuum centrifugation, and re-suspended in solution A.

3.12 Liquid-chromatography-tandem mass spectrometry

Liquid-chromatography-tandem mass spectrometry (LC-MS/MS) analysis was performed using a Dionex Ultimate 3000 nano-ultra high pressure reverse-phase chromatography coupled on-line to an Orbitrap Fusion Lumos mass spectrometer (Thermo Scientific) (REF: adan's 3-5 dropbox). In brief, samples were separated on an EASY-Spray PepMap RSLC C18 column (500mm × 75 μ m, 2 μ m particle size; Thermo Scientific) over a 60 min (120 min in the case of the matching proteome) gradient of 2–35% acetonitrile in 5% dimethyl sulfoxide (DMSO), 0.1% formic acid at 250nl min⁻¹. MS1 scans were acquired at a resolution of 60000 at *m/z* 200 and the top 12 most abundant precursor ions were selected for high collision dissociation (HCD) fragmentation.

3.13 CyTOF

Get data off ADAM

3.14 Data Processing

3.14.1 Bulk RNA-seq

Fasta files were processed using a CGAT-flow[120] pipeline, the workflow can be found at: https://github.com/cgat-developers/cgat-flow/blob/master/cgatpipelines/tools/pipeline_rnaseqdiffexpression.py. The pseudo-alignment tool, Kallisto[121], was implemented to pseudo-align reads to the reference human genome sequence (GRCH38 (hg38) assembly) and to construct a counts matrix of samples against transcripts. DESeq2[122] was used for differential expression analysis of counts matrices (using negative binomial generalized linear models) within the R statistical framework (v3.5.1). XGR[123], Reactome[124] and KEGG[125] were used

to perform pathway analysis, within R. Org.Hs.eg.db[126], AnnotationDbi[127] and biomaRt[128] were used for converting between Ensembl IDs, HGNC symbols and ENTREZ IDs.

3.14.2 ATAC-seq

Raw ATAC reads (in fasta file format) were mapped to the GRCh38 reference genome using the CGAT-flow mapping pipeline (https://github.com/cgat-developers/cgat-flow/blob/master/cgatpipelines/tools/pipeline_mapping.py), using the mapper Bowtie. The mapped bam files were then used as input for the CGAT-flow peak calling pipeline ((https://github.com/cgat-developers/cgat-flow/blob/master/cgatpipelines/tools/pipeline_peakcalling.py)). Filtering was performed to filter out [what is filterd out!!] and peak calling was implemented using macs2 (v2.2.7)[129].

3.14.3 Single-cell RNA-seq

The computational pipeline outlined in section 4.2 was used to process scRNA data.

3.14.4 LC-MS/MS

Mass-spectrometry raw data were searched against the UniProtKB human sequence data base and label-free quantitation (LFQ) was performed using MaxQuant Software (v1.5.5.1). Digestion was set to trypsin/P. Search parameters were set to include carbamidomethyl (C) as a fixed modification, oxidation (M), deamidation (NQ), and phosphorylation (STY) as variable modifications. A maximum of 2 missed cleavages were allowed for phosphoproteome analysis and 3 for the GlyGly peptidome analysis, with matching between runs. LFQ quantitation was performed using unique peptides only. Label-free interaction data analysis was performed using Perseus (v1.6.0.2). Results were exported to Microsoft Office Excel and imported into the R statistical framework (v3.5.1) for further analysis.

4

Computational method development

4.1 Introduction

4.1.1 Reproducible workflows

In data analysis, particularly in bioinformatics, many users often create simple bash or R scripts to execute the specific task at hand. However, if this is done frequently, the user will have an accumulation of these single-use scripts, which are often named uninformatively and never used again. This may mean the user creates numerous scripts which perform the same function. Another example of a bad practice is using the command line alone to perform tasks. This means that exactly how the analysis was performed is not recorded and may be lost or difficult to find later. These are bad practices in terms of efficiency and reproducibility. It is much better practice to create well-documented, generalised workflows which can then be applied to multiple different experiments. This enables the user to reuse their code more easily and reproduce results, if need be. This also allows other researchers to reproduce results or apply the code to their own research.

In addition to creating generalised, reproducible workflows, it can be beneficial to create more extensive computational pipelines for jobs which require multiple tasks or actions to be performed sequentially.

4.1.2 Computational pipelines

A computational pipeline consists of a series of manipulations and transformations, where the output of one element is the input of the next. Often these elements are executed in parallel. Pipelining ‘omics’ data-processing means that tasks that are not interdependent can be executed simultaneously. Additionally, multiple samples can be processed in parallel, thereby reducing run time. There are many available pipelining frameworks, for example Snakemake[130], Luigi and Ruffus[131].

For this work, a series of computational pipelines and workflows were generated. Ruffus and CGAT-core[132] were used as the backbone for the pipelines developed.

4.2 scRNA-Seq pseudoalignment pipeline

Fewer pipelines exist for single-cell RNA-Seq compared to bulk RNA-Seq. For the Chromium 10X Genomics platform, most of the processing and analysis is automated by Cell Ranger; however for other technologies, the workflow is not as well defined. A single-cell analysis pipeline was constructed with the aim to produce an easy-to-use, robust and reproducible workflow that works for Drop-Seq as well as 10X technology, which utilises pseudoalignment rather than traditional mapping methods.

4.2.1 Psuedoalignment

Traditional mapping techniques such as Tophat[133] or STAR[134], rely on aligning each read to a reference genome. This is generally very time consuming and computationally expensive. Another challenge that arises with traditional mapping is the occurrence of multi-mapping, whereby a read cannot be uniquely aligned as it could map equally well to multiple sites in the genome[135]. More recently, a series of methods called pseudoaligners have been developed that overcome some of the issues associated with traditional mapping approaches. Pseudoalignment (sometimes referred to as quasi-mapping) methods provide a lightweight, alignment-free alternative to traditional mapping. It has been shown that information on where exactly inside transcripts sequencing reads may have originated is not required for

accurate quantification of transcript abundances[136]. Rather, only which transcript the read could have originated from is needed and transcript abundances are calculated by computing the compatibility of reads with different transcripts. This negates the need for alignment to a reference genome, alleviating the issue of multi-mapping and reducing the computational load. Pseudoaligners have been shown to complete data processing of RNA-seq datasets up to 250-times faster than traditional alignment and quantification approaches[121]. Kallisto[121] and Salmon[137] are tools which implement pseudoalignment. They have similar speed and accuracy for bulk RNA-seq data¹.

Pseudoalignment of scRNA-seq

Pseudoalignment tools have recently been developed for droplet-based scRNA-seq analysis (dscRNA-seq). Additional challenges come with dscRNA-seq data processing, having the extra complication of cellular barcodes (CBs) and unique molecular identifiers (UMIs). These tools must handle transcript abundance estimation, as with bulk RNA-seq analysis, but also perform CB detection, collapsing of UMIs (arising from PCR duplication of molecules) and barcode error correction. Kallisto BUS[138] has been developed as an analysis tool and file format specifically for single-cell analysis, alongside BUStools, for processing of the resultant BUS file[139]. Salmon Alevin[140] has also been developed for single-cell RNA-seq analysis.

Pipeline outline

Kallisto BUS or Salmon Alevin performs pseudoalignment and generation of a cell-by-gene expression counts matrix. Quality control is performed using Scater[141] and alevinQC. Clustering is performed using Seurat3[142] and Monocle[143]. Clusters are projected onto tSNE and UMAP plots. Differentially expressed genes are identified by performing non-parametric Wilcoxon tests on $\log_2 TPM$ expression values and Fisher's exact test for comparing expressing cell frequency, these p values combined using Fisher's method. Multiple comparisons are accounted for by performing the Benjamini-Hochberg correction to adjust the false discovery rate.

¹<https://liorpachter.wordpress.com/2017/09/02/a-rebuttal/>



Figure 4.1: Flowchart outlining scRNA-Seq pseudoalignment pipeline- PLACEHOLDER- remake figure

4.2.2 Benchmark

Benchmarking measures the performance of a method/software relative to other methods available. Run time and the accuracy of results are often the factors considered in a benchmark. To be able to calculate the accuracy of results, the ‘true’ results must be known. This is difficult in scRNA-seq analysis as no gold standard analysis protocol exists. Instead, methods are compared against simulated results which act as the underlying ‘ground truth’.

Simulated data

Simulated reads with a know ground truth counts matrix were generated as follows: 10X (version 2) fastq files of 4k PBMCs from a healthy human donor

were downloaded ². These sequencing files were processed using Salmon Alevin. The resulting Alevin output folder was used as input for Minnow, using Minnow's alevin-mode. Minnow generates droplet-based scRNA-seq simulated reads, working backwards from a known counts matrix to generating raw sequencing files from which the counts matrix could have originated. The valid cell barcode list (whitelist) for 10X chemistry was used (*737K-august-2016.txt*³). Minnow was ran with an error rate of 0.001 and with 12 simulated PCR cycles. Minnow accounts for core experimental dscRNA-seq characteristics, such as PCR amplification bias, barcode sequencing errors, the presence of doublets and ambiguously mapped reads, to try and emulate a realistic set of sequencing reads consistent with the provided counts matrix.

The ground-truth counts matrix was converted to a Single Cell Experiment object (SCE) and the simulated reads were used as input for the scRNA-Seq pseudoalignment pipeline. The resulting count matrices outputted by Salmon Alevin and Kallisto BUS were converted into SCEs, subset and reordered so that they all contained the same cells and genes, in the same order. The Salmon Alevin and Kallisto BUS produced SCEs could then be compared to the ground truth SCE.

Run time

The simulated reads consisted of 434 million reads. Running Salmon Alevin and creating an SCE object took approximately 64 minutes; running Kallisto BUS, sorting and creating an SCE object took approximately 24 minutes. Using the bustools 'count' command to create a counts matrix may have further reduced run time, however more time would be needed to parse it into R and create an SCE object.

Cell barcode handling

The ground-truth data contained 4340 cells. Alevin determined a threshold for the initial whitelist (a set of CBs that likely represent non-empty droplets) by finding a 'knee' in the knee plot shown in Figure 4.2. This initial whitelist contained 5261 cell barcodes, each observed at least 191 times. Following barcode error correction,

²<https://support.10xgenomics.com/single-cell-gene-expression/datasets>

³<https://github.com/COMBINE-lab/minnow/blob/master/data/737K-august-2016.txt>

the final whitelist contained 4340 cells, all of which corresponded to the same CBs as the ground-truth data.

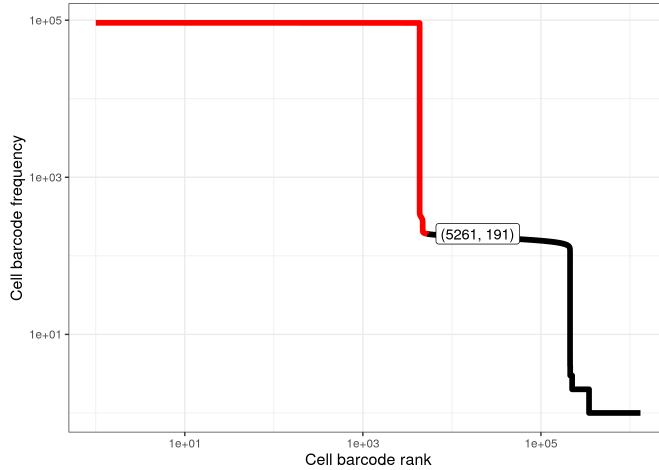


Figure 4.2: Alevin knee plot. This plot displays the number of times each cell barcode is observed, in decreasing order. Finding a ‘knee’ in this plot determines a threshold for the initial whitelist of CBs, which are unlikely to be empty droplets.

For Kallisto BUS, valid cell barcodes were determined using either `emptyDrops` (`DropletUtils`) or by using `barcodeRanks` and calculating the inflection point of a rotated knee plot (where the x- and y- axis are transposed; Figure 4.3). The inflection point method, gave a whitelist of 4339 cell barcodes (one fewer than the ground truth number), but all 4339 CBs corresponded to ground truth CBs. `emptyDrops` gave a total cell number of 12037, only 3746 of which were in the ground truth list of 4340 CBs. This was a large overestimate of number of cells present and the whitelist did not contain all of the valid CBs. Therefore, using the inflection point of the rotated knee plot was found to be the preferred method of filtering cell barcodes.

Gene expression predictive accuracy

To quantify each tool’s accuracy of gene expression, precision, recall and an F1 score were calculated for each gene. The F1 score is a measure of a test’s accuracy,

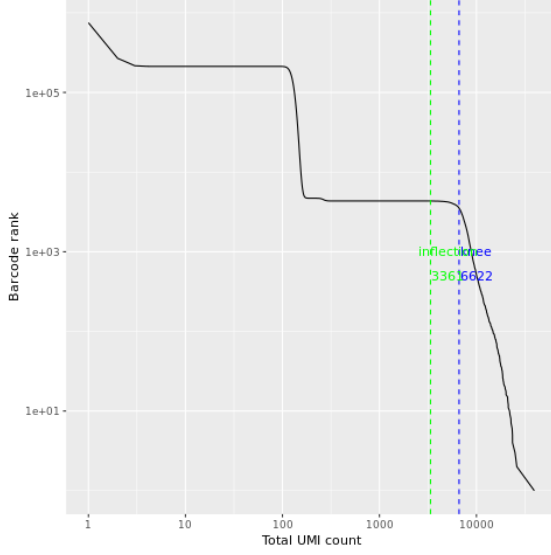


Figure 4.3: Kallisto BUS rotated knee plot. This plot shows the number of distinct UMIs against the rank of the barcode. The Pachter lab transpose the x- and y-axis on their knee plot, so that the x-axis displays distinct UMIs and the y-axis displays ranked cell barcodes, according to the number of corresponding UMIs to each CB. This is supposed to be more intuitive, having the number of distinct UMIs as the independent variable rather than cell barcode rank, as number of UMIs determine the cell barcode rank.

it is the harmonic mean of precision and recall:

$$\begin{aligned} \text{precision} &= \frac{tp}{tp + fp} \\ \text{recall} &= \frac{tp}{tp + fn} \\ F_1 &= 2 \cdot \frac{\text{precision} \cdot \text{recall}}{\text{precision} + \text{recall}} \end{aligned} \quad (4.1)$$

Where for each gene: tp = number of true positives, fp = number of false positives, fn = number of false negatives.

		Ground truth	
		Expressed	Not Expressed
Alevin /BUS	Expressed	True positive	False positive
	Not Expressed	False negative	True negative

Table 4.1: Carol diagram of true/false positives/negatives based on expression between predicted values by Alevin/BUS and the ground truth matrix.

No expression was denoted by 0, and expression by 1. When recall or precision

was undefined, i.e. a gene in Alevin/BUS matrix or the ground-truth matrix was not expressed by any cell, F score was defined as 0.

The mean F1 scores for Alevin and BUS processed data (Figure 4.4) were extremely similar to each other with scores of 0.93 and 0.95, this was due to the large number of F1 scores equal to 1. Figure 4.5 shows the distribution of F1 scores more clearly. Alevin seemed to produce more lower F1 scores than BUS.

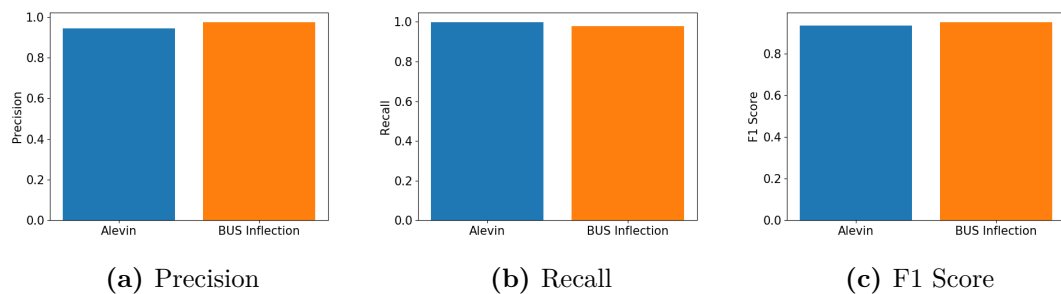


Figure 4.4: F1 score. Two times the product of precision and recall divided by the sum of precision and recall. Measure of accuracy for the tools ability to predict gene expression. Expression classified by 0 or 1. Undefined scores have been removed. F1 scores were calculated for each gene across each cell.

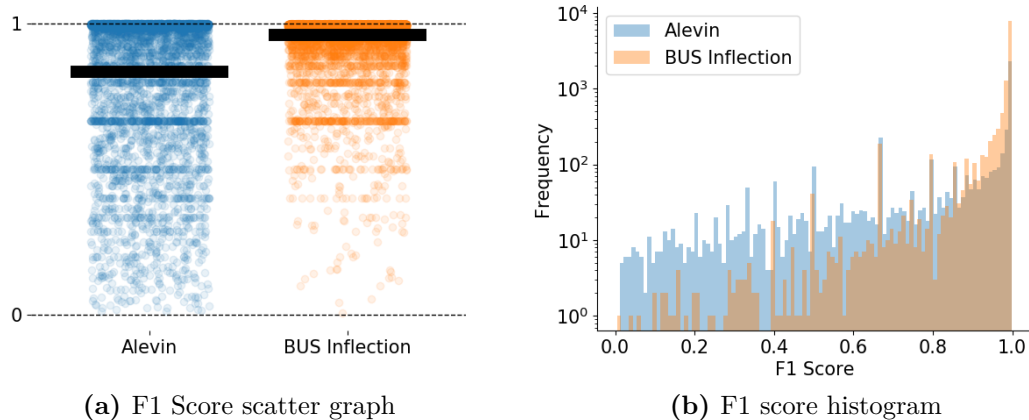


Figure 4.5: F1 score distributions. 4.5a shows the F1 score for each gene expressed across all 4339 cells. The black bar denotes the mean F1 score for each cell. F1 scores of 0 have been removed.

Clustering

Clustering analysis was performed to visualise how well the tools processed the single-cell data and how clusters compared to ground-truth data. Seurat3 integrative

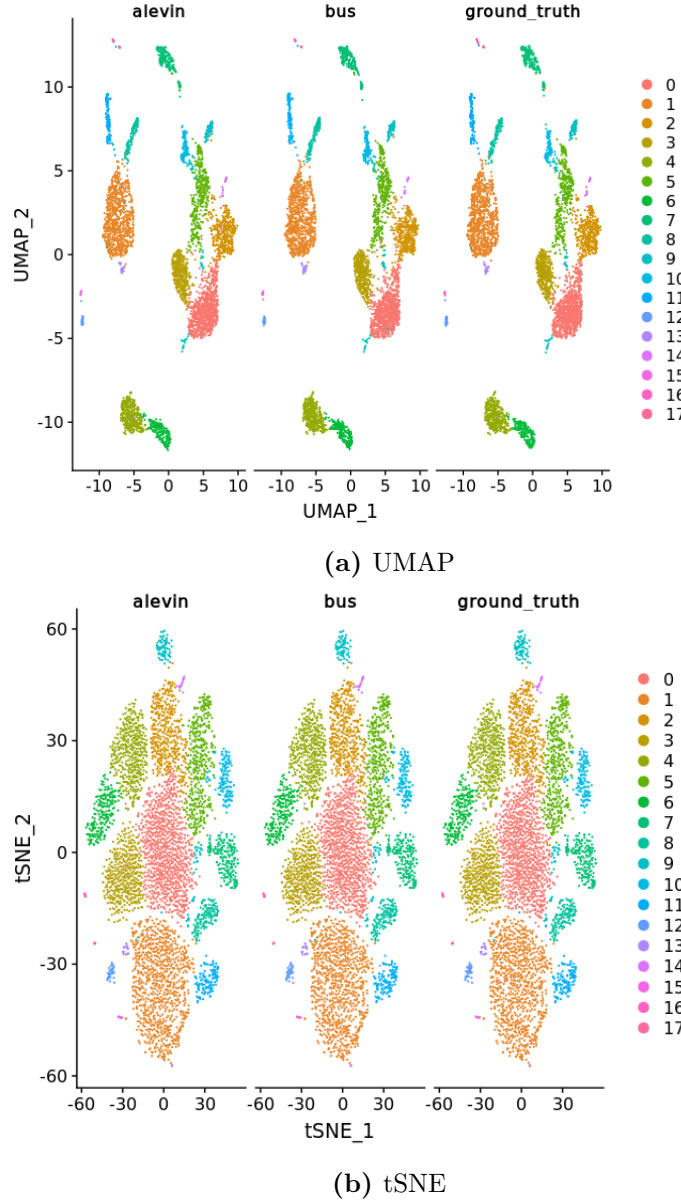


Figure 4.6: Clustering analysis of the simulated data. 18 clusters are present in the ground truth data and Alevin and BUS processed data. Integrated clustering was performed using Seruat3[142], using both Uniform Manifold Approximation and Projection (UMAP) and t-distributed Stochastic Neighbor Embedding (tSNE) dimension reduction techniques.

analysis was performed so that the clusters of each sample could be directly compared. Figure 4.6 shows clustering of Alevin, BUS and ground-truth clustered data, using UMAP and tSNE dimension reductions. 18 clusters are present in all three of the data sets. Visual analysis suggests that the two dscRNA-seq quantification tools compare well to the ground-truth and capture most aspects of the data. From the

benchmark it seems as if both tools are fit for purpose and can accurately quantify gene expression and correctly handle CBs and UMIs.

4.2.3 Updated scRNA-seq pipeline

Following the bench-mark it was decided that Kallisto BUS and BUStools would be used to analyse single-cell data. This was due to its faster run-time and higher F1 scores. The analysis pipeline has been updated continually throughout the project. The updated workflow is outlined in figure 4.7. Each major task has been split into its own pipeline, each containing multiple minor tasks. The black arrows in Figure 4.7 denote a separate processing pipeline. This allows the user to analyse the output after each step, and make changes to parameters as they see fit.

Following sequencing, raw FASTQ files are inputted into Kallisto bus. A bus file is generated along with corresponding information about equivalence classes and transcript names. Kallisto bustools is used to generate a cell by gene count matrix for each sample. These matrices are loaded into R and converted to Single cell experiment (SCE) and Seurat objects. Next, quality control (QC) is performed. Poor quality cells are removed based on numerous parameters. Cells with fewer than a set UMI number (the default is 500), cells with a very low or very high gene count (the default minimum and maximum is 300 and 6000, respectively), and cells with mitochondrial content over a certain ratio (the default is 0.1). This is a user-supervised process, which requires fine-tuning after inspecting graphs and quality metrics. Parameters can be altered and QC performed again. Following quality control, each sample is clustered individually by Seurat, then all samples are integrated together, by either Harmony or Seurat's SCTransform functionality. The integrated dataset is then annotated by cell type by the packages scClassify, singleR or clustfyr, in combination with a reference dataset or model. Often manual annotation by the user (using known biological markers) is required for finer annotation. The user can then perform further downstream analysis on the annotated integrated dataset, for example differential expression analysis and composition analysis.

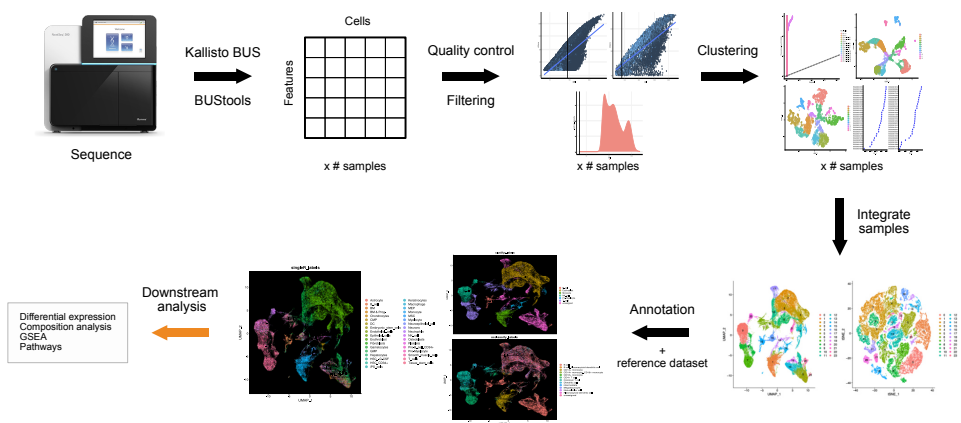


Figure 4.7: Outline of updated scRNA-seq workflow. Each black arrow denotes an independent pipeline. Following sequencing raw FASTQ files are processed by Kallisto bus and bustools and cell x gene count matrices are produced for each sample. Quality control (QC) is performed, based on a number of parameter thresholds. QC often requires user supervision to fun-tune the process. Seurat is then used to perform dimensionality reduction techniques and cluster samples individually. Harmony and Seurat are implemented to perform sample integration. The integrated dataset can then be annotated by cell type using packages cClassifiy, singleR or clustfyr, in combination with a reference dataset or model. The user can then perform further downstream analysis on the dataset, for example differential expression analysis.

4.3 scRNA-Seq velocity analysis pipeline

4.3.1 RNA velocity

"RNA velocity is a high-dimensional vector that predicts the future state of individual cells on a timescale of hours"[144]. In combination with clustering analysis, the trajectory of a single-cell can be tracked.

4.4 tRNA-seq analysis pipeline

4.4.1 Introduction

Transfer RNAs (tRNAs) are non-coding RNAs that transport amino acids to ribosomes during translation, to implement the genetic code. Most genomes contain distinct tRNAs for all 61 codons and these are encoded across multiple sites throughout the genome. Sequencing tRNAs is challenging both experimentally and computationally. The main experimental challenges arise from a stable secondary and tertiary structure, making library preparation difficult [145]. Therefore, efficient library preparation methods must be employed to overcome the ridged structure of tRNA, which usually limits the use of standard library prep methods for sequencing [146]. Computationally, the challenges come from overcoming the reverse transcription errors introduced by chemical modifications and accurately mapping reads to tRNA genomic regions, given their multiple genomic loci [147]. Typically, most mapping strategies for gene expression analysis only report read alignments with unique best matches and thus discard reads mapping to tRNA altogether. As a consequence, specialist mapping strategies to accurately map tRNAs have been proposed [148–150]. Specifically, Hoffmann et al (2018) proposed a two pass mapping strategy that first maps reads to a tRNA masked genome then secondly these unmapped reads are aligned directly to merged tRNA clusters [147].

Computational workflows for small RNA-seq data analysis have been developed previously [151]. However, there are currently a limited number specifically focused on tRNA data analysis. While there have been low-throughput implementations to aid tRNA analysis, such as tDRmapper [148], tRF2Cancer [152], SPORTS1.0 [153] and MINTmap [149], there is now a significant unmet requirement for high-throughput approaches to analyse tRNA data. Firstly, it is desirable to have a single pipeline with the flexibility to perform read quality control, mapping to both general genomic and tRNA features, and the ability to perform qualitative and quantitative analyses on tRNAs within a sample. Secondly, with decreasing sequencing costs, it is now common for small RNA-seq libraries to consist of many

biological and technical repeats and be sequenced at a much greater depth than mRNA-seq libraries. Thirdly, an appropriate level of detailed reporting output is critical for biological interpretation, which should include appropriate visualisation and publication quality figures.

4.4.2 Pipeline outline

A tRNA analysis pipeline (tRNAAnalysis) was constructed with the aim to meet the demands of tRNA-seq analysis. tRNAAnalysis is written using the Computational Genomics Analysis Toolkit (CGAT) core workflow manager [132]. All analysis modules are seamlessly integrated and tRNAAnalysis runs from a single launch command, whilst still also being able to maintain the modularity of running any individual task within the pipeline, if so desired. tRNAAnalysis implements best practice mapping strategies to allow accurate mapping of tRNA reads [147]. The pipeline is optimised so that it can process all input fastq files in parallel, produce detailed logging information, and can be locally run or executed in parallel across a high-performance cluster. tRNAAnalysis can be installed using Conda and a Docker image is also provided with all of the software and packages installed. Users can therefore use the pipeline without having to install numerous dependencies. Finally, tRNAAnalysis provides a user-friendly html report to visualise qualitative and quantitative outputs. Given that the report is written in R using Rmarkdown, the report is easily extensible and customisable, thus providing a user-friendly approach for visualising the analysis.

The workflow is written predominantly in Python and R, using Ruffus decorators [131], and the CGAT-core workflow manager [132], allowing for automatic cluster submission and parallelisation. The pipeline runs via a single command line interface, providing appropriate default settings, with the option to customise configuration parameters and job resources as required.

The main steps in the analysis are:

- Read pre-processing and quality control

- Mapping of reads
- tRNA quantitative and qualitative analysis
- Downstream analyses and report generation

Read pre-processing and quality control

tRNAnalysis accepts raw sequencing data (single-end fastq files) as input and integrates several tools for read quality checking and filtering, including CGAT tools [120], FastQC [154] and FastQ Screen [155]. Given the short length nature of small RNA library preparation, Trimmomatic [156] is used for adapter removal and to filter reads that fall short of quality thresholds. The quality metrics for each sample are then summarised with MultiQC [157].

Mapping of reads

The pre-processed fastq files and a list of gencode annotations are used as the input for mapping. The gencode annotations are supplemented with automatically downloaded RNA repeats (including RNA, tRNA, rRNA, snRNA, srpRNA) from the UCSC database [158]. Firstly, mapping is performed against the genome using Bowtie [159] to obtain a global representation of RNA types. For effective tRNA mapping, tRNAnalysis implements the best-practice mapping strategy proposed by Hoffmann et al (2018), in which tRNA loci are masked from the genome and instead, intron-less tRNA precursor sequences are appended as artificial chromosomes [147]. In first-pass mapping, reads that overlap the boundaries of mature tRNAs are extracted. In a subsequent round of mapping, the remaining reads are mapped to a tRNA-masked target genome that is augmented by representative mature tRNA sequences.

tRNA qualitative and quantitative analysis

tRNAnalysis provides RNA profile analysis that summarises the output of the read alignments mapping to various RNA sequences (e.g. miRNA, piRNA, snoRNA, lncRNA) by counting reads to features with featureCounts [160]. Plots are then generated for each sample, where the positional coverage counts are plotted relative

to the exon, upstream and downstream regions of the tRNAs. These plots can then be used to infer the levels of tRNA fragments within a sample. Using the nomenclature first proposed by Selitsky et al. (2015), if the primary tRNA sequence is < 41 nts and ≥ 28 nts, then it is defined as a tRNA-half, while if it is > 14 nts and < 28 nts then it is defined as a tRNA-fragment [161]. The frequency of read end site relative to the tRNA length is calculated and plotted as bar graphs for each tRNA cluster type. For quantitative measurement of tRNA differences between groups of samples, tRNAAnalysis implements DESeq2 to perform differential expression analysis [122]. Finally, given there can be large sequence variations between tRNAs from different tissues as a consequence of RNA modifications [147, 162, 163], any nucleotide misincorporations in the mapped reads are determined. In order to accurately distinguish sequencing errors for true mismatches, *samtools mpileup* is employed to collate summary information in the mapped bam file and then likelihoods of misincorporation are calculated. This information is stored in a bcf file that is then parsed by *bcftools call*, which performs variant calling for each tRNA sequence [164]. The output is then stored as a vcf file and normalised for indels, then filtered for sequencing depth.

Visualisation

In order to visualise the output of tRNAAnalysis, summary statistics are generated and then reports of these summaries are rendered in html format for visualisation using the Rmarkdown framework. Publication ready reports include figures such as coverage plots, volcano plots of differential expressed tRNAs, tables detailing tRNA modifications and bar graphs of tRNA frequencies. Once these standard reports have been generated, exploratory data analysis can be performed by modifying the Rmarkdown code, allowing for bespoke analysis, tweaking some parameters or to generate more specific publication-quality images.

4.4.3 Simulated data

The accuracy of tRNAAnalysis was assessed using simulated read data. Splatter [165] was used to generate a simulated counts matrix, sampling over a negative binomial distribution with two conditions and three replicates. Biostrings [166] was used to generate a fasta file containing sequences from miRNAs, full-length coding mRNAs and tRNA clusters. The simulated counts matrix from splatter and the mixed RNA-species fasta file were used as input for Polyester [167]. Polyester was ran with an error rate per base of 0.003 to generate simulated RNA-seq read fasta files. Next, CGAT-apps [120] was used to convert the output fasta files into fastq files with uniform quality scores. The simulated fastq files were then used as input for tRNAAnalysis to assess its performance.

Performance metrics

The accuracy of tRNAAnalysis was evaluated using the simulated data above. Receiver operating characteristic (ROC) curves, precision, accuracy and F1 scores were generated, comparing differential expression between the tRNAAnalysis processed simulated data to the known ground truth count matrices. The area under the curve (AUC) was found to be 0.91. The precision was 0.92, recall 0.64 and F1 score 0.76 using a p-value of 0.05, demonstrating a high level of accuracy (figure 4.8).

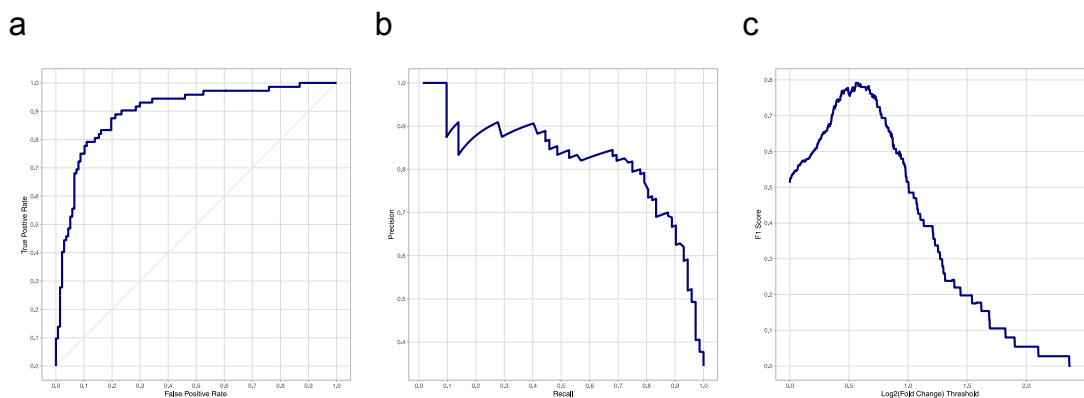


Figure 4.8: tRNAAnalysis performance metrics. Simulated data ground truth counts matrix compared to tRNAAnalysis-processed data. Classifying for whether a feature is differentially expressed or not by varying the $\log_2 FC$ threshold. a) Receiver Operator Curve (ROC) showing the accuracy of tRNAAnalysis to perform differential expression. (Area Under the Curve (AUC), 0.91). b: Recall and precision. c) F1 score.

4.4.4 Reproduce published tRNA-seq analysis

Next, tRNAnalysis was applied to real data originating from a published paper. tRNA-seq data from Chiou et al. (2018) [168] was used to illustrate the functionality of our pipeline.

This dataset comprises samples isolated from activated T cells and their associated extracellular vesicles. Using tRNAnalysis, we were able to confirm the main findings of Chiou et al. (2018), mainly that specific tRNA fragments are enriched in extracellular vesicles and released by activated T cells (Figure 2). Furthermore, we were able to present a more comprehensive evaluation of the different tRNA types within the extracellular vesicles (Figure 2). For each set of input files, interactive plots were created to show the coverage of reads over different types of tRNA fragments. For example, we plot coverage of our reads collapsed across the full length of all tRNAs (Figure 2A), which is further divided into coverage across codons (not shown) and amino acids (Figure 2B). Furthermore, we also report the relative proportion of tRNA fragments (Figure 2C) and tRNA halves (Figure 2D) in each of the input samples. Graphical comparisons between samples and relative controls are shown wherever possible. For example, the differential expression analysis is fully customisable by the user and generates comparative reports depending on the specific contrast supplied (Figure 3).

4.5 Myeloma bone marrow classifier

Using the two scRNA-seq MM BM datasets from chapter 6, cell classifiers were constructed for MM patient BM.

In chapter 6, two scRNA-seq MM BM datasets are investigated. For these datasets, references based on healthy tissue were used to inform annotation packages: clustifyr, scclassify and singleR. As the references originated from healthy tissue, they were unable to label pathological myeloma cells, and MM cells had to be identified manually using known biological markers.

Many studies separate MM cells from other immune types before sequencing, as in ... <List papers>. Using CD45 and CD138 as markers. However, often many MM cells no longer express CD138 as the disease progresses... Lose information on the surrounding microenvironment. etc. etc.

4.5.1 Classifier testing

In order to test the performance of the MM classifiers, publically available MM scRNA-seq data was downloaded from GEO. The test data comprises one PBMC sample from a relapsed and refractory MM patient (GEO accession number *GSE188632*). The deposited counts matrix was processed using the clustering and annotation modules of the scRNA-seq analysis workflow outlined in sectioned 4.2.3.

5

Bulk RNA-seq analysis of PRS inhibitors

5.1 Introduction

Although MM treatment has improved significantly in the last 10-20 years, MM remains an incurable disease. Most MM patients relapse and become resistant to drugs they have previously been treated with. Therefore, research into novel therapeutics that can overcome multi-drug resistance and can be used to treat relapsed patients is of great importance.

There is a new exciting class of compounds for the treatment of many diseases, including numerous cancers. This class of compounds are analogues of, and have been derived from the drug Febrifugine. Febrifugine is the biologically active component of the herb *Dichroa febrifuga*, which is considered one of the fundamental herbs in traditional Chinese medicine. Febrifugine was shown to have strong anti-malarial effects. One such Febrifugine derivative, Halofuginone (HF), has been shown to inhibit T Helper 17 (TH17) cell differentiation, by activating the amino acid response (AAR)[103]. Halofuginone inhibits the enzyme glutamyl-prolyl tRNA synthetase (EPRS). EPRS is a bifunctional aminoacyl-tRNA synthetase (AARS) and catalyses the the aminoacetylation of glutamic acid and proline tRNA species (such that it charges its cognate tRNAs with glutamic acid and proline). Halofuginone and Febrifugine compete with proline at the prolyl-tRNA synthetase active site of EPRS, specifically targeting utilisation of proline during translation[104]. This results in an accumulation of uncharged tRNA^{Pro}s, giving the same cellular environment as if

the cell were proline deficient, triggering the AAR to respond to the apparent proline deprived state.

AARSs are essential in protein synthesis, aiding in building chains of amino acids. Human cancer cells often have an increased rate of protein synthesis, this is especially true in multiple myeloma, creating huge amounts of non-functional paraprotein, therefore are more reliant on aaRSs. As discussed in chapter 2, HF has previously shown anti-MM activity in vitro and in-vivo. HF induced cytotoxicity and apoptosis in numerous MM cell-lines and primary MM cells. HF was also shown to inhibit MM growth and prolong survival in a mouse xenograft MM model. However, the mechanism by which HF exerted its affect was not elucidated, and it is not clear if the AAR plays a role in HF's effectiveness in MM.

It has also been shown that HF's anti-MM effect can be reduced in the presence of excessive proline. Tumours have more proline than healthy cells <REF>. This means that HF has a very narrow therapeutic window, and exhibits many side effects.

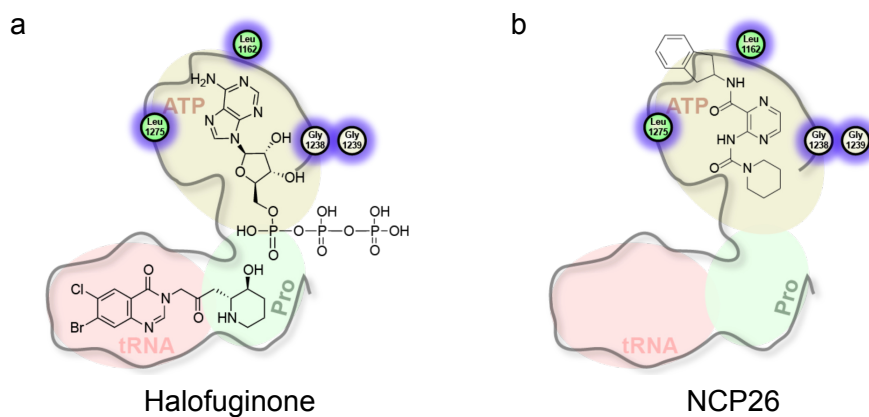


Figure 5.1: Diagrams of Halofuginone/MAZ1392 (a) and NCP26 (b) and their chemical structures. Halofuginone is an ATP dependent, proline and tRNA competitive ProRS inhibitor. NCP26 is an ATP competitive and proline uncompetitive ProRS inhibitor. Aminoacylation is an ATP-dependent process, requiring ATP to activate amino acids. Figure by Ralph Mazitschek.

Recently, The Mazitschek group have synthesized numerous other compounds which target the ProRS site of EPRS. One such example, NCP26 (figure 5.1b), does not compete with proline for the active site of ProRS, unlike Febrifugine and Halofuginone. NCP26 binds to the ATP binding site of ProRS, inhibiting utilization

of ATP. Aminoacylation is an ATP-dependent process, therefore blocking ATP binding inhibits this process, and also leads to an accumulation of uncharged tRNA^{pros}. NCP26 will hopefully alleviate some of the issues associated with HF treatment. More ProRS inhibitors have been synthesized by the group, including NCP22.

This chapter uses MM cell-line models to assess the effectiveness of these ProRS inhibitors and uses bulk-RNA-seq to capture the transcriptional landscape following treatment, and determine their mechanism of action in MM.

5.2 Cell-based assay results

5.2.1 Cell line validation

In this chapter, two MM cell lines are used. Low-passage-number AMO-1 (WT) cells, which are sensitive to various MM treatments, and carfilzomib-resistant L363 cells (CFZr). Dose response curves for the two cell lines treated with proteasome inhibitors (PIs), carfilzomib and bortezomib, were generated to confirm the PI-resistance of CFZr cells and PI-sensitivity of WT cells (figure 5.2). The IC₅₀s for

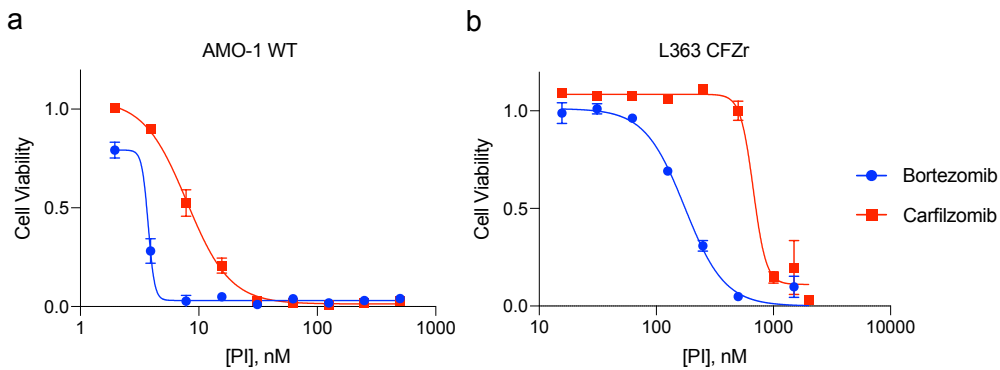


Figure 5.2: Multiple myeloma (MM) cell lines, AMO-1 (WT) and Carfilzomib-resistant L363 CFZr, treated with carfilzomib (CFZ) and bortezomib (BTZ). MM cell lines were treated for 48 hours with a range of concentrations (approximately 1nM-1μM) of proteasome inhibitors (PI). a) AMO-1 (WT cells). b) Carfilzomib-resistant L363 cells (CFZr).

AMO-1 (WT) cells were approximately 3.74nM and 7.93nM for bortezomib and carfilzomib, respectively. For L363 CFZr cells, the IC₅₀s were approximately 174.8nM and 673nM for bortezomib and carfilzomib, respectively. This confirms that WT cells are very sensitive to proteasome inhibition, with IC₅₀s in the low nanomolar range. It also demonstrates that CFZr cells are much more resistant to proteasome inhibition, with the IC₅₀ of CFZr cells almost 85 times higher than WT cells for carfilzomib.

As expected, CFZr cells are more resistant to carfilzomib treatment than bortezomib treatment, as their resistance was developed by exposure to increasing carfilzomib concentrations. However, it is clear that some cross-resistance to bortezomib is also acquired (the IC₅₀ for BTZ treatment is 46 times greater in CFZr cells than WT cells.) This is likely due to the similar mechanism of actions of the

two drugs, conferring resistance to bortezomib treatment too. It could also be in part due to increased expression of general multi-drug resistant genes, such as ATP Binding Cassette Subfamily B Member 1 (*ABCB1*). This mimics clinical MM well, as patients are treated with carfilzomib as a second-line treatment, once they have already been treated with bortezomib, developed some resistance to it and relapsed.

Together, this validates that AMO-1 (WT) cells and L363 CFZr cells are sufficient cell models for PI-sensitive and PI-resistant MM.

5.2.2 Halofuginone and NCP26 are cytotoxic to drug sensitive and drug resistant MM cell lines in a dose-dependent manner

The effect of the PRS inhibitors NCP22, NCP26, MAZ1805 (Halofuginol) and MAZ1392 (Halofuginone; HF) on cell viability was investigated using the MM cell lines AMO-1 and L363 CFZ-r. The cell lines were treated with a range of concentrations of each compound (3.9nM- 1 μ M). Cell viability was assessed using presto-blue assays (section 3.3.2), and dose response curves were generated (Figures 5.3a and 5.3c). Halofuginone and NCP26 reduced cell viability of PI-sensitive AMO-1 cells and carfilzomib resistant L363 cells in a dose-dependent fashion. For this concentration range, MAZ1805 and NCP22 seemed to have little effect on cell viability of WT or CFZ-r cells, and IC₅₀ values were unable to be calculated. Halofuginone was found to be more potent than NCP26.

For WT AMO-1 cells, Halofuginone had an IC₅₀ of 141.8nM and NCP26 an IC₅₀ of 502nM. For CFZ-r cells, Halofuginone had an IC₅₀ of 1185nM and NCP26's IC₅₀ was ambiguous, a higher stock concentration would be required for calculation. Figure 5.3b and 5.3d show the proportion of viable cells following 48 hours of treatment of the PRS inhibitors. WT AMO-1 cells were found to be more sensitive to NCP26 and Halofuginone treatment than carfilzomib resistant L363 cells. This may indicate some acquired cross-resistance built up from carfilzomib exposure or inherent resistance in the L363 cell line over the AMO-1 cell line.

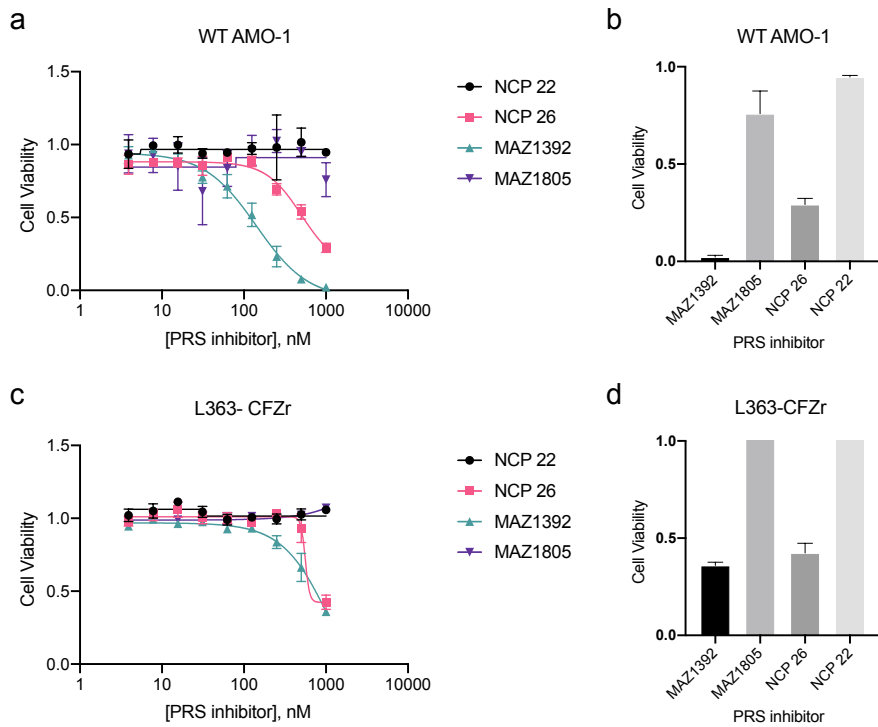


Figure 5.3: Multiple myeloma (MM) cell lines treated with PRS inhibitors- MAZ1392 (Halofuginone), MAZ1805, NCP26 and NCP22. MM cell lines were treated for 48 hours with a range of concentrations (3.91nM-1μM) of PRS inhibitors. a) and c) Dose response curves. a and b) WT AMO-1 cells. c) and d) Carfilzomib resistant L363 cells. b) and d) Proportion of cells viable after 48 hours of 1μM treatment with each agent.

5.2.3 Carfilzomib and NCP26 have an additive or mild antagonistic effect together

Drug combinations have proved effective in MM in recent years, for example the combination of bortezomib, lenalidomide, and dexamethasone (VRd) is used extensively for newly diagnosed MM patients. Drugs are often used in combination so that outcomes are improved (synergistic efficacy) or to reduce off-target effects and toxicity by minimizing the doses of the drugs (synergistic potency) [169].

To assess if NCP26 and carfilzomib work together synergistically, AMO-1 cells were treated with varying concentrations of NCP26 and Carfilzomib for 72 hours, then presto blue assays were performed to determine cell viability (see figure 5.4). SynergyFinder [170] was used to calculate the compounds' synergy scores (-4.66 ZIP; -4.18 Loewe; -5.53 Bliss). From these values it is unlikely that NCP26 and

Carfilzomib work together synergistically. NCP26 and carfilzomib seem to have an additive effect together, or slight antagonistic effect. This reflects a previous result

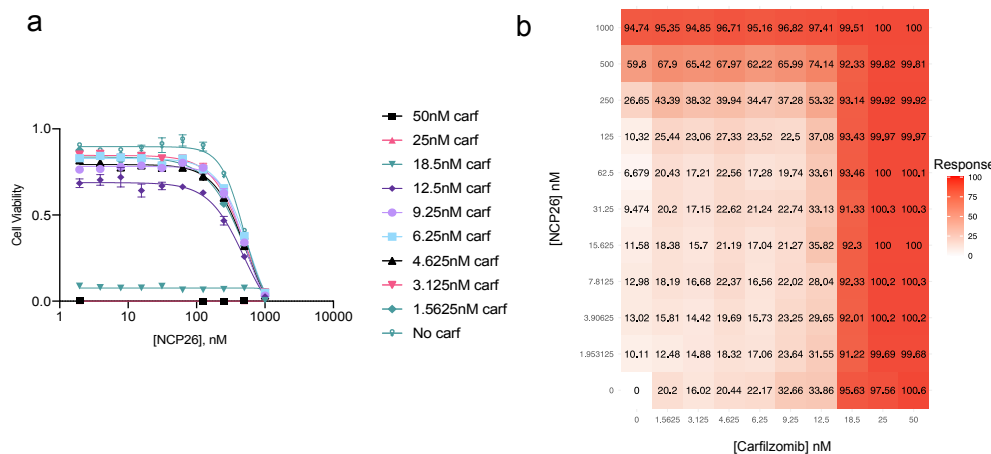


Figure 5.4: Investigating potential synergy between NCP26 and carfilzomib. AMO-1 cells were treated with varying concentration combinations of NCP26 and carfilzomib for 72 hours, then cell viability was determined using presto blue assays. **A)** Dose response curves for NCP26 with different carfilzomib concentrations. **B)** Matrix view of NCP26 and Carfilzomib concentration responses.

where HF demonstrated moderate antagonism with the bortezomib [117]. Although, this study only used a single concentration of BTZ (5nM) and 3 concentrations of HF (25, 50 and 100nM), so may not fully represent the full range of combinations of these two agents. This may indicate that HF and NCP26 may not work very well in combination with proteasome inhibitors in MM. In the same study HF was shown to exhibit moderate synergistic cytotoxicity with other anti-MM agents, including the IMiD lenalidomide and the corticosteroid dexamethasone. This may hint to the potential of ProRS inhibitors used in combination with IMiDs and corticosteroids, perhaps in place of proteasome inhibitors, once they become less effective for patients. Indeed, with the promising results in figure 5.3d, HF and NCP26 have shown cytotoxicity against PI-resistant cells and therefore show potential against relapsed and PI-resistant MM.

5.3 Bulk RNA-seq

5.3.1 Experiment overview

The transcriptome of drug-sensitive and PI-resistant MM cells following treatment with four ProRS inhibitors was investigated using bulk RNA-seq. PI-sensitive WT AMO-1 cells and carfilzomib resistant L363 cells (CFZr) were used. Cells were treated for 6 and 24 hours with a DMSO control, or 1 μ M of a ProRS inhibitor (MAZ1392 (Halofuginone), NCP26, NCP22 and MAZ1805 (Halofuginol)), or 100nM carfilzomib (AMO-1 WT cells only). CFZr cells were treated in the presence of 100nM carfilzomib. Samples were prepped for sequencing as in section 3.4. The computational workflow for bulk RNA-seq analysis is outlined in section 3.14.1.

5.3.2 Clustering

Figure 5.5 shows clustering analysis. Figure 5.5a and 5.5b show the samples distinctly separate into respective cell types, and this makes up the majority (69%) of the variance in the dataset. Therefore, the different cell types (WT and CFZr) were separated and analysed individually (figures 5.5 c-f).

The less active compounds from the ProRS inhibitor dose response curves, MAZ1805 and NCP22, cluster closely with the DMSO-treated controls. The more active inhibitors, NCP26 and Halofuginone, cluster separately from DMSO controls and less active inhibitors, indicating they have elicited a stronger transcriptional response. At 6 hours NCP26 and Halofuginone cluster closely together. At 24 hours NCP26 and Halofuginone separate more. This may suggest a distinction in their mechanism of action, or this could just be reflecting the differences in their potency.

In WT cells, carfilzomib clusters separately from DMSO and the less active inhibitors. At 6 hours, carfilzomib samples are very separate from the NCP26 and Halofuginone cluster, but they cluster together more closely at 24 hours. This could suggest an initial difference in mechanism of action and transcriptional response to the ProRS inhibitors, but culminating in a similar response as time progresses, such as cell stress and cell death pathways.

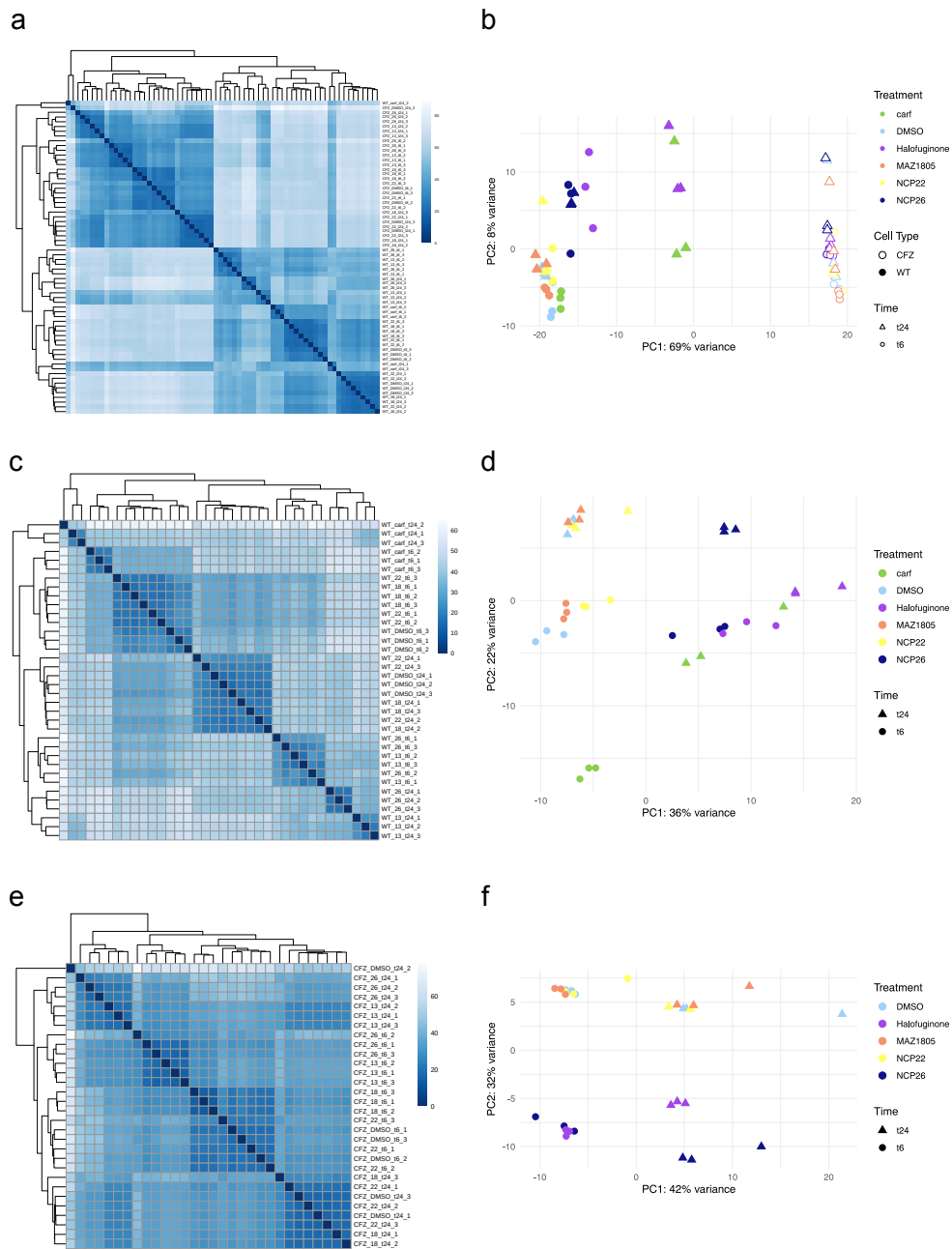


Figure 5.5: Bulk RNA-seq sample clustering. WT AMO-1 cells and CFZr L363 cells treated for 6 or 24 hours with a DMSO control, 100nM carfilzomib or 1uM of a PRS inhibitor (MAZ1392/Halofuginone, MAZ1805/Halofuginol, NCP26 and NCP22). Clustering analysis of sample-sample distances (a, c and e) and principal component analysis (PCA; b, d and f). a) and b) both cell types (WT and CFZr) displayed; c) and d) WT AMO-1 only; e) and f) CFZr samples only.

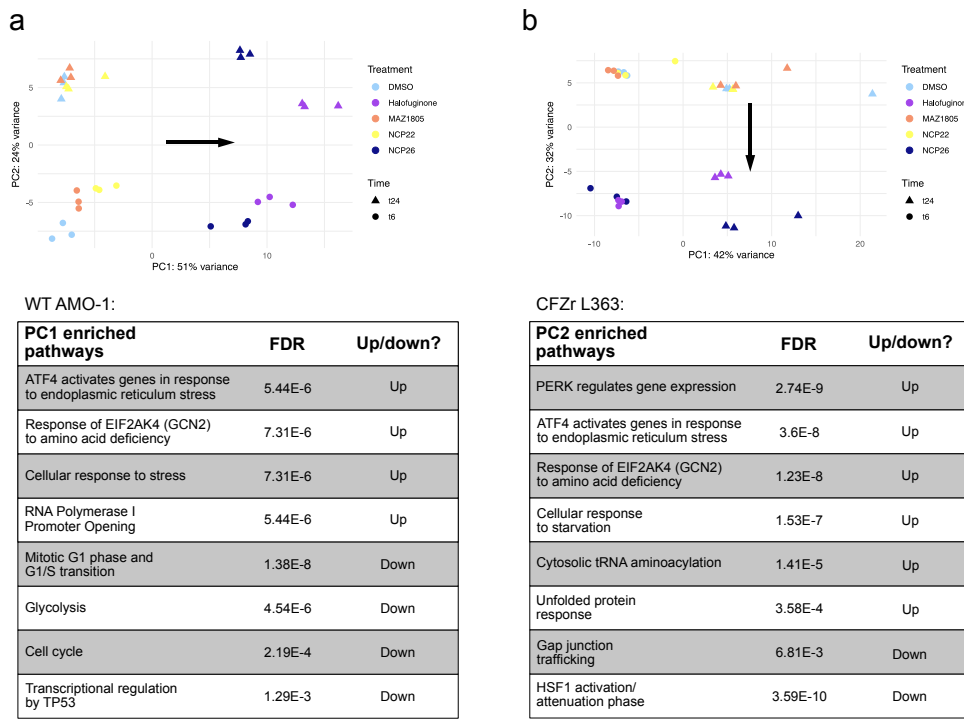


Figure 5.6: PCA pathway enrichment. Pathway enrichment analysis performed using REACTOME of top contributing genes from principal component analysis (PCA). A) WT AMO-1 dataset with carfilzomib-treated samples removed. PC1 seems to account for the separation between DMSO controls/ less active compounds and the more active ProRS inhibitors (NCP26 and Halofuginone). Genes contributing positively towards PC1 are upregulated in NCP26/ Halofuginone compared to controls/ less active compounds. Enriched pathways from top genes in PC1 shown beneath PCA plot. B) CFZr L363 dataset. PC2 seems to account for the separation between DMSO controls/ less active compounds and the more active ProRS inhibitors (NCP26 and Halofuginone). Genes contributing negatively towards PC2 (down arrow), are upregulated in NCP26/ Halofuginone compared to controls. Pathways enriched from top genes in PC2 shown beneath PCA plot.

The top principal components of the WT AMO-1 and CFZr datasets were examined more closely. Carfilzomib-treated samples were removed from the WT dataset, to ensure that the difference between controls and active ProRS inhibitors was captured in PC1 or PC2 following dimensionality reduction. Differential expression, variance stabilising transformation and PCA was re-performed to the CFZ-less dataset. Pathway enrichment analysis was performed for the top genes from the principal component accounting for the difference between controls and Halofuginone/NCP26 treatment (PC1 for WT cells, PC2 for CFZr cells; Figure 5.6). The pathways ‘ATF4 activates genes in response to ER stress’, ‘Response

of GCN2 to amino acid deficiency' and 'cytosolic tRNA aminoacylation' were all enriched, suggesting that the amino acid response is activated. The 'unfolded protein response' (UPR) was also enriched, as well as 'PERK regulates gene expression'. The UPR is a member of the integrated stress response (ISR), so many genes involved in the UPR overlap with genes involved in the AAR. *PERK* regulates the translation response of the UPR. Additionally, genes involved in the cell cycle and G1/S transition are negatively enriched. Previously, HF has been shown to induce the accumulation of cells in the G₀/G₁ cell cycle [117].

5.3.3 Drug sensitive MM

Differential expression

For WT cells at 6 hours, 2119 genes were differentially expressed ($|log_2FC| > 0.5$ and $p_{adj} < 0.05$; DE) for NCP26 treated-samples, 3019 DE genes (DEGs) for Halofuginone, 33 DE genes for MAZ1805, 218 DE genes for NCP22, and 983 DE genes for carfilzomib-treated samples compared to DMSO controls. At 24 hours, NCP26-treated samples had 3323 DE genes, Halofuginone 3426, MAZ1805 2, NCP22 1, and carfilzomib 2260 DE genes compared to DMSO controls. DEGs for NCP26 and Halofuginone treatment are shown in figure 5.7. Genes highly differentially expressed by the ProRS inhibitors are coloured and annotated with their gene symbol. Numerous genes involved in stress response pathways can be seen to be upregulated following NCP26 and Halofuginone treatment, including *TRIB3*, *JUN* and *ATF3*. Additionally, various histone genes are upregulated. Some genes involved in cell cycle progression, such as *CDKN1A*, are downregulated following ProRS inhibition.

The transcriptional changes for 6 hours of exposure to NCP26 were compared with proteomic changes for the same treatment condition (figure 5.8). The proteomic data was supplied by collaborators <COLLABORATORs names here>. Consistent with a mechanism of a global reduction of protein synthesis (with the exception of preferential translation of *ATF4*) upon ISR induction and eIF2 α phosphorylation, very few proteins with an increased abundance were identified (52 proteins with a log_2 fold-change between 0.2 and 0.64). Additionally, a larger shift of proteins with

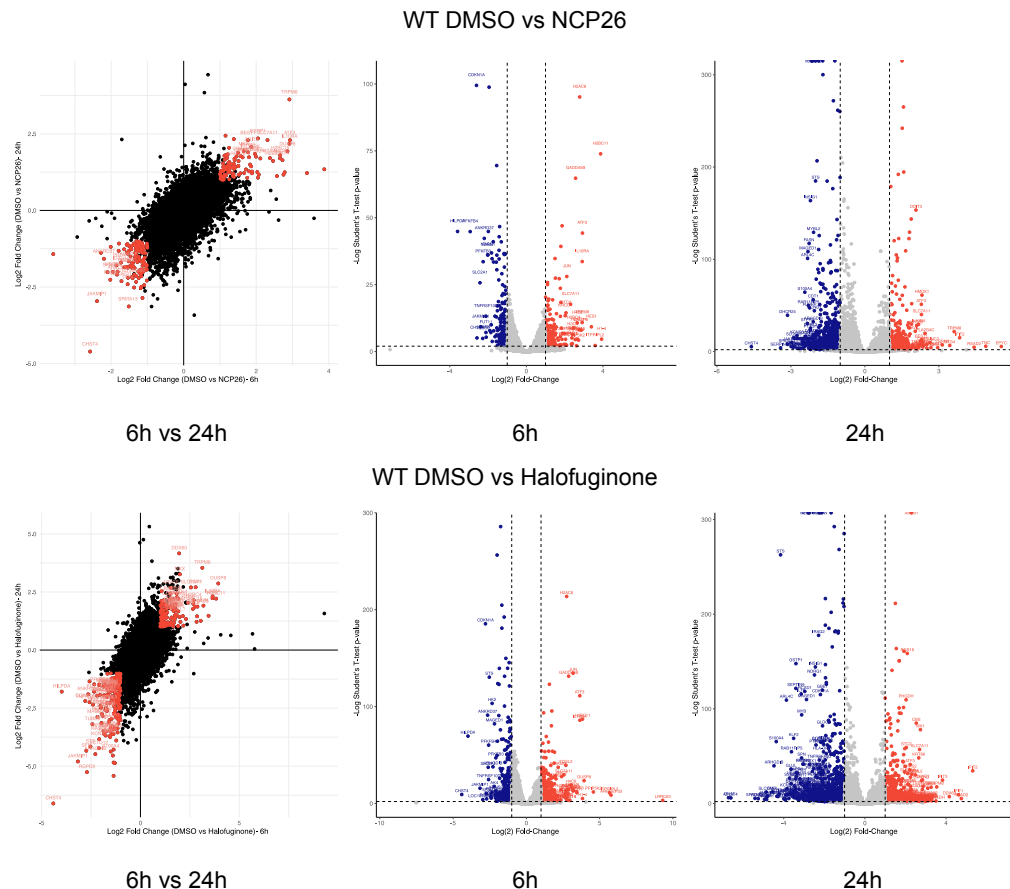


Figure 5.7: Differentially expressed genes (DEGs) for Halofuginone and NCP26 treated WT AMO-1 cells at 6 and 24 hours. Scatter plot of genes for 6 hours treated vs 24 hours treated. Red points indicate genes which are differentially expressed ($p_{adj} < 0.01$) at both 6 and 24 hours. Volcano plots are also shown. Blue points indicate downregulated DEGs ($p_{adj} < 0.01$ & $\log_2 FC < -1$). Red points indicate upregulated DEGs ($p_{adj} < 0.01$ & $\log_2 FC > 1$). Top DEGs are annotated with HGNC symbols.

lower abundance in NCP26 samples compared to DMSO was found. This correlates with transcriptional data where more DEGs were downregulated than upregulated for all ProRS treatment conditions. Also, in-fitting with this mechanism, selective *ATF4* target genes are upregulated, such as *TRIB3* and *INHBE* (as seen highlighted in figure 5.8). This shows a dominant role of the integrated stress response at 6 hours on the transcriptomic and proteomic level.

The effects of NCP26 and Halofuginone treatment on the amino acid starvation response was examined in more detail. AAR genesets ‘GOBP response to amino acid starvation’, ‘REACTOME response of EIF2AK4/GCN2 to amino acid deficiency’ and ‘KRIGE amino acid deprivation’ were collated from the Molecular Signatures

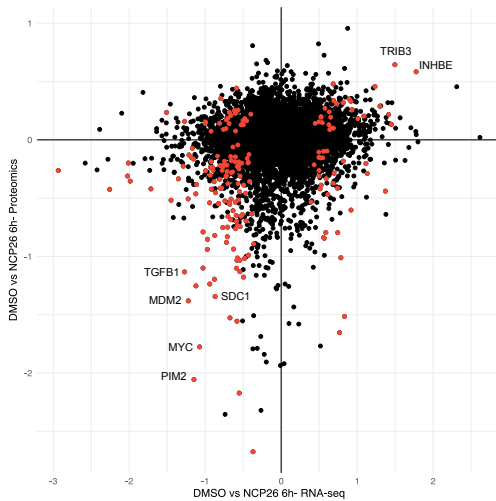


Figure 5.8: Scatterplot of proteomic and RNA-seq datasets depicting changes after 6 hr NCP26 exposure to AMO-1 cells. Red points indicate genes which are differentially expressed ($p_{adj} < 0.01$) in both RNA-seq and proteomic datasets. Following NCP26 treatment, TRIB3 and INHBE (both *ATF4* targets) were the proteins with the highest increase in abundance ($\log_2FC = 0.64$ and INHBE $\log_2FC = 0.58$) in the proteomic dataset and both were significantly upregulated in the RNA-seq dataset.

Database (MSigDb), making up a list of 166 unique genes.

This list of AAR genes was intersected with DEGs for ProRS inhibitor-treatment (i.e NCP26 vs DMSO and Halofuginone vs DMSO at 6 or 24 hours) and a heatmap was constructed (figure 5.9). The AAR transcription factors *ATF3*, *DDIT3* (CHOP), *CEBPB* and *CEBPG* are all markedly upregulated following NCP26 and Halofuginone treatment. Genes coding for aminoacyl tRNA synthetases are also shown to be upregulated, such as *WARS1*, *SARS1* and *CARS1*. Amino acid transporters, such as *SLC7A11*, were also upregulated following ProRS treatment.

A list of 287 unique genes activated by *ATF4* was compiled from the genesets: 'REACTOME ATF4 activates genes in response to endoplasmic reticulum stress' and 'ATF4 Q2' (genes having at least one occurrence of the transcription factor binding site V.ATF4 Q2 in the regions spanning up to 4 kb around their transcription starting sites). Figure 5.10 shows a heatmap for *ATF4* activated genes and genes DE by NCP26 and Halofuginone. Numerous *ATF4* targets can be seen to be differentially expressed following ProRS inhibition with NCP26 and Halofuginone.

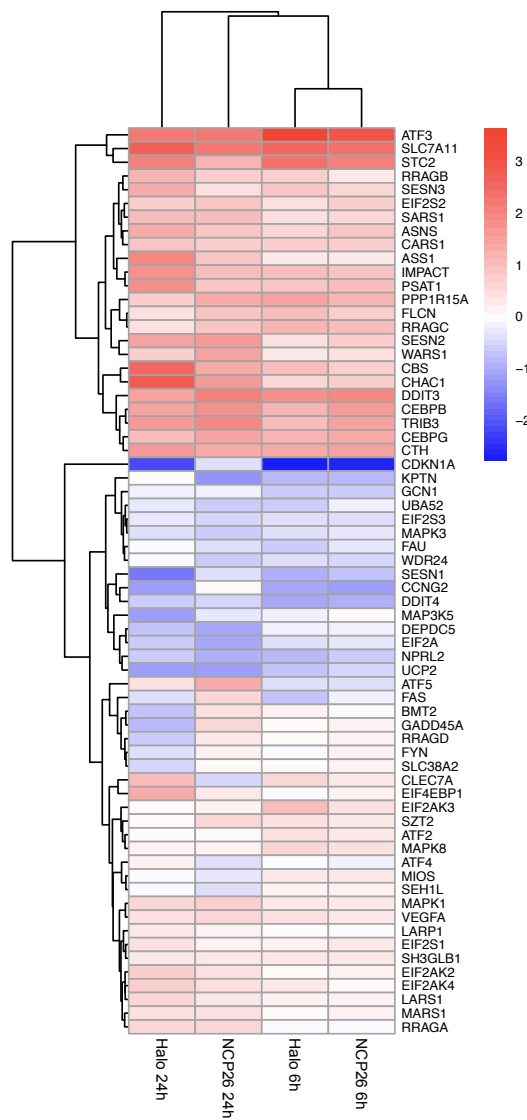


Figure 5.9: Amino acid starvation response (AAR) genes heatmap for WT cells. Differentially expressed genes (DEGs) from WT AMO-1 cells intersected with genes involved in the AAR. A list of known AAR genes was compiled by collating AAR genesets from the Molecular Signatures Database (MSigDb). The colour scale shows \log_2 fold change of expression for each treated sample, compared to its DMSO time control. Red indicates an upregulated gene and blue a downregulated gene.

Of the 287 genes in the list, 165 genes were differentially expressed with NCP26 or Halofuginone treatment.

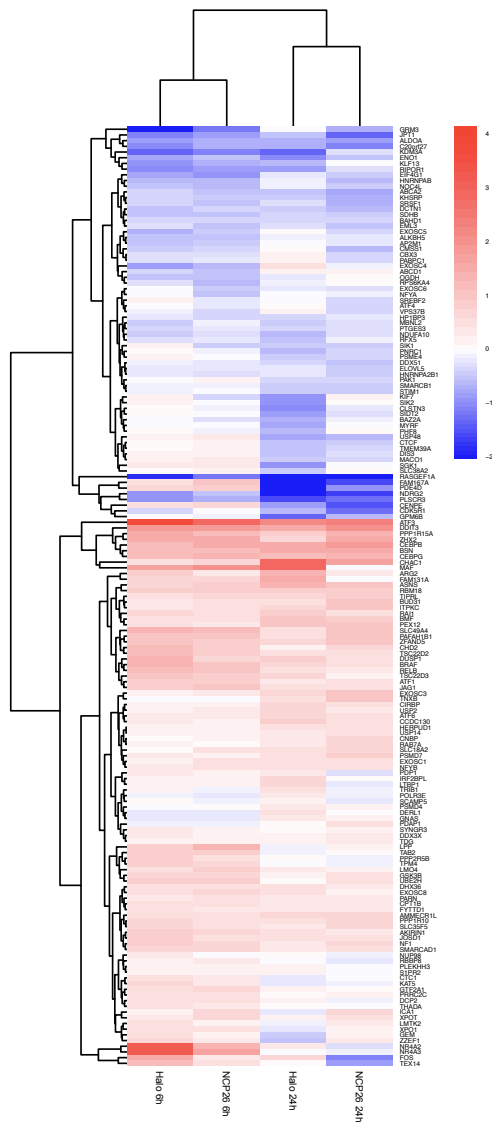


Figure 5.10: Heatmap of *ATF4* activated genes for WT cells treated with NCP26 and Halofuginone. Differentially expressed genes (DEGs) from WT AMO-1 cells intersected with *ATF4* activated genes. A list of genes activated by the transcription factor *ATF4* was compiled by collating genesets from the Molecular Signatures Database (MSigDb). The colour scale shows \log_2 fold change of expression for each treated sample, compared to its DMSO time control. Red indicates an upregulated gene and blue a downregulated gene.

Taken together, it is clear that following HF and NCP26 treatment of AMO-1 MM cells, the amino acid starvation response is activated, mediated via the transcription

factor *ATF4* and culminates in ER stress and downstream apoptotic mechanisms.

ProRS inhibitors vs carfilzomib

The mechanism of action of proteome inhibitors has been extensively studied and well described. Figure 5.5d demonstrates similarities at 24 hours between the transcriptional effects of ProRS inhibitors and the proteasome inhibitor carfilzomib on AMO-1 cells. It also demonstrates ProRS and carfilzomib-treated samples separation at 6 hours, highlighting differences in their initial mechanism of action.

The similarities and differences between ProRS inhibitors and carfilzomib were studied in more detail. As seen by x-y trends of scatter plots and number of overlapping genes in venn diagrams (figure 5.11a and b), ProRS inhibitors and carfilzomib treatment are more similar at 24 hours, and share more DEGs (compared to DMSO controls), than at 6 hours. Somewhat obviously, NCP26 and HF share more overlapping DEGs with eachother than with CFZ. HF shares more overlapping DEGs with CFZ than NCP26. This is likely to do with potency and dosing, where both HF and CFZ were used at approximately 10 times their IC₅₀ value.

Carfilzomib treatment of AMO-1 cells resulted in a pronounced induction of the heat shock response, changes to ubiquitin mediated processes and protein folding, in line with the well-defined cellular changes of proteasome inhibition. At 6 hours, some ISR effectors, such as *ATF3* and *JUN*, are upregulated by both carfilzomib and NCP26/Halofuginone treatment.

At 24 hours, CFZ and ProRS inhibitor treatment seem to culminate in similar end-stage stress, cell cycle changes and apoptotic mechanisms. Both ProRS inhibition and CFZ treatment result in pathway enrichment of ‘cell cycle arrest’, ‘negative regulation of cell proliferation’, ‘positive regulation of apoptosis process’ and ‘type 1 interferon signalling pathway’. Figure 5.11d shows a heatmap of NCP26, Halofuginone and CFZ for log₂ FC compared to DMSO controls at 6 and 24 hours for selected genes, belonging to various pathways/ classes. NCP26 and Halofuginone demonstrate similar effects on myeloma markers as CFZ, such as downregulating the MM pathological marker (*SDC1/ CD138*). CFZ is an established anti-MM

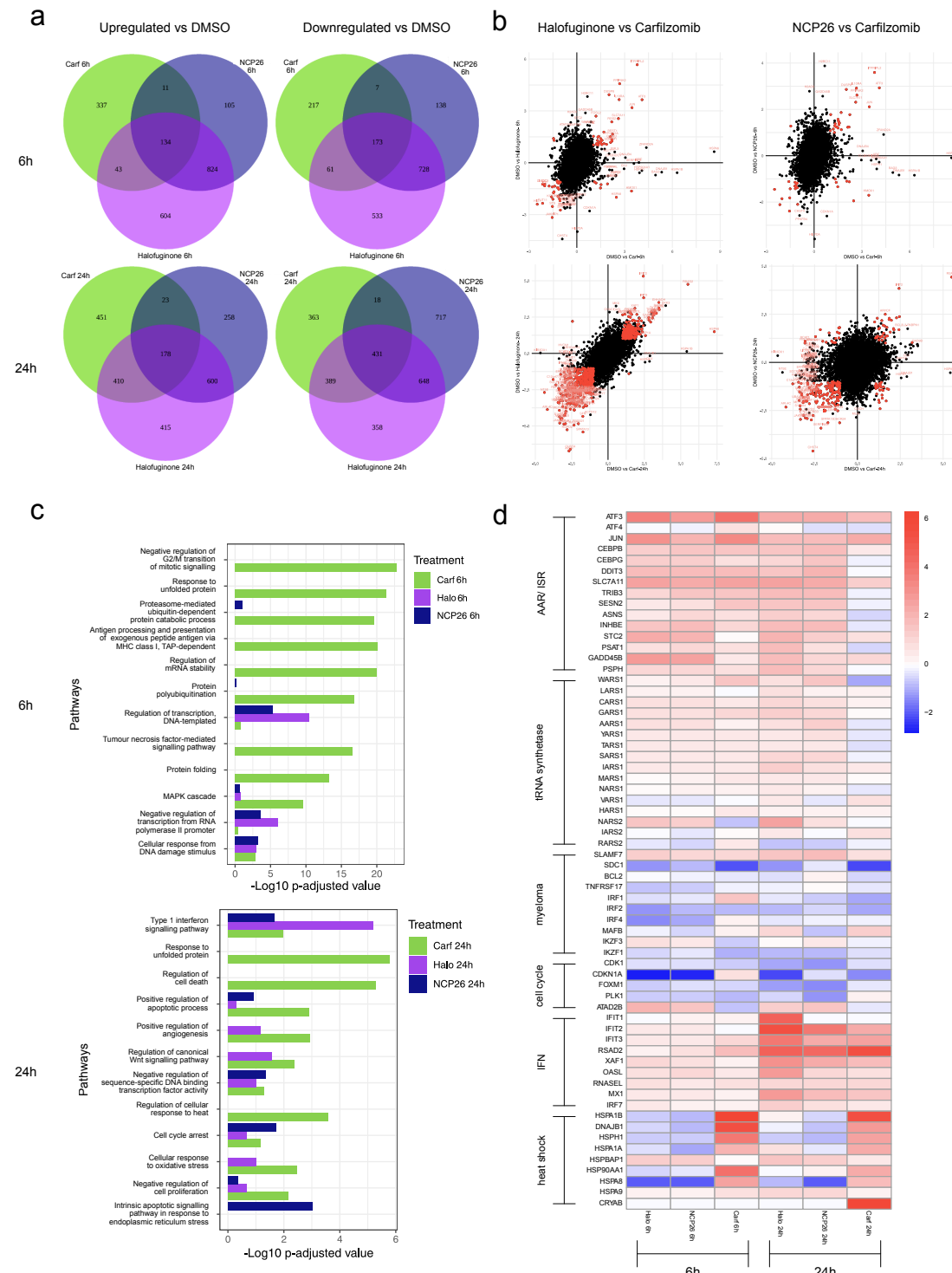


Figure 5.11: ProRS inhibitors compared with Carfilzomib's mechanism of action. a) Venn diagrams showing overlapping differentially expressed genes (DEGs; upregulated or downregulated following treatment) at 6 and 24 hours. b) Scatter plots showing correlation of carfilzomib DEGs against Halofuginone or NCP26 DEGs. c) Pathway analysis (Gene ontology biological processes; GOBP) for top upregulated genes. d) Heatmap of selected differentially expressed genes upon carfilzomib, Halofuginone and NCP26 treatment.

therapy approved in the clinic, therefore this is promising for the effectiveness of ProRS inhibitors in MM.

5.3.4 Carfilzomib-resistant cells

Differential expression

For CFZr L363 cells at 6 hours, 1165 genes were DE for NCP26 treated-samples, 2424 DEGs for Halofuginone, 222 DEGs for MAZ1805, and 0 DE genes for NCP22-treated samples compared to DMSO control samples. At 24 hours, NCP26-treated samples had 852 DE genes, Halofuginone 256 DE, and MAZ1805 and NCP22 no DE genes compared to DMSO controls. DEGs for NCP26 and Halofuginone treatment

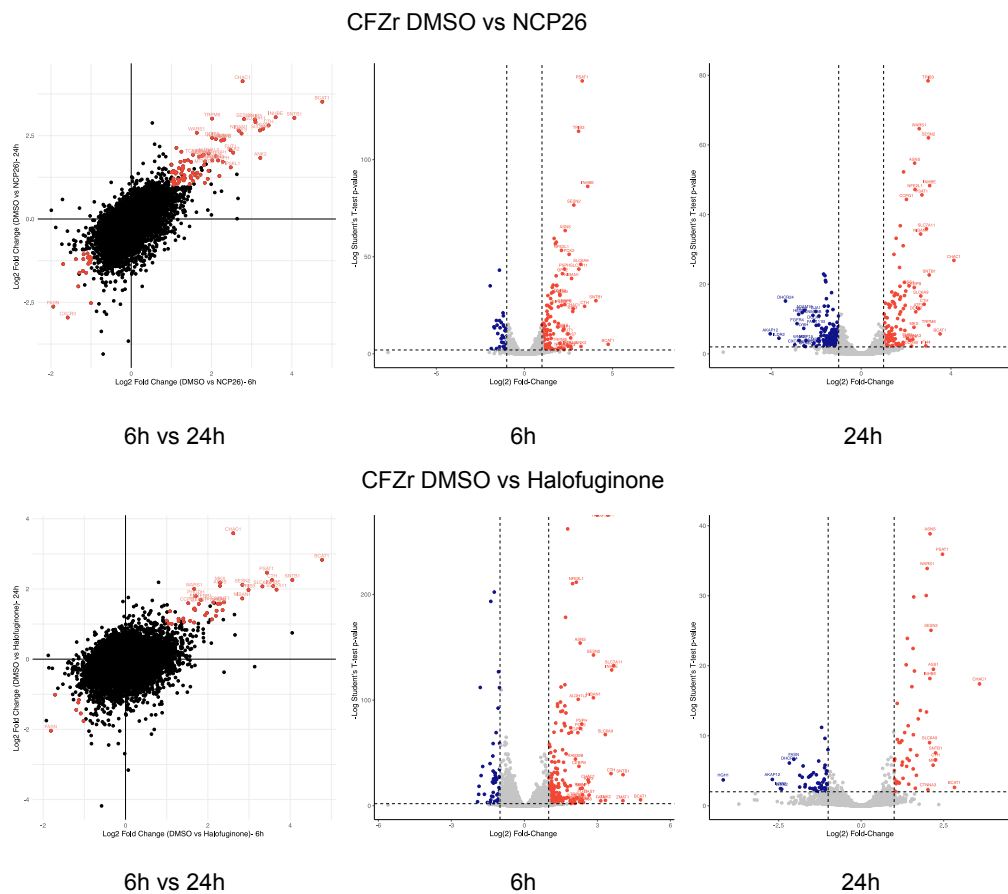


Figure 5.12: Differentially expressed genes (DEGs) for Halofuginone and NCP26 treated CFZr L363 cells at 6 and 24 hours. Scatter plot of genes for 6 hours treated vs 24 hours treated. Red points indicate genes which are differentially expressed ($p_{adj} < 0.01$) at both 6 and 24 hours. Volcano plots are also shown. Blue points indicate downregulated DEGs ($p_{adj} < 0.01 \& \log_2 FC < -1$). Red points indicate upregulated DEGs ($p_{adj} < 0.01 \& \log_2 FC > 1$). Top DEGs are annotated with HGNC symbols.

are shown in figure 5.12. Numerous genes involved in the AAR can be seen to be upregulated following NCP26 and Halofuginone treatment, including *TRIB3*, *ATF3*, *CHAC1*, *INHBE*, *PSAT1*, *ASNS* and *SESN2*; Amino acid transporters *SLC7A11* and *SLC6A9*, and tRNA aminoacyl synthetase *WARS1* are also upregulated.

Pathway enrichment analysis

Pathway enrichment analysis was performed for the top DE genes for NCP26 and Halofuginone treated samples compared to DMSO controls (figure 5.13). Multiple pathways relating to endoplasmic reticulum stress and apoptosis were enriched following ProRS inhibition. ‘Cellular response to amino acid starvation’ was enriched, along with ‘amino acid transport’, indicating that NCP26/Halofuginone likely activate the amino acid starvation response in the CFZ resistant MM cells. Additionally the pathway ‘response to unfolded protein’ was enriched. The unfolded protein response is part of the integrated stress response so shares many of the same effectors as the AAR, such as *DDIT3/CHOP*.

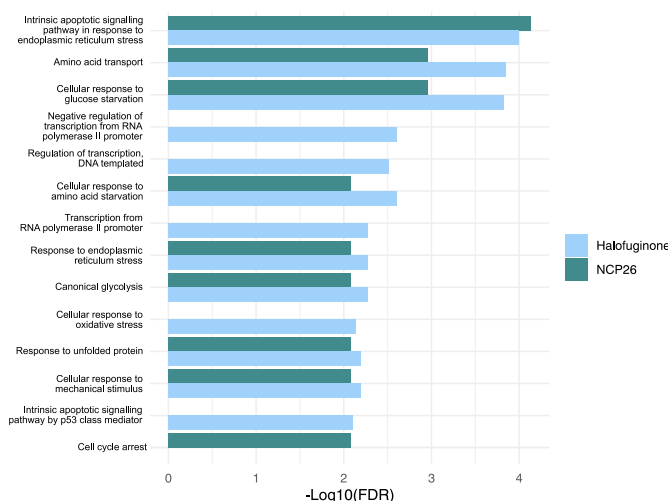


Figure 5.13: Pathway analysis for ProRS treated CFZr cells at 24 hours [CHECK]. Gene ontology biological processes (GOBP) was performed for the top DE genes.

The amino acid response was investigated more closely for the PI-resistant cell line. Figure 5.14 shows a heatmap of DEGs for NCP26/Halofuginone treated CFZr cells at 6 or 24 hours, which are known members of the AAR (collated from MSigDbl- as above).

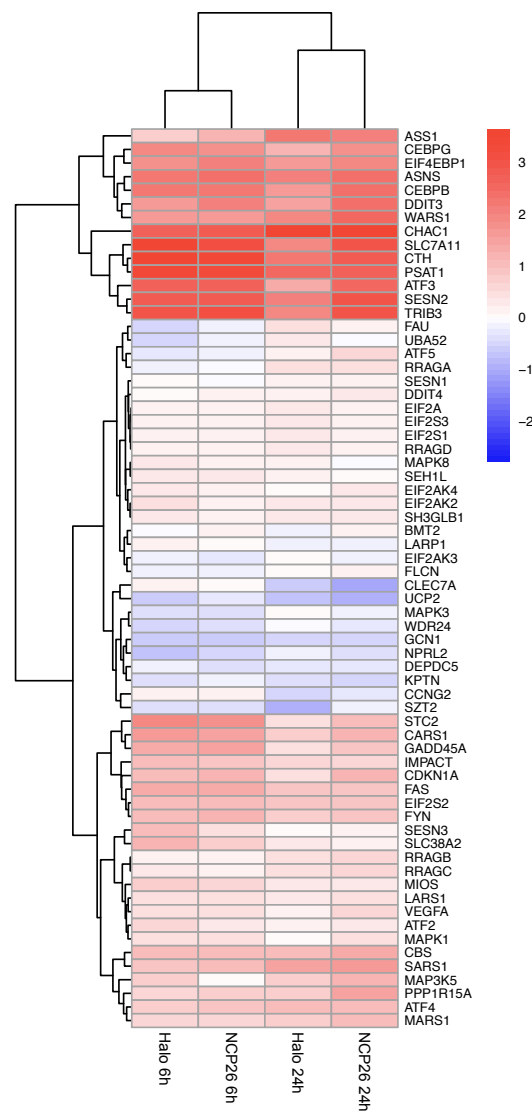


Figure 5.14: Amino acid starvation response (AAR) genes heatmap for CFZr cells. Differentially expressed genes (DEGs) from CFZr AMO-1 cells intersected with genes involved in the AAR. A list of known AAR genes was compiled by collating AAR genesets from the Molecular Signatures Database (MSigDb).

A strong elicitation of the AAR can be seen following NCP26 and Halofuginone treatment. The transcription factors *CEBPG*, *CEBPB*, *ATF3*, *DDIT3* are strongly upregulated, along with amino acid transporter *SLC7A11* and tRNA aminoacyl synthetase *WARS1*. Other AAR genes are strongly upregulated with ProRS inhibitor treatment, such as *ASS1*, *ASNS*, *CHAC1*, *PSAT1*, *PPP1R15A/GADD34* and *TRIB3*.

A list of 287 unique genes activated by *ATF4* was compiled from the genesets: 'REACTOME ATF4 activates genes in response to endoplasmic reticulum stress' and 'ATF4 Q2' (genes having at least one occurrence of the transcription factor binding site V.ATF4 Q2 in the regions spanning up to 4 kb around their transcription starting sites). Figure 5.15 shows a heatmap for ATF4 activated genes and genes DE by NCP26 and Halofuginone in CFZr cells. Numerous *ATF4* targets can be seen to be differentially expressed following ProRS inhibition with NCP26 and Halofuginone. Of the 287 genes activated by *ATF4* in the list, 157 genes were differentially expressed with NCP26 or Halofuginone treatment. This indicates that many *ATF4* downstream targets are being mediated, indicating that *ATF4* has high transcriptional activity.

CFZ-resistant L363 cells show a similar engagement of the AAR and activation of *ATF4*-mediated genes as PI-sensitive AMO-1 cells, following NCP26/HF treatment. Additionally many downstream apoptotic mechanisms are enriched, indicating the initiation of cell death by AAR activation. Taken with the cytotoxicity dose response data, this demonstrates that ProRS inhibitors are capable of killing PI-resistant MM cell lines. Thus targeting the ProRS could be a potential effective strategy in overcoming PI drug resistance in clinical MM.

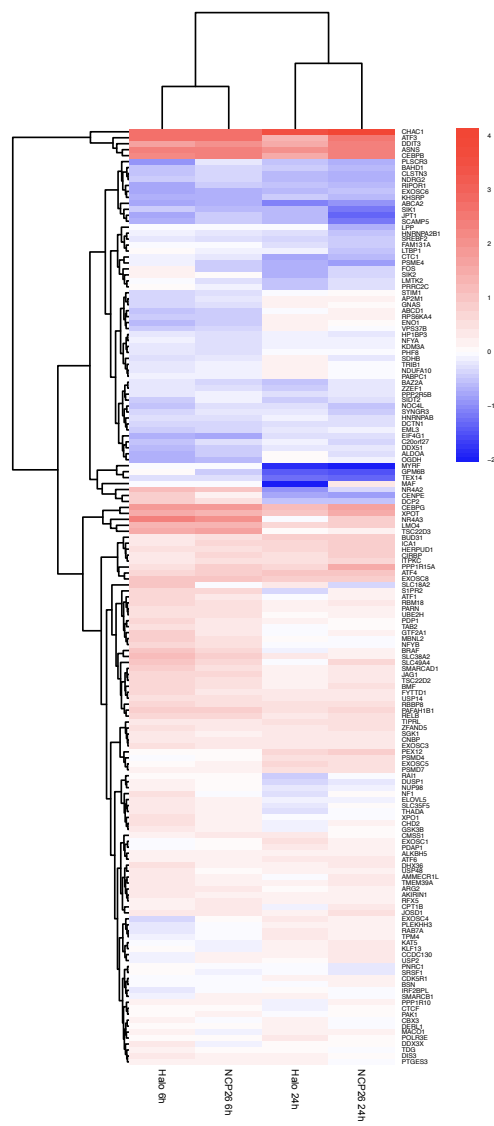


Figure 5.15: Heatmap of *ATF4* activated genes for CFZr L363 cells treated with NCP26 and Halofuginone. *ATF4* activated genes, which are differentially expressed following NCP26 or Halofuginone treatment at 6 or 24 hours. A list of genes activated by the transcription factor *ATF4* was compiled by collating genesets from the Molecular Signatures Database (MSigDb). The colour scale shows \log_2 fold change of expression for each treated sample, compared to the corresponding DMSO control. Red indicates an upregulated gene and blue a downregulated gene.

5.4 Summary

NCP26 and Halofuginone have been shown to reduce cell viability of drug sensitive and PI-resistant MM cell lines in a dose-dependent manner. It has been shown that NCP26 and carfilzomib are not synergistically cytotoxic together. This reflects a previous study where HF and bortezomib were found to be moderately antagonistic in combination. Therefore, administering a PI and ProRS inhibitor in combination would likely offer no increased benefit to patients. This is likely due to similar mechanisms of activating ER stress pathways. However from the dose response data, it is clear that ProRS inhibitors are cytotoxic against PI-resistant MM cell lines. Indicating that their mechanism of actions are still distinct enough that NCP26 and HF have an effect on CFZ resistant cells. Together with the previous result of HF interacting synergistically with lenalidomide and dexamethasone[117], this could support a case for using ProRS inhibitors as a replacement for PIs once patients have relapsed and stop responding to proteasome inhibition.

From omics data, a general shift towards lower protein abundance and downregulation of gene expression has been demonstrated following exposure of MM cells to ProRS inhibitors. This indicates activation of components of the ISR to halt global protein synthesis (with the exception of *ATF4* target genes). *ATF4* target genes were shown to be upregulated and larger in abundance following NCP26 and Halofuginone treatment. *ATF4* is the master regulator of amino acid metabolism. It is activated by amino acid deprivation. Halofuginone and NCP26 treatment caused increased expression of *ATF4* in both drug sensitive and PI-resistant MM cell lines. Expression of numerous genes involved in the AAR and ISR were markedly increased following Halofuginone and NCP26 treatment. *DDIT3* and other downstream pro-apoptotic genes were over expressed following NCP26 and HF treatment. Western blot data from collaborators has demonstrated that NCP26 and Halofuginone elicit canonical ISR activation with GCN2 and eIF2 α phosphorylation in a dose-dependent manner (figure A.1). Together this data shows that the ProRS inhibitors NCP26 and Halofuginone activate the amino acid starvation response in MM cell lines. It also demonstrates that apoptotic pathways are activated following AAR activation,

indicating that the cytotoxic effects of NCP26 and Halofuginone in MM cell lines are attributable (in part) to AAR activation and its downstream apoptotic mechanisms.

6

Single-cell RNA-seq analysis of ProRS inhibitors

6.1 Introduction

MM cells grow within the bone marrow and are supported as they grow by their microenvironment. The MM microenvironment comprises a cellular compartment (composed of immune cells, endothelial cells, osteoblasts, osteoclasts and stromal cells) and a non-cellular compartment (composed of the extracellular matrix (ECM), cytokines, chemokines and growth factors)[171, 172]. There are interactions between malignant plasma cells and the surrounding microenvironment. The bone marrow microenvironment has been indicated to play a supportive role in migration, proliferation, differentiation and drug resistance of malignant plasma cells. There is evidence linking the tumour microenvironment to progression of MGUS to active MM, for example significant matrix remodelling has been seen between the bone marrow of healthy individuals, MGUS and MM patients[172]. Therefore, to get an accurate picture of MM, information must be acquired about the surrounding niche.

Historically, the tumour environment has been investigated following the isolation of populations of cells sorted from the tumour and then sequenced using traditional microarray or bulk RNA-seq techniques. Bulk techniques measure the average expression across a sample, which is the sum of cell type specific expression weighted by cell type proportions. Single-cell techniques measure expression for each individual cell and therefore provide information on clonal diversity that may

be lost when pooling cells into bulk samples. Furthermore, multiple myeloma is an extremely heterogeneous disease, this is seen both between patients and within an individual's own tissue. Applying single-cell techniques to capture the inter- and intra-individual heterogeneity is fundamental to identifying molecular and cellular signatures that define MM.

The advent of single-cell technologies has led to a better understanding of the complexity and diversity of the tumour microenvironment. Seminal papers from Melnekoff et al. (2017)[173] and Ledergor et al. (2018)[174] use scRNA-seq to reveal clonal transcriptomic heterogeneity in MM samples. Melnekoff et al. (2017) demonstrated the clonal heterogeneity within MM using samples that were collected from eight relapsed MM patients. The group performed t-SNE clustering analysis and the samples separated into eight transcriptionally distinct clones, each corresponding to a different patient. This highlights the inter-patient differences of MM. Ledergor et al. (2018) performed a similar study to evaluate clonal heterogeneity within MM but also had a set of controls with which to compare the MM group. They found that MM patients have greater inter-individual transcriptional variation, where each MM patient possessed a unique and individual plasma cell transcriptional program. They also showed substantial intra-tumour heterogeneity (subclonal structures) of plasma cells in a third of their MM patient cohort. These papers established the importance of using single-cell techniques to study MM, as to not miss the underlying clonal heterogeneity. However both of these papers focussed solely on plasma cells and did not look at the surrounding bone marrow microenvironment. To truly understand the complexities of MM and treatment of MM, interactions between plasma cells and the bone marrow niche must also be explored using single-cell techniques.

This chapter uses MM patient-derived bone marrow (BM), from both newly-diagnosed and relapsed patients, to investigate the transcriptional effects of ProRS on MM cells and their surrounding immune microenvironment.

6.1.1 Experiment overviews

Three single-cell experiments, comprising samples from four MM patients, were performed to explore the effect of various compounds (including Halofuginone and NCP26) on MM patient tissue. The BM samples for experiments 1 and 2 were obtained from two treatment-naive, newly-diagnosed MM patients. Experiment 3 comprised samples from two patients, both with relapsed MM, therefore both presenting with a degree of acquired anti-cancer drug resistance. For experiment 1, BM samples were treated for 24 hours with 1 μ M Casin, GSK-J4, Halofuginone, NCP26, SGC-GAK, Verteporfin or a DMSO control, totalling 7 samples. For experiment 2, BM samples were treated with 1 μ M CAMKK2, CLK or CSNK2 for 24 hours; 1 μ M SGC-GAK, Halofuginone, NCP26 or a DMSO control for 24 and 48 hours, totalling 11 samples. For experiment 3, BM samples from patient 3 and 4 were treated for 24 hours with either a DMSO control, 1 μ M Halofuginone, 1 μ M NCP26 or 5 μ M NCP26, totalling 8 samples.

Following compound treatment, scRNA-seq library preparation was performed by Dr Martin Philpott as outlined in section 3.5.2. <GENEWIZ experiment 3 >

6.2 Data processing

Initially all four patient samples were processed and integrated together. However, integration was found to be poor between treatment-naive patients and the relapsed patients. This was expected as MM patients' transcriptome has been shown to change considerably following <XYZ> rounds of treatments REF. Therefore, samples from experiments 1 and 2 (treatment naive patients 1 and 2) were integrated together and samples from experiment 3 (relapsed patients 3 and 4) were integrated together.

Experiments 1 and 2 contained treatment samples that were not of interest to this project. However, all samples originated from MM patients, therefore all 18 samples were included in the analysis up to and including the integration and annotation stage of data processing. This was to increase the granularity of the

data, and allow for easier annotation of clusters. However, only the downstream analyses of DMSO, Halofuginone and NCP26 samples are included in this work.

6.2.1 Analysis overview

The single cell analysis pipeline outlined in section 4.2.3 was used to process the scRNA-seq data. Kallisto BUS/ BUStools was used to pseudoalign reads and quantify gene expression. Next, quality control and filtering of the samples was performed. Poor quality cells are likely to have a low number of genes and UMIs per cell. So any cells with fewer than 500 UMIs were removed. Cells with a gene count below 300 or above 6000 were also removed. Cells with a mitochondrial ratio higher than 0.1 were removed (a high proportion of mitochondrial genes indicates mitochondrial contamination from dead or dying cells).

After quality control and filtering of the data, clustering was performed using Seurat v3, followed by integration of all samples, using Seurat and Harmony. Using the Seurat SCTransform normalisation method of integration, the two experiments were too large to integrate across all samples. Therefore, a reference-based approach was taken, whereby a subset of samples were selected (based on their cell richness and relevance to the research question) and listed as ‘reference datasets’ for SCTransform normalisation. Harmony integration was also implemented, using patient and experiment as additional covariates.

Cell type annotation was performed by using several automated packages (singleR, clustifyr and scClassify), then fine-tuned manually using a list of known biological markers. The HumanCellAtlas database was used to inform singleR and clustifyr annotation. scClassify was performed using a pre-trained scClassify model, based on seven PBMC single-cell datasets (including 10X V2, 10X V3, smartSeq, celSeq, dropSeq and inDrops datasets). As the reference datasets were based on healthy tissue, they were unable to label pathological cells, like myeloma cells. Myeloma cells were identified manually using a range of known markers, for example: *CD38*, *CD138*, *SLAMF7* and *BCMA* (see table 6.2).

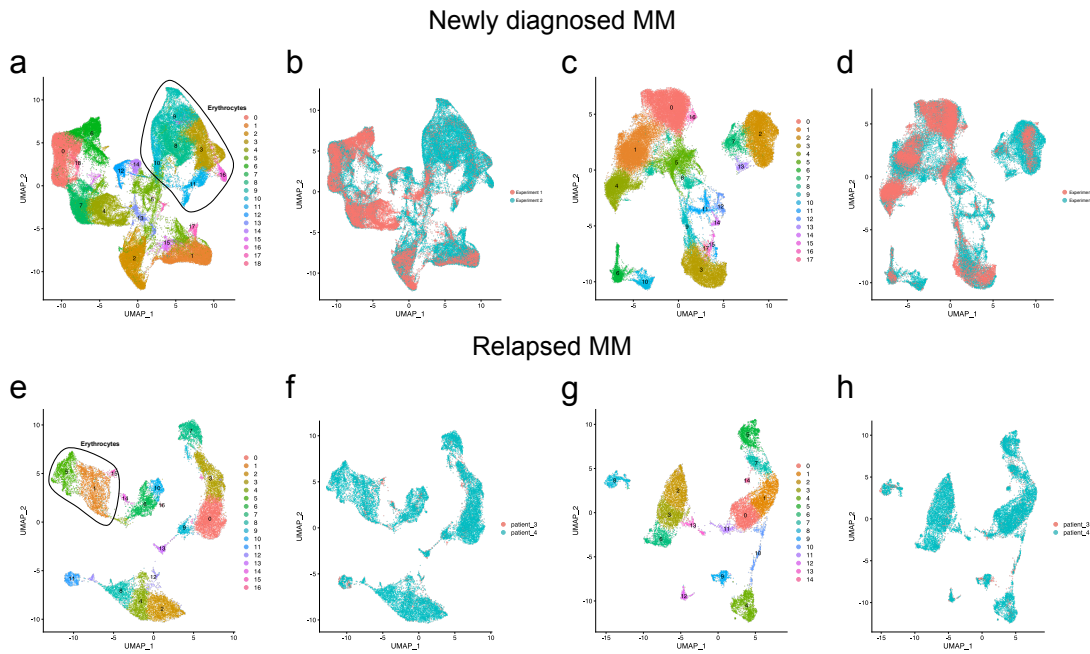


Figure 6.1: UMAP dimension plots following integration of samples from experiment 1 and 2 (treatment naive patients), and samples from patients 3 and 4 in experiment 3 (relapsed MM). [a-d] Experiment 1 and 2-newly diagnosed MM patients. [e-h] Experiment 3, patients 3 and 4- relapsed MM patients. [a, b, e, f] Integrated UMAP plots before erythrocyte cell and gene removal. Erythrocyte clusters are circled in a) and e). [c, d, g, h] UMAP plots following removal of erythrocyte cell clusters and genes and re-integration of samples. [b, d, f, h] show the composition of each dataset by experiment or patient.

Experiment 2 was found to have an extremely high erythrocyte population (figure 6.1a and b). In addition, many other cell populations were expressing erythrocyte specific genes, where we would not expect to see them being expressed, for example MM cells expressing numerous haemoglobin subunit genes. Many of the variable features that Seurat uses for clustering analysis and dimension reduction were made up of these erythrocyte-specific and haemoglobin genes. The high expression of these genes was thought to be affecting the integration of the two experiments together. A theory for the presence of the large number of erythrocytes and un-localised erythrocyte gene expression is that perhaps the BM sample taken for experiment 2 was one of the later samples taken from the patient and contained a large amount of blood. Library prep clean-up may have missed many of these cells, leading to ambient erythrocyte RNA being present within many droplets.

It was decided to remove the erythrocyte clusters (clusters 3, 8, 9, 10, 11 and

16 in newly diagnosed; clusters 1, 5 and 15 in relapsed) and haemoglobin related genes or erythrocyte specific genes that seemed to be dominating expression in the integrated dataset. After the integrated Seurat object had the erythrocyte genes and cells removed, it was split back up into separate Seurat objects for each sample, and integration and clustering was performed again. Seurat's SCTransform with reference datasets and Harmony (using a multi-covariate model, accounting for each different sample and the two different experiments) were used to re-integrate all samples. Harmony integration was found to integrate clusters across patients and experiments better than using Seurat's SCTransform. The Harmony-integrated datasets were taken forwards and used for cell type annotation. Better integration was achieved after removing erythrocytes and erythrocyte-specific genes (see figure 6.1d).

A large erythrocyte component was also found for patient 4 for in experiment 3 (figures 6.1e and 6.1f). The same analysis workflow was applied to experiment 3, removing the erythrocyte cluster and erythrocyte specific genes and re-integrating using Harmony with samples and patients as covariates.

Table 6.1 shows for each experiment: the total number of cells, the number of cells passing filter, and the number of cells remaining following erythrocyte removal. Patients 1, 2, 3 and 4 have a final cell number of 23915, 37161, 2257 and

Experiment	Patient	Total cells	Cells passing filter	Cell number after erythrocyte removal
Experiment 1	Patient 1	112452	25779	23915
Experiment 2	Patient 2	462560	61059	37161
Experiment 3	Patient 3	4894	2625	2257
	Patient 4	21682	18674	14934

Table 6.1: Total cells, the number of cells passing filter, and the number of cells passing filter once erythrocyte clusters were removed across all samples for each patient.

14934 cells, respectively, remaining for downstream analysis. The low number of cells for patient 3 may be a barrier to interpreting meaningful results. However, patients 1, 2 and 4 have more than sufficient cell numbers to identify MM cells and perform differential expression.

6.2.2 Annotation of re-integrated data

R packages clustifyr and scClassify were used to aid in cell type annotation of the integrated datasets, the result of this annotation can be seen in figure 6.2.

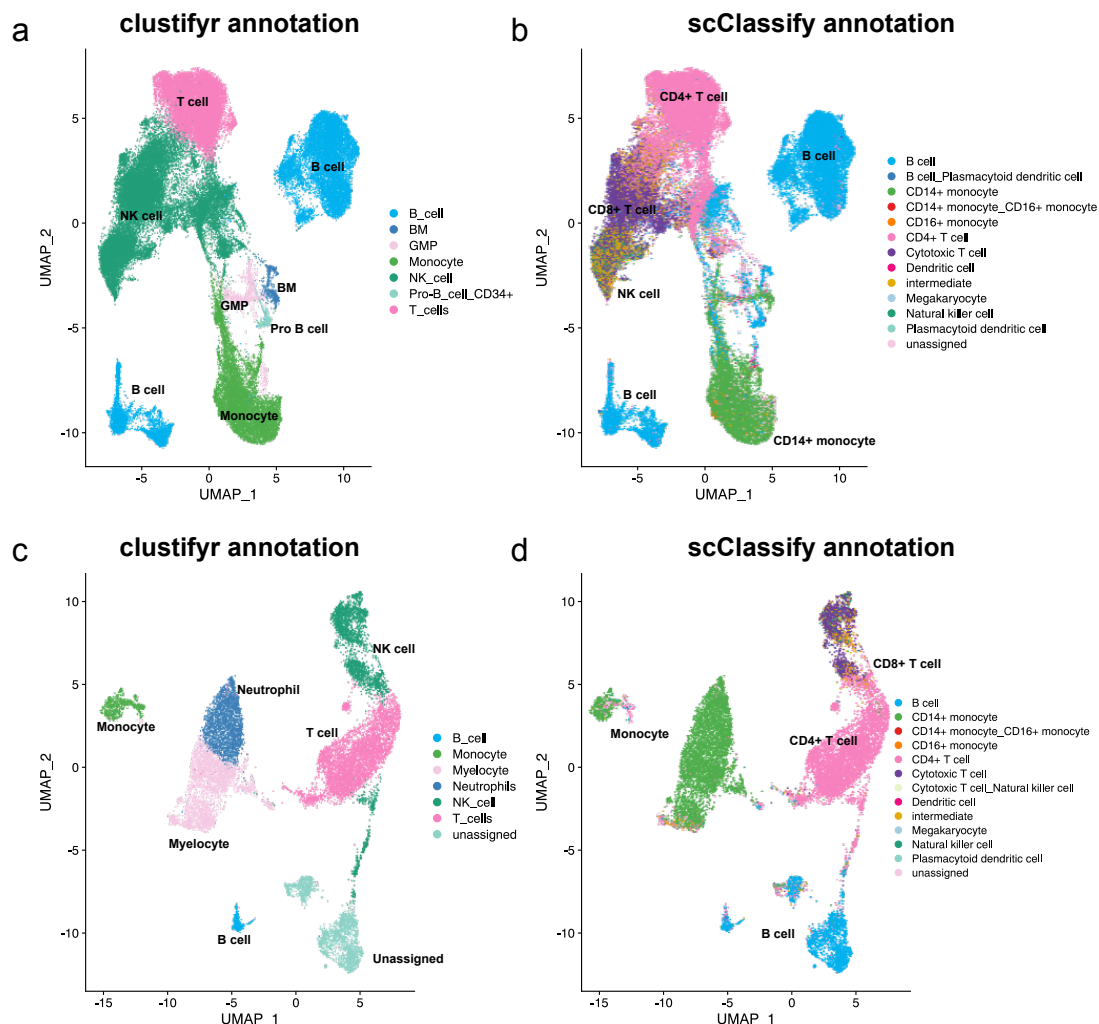


Figure 6.2: Automated annotation of scRNA-seq cell clusters, using the R packages clustifyr and scClassify in combination with reference datasets. [a, b] newly-diagnosed MM, [c, d] relapsed MM. The output of automated packages clustifyr and scClassify is used to aid cell type annotation. Clustifyr assigns a cell type to each cell cluster, whilst scClassify assigns a cell type to each individual cell. Clustifyr was used in conjunction with the HumanCellAtlas reference. scClassify was ran using a pre-trained model trained on seven PBMC single cell RNA-seq datasets. Both references originate from healthy data, therefore neither are able to identify the myeloma cell population.

These packages give a good starting point for more detailed manual annotation. However, currently they are not a complete substitute for manual annotation

using underlying biological knowledge of cell markers. For example, both references originate from healthy tissue, therefore the myeloma cluster could not be identified using clustifyr, scClassify and these healthy references alone. Both packages either incorrectly labelled the MM cluster as B cells, or were unable to assign any cell type to the myeloma cluster with any confidence and left it unassigned. Thus, the MM cluster had to be identified manually using known MM pathological markers.

Figures 6.3 and 6.4 show how the MM cell clusters were identified. From MM cells, you would expect to see expression of CD38, but lower expression than in normal plasma cells. You would expect to see high expression of the pathological marker CD138 in MM cells, as well as high expression of SLAMF7, BCMA, KRAS, IGKC and IGL2, however these are not exclusive to the MM cluster. You would expect to see little or no expression of CD45 and CD19 in the MM cluster and reduced expression of CD20 in MM cells compared to normal B cells. Using the expression of these markers, clusters 2, 7 and 13 were identified as the MM cell population in the newly diagnosed dataset; and cluster 4 was identified as the MM cell population in the relapsed MM dataset.

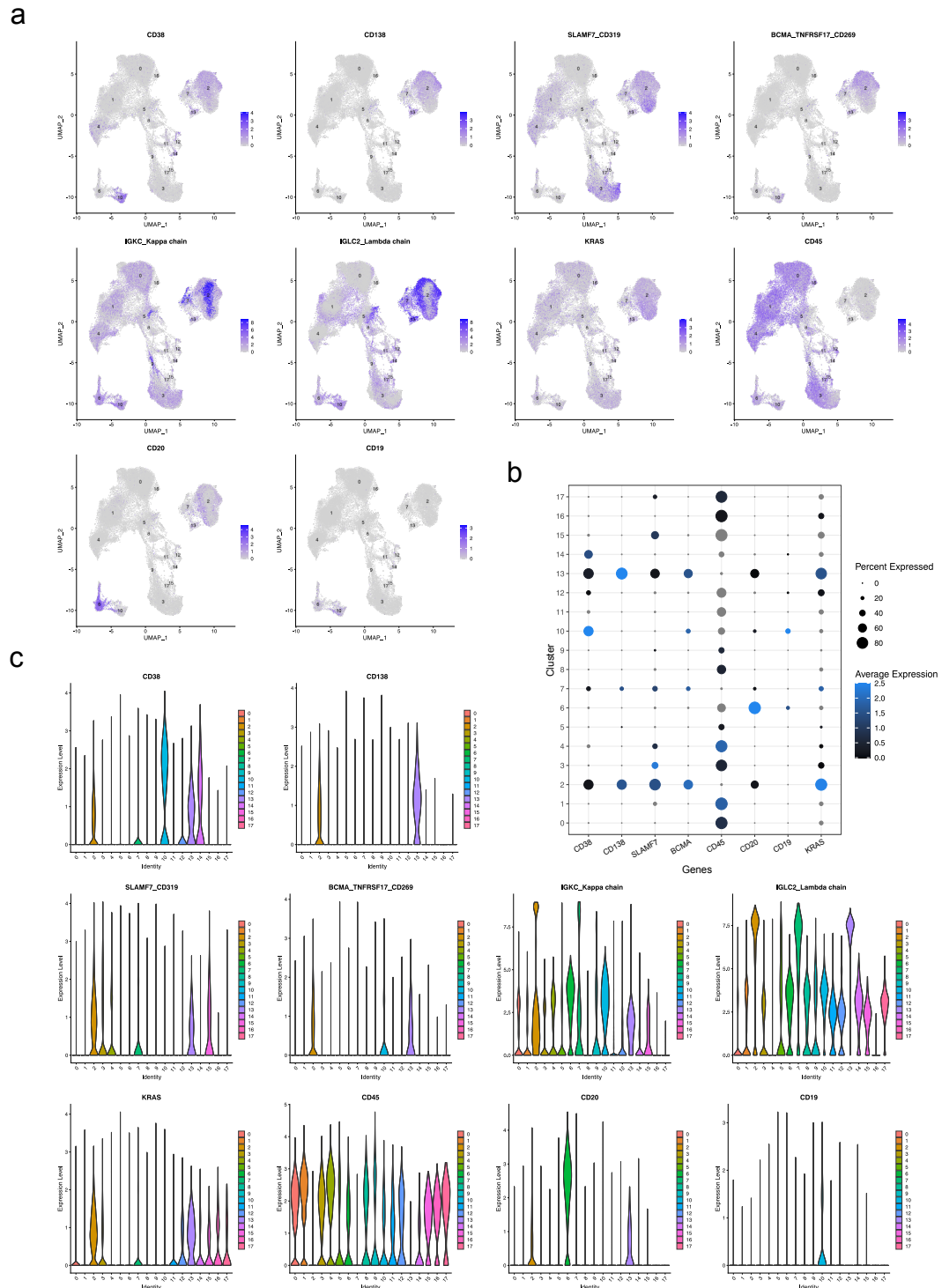


Figure 6.3: Manual annotation of multiple myeloma (MM) cell clusters in experiment 1 and 2 (newly-diagnosed patients) using known MM biological markers. a) Seurat UMAP feature plots showing MM marker expression. Grey indicates no expression and purple indicates expression. b) A dot plot showing for each cluster the percentage of cells expressing a given marker and the average expression. c) Violin plots showing gene expression for each cluster.

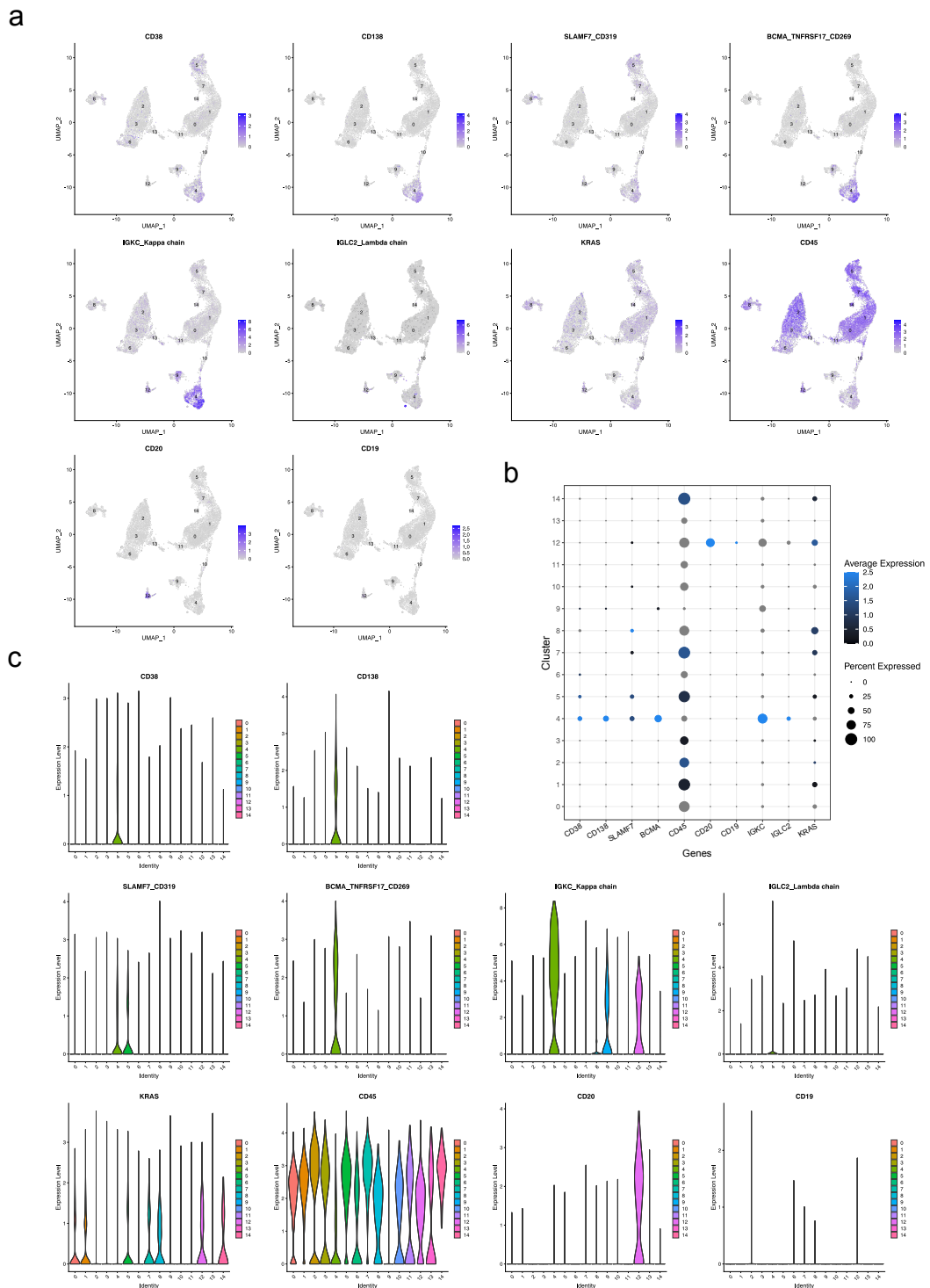


Figure 6.4: Manual annotation of MM cell clusters in experiment 3 (relapsed MM) using known biological markers. a) Seurat UMAP feature plots showing MM marker expression. Grey indicates no expression and purple indicates expression. b) A dot plot showing for each cluster the percentage of cells expressing a given marker and the average expression. c) Violin plots showing gene expression for each cluster.

Marker type	Expressed/ over-expressed	Not expressed/ reduced expression
Multiple myeloma cells	CD138, CD38 (lower than plasma cells), SLAMF7, BCMA, KRAS, IGKC, IGCL2	CD20, CD19, CD45
Normal plasma cells	CD38, CD19, some BCMA	place
B cells	CD20, some CD19	b down
T cells	TRAC, CD3D	cd4 down
Cytotoxic cells	GZMH, GZMB, GZMA, PRF1	cd4 down
CD4+ T cells	CCR7, SELL, TCF7 and T cell markers	cd4 down
CD8+ T cells	CD8A, cytotoxic markers and T cell markers	cd 8 down
NK cells	KLRB1, KLRF1 and cytotoxic markers	nk down
Dendritic cells	dc up	dc down
Monocytes	mono up	mono down
etc...	else	else down

Table 6.2: Manual annotation markers for cell types originating from transcriptomic profiles of bone marrow samples. SLAMF7, BCMA, KRAS, IGKC and IGCL2 are very highly expressed by MM cells, but are not exclusive to this cluster.

6.3 Results

6.3.1 Newly-diagnosed MM

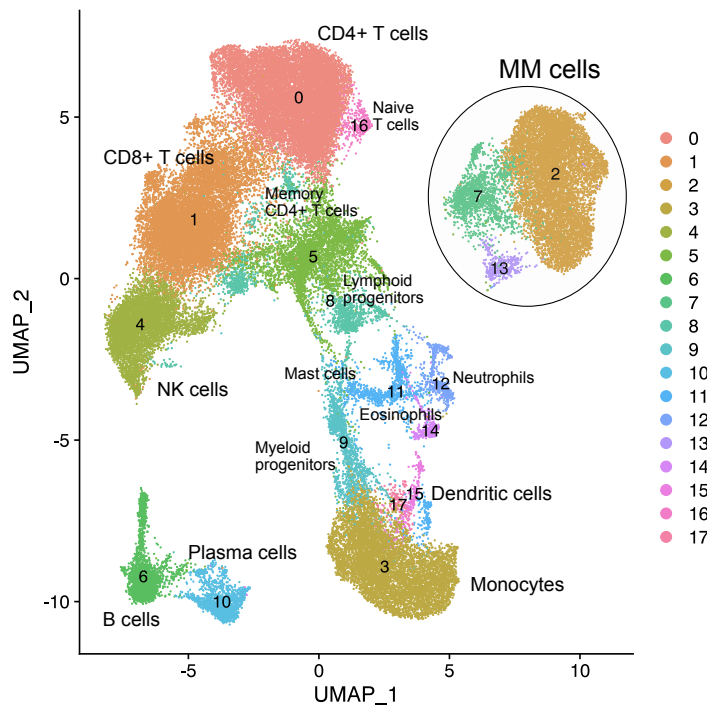


Figure 6.5: Fully annotated UMAP clustering analysis of two newly-diagnosed multiple myeloma (MM) patients. The MM cell population (circled), consists of three distinct clusters.

Figure 6.5 shows final cell-type annotation for the newly-diagnosed MM integrated dataset. 18 distinct clusters were identified using Seurat embeddings. The expected major immune clusters were identified (such as B cells, T cells, NK cells and myeloid cells.) Using the established MM biological markers (shown in figure 6.3), three distinct MM clusters (2, 7 and 13) were identified.

Composition

Cluster composition analysis by treatment condition is shown in figure 6.6. Halofuginone treatment at both 24 and 48 hours reduced the proportion of cells in the MM cluster ($p<0.00001$) compared to DMSO, for both experiment 1 and 2. NCP26 treatment at 24 hours reduced the proportion of cells in the MM cluster ($p<0.00001$), for both experiment 1 and 2. HF and NCP26 treatment increase the proportion

of cells in the CD4+ T cell, CD8+ T cell and B cell clusters. Together with dose response curves and cell death assays, this suggests that Halofuginone and NCP26 are selectively killing MM cells to a higher degree than other cell types.

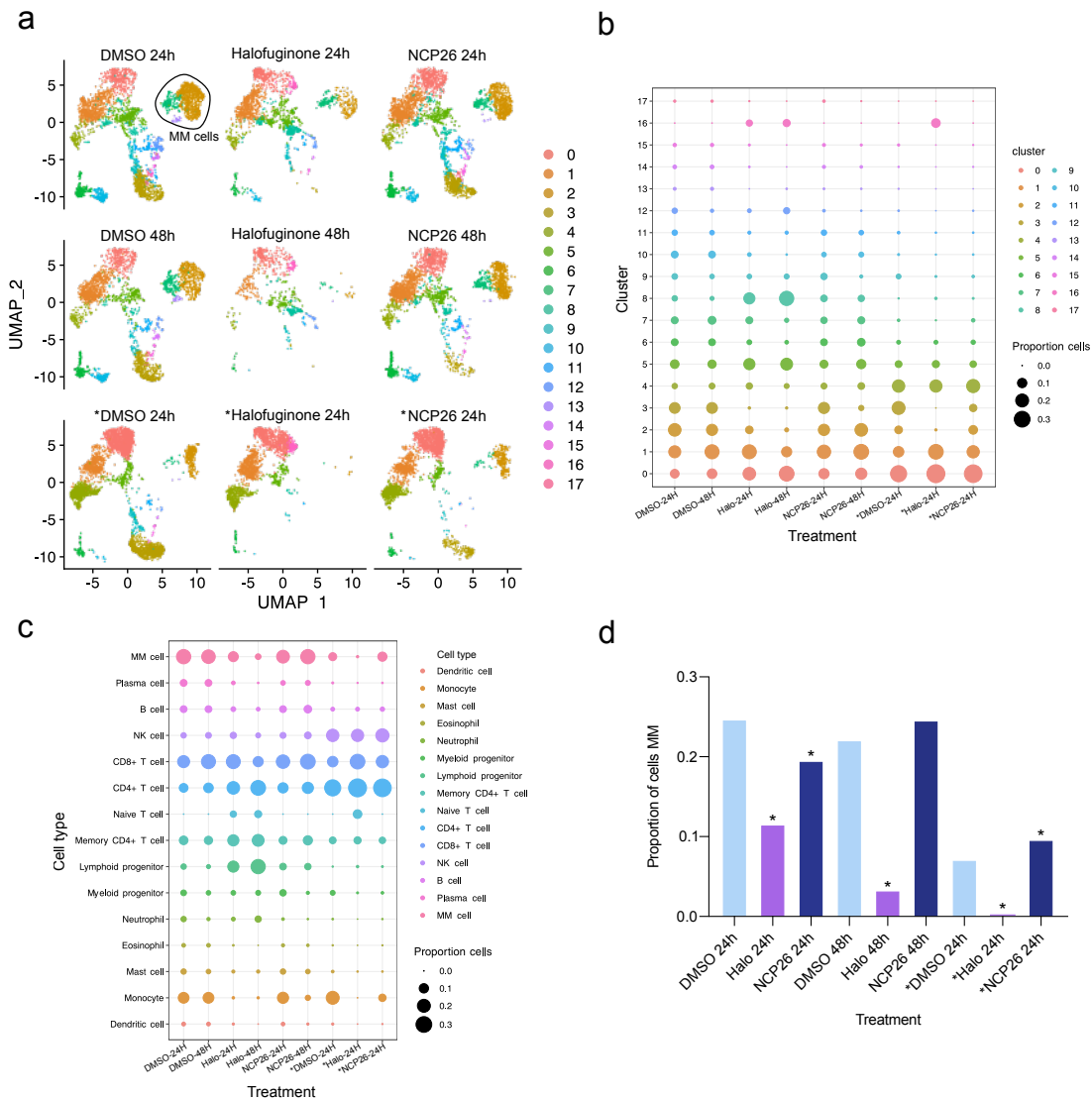


Figure 6.6: Composition analysis of newly diagnosed MM. a) UMAP cell composition plots separated by treatment condition. b) Dot plot showing proportion of cells in each cluster for each sample. c) Dot plot showing proportion of cells in each cell class for each sample (as labelled in Figure 6.5). d) The proportion of cells in the MM cluster only (stars above bars indicate significant at $p<0.01$ compared to corresponding control). Sample names starting with asterisks originate from experiment 1, no asterisk indicates experiment 2 origin.

Halofuginone treatment and 48-hour NCP26 treatment were also found to reduce the proportion of cells in the monocyte and neutrophil cluster ($p<0.00001$),

indicating they may have some off-target effects on myeloid cells.

All of the compounds were used at a concentration of 1 μ M. This is approximately 10 times the concentration of Halofuginone's IC₅₀ value on MM cell lines and two times NCP26's IC₅₀ value. As you can see, large number of cells were killed by Halofuginone treatment at both 24 and 48 hours. This could be a dosing issue

Differential expression

Next, differential gene expression was investigated using Seurat's FindMarkers function. At 48 hours there is substantial cell death, therefore the 24-hour time point will be mainly focussed on.

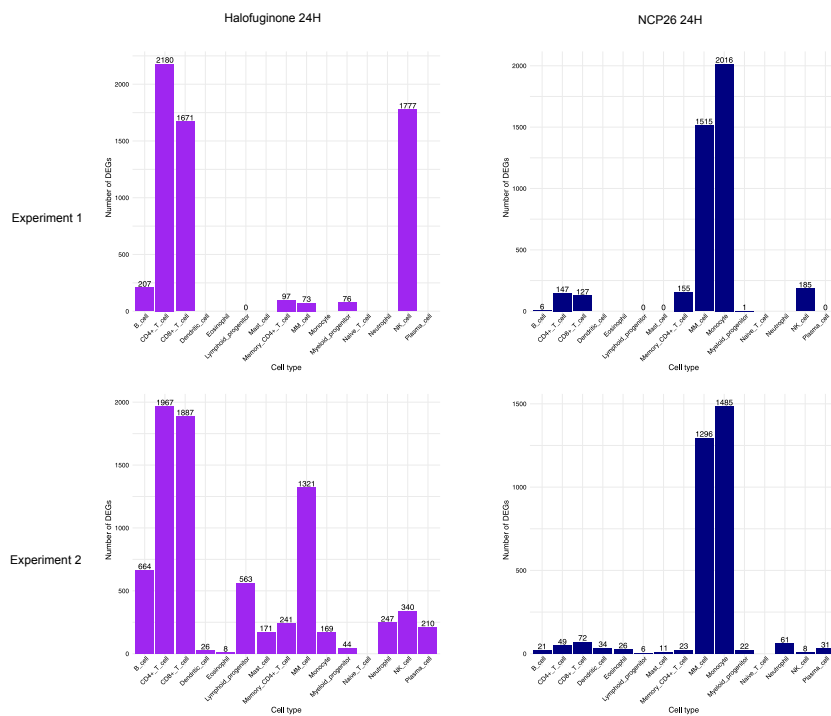


Figure 6.7: Number of differentially expressed genes (DEGs; $p_{adj} < 0.05$) broken down by cell type for two newly-diagnosed MM patients treated with ProRS inhibitors (Halofuginone and NCP26) for 24 hours. Cell type annotation corresponding to figure 6.5. Experiment 1 and experiment 2 denote separate experiments, each containing BM samples from different newly-diagnosed MM patients.

Following 24-hour 1 μ M NCP26 treatment, 1515 genes were differentially expressed (DE; $p_{adj} < 0.05$) in the myeloma cell population in experiment 1, and 1294 genes in experiment 2, compared to DMSO. Figure 6.7 shows the breakdown of

DEGs per cell type for NCP26 and HF treatment. 24 hour 1 μ M NCP26 treatment had very little transcriptional effect on many of the other immune cell types, for example T, B and NK cells, where the number of DEGs is more than 10 times smaller than for MM cells. This corroborates the composition analysis, where MM cells seem more sensitive to ProRS inhibition than other immune cell types. However, there were 2016 and 1485 genes DE in the monocyte cluster following NCP26 treatment in experiment 1 and 2, respectively. This reflects the composition analysis, where the proportion of cells in the monocyte cluster was markedly reduced. This could indicate that NCP26 is not be selective for MM cells over myeloid cells.

In MM cells, 73 genes were DE following Halofuginone treatment in experiment 1, and 1318 genes in experiment 2. This reflects results seen in the composition analysis (figure 6.6), whereby we saw very few cells remaining in the MM cluster for experiment 1. Due to the low cell number, there is less statistical power and diversity across the cells, therefore you would likely see fewer statistically significant DEGs for this cluster. Unlike NCP26 treatment, for 24 hour 1 μ M HF treatment we see a large number of DEGs in B, T and NK cell clusters. This likely reflects the differences in potency between NCP26 and HF. HF is approximately five times more potent than NCP26. This fits with composition analysis, where HF showed a great deal of cell killing in the myeloid and MM cluster.

At lower doses (i.e. NCP26 at 1 μ M) of ProRS inhibition we see clear evidence of greater transcriptional effects on MM cells and monocytes over other immune subtypes. However at higher doses (i.e. Halofuginone at 1 μ M) we see substantial cell killing of MM cells and monocytes and larger transcriptional effects on other immune subtypes. This together with the composition analysis, proves NCP26 and Halofuginone are selective for MM cells over most healthy immune cells, except for monocytes. However, this could be a problem with the doses used for treatment. 1 μ M is almost 10 times greater than Halofuginone's IC₅₀ and two times greater than NCP26's IC₅₀. This experiment should be performed at a lower concentration to ascertain if NCP26 and Halofuginone are more selective for MM cells or monocytes. Ideally, the experiment would be performed over the course of

three days at a lower concentration, as in our cell line studies. However, human bone marrow samples do not last very well in extended culture, therefore acute treatment was the preferred method.

Pathway analysis

Pathway enrichment analysis was performed for the top 200 upregulated and top 200 downregulated DEGs in the MM cluster, following Halofugininone and NCP26 treatment (figure 6.8).

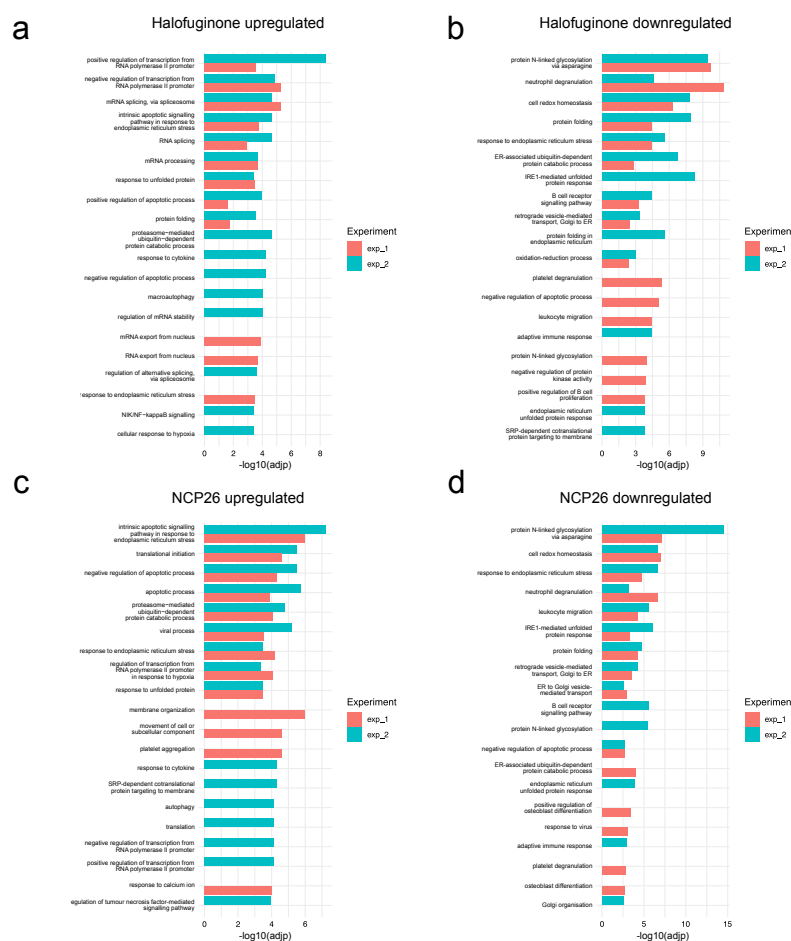


Figure 6.8: Pathway analysis (Gene ontology biological processes; GOBP) of the MM cluster. a) and b) Halofugininone treatment. c) and d) NCP26 treatment. a) and c) GOBP pathway analysis performed using top 200 upregulated DEGs ($p_{adj} < 0.05$ and $\log_2FC > 0$) ranked by fold change. b) and d) GOBP pathway analysis performed using top 200 downregulated DEGs ($p_{adj} < 0.05$ and $\log_2FC < 0$) ranked by fold change.

For NCP26 and Halofuginone treatment of MM cells, pathways related to ER-stress and apoptosis are enriched. The unfolded protein response- another member

of the ISR, which shares many effectors with AAR- is also enriched. This supports our bulk RNA-seq results that the AAR response is activated following NCP26 and Halofuginone treatment.

Figure 6.9 shows a more detailed exploration of AAR genes and their expression in MM cells, as well as the immune microenvironment. Figure 6.9a shows *ATF4*/

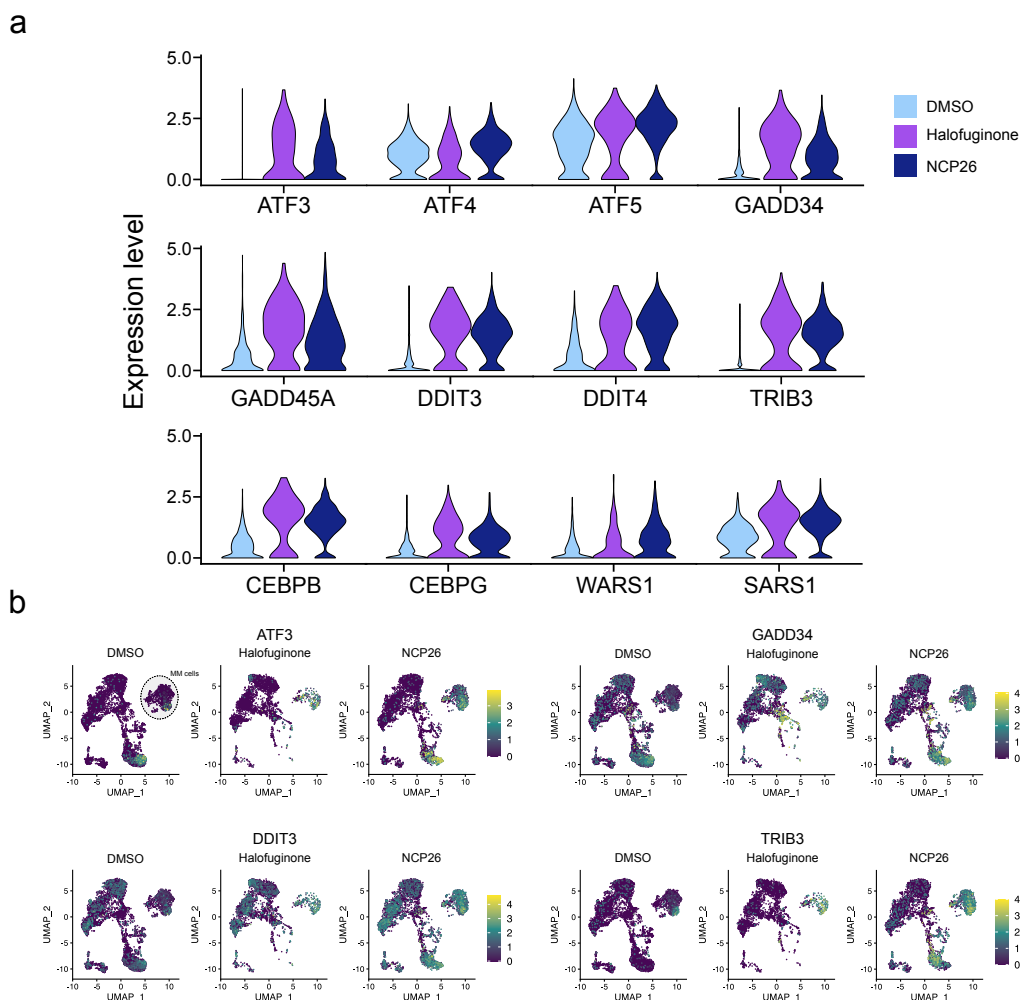


Figure 6.9: A selection of differentially expressed amino acid starvation response (AAR) genes following 24 hour treatment with ProRS inhibitors (newly-diagnosed MM patients). a) Violin plots showing expression of selected AAR genes in the MM population following DMSO, Halofuginone and NCP26 treatment. b) Feature plots of UMAP clustering, showing gene expression of AAR genes *ATF3*, *GADD34*, *DDIT3* and *TRIB3*. Feature plots are split into three panels based on treatment condition (DMSO, Halofuginone and NCP26). The myeloma cell population is circled in the first panel.

target genes' expression increased in HF and NCP26 treated samples compared

to DMSO in MM cells. tRNA aminoacyl synthetase genes (*WARS1* and *SARS1*) are also over-expressed in MM cells following ProRS inhibitor treatment. Figure 6.9b shows selected AAR genes' expression for the entire UMAP clustering plot, separated by treatment condition. Markedly increased expression of *ATF3* and *TRIB3* can be seen localised to MM cells and monocytes for HF and NCP26 treatment. Increased expression of *DDIT3* and *GADD34* can be seen across a few different clusters, however it is most pronounced in the MM and monocyte clusters. Additionally, the apoptotic mediator *TNFRSF10B*, positively regulated by *DDIT3*, is upregulated in MM cells and monocytes (figure 6.10b). Indicating that, although aaRSs are ubiquitous enzymes and ProRS inhibitors cause some degree of activation of the AAR in all cells, they are cytotoxically more selective for MM cells, and lead to an apoptotic cascade of events, preferentially over other cell types. *CDKN1A* is also overexpressed in MM cells following NCP26 and HF treatment. *CDKN1A* is a target gene of *ATF4* and arrests cell cycle progression by inhibiting the activity of cyclin-dependent kinases. HF has previously been shown to cause cell cycle arrest and accumulating cells in the G₀/G₁ phase of cell cycle. Moreover, figure 6.10 demonstrates that NCP26 and HF have profound anti-MM effects on MM cells. NCP26 and HF treatment markedly reduce expression of MM pathological marker *CD138* in MM cells. Additionally NCP26 and HF treatment reduced expression of *IGKC* and *IGLC2*, the genes coding for the constant regions of immunoglobulin light chains, Kappa and Lambda. Previously these genes have been implicated in MM outcome.

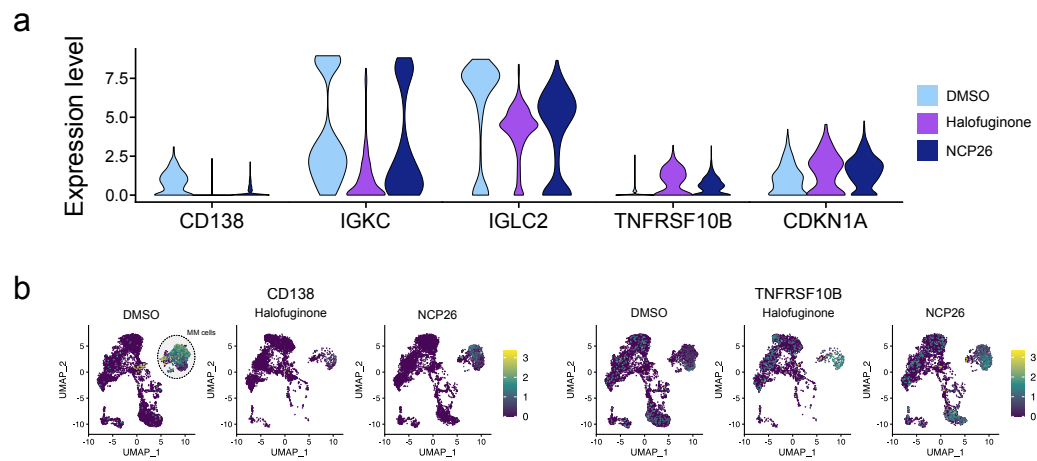


Figure 6.10: A selection of differentially expressed multiple myeloma (MM) and cell cycle/apoptotic markers in newly diagnosed patients treated with ProRS inhibitors for 24 hours. a) Violin plots showing expression of selected genes in the MM population following DMSO, Halofuginone and NCP26 treatment. b) Feature plots of UMAP clustering, showing gene expression of MM pathological marker *CD138* and apoptotic marker *TNFRSF10B*. The myeloma cell population is circled in the first panel. Halofuginone and NCP26 treatment reduce *CD138* expression and increase *TNFRSF10B* expression in the MM clusters.

6.3.2 Relapsed MM

Figure 6.11 shows final cell-type annotation for the integrated relapsed MM patients. 15 distinct clusters were identified. The expected major immune clusters were identi-

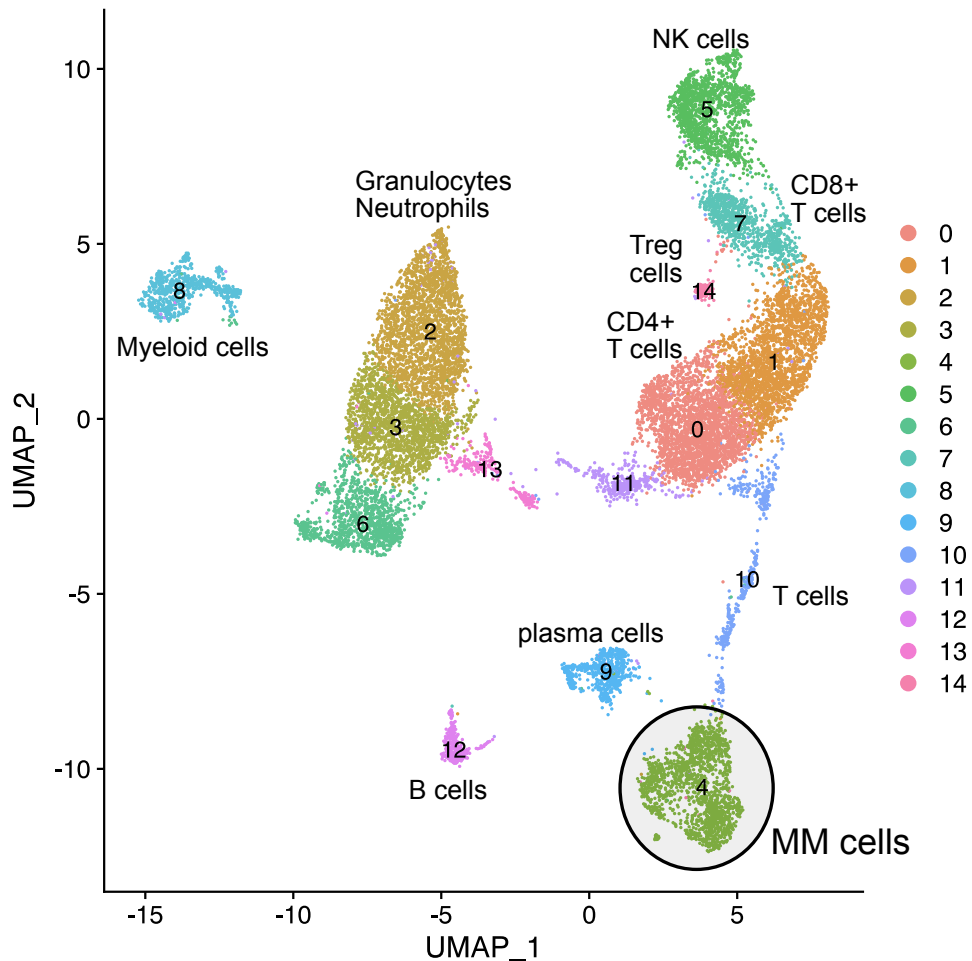


Figure 6.11: Fully annotated UMAP clustering analysis of two relapsed multiple myeloma (MM) patients. 15 distinct clusters were identified, one of which was identified as the MM population (circled).

fied (such as B cells, T cells and myeloid cells.) Using the established MM biological markers (shown in figure 6.4), one distinct MM cluster (cluster 4) was identified. The relapsed patients show substantial transcriptional differences to treatment-naïve patients, in both their myeloma cells and in their normal immune cells.

Composition

Cluster composition analysis for relapsed MM by treatment is shown in figure 6.12. 1 μ M and 5 μ M NCP26 treatment reduced the proportion of cells in the MM

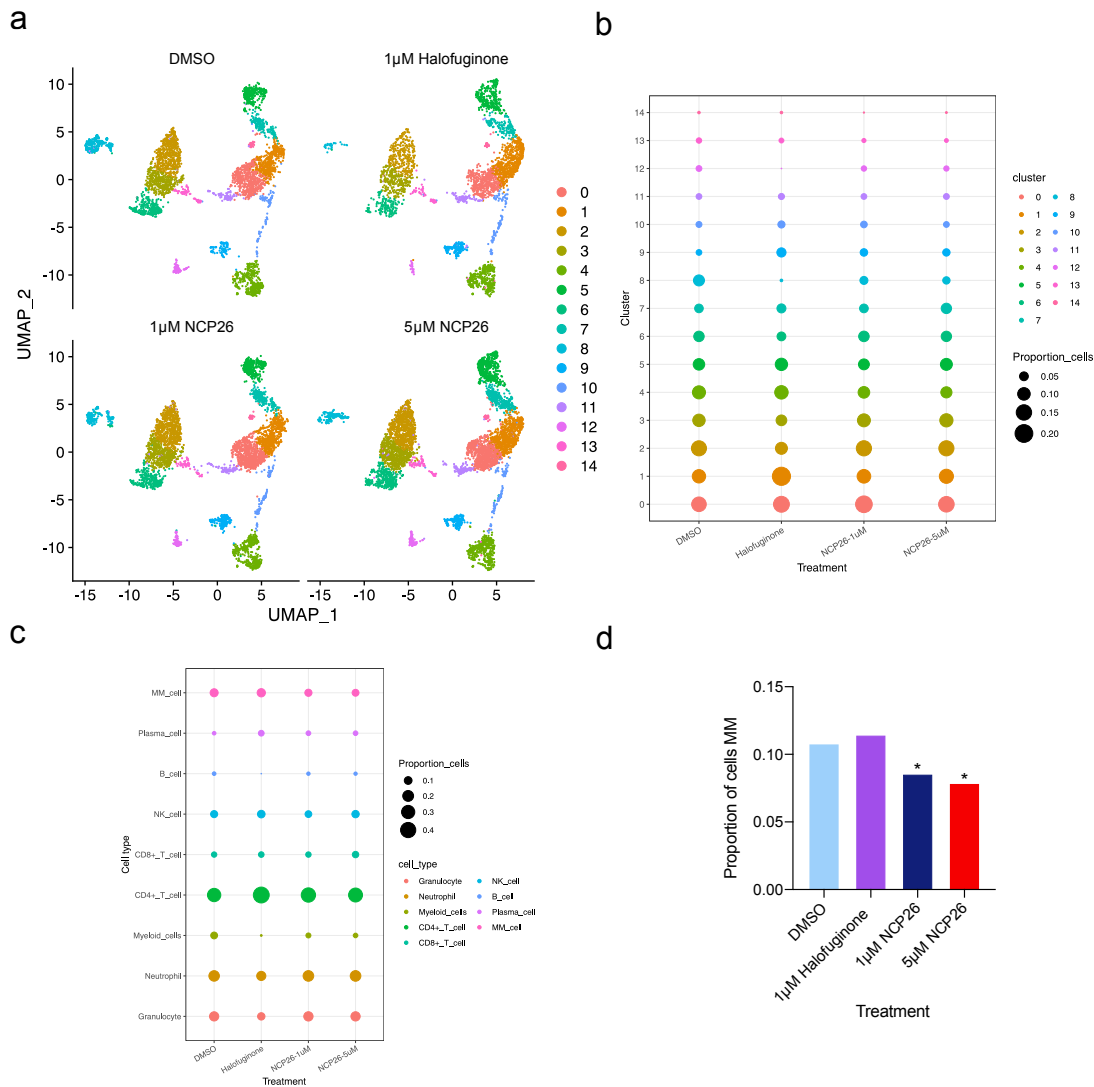


Figure 6.12: Composition analysis of relapsed MM cells treated for 24 hours with ProRS inhibitors. a) UMAP cell composition plots separated by treatment condition. b) Dot plot showing proportion of cells in each cluster for each sample. c) Dot plot showing proportion of cells in each cell class for each sample (as labelled in Figure 6.11). d) The proportion of cells in the MM cluster only (stars above bars indicate significant at $p < 0.01$ compared to DMSO control). NCP26 treatment reduces the proportion of cells in the MM cluster ($p < 0.01$).

cluster compared to the DMSO control. There is insufficient evidence ($p > 0.05$) to conclude if Halofuginone treatment affected the proportion of cells in the MM cluster.

However, Halofuginone significantly reduced the proportion of cells in the myeloid and B cell clusters. This perhaps indicates that in relapsed MM, Halofuginone is not selective for myeloma cells over other immune cells. This may indicate an advantage of NCP26 and proline non-competitive ProRS inhibitors in relapsed myeloma.

Differential expression

Next, differential gene expression was investigated for the relapsed MM patients.

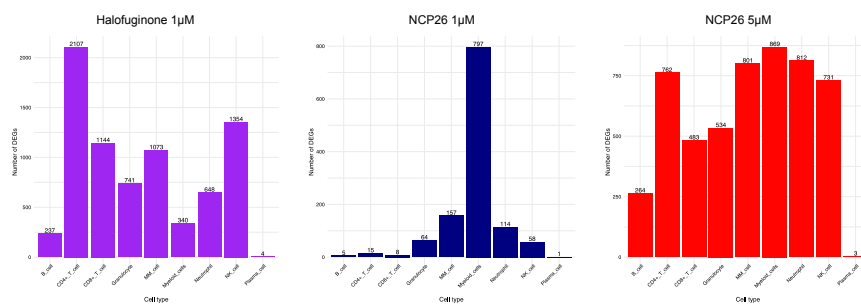


Figure 6.13: Number of differentially expressed genes (DEGs; $p_{adj} < 0.05$) broken down by cell type for a relapsed MM patient (patient 4 only) treated with ProRS inhibitors (Halofuginone and NCP26) for 24 hours. Cell type annotation corresponding to figure 6.11.

1073 genes were DE in the MM cell cluster following 24-hour 1 μ M HF treatment. 157 and 801 genes were DE in the MM cell cluster following 24-hour 1 μ M and 5 μ M NCP26 treatment, respectively. The breakdown of DEGs per cell type for NCP26 and HF treatment is shown in figure 6.13.

In contrast to the newly-diagnosed MM data, many of the other immune subtypes are differentially expressed to similar degrees as MM cells by ProRS inhibition. Additionally, the myeloid cell cluster has more DEGs than MM cells for 1 μ M and 5 μ M NCP26 treatment. Myeloid cells have fewer DEGs than MM cells for HF treatment, however figure 6.12 demonstrates substantial cell killing of myeloid cells by HF treatment, therefore you may not expect as many statistically significant DEGs for the few cells remaining in the cluster. This may indicate that NCP26 and HF are not as selective for MM cells for relapsed patients as they are for newly-diagnosed MM.

6.3.3 Monocyte/macrophage analysis

6.4 Myeloma bone marrow classifier

7

tRNA-seq analysis of PRS inhibitors

7.1 Introduction

7.1.1 Experiment overview

In parallel with mRNAs

7.2 Data processing

7.2.1 Quality control and filtering

7.3 Results

7.3.1 tRNA results

7.3.2 mRNA results

7.3.3 Multi-omics factor analysis

7.4 PRS inhibition general discussion

8

Epigenetic reversal of drug resistance in MM

8.1 Introduction

8.1.1 Experiment overviews

8.2 Dose response curves

8.3 Bulk RNA-seq

8.4 Single-cell RNA-seq

8.5 Proteomics

8.5.1 CyTOF

8.5.2 Total proteomics (LC-MS/MS)

8.5.3 Phosphoproteomics

8.5.4 Ubiquitinomics

8.6 ATAC-seq

8.7 Multi-omics factor analysis

8.8 TRIM24i general discussion

Appendices

A

Supplementary figures

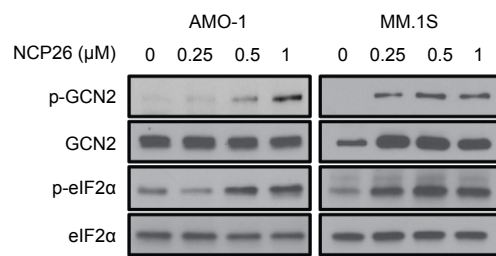


Figure A.1: Western blot demonstrating dose-dependent NCP26 canonical ISR activation with GCN2 and eIF2 α phosphorylation. AMO-1 and MM.1S myeloma cell lines used.

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