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Elucidating the Novel Role for Core Binding Factor beta in Osteosarcoma Protein
Translation

By

NICHOLAS ALEXANDER OLDBERG

DISSERTATION

Submitted in partial satisfaction of the requirements for the degree of

DOCTOR OF PHILOSOPHY

in

Pharmacology and Toxicology

in the

OFFICE OF GRADUATE STUDIES

of the

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Approved:

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ABSTRACT OF DISSERTATION

Osteosarcoma (OS) is the most common primary bone malignancy in humans and canines, and in humans primarily affects younger patients 10-14 years of age. While considerable efforts have been put forth in new therapeutic approaches to this disease, the treatment and prognosis for OS has changed very little since the 1980s. Targeted therapeutics have made considerable progress in other cancer types, leveraging characteristics of cancer cells which differentiate them from that of normal healthy cells. In comparison to other cancers, OS is highly heterogeneous, and no single unifying driver mutation has yet been found. Development of a therapy which could overcome the high degree of heterogeneity amongst OS tumors could go a long way in improving the lives of OS patients. Targeting protein translation has been proposed as one mechanism by which to overcome tumor heterogeneity, and this dissertation focuses on studying a noncanonical role of core binding factor beta (CBF β) as a regulator of protein translation, and elucidating whether this could represent a potential therapeutic target for OS.

Utilizing a wide array of *in vitro* assays, we have been able to demonstrate that loss of CBF β reduces protein expression of RUNX2 in a post-transcriptional manner, and this decrease in RUNX2 protein level is not fully explained by alterations in RUNX2 stability brought about by loss of its binding partner CBF β . Additionally, we demonstrate that loss of CBF β also causes a decrease in global protein translation, and confirmed an interaction between CBF β and hnRNPk which has thus far only been observed in breast cancer cells. Importantly, this interaction with hnRNPk is said to be the mechanism by which CBF β influences protein translation, and our results corroborate those observed in breast cancer cells and suggest CBF β may also perform this role in OS.

Reports of the interactions between CBF β and hnRNPk or RUNX2 allude to mutual exclusivity in interaction, and with the transcriptional role of CBF β accomplished via binding to RUNX proteins, and the translational role of CBF β accomplished via binding to hnRNPk, it is entirely possible these two roles are antagonistic in some fashion. To investigate the relevance of certain CBF β residues in terms of this translational role of CBF β , and avoid confounding variables from the transcriptional role of CBF β , we utilized point mutations to interrupt CBF β -RUNX2 interaction. Using various in vitro assays, we validated key residues of CBF β which are involved in its interaction with RUNX2, re-introduced this mutant form into CBF β knockout cells, and measured alterations to RUNX2 interaction and nuclear shuttling. We confirmed that our mutant displays reduced binding to RUNX2, and drastically reduced nuclear shuttling.

Lastly, we expanded our studies from RUNX2 to the entire genome and proteome. Encouraging data thus far had suggested CBF β may play a role in protein translation, and necessary next steps were to assess which proteins CBF β may be interacting with in performance of this role, and elucidate which proteins may be under the translational purview of CBF β . Using immunoprecipitation mass spectrometry we identified numerous specific interactors of CBF β , with high enrichment in pathway analysis terms associated with protein translation. Additionally, using two different methods we generated a list of proteins which may be under the translational purview of CBF β , and found strong enrichment of numerous cancer-associated terms among this list.

These studies establish that CBF β participates in protein translation in OS, with many genes under its purview associated strongly with cancer in general, and OS specifically. This provides justification for future studies delving deeper into this novel role of CBF β , and opens up another mechanism by which protein translation could be targeted therapeutically in OS.

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I would like to thank my mom for being so supportive and involved at every step of my educational career. She worked tirelessly to find scholarships I could qualify for, which facilitated my travel to COSMOS summer camp at UC Irvine, attendance at SAT prep programs, and helped pay for my college tuition. Her commitment to me getting the best education possible played a tremendous role in my development as a scientist and as a person overall. I would also like to mention how much of a boost the frozen meals she always sent home with me were. There were many times during my graduate studies when her homemade cooking was what gave me that extra push to keep going.

I would like to thank my dad for his encouragement throughout my time in graduate school. He has made a strong effort to understand the specifics of my research to fully grasp the weight of the triumphs and pitfalls in my journey. This has meant the absolute world to me, and it has been a joy sharing updates with him as I have progressed through graduate school.

While growing up, I faced several health challenges and was diagnosed with juvenile rheumatoid arthritis at a very young age. My dad found a program called Healthy Families, which allowed us to afford my prescription for a new drug that had just come on the market, Enbrel. This health program and Enbrel changed my life in two significant ways: health-wise, they gave me the ability to move my joints freely and return to the physical activities I loved so dearly; and career-wise, witnessing the miraculous transformation in my body brought about by Enbrel inspired me to pursue pharmacology as a field of study.

I would also like to thank my grandma, Sara Oldberg, for supporting my passion for science and providing me with such interesting reading material when I was growing up. She was an avid reader and would always give me her *Discover* magazines after she was done reading them. Discussing these articles with her are memories I will forever cherish, and they deepened my resolve that becoming a scientist was the career path for me.

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DEDICATION

I would like to dedicate this work to all scientific researchers—past, present, and future—and celebrate their tenacity in pushing the envelope and bringing new discoveries that benefit humanity. As scientists, we truly stand on the shoulders of giants, and the work of today is only possible because of the strong commitment of those who came before us.

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Chapter 1

Literature Review and Project Rationale

Osteosarcoma

Incidence and Prognoses

Osteosarcoma (OS) is the most common form of primary bone cancer in humans and canines (1), with ~1,000 new cases diagnosed per year in the United States (2,3). The age of diagnosis with OS has a bimodal age distribution, with the first, and largest, peak in the 10-14-year-old age group, with the second for patients aged 65+ (4). The early age of presentation in patients makes OS somewhat of an outlier in the field of oncology, as the average age at diagnosis for all cancer patients is 66 years (3). OS can present in a variety of bones in the body, however the lower long bones are the most common, comprising $\frac{3}{4}$ of all early-onset diagnoses (2). Within the bone, OS most often occurs at the metaphysis (4), the portion of bone between the diaphysis (middle shaft) and epiphysis (rounded end) which contains the growth plate.

The lung is far and away the most common site of metastasis in OS patients, and 20-30% of patients already harbor clinically detectable metastatic lesions at time of diagnosis (5,6). Importantly, for metastases to be clinically detectable they must be large enough to be seen by imaging techniques, resulting in small lesions often remaining undiscovered. This presents a clear limitation, and the fraction of OS patients harboring metastatic lesions at time of diagnosis is likely far higher, closer to ~50% by some estimations (7). When presenting with localized disease at time of diagnosis, patients with OS have an estimated five-year survival rate of 60-80% (8), but for those with clinically detectable metastases this plummets to ~20% (6,9).

Treatment Options

Advancements in mechanisms to treat OS have lagged behind that of cancer in general, and the current treatment regimen has remained essentially unchanged over the past 30 years (10). The standard treatment of OS is so-called MAP therapy, standing for Methotrexate, Adriamycin, and cisPlatin, with or without ifosamide (11). When introduced in the 1980's, MAP therapy represented a considerable improvement in treatment of OS. Prior to this, the standard approach to OS was amputation, which was not only gruesome, but sadly yielded a paltry 5-year survival rate of ~17% for patients with OS (12). Although nearly four decades have passed since its introduction, MAP therapy remains as the most effective treatment for OS (13).

This lack of new approved therapies for OS is not due to a lack of effort, as numerous approaches leveraging a wide range of mechanisms have been attempted to treat this disease. These include proteasome inhibition (14–16), immunotherapies (17,18), T cell engaging bispecific antibodies (19), anti-IL-2 monoclonal antibodies (mAb) (20), oncolytic viruses in solo (21) or in combination with another therapy (22), or modulation of innate immunity (18,23). Additionally, drugs synergistic or additive to existing MAP therapy have been evaluated, such as anti-CD47 mAb (24) or Sorafenib (25) in combination with Doxorubicin, and PPAR γ agonists (26), natural products such as curcumol (27), or Eicosapentaenoic acid (28) in combination with Cisplatin.

Disease Profile

In contrast to other sarcomas, OS is not typified by a specific translocation or genetic mutation, and instead presents as an assortment of widespread chromosomal abnormalities (29). Osteosarcoma tumors can possess a large range of chromosome number, ranging from haploid all the way up to hexaploid (30,31). In lieu of gain of function mutations in oncogenes, or loss of

function mutations in tumor suppressors, OS is driven more so by changes in copy number of the genes themselves. This occurs by way of genomic amplification occurring in chromosomal regions containing oncogenes, and copy number losses occurring in regions harboring tumor suppressor genes (29).

In terms of common alterations observed in OS, these include amplification of oncogenes MYC (32–35) and mouse double minute 2 homolog (MDM2) (33,36–38) and loss of tumor suppressors tumor protein p53 (TP53) (36,37), retinoblastoma protein (RB1) (32,36,37), and cyclin-dependent kinase inhibitor 2A (CDN2A) (32,36,37). Acquisition of these genetic abnormalities in OS is likely to occur by a variety of mechanisms, such as point mutation during DNA replication (39) or aneuploidy due to errors in cell division (40,41). Additional to these canonical methods, a new mechanism, chromothripsis, has recently been identified (29). In chromothripsis, tens to hundreds of individual genomic rearrangements can occur in a single event, resulting in a genetic catastrophe for the cell (29,42). This mechanism was estimated by the authors to occur in at least 2-3% of all cancers, but in the words of the authors, is “particularly common in bone cancers”, reaching a prevalence of ~25% (42).

This genomic landscape makes development of therapies for OS challenging, as the lack of *bonafide* driver mutations in OS (43) leaves few specific differences between tumor and healthy tissue which could be leveraged therapeutically. The heterogeneity of OS places it at a relative disadvantage when compared to other cancers which possess strong genetic links to disease. Salient examples of targeted therapeutics in other cancers which have utilized disease-specific mutations include imatinib (44,45) which targets the BCR-ABL fusion gene in chronic myelogenous leukemia (46), EGF/human EGFR (HER) targeting therapies such as osimertinib in

non-small cell lung cancer and breast cancer (47–49), and the wide variety of development targeting RAS mutations in many cancers (50–54).

Cancer therapeutics do not have to target a specific mutation in order to be effective, and could instead target proteins which cancer cells are highly reliant upon for survival. This has been employed to great success with the IMiD drugs Lenalidomide and Mezigdomide, which function by co-opting the ubiquitin proteasome system (UPS) to degrade Ikaros and Ailos (55–57). These two transcription factors are necessary for the growth and survival of multiple myeloma cells (58,59), and their degradation by these drugs results in the selective killing of myeloma cells. Therapeutics leveraging the UPS have been explored in OS (60), but studies on this are very limited. This relatively new modality to treat disease could provide potential in OS, however before debating between mechanisms by which to attack a target, it is of course necessary to identify a viable target in the first place.

Core Binding Factor Transcriptional Complex

RUNX2 and CBF β Overview

RUNX2 is a member of the RUNX family of DNA-binding transcription factors, which are co-activated by interaction with another protein, CBF β . RUNX2 is known as the master regulator of bone growth and differentiation, and has critical roles in chondrocyte maturation and osteoblast differentiation (61). Core binding factor beta (CBF β) acts as a binding partner to the RUNX family of transcription factors, and serves as a transcriptional co-activator. When bound to CBF β , RUNX proteins have a higher affinity for DNA (62,63).

In normal conditions, RUNX2 plays a pivotal role in the formation of bones and teeth. RUNX2^{-/-} mice are devoid of intramembranous bones and bone collar formation (64), and RUNX2^{-/-} calvarial cells are incapable of differentiating into osteoblasts (65). In addition to playing a pivotal role in development of the skeleton and teeth, RUNX2 is also expressed in other tissues such as the ovaries, testis, brain, and B cells in the blood (66,67). CBF β is also crucial for skeletal development, a discovery only possible via a CBF β -GFP ‘knock-in’ approach(68–70). Homozygous disruption of CBF β is embryonic lethal in mice, with death occurring ~12 days post coitum from central nervous system hemorrhaging and disruption of liver hematopoiesis (71,72).

RUNX2 is de-regulated in OS (73), and high expression levels of RUNX2 (74,75) or CBF β (76) are each correlated with poor response to chemotherapy, and poor overall prognosis, in OS. Additionally, many of the gene targets of RUNX2 have been linked to poor therapeutic outcome in OS (77,78), such as vascular endothelial growth factor (VEGF) (79,80), alkaline phosphatase (ALP) (81), and matrix metalloproteinase 9 (MMP-9) (82). These data brought RUNX2 and CBF β into our focus in studying OS, and spurred questions about whether the actions of these proteins could be interrupted as a therapeutic target.

Mutations in RUNX2 and CBF β

Mutations in the RUNX proteins and/or CBF β can have significant consequences to health. Alterations in RUNX2 are strongly implicated in cleidocranial dysplasia (CCD) (83), a rare condition which presents as developmental difficulties in teeth and bone. CCD is predominantly caused by haploinsufficiency of RUNX2 (84–86), and mutations of RUNX2 are found in 60-70% of CCD cases (87). These mutations are commonly found in the Runt Homology Domain (RHD) (88), and typically result in inhibited binding of RUNX2 to DNA (89) or to CBF β (83). Many

cases of CCD with no genomic alteration to RUNX2 were found to be caused by pathogenic variants of CBF β , underscoring how critical RUNX2 and CBF β are in bone formation (90). Mutations in RUNX2 and CBF β are also seen in ~25% of all *de novo* acute leukemias, and are the most commonly disrupted genes in leukemia in humans (91). RUNX1 and CBF β are mutated in a form of acute myeloid leukemia (AML) known as CBF AML, in which chromosomal rearrangements lead to the RUNX1/RUNX1T1 and CBF β -MYH11 (also known as CBF β -SMMHC) fusion proteins (92). CBF β -SMMHC is formed by the fusion of the heterodimerization domain of CBF β to the coiled-coil domain of Smooth Muscle Myosin Heavy Chain (93,94). Expression of this fusion protein leads to dysregulation of hematopoietic development, likely resulting from CBF β -SMMHC binding RUNX1 with higher affinity than wild-type CBF β (93). RUNX1/2 and CBF β are also found to be frequently mutated in breast cancer (95,96).

Therapeutic Targeting of RUNX2 and CBF β

Efforts to target RUNX2 and CBF β in the context of OS have generated encouraging data, as siRNA knockdown of RUNX2 in OS cells decreases colony formation and cell invasion (97), and knockdown of CBF β in OS cells reduces cell proliferation, migration, and invasion (76). RUNX2 has been targeted therapeutically via inhibition of DNA binding by small molecule, with clinical trials currently underway (98). Inhibition of the RUNX proteins has also been proposed as a way to enhance the sensitivity of myeloma drugs which leverage the UPS (99). Ro5-3335 is an inhibitor of CBF β -RUNX1 interaction and has shown promise *in vitro* and *in vivo* against leukemias containing translocations or inversions at CBF β or RUNX1 genes (100), and siRNA knockdown of RUNX2 in OS cells increases sensitivity to Doxorubicin (101). Pharmacological inhibition of interaction of CBF β with the RUNX proteins has been explored in leukemia and

breast cancer (102,103) and inhibition of CBF β -RUNX2 interaction has been proposed as a novel therapeutic target in osteosarcoma (104).

Structural Biology of RUNX2

RUNX proteins are typified by the Runt homology domain (RHD) (105), a conserved domain key in their interaction with DNA (106). Located close to the N terminus, this domain contains the PyGPyGGTPy consensus sequence, which is essential for their binding with DNA as well as CBF β (107). Next to the RHD is the glutamine-alanine (QA) repeat region, important in transactivation (108). The nuclear localization signal (NLS), a sequence of nine amino acids next to the RHD, governs translocation of RUNX proteins between cytoplasmic and nuclear compartments (67), and RUNX2 is predominantly found in the nucleus (109). The C-terminus of RUNX2 is also crucial in RUNX2 function, with deletion of the C-terminus phenocopying that of RUNX2 null and resulting in complete bone loss (110,111). Within the C-terminus lies the proline-serine-threonine-rich (PST) region, which possesses both transactivatory and transinhibitory functions (67,111). The matrix targeting signal (NMTS) can be found within the PST region, and works together with the NLS to govern RUNX2 nuclear localization (112). The final five amino acids within the PST region, known as VWRPY, function as a transcriptional repression domain (113,114).

Regulation of RUNX2 and CBF β Activity

Numerous proteins and processes are implicated in regulation of RUNX2 expression and activity. In terms of expression level, RUNX2 levels are controlled by a combination of

acetylation, deacetylation, and ubiquitination (115). RUNX2 levels normally oscillate with the cell cycle, but this oscillation is disrupted in OS cells (116) due to de-regulation of proteolytic degradation (117). The regulation of RUNX2 with the cell cycle is thought to be mediated by cdc2-mediated phosphorylation of RUNX2, and RUNX2 has been implicated in entry in and exit out of cell cycle phases G2/M (118).

RUNX2 is normally degraded by the proteasome, however under stressful conditions it can also be trafficked to the lysosome by SOX9 (119). SMURF1, an E3 ligase, has been reported to regulate RUNX2 levels (120,121), and RUNX2 levels are also regulated by the CK2/HAUSP pathway (122). HDAC4/5 deacetylate RUNX2, allowing Smurf1-mediated degradation of RUNX2 to take place (115), and Akt enhances the stability of RUNX2 by regulating Smurf1 (123). WW domain-containing oxidoreductase (WWOX) has been implicated as a suppressor of RUNX2 levels and activity (124), and through this mechanism may determine the aggressive phenotype of OS (125). Hes1 stabilizes RUNX2 (126), and CBF β has also been reported to also stabilize the RUNX proteins (127). According to published data, binding of CBF β to RUNX proteins and subsequent protection from degradation is less pronounced in RUNX2 than in RUNX1 or RUNX3, possibly due to it binding RUNX2 with lower affinity (128). Hif-2 α has been proposed to compete with CBF β for interaction with RUNX2, and thereby downregulate RUNX2 activity via removing stabilization by CBF β (129).

As the canonical role of CBF β is that of a co-activator to the RUNX proteins, little is published about the regulation of CBF β activity specifically. While RUNX2 levels oscillate during the cell cycle in healthy cells, levels of CBF β remain more consistent (117). CBF β can be found in both the nucleus and the cytoplasm (130), but cannot freely move between compartments on its own. CBF β does not possess a nuclear localization signal (NLS) (131,132) and is carried into the

nucleus via piggybacking onboard the RUNX proteins (130,133). Binding of CBF β to Filamin A prevents binding to the RUNX proteins, and thereby retains it in the cytoplasm (134). Subcellular localization of CBF β has also been postulated to involve CRLZ-1 (also known as SAS10 or UTP3) which interacts with CBF β (135) and may play a role in nuclear shuttling (136,137) and OS development at large (138).

As it stands today, the heterogeneity of OS patient tumors remains a significant hurdle in developing therapies to treat this population (43). In order for the wide array of mutations in OS to manifest as advantageous to cancer cells, these mutated genes must be expressed as protein. Targeting protein translation has been proposed as a way of overcoming tumor heterogeneity, as protein translation acts as a convergence point of multiple signaling pathways (139). A useful approach, in theory, would be to affect this mechanism in such a way that is preferential towards proteins implicated in cancer, so as to avoid inhibiting translation of beneficial proteins and thereby injuring healthy cells. Recently, CBF β has been proposed to participate in protein translation (130), which may provide a new mechanism to affect protein translation in OS.

Protein Translation

Overview

Proteins are translated by two main mechanisms; cap-dependent, and cap-independent translation. Cap-independent translation relies on an internal ribosomal entry site (IRES), and is utilized under conditions where cap-dependent translation is compromised (140–142). These conditions include endoplasmic reticulum (ER) stress, hypoxia, mitosis, or nutrient limitation, and mRNAs containing IRES elements often encode proteins which are involved in recovering from

or maintaining function during these states (143). Outside of these conditions, cap-dependent translation is by far the most commonly used, and is estimated to be employed in the translation of ~95% of total cellular mRNAs (144). RUNX2, like most proteins in the cell, is expressed via cap-dependent translation under normal conditions (145), however it may rely on IRES-mediated cap-independent translation during cellular stress (146).

Cap-dependent translation is used in the expression of mRNAs with highly structured 5' un-translated regions (UTR's), as these structures must be broken down to allow ribosomes to bind to mRNA and begin protein translation (147,148). During cap-dependent translation initiation, Eukaryotic Translation Initiation Factor 4E (eIF4E) binds to the 7-methylguanosine (m7G) cap of mRNA's and initiates assembly of the translation initiation complex and subsequent direction of ribosomes to mRNA. Following binding of eIF4E, eIF4G binds and stabilizes eIF4E at the 5' cap (149). This is followed by binding of eIF4A, a helicase which unwinds 5' UTR structures (150). Together, these proteins comprise the eIF4F translation initiation complex (TIC). Although least abundant of the translation initiation factors, eIF4E is indispensable to this process as recruitment of complex proteins to the m7G cap is the rate-limiting step of the translation process (151).

Also important in cap-dependent translation are eIF4E binding proteins (4E-BPs), proteins which bind to eIF4E and function to inhibit protein translation (149,152). 4E-BPs are regulated by the mechanistic target of rapamycin complex 1 (mTORC1), which causes their phosphorylation. These proteins compete with eIF4G for a binding site on eIF4E, and phosphorylation of 4E-BPs prevents their interaction with eIF4E (152). 4E-BPs are important in translational regulation, and some repress translation of a large range of transcripts while others are restricted to smaller subsets of mRNAs by also associating with RNA binding proteins (153).

Cap-Dependent Translation in Cancer

Cap-dependent translation has garnered interest in the context of cancer, and is employed in the translation of many oncogenes implicated in OS such as MYC, Fibroblast Growth Factor 2 (FGF-2), Platelet Derived Growth Factor (PDGF), Vascular Endothelial Growth Factor (VEGF), Survivin and cyclins (154). In fact, nearly all potentially oncogenic pathways intersect at the eIF4F complex (155,156), and cap-dependent translation is commonly upregulated in most human malignancies (147). In addition to the machinery of the cap-dependent complex being essential to synthesis of many identified oncogenes (154), de-regulated expression patterns in the translation components themselves have also been implicated in OS, and cancer in general.

eIF4E expression is elevated in OS cell lines and patient samples compared to nonmalignant control cell lines and healthy bone tissue (157,158), and is further elevated in OS patients harboring distant metastases compared to those with local disease (158). This correlation between eIF4E expression and poor outcome was challenged by another study (159), although these data relied upon a four point scale of expression level rather than a more quantitative metric such as western blotting or qRT-PCR. As eIF4E is expressed at a low level, and this protein is the rate-limiting factor for translation initiation, a small shift in its expression could have drastic effects on translation activity. Importantly, an increase in quantity or activity of eIF4E does not necessarily lead to globally elevated protein translation, but rather an increase in translation of a subset of mRNAs (153). This subset includes mRNAs possessing extensive secondary structure (160) and oncogenes of particular interest in OS such as MYC, FGF and VEGF (161,162). Additionally, 4E-BPs, the negative regulators of eIF4E and therefore protein translation, are reported to exert tumor suppressive activity (163), further implicating the EIF4F complex in cancer.

Therapeutic targeting of members of the cap-dependent translational complex has shown encouraging results in OS, although the total number of studies are limited. When used in conjunction with conventional chemotherapeutics or as solo therapies, small molecule inhibitors (157) or micro RNA's (158) affecting eIF4E have shown promising data in decreasing OS cell proliferation, migration, and invasion *in vitro* and *in vivo*. Positive results have also been shown in OS by inhibitors targeting eIF4A (164,165), the helicase responsible for unwinding 5' UTR mRNA secondary structures. Additionally, use of rapamycin has demonstrated effectiveness as an inhibitor of metastasis of OS cells *in vivo* by interrupting mRNA translation (166).

Numerous papers mention that despite decades of research revealing much of the mechanism of protein translation in eukaryotes, our understanding of this process as a whole remains incomplete (167). Recently, Malik et al put forth evidence from breast cancer cells reporting that CBF β may perform a noncanonical role, as a regulator of cap-dependent protein translation (130). CBF β reportedly exerts this role by binding to mRNA's via hnRNPk, a multifunctional protein which participates in transcription and translation, and enhances the translation of these mRNA's through eIF4B (130). Additionally, CBF β interaction with RUNX proteins or hnRNPk may be mutually exclusive, since cytoplasmic CBF β did not bind to RUNX1, and nuclear CBF β did not bind to hnRNPk, although all three proteins were present in both the nucleus and cytoplasm (130). CBF β performs its transcriptional role in the nucleus of the cell, while this translational role is reported to occur in the cytoplasm, suggesting that shuttling of CBF β into the nucleus by the RUNX proteins (131) could in some way alter its activity in the cytoplasm. If CBF β also regulates cap-dependent protein translation in OS cells, this could provide a new therapeutic target to affect protein translation in OS. As protein translation acts as a convergence

point of multiple signaling pathways (168), targeting CBF β could allow a mechanism to overcome the high degree of heterogeneity among OS patient tumors (43).

Project Rationale and Justification

OS is a highly heterogeneous cancer type, making development of targeted therapies challenging. Enhancing our understanding of mechanisms to overcome OS patient heterogeneity could greatly advance OS therapeutic development, and targeting protein translation has been put forth as one method of accomplishing this. Protein translation is an indispensable process in cells, and decades of research into this process have provided an understanding of the machinery which regulates and facilitates this process. Recent reports of CBF β performing a non-canonical role, as a regulator of protein translation, are intriguing as they may represent a new target through which protein translation could be affected. Thus far, this noncanonical role of CBF β has only been observed in breast cancer cells, and many questions still remain in understanding the broad and specific impacts of this role. This dissertation aims to study whether CBF β also performs this role in OS cells, and to elucidate the significance of this role in the context of OS at large. In generating CBF β knockout cell lines, we noticed a decrease in RUNX2 protein expression, similar to the CBF β knockout-induced decrease in RUNX1 expression observed in breast cancer cells. We hypothesized that CBF β may regulate RUNX2 levels in a post-transcriptional manner, and loss of CBF β may cause alterations in protein translation within the cell. This hypothesis is tested in **Chapter 2 (CBF β regulates RUNX2 protein levels in osteosarcoma cells)**. This chapter evaluates changes in RUNX2 expression induced by loss of CBF β via qPCR and western blotting, and studies whether re-introduction of CBF β can rescue low RUNX2 levels. Next, it utilizes western blotting coupled with proteasome and translation inhibitors to study potential changes in

RUNX2 stability brought about by loss of CBF β . Finally, this chapter describes the influence of CBF β loss on global protein translation, and investigates a potential interaction with hnRNPk that underpins this novel role of CBF β in breast cancer cells.

Based upon the results obtained from Chapter 2, we next attempted to interrupt the binding of CBF β to RUNX2. Previous data had suggested stabilization of RUNX2 by interaction with CBF β did not fully explain the reduction in RUNX2 protein level, but we could not rule stabilization out as contributing somewhat to this reduction. Additionally, as interaction of CBF β with RUNX proteins or hnRNPk may be mutually exclusive and occur in separate cellular compartments (130), it's possible the translational and transcriptional roles of CBF β are in some way antagonistic. Possible competition for the existing pool of CBF β taking place between RUNX proteins and hnRNPk would make individual study of each role challenging. Furthermore, knockout of one of the RUNX proteins leads to compensatory up-regulation in other members of the RUNX family (169,170). Due to these confounding effects, we opted to generate a mutant form of CBF β , with inhibited RUNX protein binding, and use this as a tool to further study the translational role of CBF β . This is tested in **Chapter 3 (Mutations in the RUNX2 binding pocket of CBF β alter its cellular activity in osteosarcoma cells)**. This chapter details recombinant protein studies used to validate residues of CBF β key in RUNX2 interaction, and subsequently generate a mutant form of CBF β incorporating mutations at these residues. Following this, we re-introduced our mutant back into CBF β knockout cells, and used western blotting, cellular electro-thermal shift assay (CETSA), and nuclear and cytoplasmic fractionation to understand the influence of our mutations on the behavior of CBF β in OS cells. We demonstrate that CBF β residues G61, N63, and N104 are important for interaction with RUNX2, and mutation of these residues to alanine results in decreased interaction with RUNX2. Additionally, we showcase that

this reduced interaction with RUNX2 prevents shuttling of CBF β into the nucleus. Finally, we demonstrate that re-introduction of our mutant form of CBF β into CBF β knockout cells was unable to recover low RUNX2 protein expression. This could be due to decreased stability of RUNX2 as a result of inhibited CBF β binding, or perhaps that these same residues utilized for RUNX2 binding are also important for the translational role of CBF β , and through this role CBF β stimulates production of RUNX2 protein.

Up until this point, we had demonstrated that CBF β regulates RUNX2 expression in a post-transcriptional manner, and loss of CBF β leads to reduced global protein translation. Furthermore, we had validated residues of CBF β key in interaction with RUNX2, which led to the generation of a CBF β mutant with reduced RUNX2 binding which can be used as a tool in future studies. With encouraging data thus far, we had two main questions: 1) which proteins does CBF β interact with in performance of this noncanonical role? and 2) which genes/proteins are under the translational purview of CBF β ? Data generation until this point had been in a targeted manner, measuring RUNX2 specifically, but in our quest to reveal the full scope of CBF β approaches casting a wider net were needed. **Chapter 4 (Profiling the interactome and translome of CBF β in osteosarcoma cells)** delves into these questions, describing our findings using more comprehensive assays. We utilized immunoprecipitation mass spectrometry (IP-MS) of CBF β , both wild-type (WT) and mutant (3xMut), to reveal specific interactors of CBF β , and understand which of these are dependent upon G61, N63, and/or N104 residues of CBF β for this interaction. IP-MS followed by gene ontology (GO) revealed CBF β to potentially interact with many proteins associated with protein translation, the vast majority of which have not been previously reported. We next used direct detection of biotinylated proteins (DiDBiT) and Ribo-Seq to understand which proteins experience an increase in transcription, translation, overall production, and translational

buffering in the presence of CBF β . Encouragingly, not only did these assays identify a multitude of proteins under the translational purview of CBF β , these proteins demonstrated high enrichment of terms and pathways linked to cancer such as Rho GTPase-associated proteins, VEGF signaling, PD-L1 and MAPK, as well as enrichment of the specific term “pathways in cancer”.

This dissertation has two main goals: the first is to expand our understanding of the interplay between loss of CBF β and reduction in RUNX2 protein expression, and whether this suggests CBF β involvement in a post-transcriptional mechanism. The second goal is to investigate this post-transcriptional mechanism, and gain insight into what proteins CBF β interacts with, and regulates, in OS cells.

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Chapter 2

CBF β regulates RUNX2 protein levels in osteosarcoma cells

Abstract

Development of novel therapies to treat OS has lagged behind much of the field of cancer research, due in large part due to the heterogeneity of OS and lack of identified driver mutations. High expression of RUNX2, the master regulator of bone growth and differentiation, is implicated in OS, and the transcriptional function of RUNX2 is enhanced by CBF β . In the process of profiling a CBF β knockout (KO) cell line developed for another project, we identified a reduction in RUNX2 expression upon loss of CBF β . We sought to unravel the mechanism underlying this and enhance our understanding of the biology of OS at large. To this end, we evaluated the loss of CBF β and how this effects RUNX2 mRNA and protein levels, RUNX2 stability, and global protein translation. Additionally, we assessed a protein-protein interaction between CBF β and hnRNPK previously demonstrated in breast cancer, and proposed to be key to a putative noncanonical role performed by CBF β . Finally, we conducted a rescue experiment reconstituting CBF β back into CBF β KO cells and evaluated alterations to RUNX2 levels. Loss of CBF β caused a reduction in RUNX2 protein expression, however RUNX2 mRNA levels were unaffected. This decrease in RUNX2 protein levels was not sufficiently explained by a change in RUNX2 stability from loss of its binding partner. CBF β was confirmed to interact with hnRNPK and loss of CBF β resulted in a decrease in global protein translation. Lastly, reconstitution of CBF β into CBF β KO cells rescued low RUNX2 protein expression. These results together demonstrate that CBF β may play a role in protein translation in OS, and could present a novel therapeutic target in treatment of this disease.

Introduction

Osteosarcoma (OS) is a highly aggressive primary bone cancer, and the most prevalent of this cancer type in humans and canines (1). OS most frequently presents early in life, affecting primarily children and adolescents (2), which places it in stark contrast to the average age of all cancer diagnoses of 66 years (3). OS Patients presenting with localized disease at time of diagnosis have an estimated 5-year survival rate of 60-80% (4), yet this plummets to ~20% for those presenting with clinically detectable metastases (5,6). Despite numerous advances in the field of cancer research as a whole, the current treatment outlook for osteosarcoma has remained essentially unchanged over the past 30 years (7). The highly aggressive nature of osteosarcoma combined with its early age of presentation and lack of therapeutic advancement represents a critical unmet need in cancer research today.

Core binding factor beta (CBF β) functions as a binding partner to the RUNX family of DNA-binding transcription factors (RUNX1-3) and acts as a transcriptional co-activator by allosterically increasing their affinity to DNA (8,9) . RUNX2 is known as the master regulator of bone growth, playing a critical role in osteoblast proliferation and differentiation (10). In the context of OS, high expression levels of RUNX2 (11,12) or CBF β (13) have each demonstrated correlation with poor disease prognosis. siRNA knockdown of RUNX2 in OS cells increases sensitivity to Doxorubicin (14) and leads to reduced colony formation and cell invasion (15), while siRNA knockdown of CBF β in OS cells causes reduced cell proliferation, migration, and invasion (13). RUNX2 modulates the expression of genes such as vascular endothelial growth factor (VEGF) (16,17), alkaline phosphatase (ALP) (18), and matrix metalloproteinase 9 (MMP-9) (19), all of which have also been linked to poor therapeutic outcome in OS (20,21). The array of data

implicating RUNX2 in OS, combined with CBF β functioning as a co-activator to RUNX2, led to the proposal of the CBF β -RUNX2 interaction as a novel therapeutic target in OS (22).

This project began in simply studying this interaction, but as additional data was gathered, the scope expanded from studying this interaction to evaluating the roles of CBF β itself. A recent study in breast cancer proposed a novel role for CBF β as a regulator of protein translation (23), and as we advanced in profiling our CBF β KO cell line, our data began to corroborate this possibility in OS as well. Protein translation acts as a convergence point of multiple signaling pathways (24), and modulating this process therapeutically could provide a mechanism to target cancers with wide mutational profiles. This is especially interesting in the context of OS, being one of the most heterogeneous forms of cancer studied (25). These parallels between our data and that in breast cancer inspired us to intensify our focus on investigating if CBF β performs this non-canonical role in OS, and if so, to what extent does this influence the malignant phenotype of OS overall.

Proteins can be translated via a cap-dependent or cap-independent mechanism, with the former garnering particular interest due to oncogenes identified in OS such as MYC, fibroblast growth factor 2 (FGF-2), platelet derived growth factor (PDGF), vascular endothelial growth factor (VEGF), survivin and cyclins being translated in this manner (26). This has led to research targeting cap-dependent translation in OS (27–29), which is the same pathway CBF β had been suggested to regulate in breast cancer cells (23). It is important to note that the study of this novel role of CBF β in breast cancer is still in its infancy, and the overall impact of CBF β on breast cancer is still a point of contention. Loss of CBF β transformed breast cancer cells and led to increased tumor formation in one study (23), while loss of CBF β was found to inhibit formation of metastases in another study (30). While there may not be as much conflict regarding the broad impact of CBF β

in OS, as high expression is known to correlate with poor overall survival (13,31), there is currently no data available regarding the role of CBF β in protein translation in OS. This study represents the first of its kind interrogating this novel function, with the goal of enhancing our understanding of OS overall and potentially leading to identification of a novel therapeutic target.

Materials and Methods

Cell Lines and Culture Conditions

Parental U2OS (American Type Culture Collection [ATCC], Manassas, Virginia) and U2OS-derived cell lines were maintained in McCoy's 5A (Iwakata & Grace Modification) media supplemented with 10% FBS (Corning Inc., Glendale, Arizona) and U/mL Penicillin-Streptomycin (P/S) (Corning) at 37C in a humidified atmosphere with 5% CO₂ unless otherwise specified. HEK293T cells (ATCC) were maintained in DMEM media (Corning) supplemented with 10% FBS and 1% P/S in the same atmosphere as previously listed cells. Cell populations were maintained in T75 flasks (Corning) and were split twice weekly with Trypsin (Gibco, Thermo Fisher Scientific, Waltham, Massachusetts) upon reaching 70-80% confluence and discarded after 20 passages.

Knockout Generation

CBF β knockout (KO) cell lines were generated in the parental human U2OS wild-type (wt) cell line. Briefly, the Synthego multi-sgRNA CRISPR/Cas9 ribonucleoprotein complex (Gene Knockout Kit v2, Synthego, Redwood City, CA) was transfected into 80,000 cells with Lipofectamine CRISPRMAX (Invitrogen, Carlsbad, CA) for 24-hours. The media was replaced with fresh supplemented media and cells were incubated for two additional days. A limited dilution (0.8 cells/well) was performed in a 96-well plate format and monitored for individual cells to produced individual colonies that were selected for sequencing. DNA was isolated using a genomic DNA purification kit (Promega, Madison, WI) from each viable clone and was submitted to Genewiz (Azenta Life Sciences, South San Francisco, CA) for Sanger sequencing. Sequencing

trace files were analyzed using the Synthego Interference of CRISPR Edits (ICE) Analysis Tool (Synthego Performance Analysis, ICE Analysis. 2019. v3.0) to determine the transfection efficiency, indel size, and knockout score.

Western Blotting

Lysate Harvesting: Cells were dissociated with Trypsin then quenched by addition of normal growth media. Cell suspension was spun down 500g x 5 min and washed twice in PBS on ice. Pellets were then resuspended in varying amounts of RIPA lysis buffer (50 mM Tris pH 7.5, 150 mM NaCl, 0.1% Triton X-100, 0.50% sodium deoxycholate, 0.1% SDS) supplemented with 1x cOmplete EDTA-free protease inhibitor cocktail (Thermo Fisher Scientific), 2mM NaVO₃, 5 mM NaF and 1mM phenylmethylsulfonyl fluoride (PMSF) (all sourced from Sigma-Aldrich, St. Louis, Missouri). Cells were incubated in RIPA buffer on ice for 5 min, passed through a 26 gauge syringe 6x, then vortexed 2,000 rpm for 30 min. at 4C and centrifuged 15,000g for 20 min. at 4C. The pellet was discarded and supernatant was transferred to a fresh tube for BCA analysis.

Protein Content Normalization: Protein content of harvested lysates was analyzed using Bicinchoninic Acid (BCA) Protein Assay Kit (Thermo Fisher Scientific) following manufacturer's recommendations. Assays were read using a Synergy H1 plate reader (Biotek Instruments, Winooski, Vermont) measuring excitation/emission of 560 and 590 nm, respectively. Acceptable calibration curves were defined as those with an R squared value ≥ 0.95 . Protein concentration of lysates was normalized by addition of varying volumes of RIPA lysis buffer.

SDS-PAGE: Normalized samples were mixed with NuPage Laemmle Buffer (Thermo Fisher Scientific) and DTT (F. Hoffmann-La Roche AG, Basel, Switzerland) to achieve final

concentrations of 1x Laemmle Buffer and 50 mM DTT, then heated at 70°C for 10 min. with occasional vortex mixing. Samples were cooled on ice for 5 minutes and then loaded into 4-12% Bis-Tris Thermo Fisher NuPage mini gels and run at 200V constant voltage for 30-50 min using a 300V Enduro power supply (Labnet, Edison, New Jersey), then transferred to PVDF (MilliporeSigma, Burlington, Massachusetts) for 37 min. at 30V constant voltage. Running and transfer buffers used were prepared according to Thermo NuPage manufacturer's recommendations. Membranes were blocked by rocking gently with 5% non-fat dry milk (NFDM; LabScientific bioKEMIX, Danvers, Massachusetts) in Tris-Buffered Saline with Tween-20 (TBST) (Sigma-Aldrich) for 1h at room temperature. Primary antibodies (Table 1) were diluted in 5% NFDM in TBST and incubated overnight (16h) with rocking at 4°C. Membranes were washed by gently rocking 3x5 min. at room temperature with TBST. Secondary antibodies were diluted in 5% NFDM in TBST and antibody solution was incubated with blots for 1.25h with gentle rocking at room temperature. Membranes were then washed via rocking 5x5min. with TBST and developed using Pico or Femto (Thermo Fisher Scientific) developing reagents. Images were taken using ProteinSimple gel imager (Bio-Techne, Minneapolis, Minnesota).

Antibodies: Antibodies were sourced from Cell Signaling Technologies (CST)(Danvers, Massachusetts), Santa Cruz Biotechnology (SCBT)(Santa Cruz, California), Sigma Aldrich (St. Louis, Missouri) and Proteintech (Rosemont, Illinois). Antibodies were diluted in 5% NFDM in TBST as described in **Table 1**.

Table 2.1: Primary and secondary antibodies used in western blot experiments

| Target | Antibody | Dilution |
|----------------|---|----------|
| RUNX2 | CST α -RUNX2 #D1L7F | 1:1,000 |
| CBF β | CST α -CBF β #D4N2N | 1:1,000 |
| GAPDH | SCBT α -GAPDH sc-166574 | 1:1,000 |
| FLAG | Sigma M2 α -FLAG F1804 | 1:1,000 |
| Actin | SCBT α -Actin SPM161 | 1:10,000 |
| hnRNPK | SCBT α -hnRNPK sc-28380 | 1:1,000 |
| Biotin | CST α -Biotin-HRP 7075 | 1:3,000 |
| eIF4B | ProteinTech α - eIF4B 17917-1-AP | 1:1,000 |
| Rabbit Primary | CST Goat α -Rabbit #7076 | 1:1,000 |
| Mouse Primary | CST Goat α -Mouse #7074 | 1:4,000 |

Membrane Stripping: Membranes were stripped and re-probed using Abcam (Cambridge, United Kingdom) mild western blot stripping protocol. Briefly, membranes were incubated by shaking for 10 min. with mild stripping buffer (0.2M Glycine, 0.1% SDS, 0.1% Tween-20, pH 2.2), then washed twice with PBS for 10 min. with shaking. Stripping was verified via incubation of membrane with Femto ECL developing reagent (Thermo Fisher) and imaged for residual signal. Any membranes showing residual signal were subjected to additional 5-10 min. of incubation with mild stripping buffer. Following verification of successful membrane stripping, membranes were subjected to 2x 5min. washes in TBST on a shaker at room temperature, and then blocked for 1.25h with 5% milk in TBST at room temperature. Following this, membranes were probed as described above.

Densitometry: Densitometric analysis was performed using ImageJ (32), with bands of interest selected using built-in gel analysis tool. Background subtraction using 80px diameter rolling ball was applied to select images.

Proteasome Inhibition Assay

U2OS wt and U2OS CBF β KO cells were plated on 10 cm dishes and grown to 50% confluence, then treated with MG132 (Sigma-Aldrich, St. Louis, Missouri) 50 μ M or equivalent DMSO control for 4h or 18h followed by western blot analysis as mentioned previously. In some experiments floating cells were also collected, spun down with harvested adherent cells and combined together before lysate collection. The extent of proteasome inhibition induced by MG132 was evaluated using Abcam Proteasome Activity kit according to manufacturer's recommendations.

Cycloheximide Pulse Assay

U2OS wt and U2OS CBF β KO cells were grown in a 6-well plate to 60% confluence, then treated with McCoy's media supplemented with 10% FBS, 1% P/S, 15 mM HEPES pH 6.95 and 150 μ g/mL cycloheximide (CHX) for up to 24h. Cells were all plated in the same batch and drugged in a staggered timeline resulting in all cells being collected at the same time. Following treatment, cells were collected and analyzed via western blot as described previously.

RNAseq

Sequencing: Batch 3' Tag-Seq RNA-seq data was collected and analysis was performed at the Bioinformatics Core (UC Davis Genome Center). The RNA sequencing data were preprocessed using HTStream, version 1.3.3 (<https://s4hts.github.io/HTStream/>) and PCR duplicates were removed using umitools, version 1.0.1 (33). Reads were aligned to GRCh38 using STAR (34). Comparison for analysis was U2OS CBF β KO vs DMSO-treated U2OS wt control.

Differential Expression Analyses: Differentially expressed genes were identified using the limma-voom Bioconductor pipeline (limma version 3.50.1, edgeR version 3.36.0) (35). In this pipeline, the TMM method is used to obtain normalization factors and adjusted library sizes for each sample (36). These adjusted library sizes are then used to calculate counts per million reads (CPMs), using the adjusted library sizes instead of the total counts as the denominator. CPMs are log₂ transformed, and voom is used to calculate variance weights for each observation, which are used in a weighted least squares model in limma (37). Empirical Bayes smoothing is used to obtain improved smoothed standard errors for log fold changes for use in hypothesis testing. The resulting p- values are adjusted for multiple testing using the Benjamini-Hochberg false discovery rate (FDR) controlling method (38).

Global Translation Assay

Global protein translation was evaluated using Thermo Fisher Click-iT OPP Alexa488 kit. Briefly, 4,500 U2OS wt and U2OS CBF β KO cells were seeded into each well of a 96-well plate 12h before labeling. Nascent proteins were labeled for 2h with McCoy's media + 6.2 μ M O-propargyl-puromycin (OPP) in DMSO, with control wells including 150 μ g/mL cycloheximide (CHX) (Sigma-Aldrich) or equivalent DMSO control. Cells were fixed and permeabilized and click conjugation was conducted, all according to manufacturer's recommendations. Nuclei of fixed cells were then stained with HCS NuclearMask Stain (Thermo Fisher Scientific) diluted 1:2,000 in PBS according to manufacturer's recommendations. Images were obtained using a Jenoptik ProgRes MF Cool CCD mounted on a Leica DMI3000B microscope with a Leica HCX PL FLUOTAR L 20X/0.40na CORR objective. Samples were illuminated by a Leica EL6000

fluorescent light source, and cell counts and nascent protein signal was quantified using ImageJ. Fluorescent signal was normalized to quantity of nuclei as stained by NuclearMask.

Lentiviral Transduction

Plasmid Design: The open reading frame (ORF) for CBF β was cloned out of OHu25418 plasmid incorporating NM_022845.3 (GenScript Biotech Corporation, Piscataway, New Jersey) using primers listed in **Table 2**, produced by Integrated DNA Technologies (IDT, Newark, NJ). CBF β ORF was then inserted into pHIV-EGFP by use of XbaI and TspMI restriction enzymes (New England BioLabs, Ipswich, MA) to generate CBF β _WT. *In silico* cloning simulations, primer design, and plasmid diagrams were made using SnapGene® software (from Dotmatics; available at snapgene.com). The vector was digested by both enzymes simultaneously in NEB Cutsmart 3.1 buffer according to manufacturer's recommendations, cleaned up by electrophoresis in 1% agarose gel (IBI Scientific, Dubuque, Iowa) in TBE run at 160V, and DNA was extracted from gel using GeneJET Gel Extraction Kit (Thermo Fisher Scientific) according to manufacturer's recommendations. Plasmid was eluted from GeneJET column using 30 μ L nuclease free water (Ambion, Thermo Fisher Scientific). Insert and vector were combined at 7:1 stoichiometric ratio and ligated using T4 Ligase (Thermo Fisher Scientific) according to manufacturer's recommendations. Vector was transformed into NEB Stable *E. coli* according to manufacturer's recommendations, then grown overnight at 37°C on LB (Luria-Bertani) (Thermo Fisher Scientific) and Agar (Sigma-Aldrich) plates containing 100 μ g/mL carbenicillin (IBI Scientific, Dubuque, Iowa). Colonies growing in presence of 100 μ g/mL carbenicillin were inoculated into 5 mL of LB broth supplemented with 100 μ g/mL carbenicillin and grown overnight at 37°C with 180rpm rotation. Following overnight growth, plasmid DNA was collected using

GeneJET Plasmid Miniprep Kit (Thermo Fisher Scientific) according to manufacturer's recommendations, with final elution from column accomplished with 50 μ L nuclease-free water. Eight hundred (800) ng of purified plasmid was sent for Sanger sequencing verification by Genewiz using primers listed in **Table 2**.

C-terminus FLAG tagged CBF β (denoted as CBF β _WT_FLAG) was generated through site directed mutagenesis of the above described vector via Q5 Mutagenesis Kit (New England BioLabs) according to manufacturer's instructions, with same transformation, miniprep, and sequence verification steps as listed above, using primers in **Table 2**.

Table 2.2: Primer sequences for Lentiviral Vector Generation

| Usage | Primer Sequence |
|--|--|
| Add XbaI restriction site and a Kozak sequence to CBF β N-term for cloning into pHIV-EGFP F | 5'-GGAGGATTCTAGAGC CACCATGCCGCGC-3' |
| Add TspMI restriction site and stop codon (snipping off FLAG tag in original pcDNA3.1 vector) to CBF β C-term for cloning into pHIV-EGFP R | 5'-GGAGGTCCCGGGTTATCAACG AAGTTTGAGGTCATCACCACC-3' |
| Mutate add C-term FLAG tag to CBF β in pHIV-EGFP F | 5'-GACGACGATAAGTGAT AACCCGGGCTAGGA-3' |
| Mutate add C-term FLAG tag to CBF β in pHIV-EGFP R | 5'-ATCCTTGTAATCACGAA GTTTGAGGTCATCAC-3' |
| Sequence T7 F | 5'- TCAAGCCTCAGACAGTGGTTC-3' |
| Begin Sanger sequencing at elongation factor-1 α promoter | 5'- TCAAGCCTCAGACAGTGGTTC-3' |

HEK293T Transfection: Wells of a 6-well plate were filled with 1 mL antibiotic-free 10% FBS DMEM and pre-incubated to reach 37C. Third (3rd) generation lentiviral plasmids (39,40), listed below, were mixed with jetPRIME (Polyplus, Sartorius, New York, New York) and added

to media according to manufacturer's recommendations, using 2:1 ratio of jetPRIME volume (μL) to DNA quantity (μg). Total quantity of each plasmid used for co-transfection is listed in **Table 3**.

Table 2.3: Plasmids and their source used in HEK293T transfection experiments

| Plasmid | Source | DNA (μg) |
|------------|----------------|-----------------------|
| pHIV-EGFP | Addgene #21373 | 0.87 |
| pMDLg/pRRE | Addgene #12251 | 0.44 |
| pRSV/REV | Addgene #12253 | 0.22 |
| pMD2.G | Addgene #12259 | 0.22 |

1.4 million HEK293T cells per well were then plated on top of 1 mL media + transfection reagents + plasmids, and cells were incubated in presence of transfection reagents for 18h. Media was changed and lentiviral particles were harvested as described below.

Harvesting of Lentiviral Particles: Viral supernatant was then removed and fresh antibiotic-free DMEM supplemented with 10% FBS was added. Supernatant containing lentiviral particles was collected 48h and 72h post-transfection and pooled. Supernatant was then cleared by centrifugation at 500g for 10 min and sterilized by passing through a 0.2 μm nylon filter (Sigma-Aldrich) and stored at -80C until lentiviral transduction.

Infection of Target Cells: Fifty-one thousand U2OS CBF β KO cells were seeded into each well of a 24- well plate and grown at 37°C for 16h prior to transduction. Previously harvested lentiviral particles were thawed at 4°C and diluted 20x, 50x, 125x, 313x, 781x, and 1953x in normal media supplemented with 4 $\mu\text{g}/\text{mL}$ Polybrene (Sigma-Aldrich) and added to target cells. Infection was allowed to continue for 3 days at 37°C, after which cells were split into 6-well plates and inspected for fluorescence using a Leica DMI3000B microscope with a Leica EL6000 fluorescent light source. The cell population from each infection used for harvesting via trypsin

and outgrowth in T75 flask was chosen based on <5% EGFP+ cells. This was done to reduce risk of double integration from lentivirus. Dilution factors of lentiviral particles in cell populations selected for outgrowth are shown in **Table 4**:

Table 2.4: Lentiviral inserts and dilution factors utilized for transduction experiments

| Lentiviral Insert | Dilution |
|---------------------------------------|----------|
| Empty vector pHIV-EGFP | 125x |
| pHIV-EGFP with CBF β (wt) | 313x |
| pHIV-EGFP with CBF β _FLAG (wt) | 781x |

FACS Sorting of Monoclonal Cell Lines: Ten million cells from each lentiviral insert-containing population were harvested via trypsin, washed 2x with PBS, then resuspended in PBS supplemented with 1% FBS and 2% P/S. Cells were passed through a sterile 40 μ m nylon mesh (Sigma-Aldrich) just before sorting. Single cells were separated into individual wells of 96 well culture plates for subsequent expansion using a 4-laser, 18-color Astrios EQ cell sorter (Beckman Coulter, Brea, CA). The Astrios was configured with its 70 μ m nozzle at 60psi to quickly and accurately deposit single EGFP+ cells into 96 well plates prefilled with McCoy's media supplemented with 10% FBS and 1% P/S. Each 96 well plate was pre-chilled to ~4C to preserve cell vitality during sorting. Following sorting into 96-well plates, media was regularly changed and cells were harvested via trypsin and transferred to successively larger vessels upon reaching ~50% confluence in each. Vessel order was: 96-well plate, 24-well plate, 6-well plate, and finally cells were seeded into a T75 flask. 5 different clonal cell lines from each of 4 different lentiviral inserts were generated.

RNA Collection

U2OS wt and U2OS CBF β KO cells were grown to 70% confluence on 10 cm plates in 10% FBS 1% P/S McCoy's. Cells were washed 2x with PBS then lysed with 1 mL TRIzol (Ambion, Thermo Fisher Scientific) per plate, which was transferred to a 1.5 mL tube, 200 μ L of Chloroform (Thermo Fisher Scientific) was added, and tubes were mixed and centrifuged at 4°C 12,000g for 15 min. The aqueous layer (top) was transferred to a fresh tube, 500 μ L of 2-propanol (Thermo Fisher Scientific) was added, and samples were mixed and allowed to sit at room temp for 5 min. Sample was pelleted at 4°C 12,000g for 10 min, supernatant was removed, and pellet was washed with 1 mL 75% ethanol (Decon Labs, King of Prussia, Pennsylvania) in water, w/v, centrifuged, washed, and centrifuged again. Supernatant was removed and pellet was dried for 10 min. and re-solvated in 30 μ L nuclease-free water and vortex mixed at 300 rpm at 55°C for 10 min. to facilitate dissolution. RNA samples were stored -80C until further analysis.

Quantitative real-time PCR (qRT-PCR)

cDNA synthesis and gDNA wipeout was performed using QuantiTect Reverse Transcription Kit (Qiagen, Hilden, Germany). Briefly, RNA was thawed on ice and all kit solutions were thawed at room temperature. gDNA wipeout and following cDNA synthesis reactions were performed per manufacturer's recommendations, using 1 μ g RNA as starting material. qRT-PCR was performed by creating a master mix comprised of: 6.5 μ L nuclease-free water, 1 μ L forward primer (2.5 μ M stock), 3 μ L reverse primer (2.5 μ M stock) and 12.5 μ L SYBR green per reaction. Twenty three (23) μ L of master mix was added to each well of a 96-well PCR plate, and 2 μ L of 5ng/ μ L cDNA was spiked into each well. qRT-PCR reaction was promptly started, with gradient

conditions of 5 min. at 95°C followed by 45 cycles of 10s at 95C and 30s at 60°C. qRT-PCR was performed on an AriaMx Real-Time PCR System (Agilent Technologies, Santa Clara, CA). Threshold was set to region of sigmoid where amplification became linear.

Table 2.5: Primers utilized for quantitative real time RT-PCR of RUNX2.

| Usage | Primer Sequence |
|--------------|-------------------------------|
| qPCR HPRT F | 5'- CCTGGCGTCGTGATTAGTGA -3' |
| qPCR HPRT R | 5'- CGAGCAAGACGTTTCAGTCCT -3' |
| qPCR RUNX2 F | 5'- TAGGCGCATTTCAGGTGCTT -3' |
| qPCR RUNX2 R | 5'- GGTGTGGTAGTGAGTGGTGG -3' |

CBFβ-hnRNP Co-Immunoprecipitation

Pulldown of CBFβ: U2OS cells were grown in 15% FBS 1% Pen-Strep McCoy's media on 2x 10 cm plates until 70% confluent. Cells were lysed according to western blot lysate collection protocol outlined above, except cells were lysed in modified RIPA buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1% NP40, 0.25% sodium deoxycholate) supplemented with 1x cComplete EDTA-free protease inhibitor cocktail, 2mM NaVO₃, NaF 5 mM and 1mM PMSF. CBFβ was immunoprecipitated from cell lysates using anti-CBFβ antibody A303-547A (Bethyl Labs, Montgomery, Texas) and Protein A Sepharose CL-4B beads (GE Healthcare, Chicago, IL) as follows: cell lysates were normalized to 1.00 mg/mL protein concentration, adjusted to a final volume 300 μL, and 3 μg anti-CBFβ antibody was added to lysates and rotated overnight at 4C. The following day agarose beads were re-hydrated and washed 3x with PBS, then finally resuspended in lysis buffer and added to samples, and rotated for 4h at 4°C. Following this, agarose beads were spun down and washed twice with lysis buffer, then twice with wash buffer (50 mM Tris pH 7.5, 0.1% NP40, 0.05% sodium deoxycholate). Washes were removed and protein was extracted from beads by heating at 70°C for 10 min. in the presence of 1x Laemmle buffer and 50

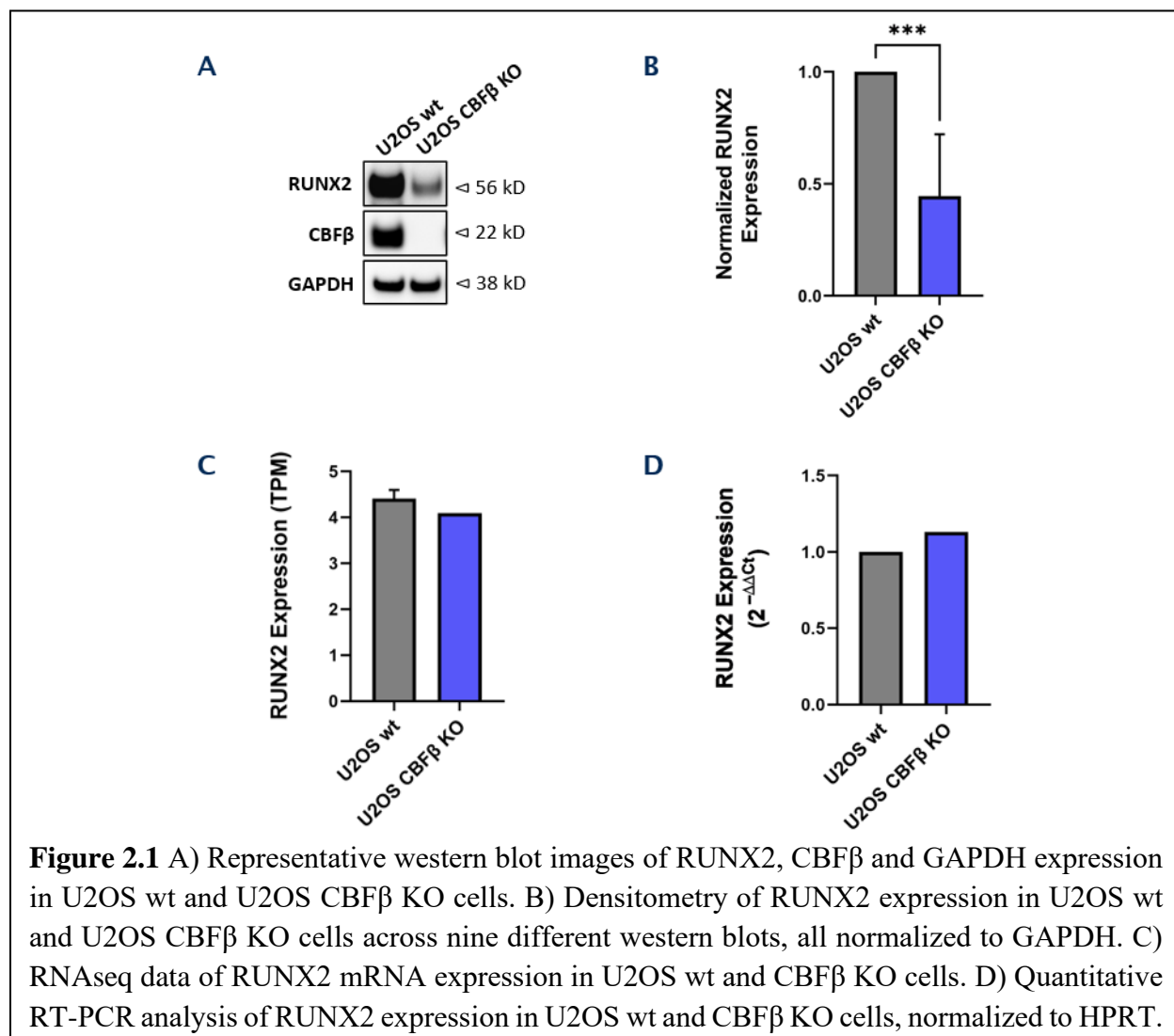
mM DTT. The eluted protein was then analyzed by western blot as described above. Membranes were stored in TBS at 4°C between antibody detection.

Pulldown of hnRNPk: Experiment was conducted the same as above except hnRNPk was immunoprecipitated from cell lysates using anti-hnRNPk antibody (Santa Cruz Biotech SC-28380) and anti-GAPDH antibody was used as isotype control (Santa Cruz Biotech SC-166574). Antibody quantities added were the same as above.

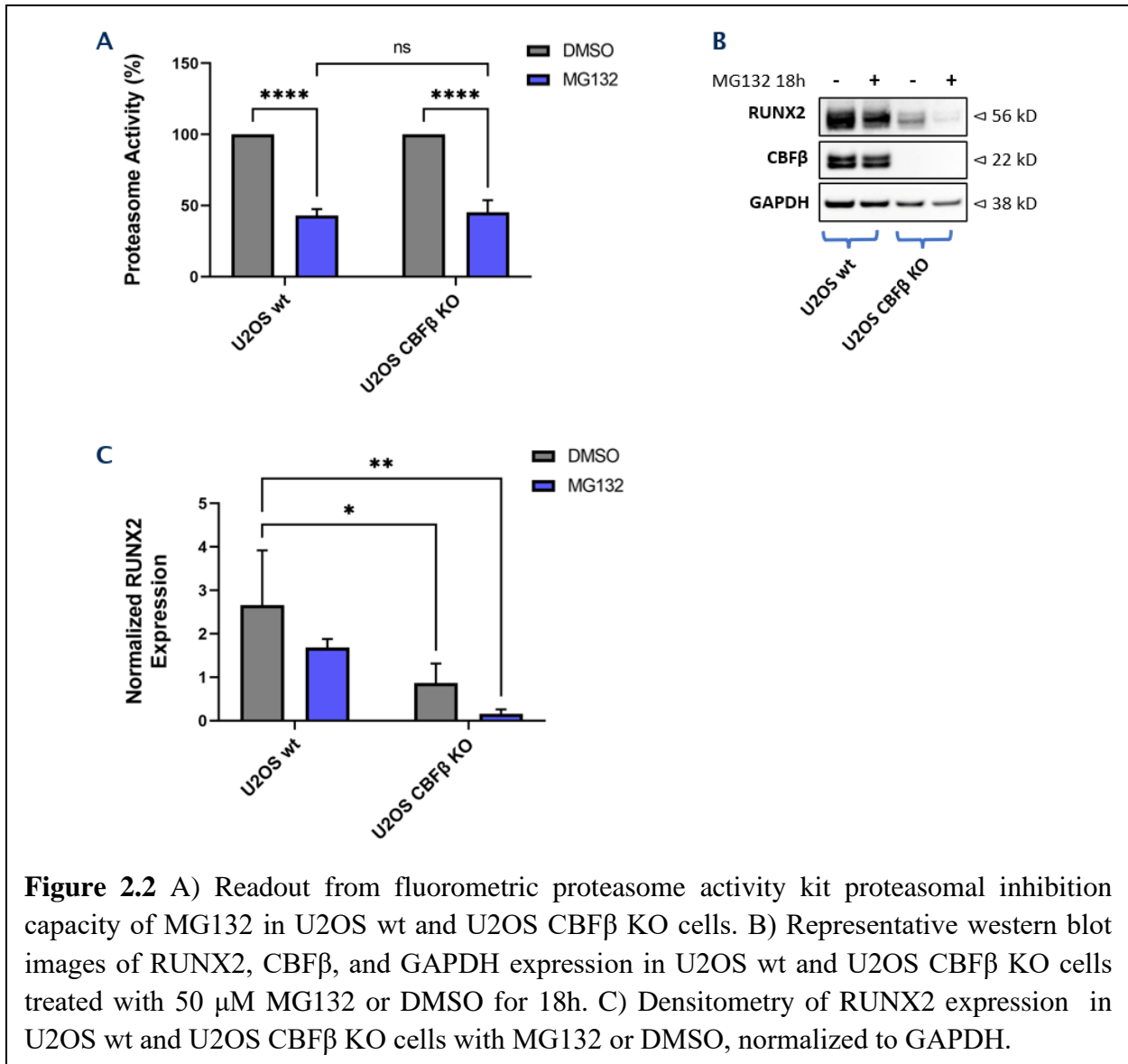
Statistical Analyses

Figure 2.1B: Error bars are standard deviation (SD), n=9 (biological), *** = p<0.0001, unpaired two-tailed Welch's t-test. ***Figure 2.2A:*** Error bars are SD, n=3 (technical); **** = p<0.0001, two-way ANOVA with Tukey's post hoc test. ns = not significant. ***Figure 2.2C:*** Error bars are SD, n=3 (biological); * = p<0.05, ** = p<0.01, two-way ANOVA with Tukey's post hoc test. ***Figure 2.4A:*** Error bars are SD, n=3 (3 biological replicates, 3 technical replicates per run); one-way t-test. ***Figure 2.6C,D:*** Error bars are SD, n=5 (biological); * = p<0.05, *** = p<0.001, one-way ANOVA with Tukey's post hoc test.

Results

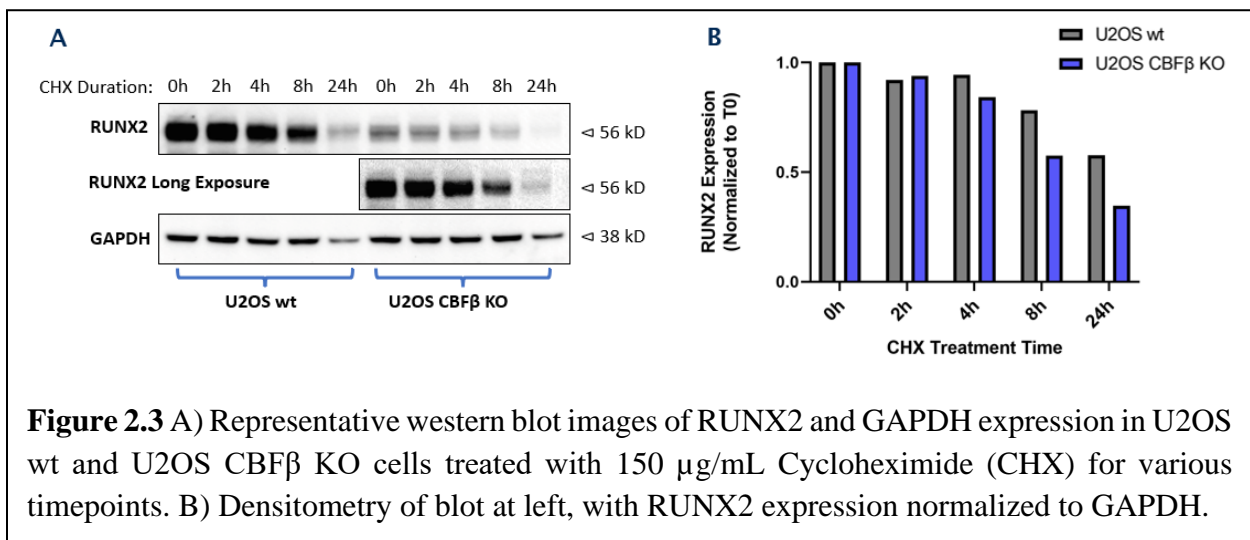


Loss of CBFβ leads to reduced RUNX2 protein expression. A U2OS CBFβ KO cell line was developed using CRISPR and profiled via western blotting. When compared to U2OS wt cells, U2OS CBFβ KO cells consistently demonstrated lower RUNX2 protein expression, normalized to GAPDH. Interestingly, both cell lines demonstrated similar relative RUNX2 mRNA expression when measured by RNAseq. qRT-PCR was then conducted to validate RNAseq data, and RUNX2 mRNA expression again appeared similar between both cell lines.



Reduced RUNX2 protein expression in U2OS CBFβ KO cells is not rescued by proteasomal inhibition. To assess if proteasomal degradation could explain this decrease in RUNX2 protein expression in U2OS CBFβ KO cells, we subjected our two cell lines to proteasome inhibition followed by western blotting. We first tested our proteasome inhibitor, MG132, to ensure it was capable of inhibiting the proteasome in our cell lines. Treatment of both cell lines with 50 μM MG132 resulted in ~45% proteasome inhibition in each, and we used this same treatment for our

western blotting experiments. Proteasomal inhibition did not rescue low RUNX2 protein expression in U2OS CBF β KO cells, and instead caused RUNX2 expression to decrease further. This was observed in both cell lines, however in terms of fold change it was more pronounced in U2OS CBF β KO cells. RUNX2 protein expression in DMSO-treated samples was again lower in CBF β KO cells than WT U2OS cells, in accordance with previous data.



Loss of CBF β does not drastically alter RUNX2 degradation in U2OS cells. To further evaluate if heightened degradation explains lower RUNX2 protein expression in U2OS CBF β KO cells, we utilized an orthogonal assay involving CHX, a protein translation inhibitor. Instead of inhibiting degradation, as before, we inhibited synthesis of new protein and observed degradation itself. Following treatment with CHX, RUNX2 protein expression did decrease over time in both cell lines, which was expected. In comparing the two cell lines, we did see a small difference in terms of quantity of RUNX2 degraded over time, with RUNX2 in U2OS CBF β KO cells seemingly having a slightly shorter half-life than that of RUNX2 in WT U2OS cells.

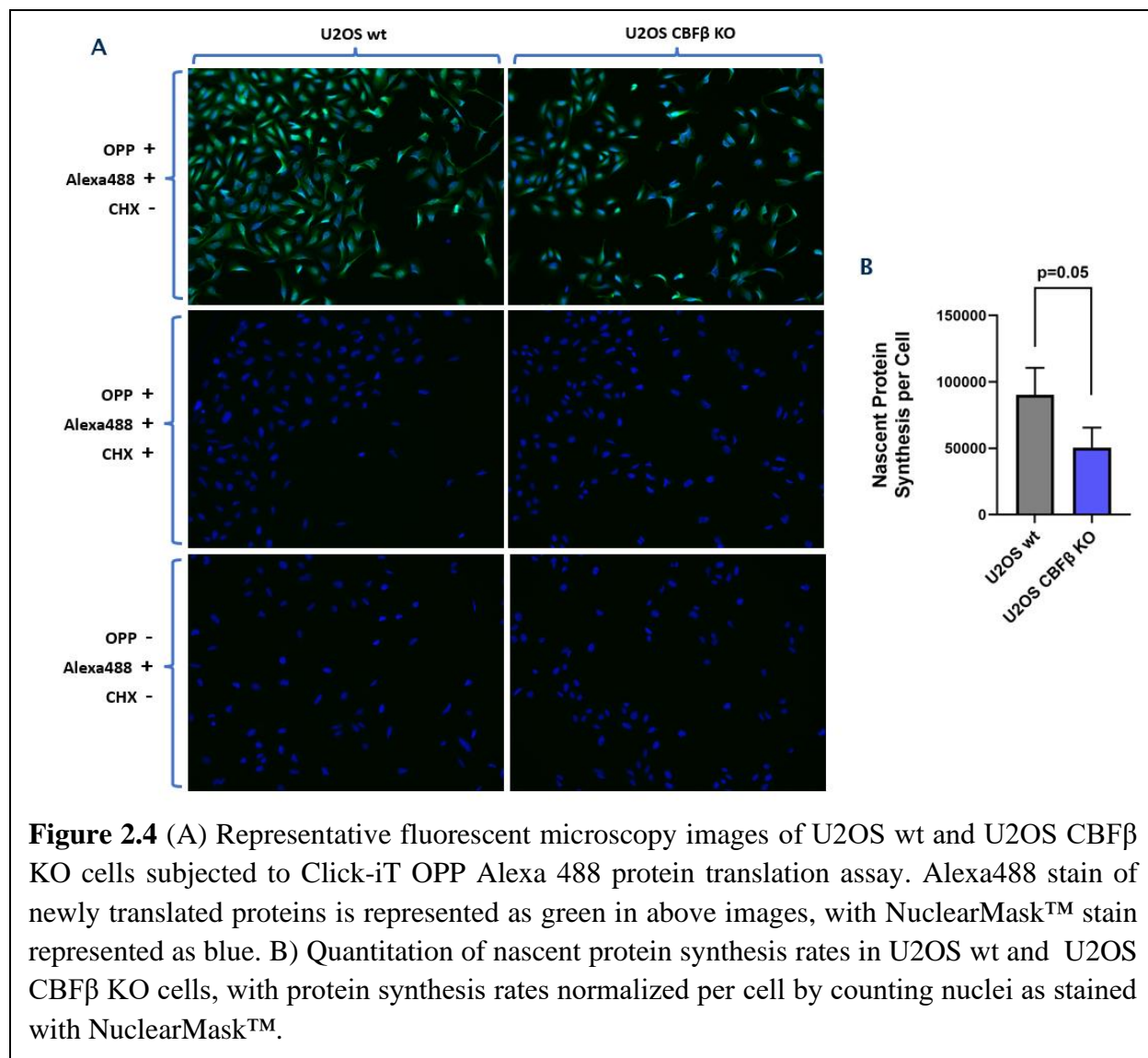


Figure 2.4 (A) Representative fluorescent microscopy images of U2OS wt and U2OS CBFβ KO cells subjected to Click-iT OPP Alexa 488 protein translation assay. Alexa488 stain of newly translated proteins is represented as green in above images, with NuclearMask™ stain represented as blue. B) Quantitation of nascent protein synthesis rates in U2OS wt and U2OS CBFβ KO cells, with protein synthesis rates normalized per cell by counting nuclei as stained with NuclearMask™.

Loss of CBFβ leads to a decrease in global protein translation in U2OS cells. In order to understand CBFβ's contribution to global protein translation, we subjected U2OS wt and U2OS CBFβ KO cells to the Click-iT OPP Alexa 488 protein translation assay. This experiment was performed in technical and biological triplicate. We observed a consistent decrease in global protein translation in U2OS CBFβ KO cells compared to U2OS WT, slightly exceeding the cutoff for statistical significance.

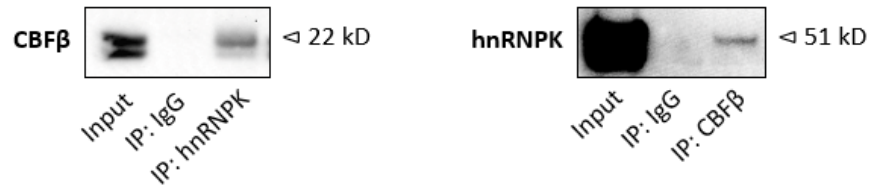
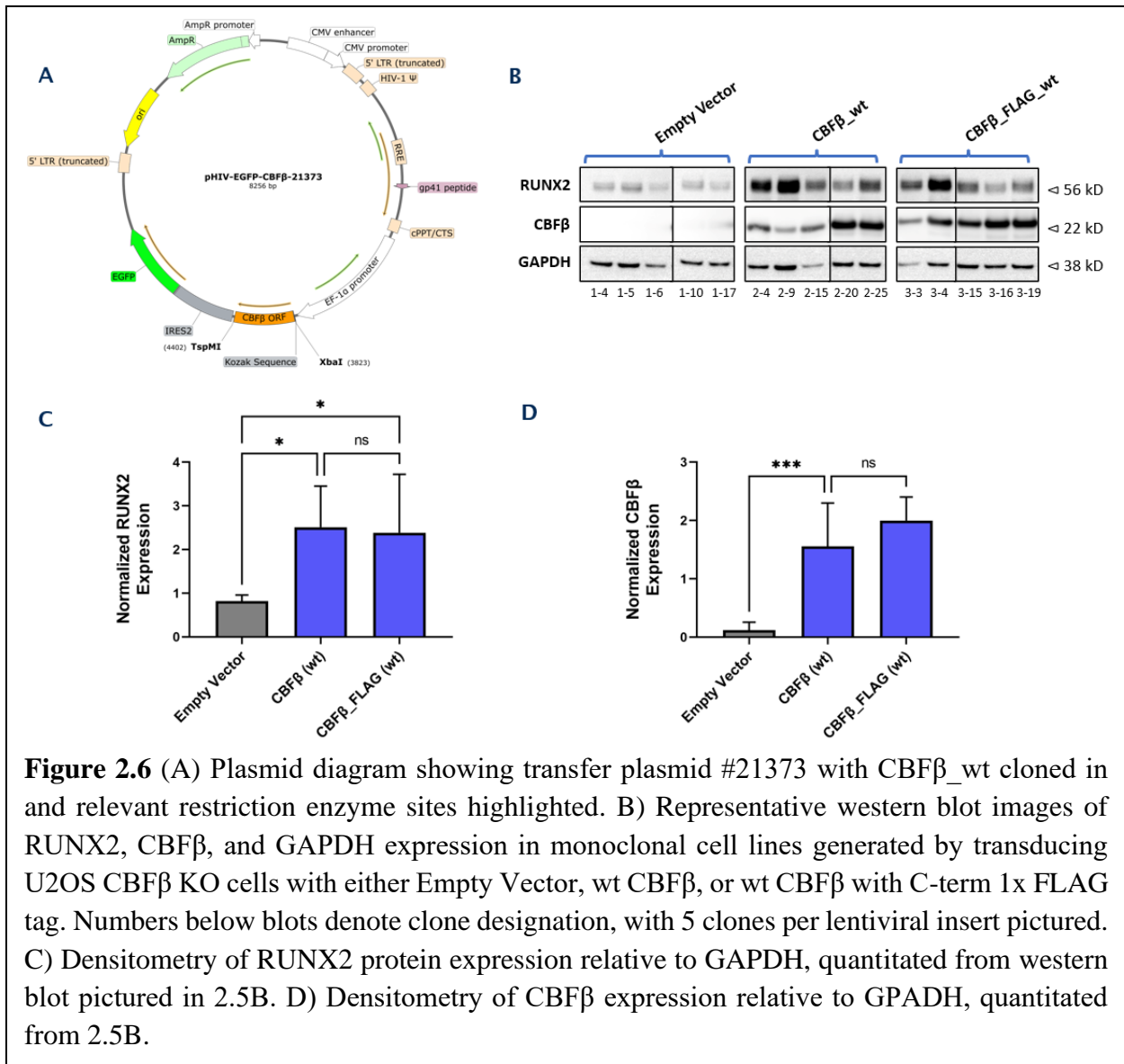


Figure 2.5 Co-IP of hnRNPk in U2OS cells followed by blotting for CBFβ, and reciprocal Co-IP pulling down CBFβ and blotting for hnRNPk.

CBFβ and hnRNPk interact in U2OS cells. Other research posited CBFβ to participate in protein translation through interaction with hnRNPk. To evaluate whether this interaction occurred in osteosarcoma cells, we subjected U2OS cells to reciprocal Co-IP's of CBFβ and hnRNPk. hnRNPk was present in CBFβ pulldown, and CBFβ was present in hnRNPk pulldown. Isotype control (IP: IgG) demonstrated no signal in either experiment.



Re-introduction of WT CBF β rescues low RUNX2 expression in U2OS CBF β KO cells. To evaluate that we were indeed studying a CBF β -specific effect, we used lentiviral transduction of CBF β KO cells to generate stable cell lines expressing Empty Vector, CBF β _WT, or CBF β _WT_FLAG. Transduction with CBF β _WT or CBF β _WT_FLAG resulted in statistically significant increases in RUNX2 protein expression relative to cells transduced with Empty Vector.

Expression levels of both endogenous RUNX2 and CBF β transgene were consistent between CBF β _WT and CBF β _WT_FLAG transduced cells.

Discussion

As measured by western blot, RUNX2 protein expression was consistently lower in U2OS CBF β KO cells than in U2OS wt cells (**Figure 2.1**). This was not accompanied by a commensurate decrease in RUNX2 mRNA expression, implying a post-transcriptional mechanism, seemingly dependent upon CBF β . Potential explanations to this include increased protein degradation, decreased mRNA stability, or decreased translational efficiency of RUNX2. We chose to focus our efforts on proteasomal degradation as this is the pathway governing RUNX2 turnover under normal conditions (41–44), and previous studies suggested CBF β protects RUNX2 from proteasomal degradation (45,46). The resulting decrease in RUNX2 levels due to the loss of CBF β was not due to proteasome degradation, because treatment with MG132 was unable to rescue low RUNX2 expression observed in CBF β KO cells (**Figure 2.2**).

It was surprising that both cell lines exhibited decreased expression of RUNX2 upon MG132 treatment, when proteasome inhibition typically causes an accumulation of proteins (47). Nevertheless, this phenomenon is not unheard of and has been observed in other studies treating U2OS cells with MG132 for a similar duration (48), as well as in other cell models (49–51). Proteasome inhibitors may lead to increased lysosomal degradation of proteins (52), and cause RUNX2 to undergo trafficking to the lysosome through interaction with SOX9 (53), which could explain the decrease in RUNX2 protein following MG132 treatment. Interestingly, RNAseq data (not shown) revealed heightened SOX9 mRNA expression in CBF β KO cells compared to U2OS wt cells, which may underly the more severe decrease in RUNX2 protein observed in U2OS CBF β KO cells compared to U2OS wt cells. Further studies of the significance of SOX9 and lysosomal degradation in RUNX2 stability would be necessary to delve into this further, and could entail

MG132 treatment in the presence of SOX9 siRNA or use of lysosomal protease inhibitors such as Bafilomycin A1 (54) or Leupeptin (55).

Treatment of our cell lines with CHX did demonstrate a slightly reduced half-life of RUNX2 in U2OS CBF β KO cells, suggesting RUNX2 may be degraded at a slightly increased rate in the absence of CBF β (**Figure 2.3**). This difference in RUNX2 half-life was small in magnitude overall, and most prominent in latter timepoints. It's important to note that CHX treatment of cell lines in other studies is typically shorter in duration, as the median half-life of human proteins hovers around 8.7 h (56). In our hands, RUNX2 appeared to have a particularly long half-life in U2OS cells, necessitating a longer treatment time to observe a significant amount of degradation. Extended treatment with CHX, and therefore shutdown of nascent protein synthesis, is likely to place cells under stress and alter degradation kinetics (57), which may confound these data at the later timepoints.

Other methods are available that could allow us to delve deeper into potential alterations in the stability of RUNX2, such as immunoprecipitation (IP) of RUNX2 followed by immunoblotting for ubiquitin (58) or pulse-chase labeling of nascent proteins followed by IP of RUNX2 (59). While these methods have been used with great success in other studies, they were not feasible for us due to a myriad of issues encountered with IP of RUNX2. These are listed in further detail in the Chapter 3 discussion, but in short, this experiment was attempted ad nauseam under numerous conditions and ultimately proved insurmountable. Ultimately, we concluded that decreased stability of RUNX2 due to loss of its binding partner, CBF β , did not sufficiently explain the discrepancy in RUNX2 levels we observed. Around the same time, similar data was reported in breast cancer cells demonstrating a reduction in RUNX1 protein upon loss of CBF β (23), and we observed a decrease in RUNX1 in our hands as well (data not shown). The authors went on to

implicate CBF β playing a critical role protein translation, which supports our results that pointed to a mechanism other than stabilization of RUNX2.

Utilizing a Click-iT OPP Alexa 488 protein translation assay, we assessed global protein translation rates of U2OS wt and U2OS CBF β KO cells. In all three biological replicates performed, U2OS CBF β KO cells consistently registered lower global protein translation rates than U2OS wt cells (**Figure 2.4**). While our result did not quite meet the criteria for statistical significance ($p=0.05$, not $p<0.05$), it's important to note that this assay evaluates translation rates of all cellular proteins collectively rather than on a case-by-case basis. The influence of CBF β on protein translation might not be broad enough to significantly skew global protein synthesis, but may strongly influence the translation of a subset of proteins within OS cells, particularly those reliant on cap-dependent translation. This consistent reduction in global protein synthesis rates between our cell lines was encouraging enough for us to move forward studying protein translation in latter, more specific assays.

Previous work postulated that the role of CBF β in protein translation is accomplished by binding to mRNA via hnRNPk, and CBF β modulates translation through binding to eIF4B, a member of the eIF4 translational complex (23). We next conducted co-IP experiments to assess if this CBF β -hnRNPk interaction also occurs in OS cells. Reciprocal Co-IP demonstrated an interaction between hnRNPk and CBF β in U2OS cells, as evidenced by CBF β being present in the pulldown of hnRNPk, and vice versa (**Figure 2.5**). We did not see strong enrichment of either protein in our pulldowns, possibly due to weak CBF β -hnRNPk interaction, or a low fraction of total CBF β and hnRNPk bound to each other at any point in time. Nevertheless, our pulldown was specific and demonstrated CBF β and hnRNPk do interact in U2OS cells as neither protein was present in IgG isotype samples.

Taken together, our results were supportive of a non-canonical role for CBF β in protein translation in OS cells. To further support this claim, it was necessary to confirm our conclusions weren't misled by off-target CRISPR activity (60) in our starting CBF β KO cell line. To this end, we conducted a rescue experiment reconstituting wt CBF β back into CBF β KO cells. Transient transfection was first attempted, but U2OS CBF β KO cells proved resistant to multiple transfection reagents and optimization panels, yielding a maximum transfection efficiency of 16%. This low efficiency precluded measure of changes in RUNX2 brought about by CBF β , as any shifts in RUNX2 expression are blunted by the abundance of un-transfected cells. Furthermore, RUNX2 demonstrated slow degradation in our previous studies, meaning its synthesis rate was commensurately slow as well (61). Transient transfection results in transgene expression lasting just a few days, which we reasoned might not be enough time for reconstituted CBF β to drive appreciable changes in RUNX2 levels.

In order to interrogate this system over a longer time period, we opted to reconstitute CBF β through lentiviral transduction. The cytomegalovirus (CMV) promoter is commonly chosen for driving transgene expression in transduced cells, however reports of it causing inconsistent expression over time are numerous (62–64). We instead selected the EF-1 α promoter due to its lauded capability to drive stable, long-term transgene expression (63,65). Cells were transduced with CBF β _WT as well as CBF β _WT incorporating a 1xFLAG tag. The FLAG tag was cloned in at the C-terminus, as the N-terminus of CBF β is key in CBF β -RUNX2 interaction (66) and RUNX protein binding would have been blocked by a FLAG tag at this location (23,67).

Lentiviral particles with transfer plasmids containing either Empty Vector, CBF β _WT, or CBF β _WT_FLAG were generated and used to infect U2OS CBF β KO cells, and multiple monoclonal cell lines for each insert were generated. RUNX2 expression in these resultant cell

lines was then evaluated via western blotting (**Figure 2.6**). We observed a statistically significant increase in RUNX2 protein expression in our CBF β -expressing transductants, indicating that CBF β alone was both necessary, and sufficient, to rescue low RUNX2 protein levels in U2OS CBF β KO cells. RUNX2 and CBF β expression was consistent between cell lines transduced with either untagged CBF β or our FLAG-tagged variant, demonstrating that our affinity tag did not alter transgene expression or hamper CBF β -mediated rescue of RUNX2.

In conclusion, we have demonstrated that loss of CBF β in U2OS cells leads to a reduction in RUNX2 protein expression while RUNX2 mRNA levels are unaffected, implying a post-transcriptional mechanism. This reduction in RUNX2 protein expression is specific to loss of CBF β , as reconstitution of wt CBF β restores low RUNX2 protein expression. The observed reduction in RUNX2 protein expression upon loss of CBF β is not fully explained by a decrease in the stability of RUNX2, and may be explained by CBF β playing a role in protein translation. This noncanonical role of CBF β has been reported in breast cancer, and, in that setting, occurs via interaction with hnRNPk. Indeed, we observed a reduction in global protein translation upon loss of CBF β , and confirmed CBF β and hnRNPk do interact in osteosarcoma cells. Further study is needed to unravel more about the significance of this noncanonical OS, and reveal which proteins or pathways may be under the purview of CBF β .

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Chapter 3

**Mutations in the RUNX2 binding pocket of CBF β alter its
cellular activity in osteosarcoma cells**

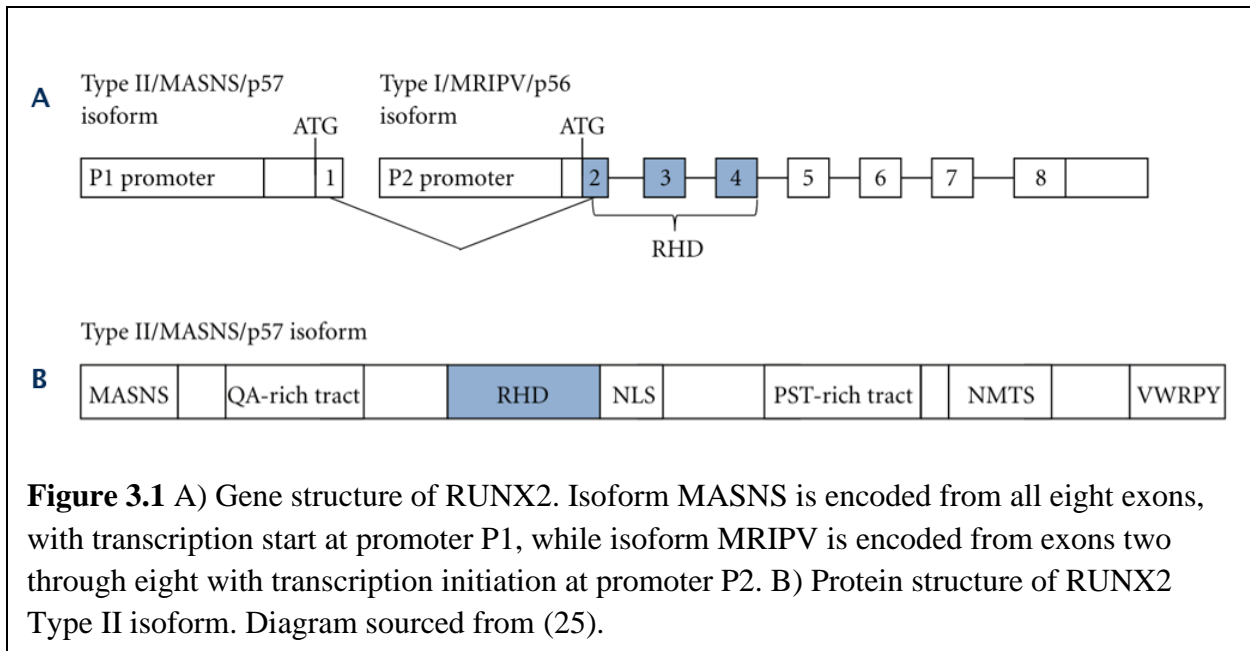
Abstract

Osteosarcoma (OS) is a highly aggressive form of bone cancer that most commonly presents early in age, and treatment options have remained unchanged in the last 30 years. OS tumors are highly heterogeneous and lack bona-fide driver mutations, hampering therapeutic development and necessitating study of methods to overcome this. Previous data in our hands and another lab have suggested that core binding factor subunit beta (CBF β), a transcriptional co-activator to the RUNX family of transcription factors, may perform a noncanonical role as a regulator of protein translation. This is of particular interest in OS, as targeting protein translation could provide a mechanism to prevail over the heterogeneity in OS tumor samples. Elucidating the relevance of this translational role of CBF β to the malignant phenotype of OS is challenging, as CBF β is shuttled away from the cytoplasm and into the nucleus by RUNX proteins, yet this non-canonical role of CBF β is suggested to occur in the cytoplasm. These two roles of CBF β may be antagonistic to each other, so we endeavored to decouple them in order to better study this putative non-canonical role of CBF β . To this end, we produced CBF β and RUNX2 in a recombinant system and used synthetic peptides to validate residues key in their interaction. We then generated stable cell lines expressing CBF β with point mutations at these residues, and assessed the influence of these mutations on CBF β -RUNX2 interaction and subsequent nuclear shuttling. In our recombinant system and in accordance with previous research, G61 and N63 of CBF β appear important to CBF β -RUNX2 interaction. Mutation of these residues and N104 of CBF β to alanine yielded a mutant form of CBF β which exhibited drastically reduced RUNX2 binding and nuclear shuttling.

Introduction

Osteosarcoma is the most common form of primary bone cancer in humans and canines, with ~1,000 new cases diagnosed per year in the United States (1). Age of diagnosis of OS has a bimodal distribution, with the first, and largest, peak in the 10-14-year-old age group, with the second for patients aged 65+ (2). The standard of care for OS is so-called MAP therapy (methotrexate, adriamycin, and cisplatin) (3), a regimen which hasn't changed in more than 30 years (4). Much of the difficulty in developing new therapies in OS has been due to the lack of identified driver mutations in this disease. In lieu of being caused by specific mutations which could be leveraged therapeutically, OS instead presents as a collection of widespread chromosomal abnormalities (5), with chromosome counts ranging from haploid up to hexaploid (6,7).

RUNX2 is a member of the RUNX family of DNA-binding transcription factors and is known as the master regulator of bone growth. RUNX2 plays a critical role in chondrocyte maturation and osteoblast differentiation (8), and high levels of RUNX2 (9,10) and many of its gene targets (11–14) have been implicated in poor therapeutic outcome in OS (15,16). CBF β acts as a transcriptional co-activator to the RUNX proteins, binding to them to form a CBF β -RUNX heterodimer. CBF β is crucial for skeletal development (17–19), and although the RUNX proteins can bind DNA on their own (20), their affinity for DNA is greatly enhanced by interaction with CBF β (21,22). In addition to RUNX2, high expression of CBF β itself is also correlated with poor prognosis in OS (23). These data have led to the aforementioned proteins being proposed as a novel therapeutic target in OS (24).



Structure of RUNX2. RUNX proteins are identified by the Runt homology domain (RHD) (25), which is required for their interaction with DNA (26). This domain is located close to the N terminus and is ~90% conserved between each member of the RUNX family, with the PyGPyGGTPy consensus sequence being essential for interaction with DNA as well as CBF β (27) (**Figure 3.1**). Upstream of the RHD is the glutamine-alanine (QA) repeat region, which plays a role in transactivation (28). Adjacent to the RHD lies the nuclear localization signal (NLS), a sequence of nine amino acids which allows translocation of RUNX proteins from the cytoplasm to the nucleus (29). The C-terminus is essential for RUNX2 function (30,31), and contains the proline-serine-threonine-rich (PST) region which has both transactivatory and transinhibitory functions (29,31). The nuclear matrix targeting signal (NMTS) is located within the PST region, and in concert with the NLS helps localize RUNX2 to the nucleus (32). Finally, the ending five residues in the PST region, referred to as VWRPY, act as a transcriptional repression domain

(33,34). RUNX2 is expressed as two isoforms, Type II/MASNS/p57 and Type I/MRIPV/p56, which are each produced via initiation of transcription at P1 or P1 promoters, respectively (35).

Mutations in RUNX2 or CBF β can have significant consequences to health. Mutations in RUNX2 are strongly implicated in cleidocranial dysplasia (CCD) (36), a disease characterized by complications with tooth and bone development. Mutations in patients with CCD are commonly found in the RHD (37), and interfere with DNA binding (38) or heterodimerization with CBF β (36). Mutations in CBF β , in the form of translocations, are found in a large percentage of human leukemias (39). These translocations are most commonly in the form of CBF β -SMMHC, a fusion protein with altered affinity to RUNX1 (40) consisting of the heterodimerization domain of CBF β fused to the coiled-coil domain of smooth muscle myosin heavy chain (40,41).

CBF β lacks an NLS (42) and relies on the RUNX proteins to be shuttled into the nucleus. While the traditional role of CBF β as a transcriptional co-activator in the nucleus is well-described, recent research has proposed that it performs a non-canonical role in the cytoplasm as a regulator of protein translation (43). This is especially interesting in the context of OS, as protein translation is a convergence point of multiple signaling pathways (44), and targeting this process therapeutically could provide a route to overcome the high degree of heterogeneity amongst OS patient tumors (45). According to observations thus far only documented in breast cancer cell lines, CBF β binds to RNAs through hnRNPK, and thereby regulates the translation of proteins via eIF4B (43).

In our hands as well as published research, knockout of CBF β leads to a reduction in RUNX protein expression (43), and knockout of one of the RUNX proteins leads to a compensatory up-regulation in other RUNX proteins (46,47). Additionally, shuttling of CBF β into the nucleus by RUNX proteins may decrease the cytoplasmic pool of CBF β available to modulate translation.

Given the apparent mutual exclusivity of CBF β binding to RUNX proteins or hnRNPk (43), and the roles accomplished through binding to each occurring in separate cellular compartments, it's possible these roles are antagonistic in some manner.

The strongly intertwined nature of RUNX proteins and CBF β makes individual study of CBF β challenging, so we endeavored to inhibit the transcriptional role of CBF β in order to isolate and study the role CBF β may play in protein translation. Inhibition of the transcriptional role of CBF β via small molecule was first attempted (24,48), however the compound we tested did not significantly interrupt CBF β interaction with RUNX2. Instead, we opted to inhibit the transcriptional role of CBF β by introducing point mutations into the RUNX2 binding face of CBF β , with the goal of interrupting CBF β -RUNX2 interaction and subsequent nuclear shuttling of CBF β . Cell lines expressing this mutant form could then allow further study of the role of CBF β in protein translation, as well as add clarity into potential competition between RUNX proteins and hnRNPk for the same binding site on CBF β .

Materials and Methods

Cell Lines and Culture Conditions

Parental U2OS and U2OS-derived cell lines were maintained in McCoy's 5A (Iwakata & Grace Modification) media supplemented with 10% FBS and 1% P/S as previously described in Chapter 2.

Recombinant RUNX2 and CBF β Production

Vector Design: RUNX2 was cloned out of Genscript pcDNA3.1 OHu22872 containing NM_001024630.4 using primers outlined in **Table 1**. RUNX2 was then inserted into pGEX 6.1 expression vector using *PasI* and *XhoI* restriction enzymes from New England Biolabs (NEB). Restriction digestion was accomplished sequentially in recommended buffers with 1% agarose gel purification cleanup between digestions. Following gel purification, DNA was extracted from gels using Thermo Fisher GeneJET gel extraction kit according to manufacturer's recommendations, with elution accomplished by 30 μ L nuclease-free water. Following this, insert and vector were mixed together in 3:1 stoichiometric ratio of insert:vector and ligated using T4 Ligase from Thermo Fisher according to manufacturer's recommendations. The ligated vector was then transformed into NEB5 α *E. coli* according to manufacturer's recommendations, then plated on LB Agar plates containing 100 μ g/mL carbenicillin and grown at 37°C overnight. Single colonies growing in the presence of 100 μ g/mL carbenicillin were selected and used to inoculate 5 mL of LB broth supplemented with 100 μ g/mL carbenicillin and grown overnight at 37°C with 180 rpm shaking. Following overnight outgrowth, plasmid was extracted using Thermo Fisher GeneJET Plasmid Miniprep system according to manufacturer's recommendations, with elution

accomplished by addition of 30 μ L nuclease-free water. Eight hundred (800) ng of purified plasmid was sent to Sanger sequencing at Azenta Labs for verification of successful cloning. pGEX vector incorporating RUNX2 was then transformed into NEB T7 Express *E. coli* according to manufacturer's recommendations, then plated on LB Agar plates containing 100 μ g/mL carbenicillin and grown at 37°C overnight. Following transformation, a single colony was selected and used to inoculate five mL of LB + 100 μ g/mL carbenicillin, which was grown overnight at 37°C with 180 rpm shaking. Following overnight growth, 500 μ L of bacterial culture was mixed with 500 μ L of 1:1 Glycerol:LB media (v/v) and frozen in cryovials at -80°C until recombinant protein production steps below.

CBF β was cloned out of Genscript pcDNA3.1 OHu25418 incorporating NM_022845.3 using primers outlined in **Table 1**. Insert was cloned into the pTXB1 expression vector, which imparted an Mxe intein/chitin binding domain (CBD) at the C-terminus. Insertion was accomplished via simultaneous digestion by XhoI and SapI (NEB) restriction enzymes in NEB Cutsmart 3.1 buffer according to manufacturer's recommendations, followed by ligation as described above. Finally, sequence verification, transformation into NEB T7 Express, outgrowth and creation of frozen glycerol stocks was accomplished as described previously.

Table 1: Primers for Recombinant Protein Production

| Usage | Primer Sequence |
|---|--|
| Add TspMI restriction site to RUNX2 N term for cloning into pGEX-6P-1 F | 5'-GGTGGTCCCTGGGCATG GCATCAAACAGCCTCTT-3' |
| Add XhoI restriction site to RUNX2 C-term for cloning into pGEX-6P-1 R | 5'-GGTGGTC'TCGAGTCAATA TGGTCGCCAAACAG-3' |
| Add NdeI restriction site to CBFb N-term for cloning into pTXB1 F | 5'-GGTGGTC'ATATGATGCCG CGCGTCGTGCCCGA-3' |
| Add SapI restriction site to CBFb C- term for cloning into pTXB1 R | 5'-GGTGGTTGCTCTTCC'GCAAC GAAGTTTGAGGTCATCAC-3' |
| Remove extra N-term Met from CBFβ-pTXB1 F | 5'-CCGCGCGTCGTGCCCGAC-3' |
| Remove extra N-term Met from CBFβ-pTXB1 R | 5'- CATATGTATATCTCCTTCTTAAAGTT AAACAAAATTATT TCTAGAGGGGAATTGTTATCCGCTC- 3' |

Recombinant Protein Expression: A glycerol stock containing NEB T7 Express *E. coli* cells harboring either recombinant RUNX2-GST or CBFβ-CBD was thawed at room temp for 1 min, then 10 μL of culture was used to inoculate 50 mL of LB broth supplemented with 100 μg/mL carbenicillin. Inoculated media was grown overnight at 37°C with 180 rpm shaking. The following day, the 50 mL culture was used to inoculate a 1L flask of LB supplemented with 100 μg/mL carbenicillin, which was then grown at 37°C with 180 rpm shaking. Optical density (OD) measurements were taken periodically and culture was induced for protein expression at OD ~0.6 by addition of isopropyl β-D-thiogalactoside (IPTG) (Millipore Sigma) to 0.5 mM final concentration. Following induction, cultures were incubated overnight at 16°C with 150 rpm shaking. The following day cultures were spun down at 3,200g in a tabletop centrifuge set to 4°C until the entire culture was pelleted. The supernatant was discarded and pellets were stored at -80°C overnight. Pellets were then thawed on ice and resuspended using lysis buffer consisting of

20 mL PBS supplemented with 1 cOmplete protease inhibitor cocktail tablet and 0.5mM PMSF final concentration. Bacteria slurry was placed on ice and sonicated for 19 min consisting of 10 seconds on and 10 seconds off, using a Fisherbrand Sonic Dismembrator sonicator with amplitude set to 43. Lysate was then spun down at 17,000 g for 20 min. at 4°C to yield clarified lysate, and pellet was discarded.

Recombinant CBF β Purification: CBF β was purified from clarified lysate by 2h rotation at 4°C with 2.5mL bed volume pre-washed NEB chitin resin. Lysate and bead mixture was then poured into a 10 mL Pierce Disposable Column (Thermo Fisher), and 200 mL ice cold PBS was poured through column to wash beads. Beads were then incubated at 4°C for 4 days in 10 mL PBS with 50mM DTT to cleave chitin tag. Free CBF β was eluted from column into a tube and PBS was added to bring volume to 30 mL total. Solution was then concentrated by centrifugation at 4°C and 3,200g with an Amicon 3 kilodalton molecular weight cutoff spin concentrator tube (Millipore Sigma). Following concentration down to 2 mL total volume, purified protein was desalted by passing through Thermo Fisher Zeba spin desalting columns twice, according to manufacturer's recommendations.

Recombinant RUNX2 Purification: RUNX2 was purified from clarified lysate by 1.5h rotation at 4°C with 1.25mL bed volume pre-washed Pierce Glutathione Agarose GSH beads (Thermo Fisher). Beads were then spun down at 700g at 4°C, then washed 3x with PBS supplemented with 0.1% Triton x-100. Beads were suspended in sufficient PBS + 0.1% Tritox-100 to achieve a 50% slurry, and kept at 4°C until time of assay start.

Recombinant Protein Expression Verification: Recombinant expression of RUNX2 and CBF β was verified by running on SDS-PAGE as described previously in Chapter 2. Gel was then

stained using GelCode™ Blue Safe Protein Stain according to manufacturer's recommendations. Gel was then imaged using ProteinSimple imaging platform set to epi white imaging settings.

RUNX2-CBF β Binding Assay

Pulldown Assay Performance: 47 femtomoles of recombinant CBF β and 378 femtomoles of recombinant RUNX2-GST bound to GSH agarose beads were spiked into 1,920 μ L of PBS supplemented with 0.1% Triton x-100 and placed on a rotator overnight at 4°C. Beads were then washed 3x with 1 mL PBS + 0.1% Triton x-100, and bound proteins were eluted by addition of 42 μ L SDS-PAGE sample buffer (5:1:14 ratios of 4x Laemmle Buffer:1M DTT in water:1x PBS) and heated at 70°C for 10 min. with occasional vortex mixing. Beads were spun down for 10 seconds at 1,000g and supernatant was diluted 81x with SDS-PAGE sample buffer, then subjected to western blotting analysis as previously described. PVDF membranes were probed with CST α -CBF β antibody #D4N2N diluted 1:10,000 and CST α -RUNX2 antibody #D1L7F diluted 1:50,000, all in 5% NFDM in TBST.

Peptide Production and Dilution: RUNX2-derived peptide (GNDENYSAEL) and CBF β -derived peptide (ATGTNLSLQFF) were synthesized by Genscript and solvated in DMSO to 30 mg/mL and stored at 4°C. As RUNX2 and CBF β -derived peptides have different properties, a separate control peptide for each was chosen. Sequence SSLTNGLFR was selected as a control peptide for CBF β -derived sequence, and sequence IATEAIENIR was selected as a control peptide for RUNX2-derived sequence. Control peptides were synthesized by Celtek Peptides (Franklin, TN) and also solvated to 30 mg/mL in DMSO and stored at 4°C.

Pulldown with CBF β -derived Peptide: 378 femtomoles of recombinant RUNX2 was pre-incubated with 3.78 or 37.8 picomoles CBF β -derived peptide or control peptide for 1h at 4°C on a rotator. Following pre-incubation, 47 femtomoles of recombinant CBF β was spiked into tube and pulldown was accomplished as described above.

Pulldown with RUNX2-derived Peptide: 47 femtomoles of recombinant CBF β was pre-incubated with 470 or 4,700 picomoles RUNX2-derived peptide or control peptide for 1h at 4°C on a rotator. Following pre-incubation, 378 femtomoles of recombinant RUNX2 was spiked into tube and pulldown was accomplished as described above.

Structural Modeling

3D structure of CBF β bound to RUNX2 (PDB #6VGE) (49) was rendered using the PyMOL Molecular Graphics System, Version 2.4.1, Schrödinger, Incorporated. ERG and 16mer DNA strand in original structure were hidden from view and sidechains of CBF β residues G61, N63, and N104 were made visible along with their respective Hydrogen bonds.

Lentiviral Transduction

Plasmid Design: CBF β 3x Mut (G61A, N63A, N104A) was generated through site directed mutagenesis of pHIV-EGFP incorporating wild-type (wt) CBF β _FLAG, generated previously. Mutagenesis was accomplished using NEB Q5 Mutagenesis Kit according to manufacturer's instructions, with primers used listed in **Table 2**. Mutagenesis was accomplished in two separate

reactions, with vector transformed into NEB Stable cells in between reactions as previously described, and plasmids were sent for Sanger sequence verification by Azenta Labs.

Table 2: Primer sequences for Lentiviral Vector Generation

| Usage | Primer Sequence |
|---|------------------------------------|
| Mutate CBF β N104A F | 5'-CATGATTCTGGCCGGAGTCTGTGTTATC-3' |
| Mutate CBF β N104A R | 5'-GGAGCCTTCAAATATACC -3' |
| Mutate CBF β G61A and N63A F | 5' CGCACTGTCTCTCCAGTTTTTTC 3' |
| Mutate CBF β G61A and N63A R | 5' GTCGCTGTGGCCACAAAAGCGAT 3' |
| Sequence T7 | 5'- TCAAGCCTCAGACAGTGGTTC-3' |
| Begin Sanger sequencing at EF-1 α promoter | 5'- TCAAGCCTCAGACAGTGGTTC-3' |

U2OS CBF β KO cells were subjected to lentiviral transduction using same protocol as outlined previously, this time using transfer plasmid pHIV-EGFP CBF β _FLAG (3xMut) generated as described below. Cells infected by 1953x dilution of lentiviral particles were chosen for outgrowth in a T75 flask based on <5% EGFP+ cells. As previously mentioned, this dilution was chosen to minimize risk of double integration from lentivirus. Sorting of individual cells and outgrowth to generate monoclonal cell lines was conducted as previously outlined.

Cellular Electrothermal Shift Assay (CETSA)

U2OS-derived cell lines were grown on 10 cm plates to 70% confluence then harvested via Trypsin. Cells were then washed with PBS twice, diluted 10x with 0.2% Trypan Blue in PBS (MP Biomedicals, LLC, Irvine, CA) and counted with a hemocytometer (Thermo Fisher). A cell suspension of 5.6 million cells/mL was prepared in PBS supplemented with 1x cComplete protease inhibitor tablet (Thermo Fisher) and 1mM PMSF (Sigma-Aldrich). 100 μ L of cell suspension was

placed into each of two PCR tubes, with one tube placed into a SimpliAmp Thermocycler (Life Technologies, Carlsbad, CA) and incubated at 40°C - 68°C (during RUNX2 melting temp assessment) or 49.5°C (during CETSA cell line panel) for 3 min while the other was incubated at room temperature for the same duration. Following incubation, cell suspensions were flash frozen by immersion in liquid nitrogen for 1 min, then thawed at 25°C for 5 min. Cell suspensions were flash frozen and thawed again, then spun down at 20,000g for 20 min. at 4°C. Supernatant was transferred to a fresh tube and mixed with Laemmle buffer and DTT to 1x and 50 mM final concentrations, respectively, and heated at 70°C with occasional vortex mixing for 10 minutes. Sample was then analyzed via western blotting as previously described.

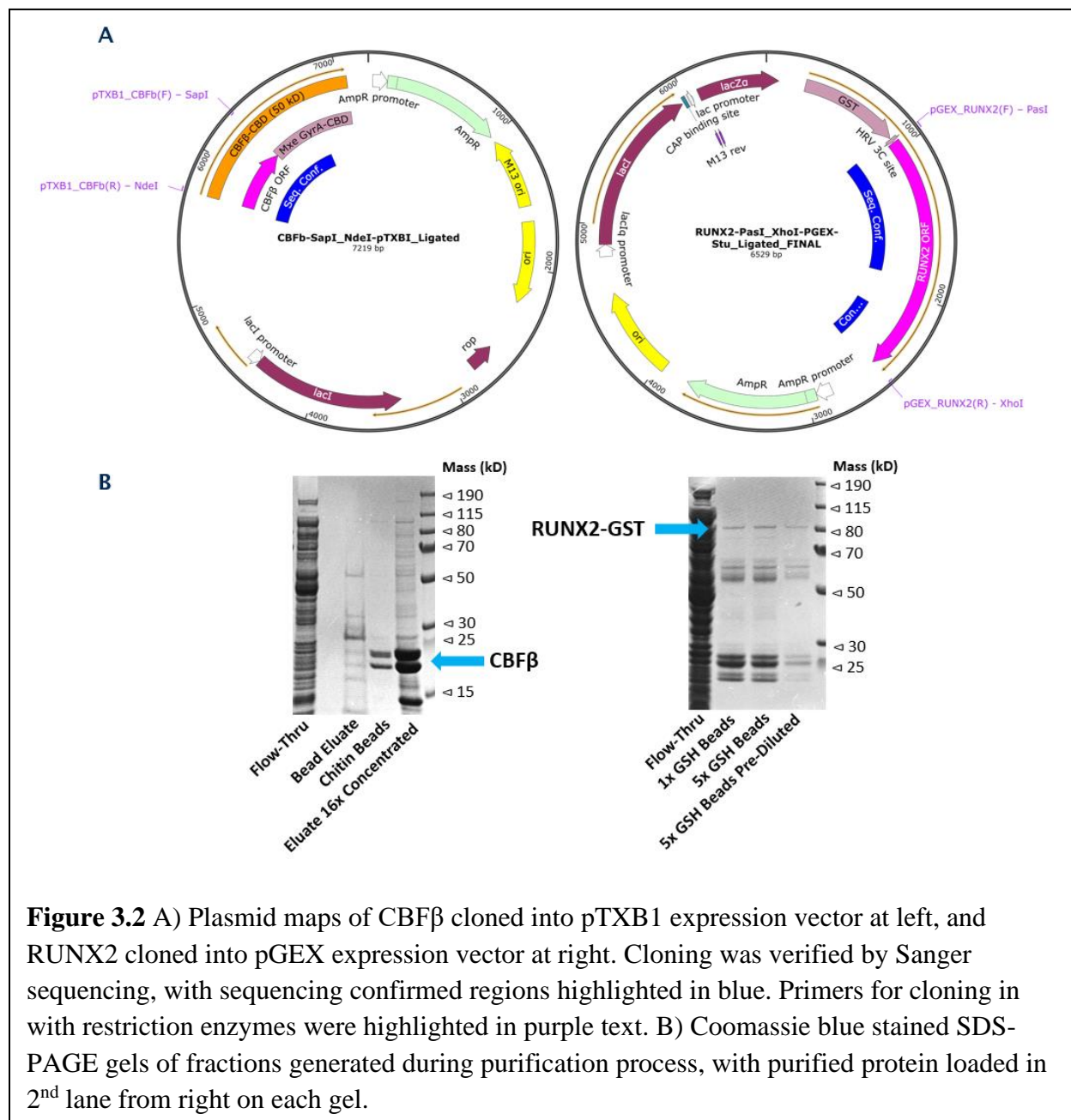
Nuclear and Cytoplasmic Fractionation

Nuclear and cytoplasmic fractionations were performed with Thermo Fisher NE-PER Kit, #78833 with slight modifications to manufacturer's recommendations. In short, the nuclear pellet was subjected to an additional wash with 200:11 (v/v) Cytoplasmic Extraction Reagent (CER) I :CER II, and pellet was vortex mixed with wash for 5 minutes. Wash was then removed with gel-loading pipette tip. Nuclear Extraction Reagent (NER) was added in manufacturer's recommended proportions, and nuclear pellet was broken up by 8 cycles of sonicating for 30s, vortexing for 15s, and placing on ice for 5 min. Samples were then centrifuged at 16,000g for 10 min. at 4°C, and placed on ice. Extracts were then mixed with NuPage Laemmle Buffer and DTT to 1x and 50mM final concentrations, respectively, and analyzed via western blotting as described previously.

Statistical Analyses

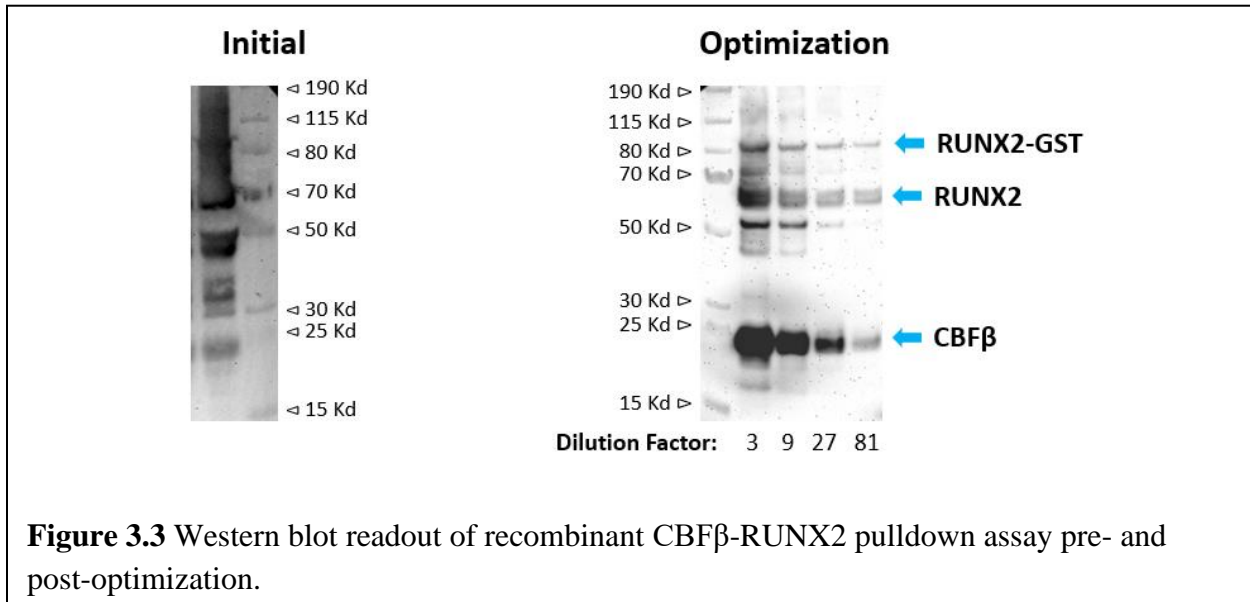
Figure 3.7C,D: Error bars are SD, n=5 (biological); * = $p < 0.05$, **** = $p < 0.001$, one-way ANOVA with Tukey's post hoc test. **Figure 3.8D:** Error bars are SD, n=3 (biological); *** = $p < 0.001$, **** = $p < 0.0001$, one-way ANOVA with Tukey's post hoc test.

Results



CBFβ and RUNX2 were successfully cloned into pTXB1 and pGEX vectors, expressed, and purified recombinantly. Cloning of CBFβ and RUNX2 into their respective expression vectors was successful as verified by Sanger sequencing. Purification endeavors of both proteins were also

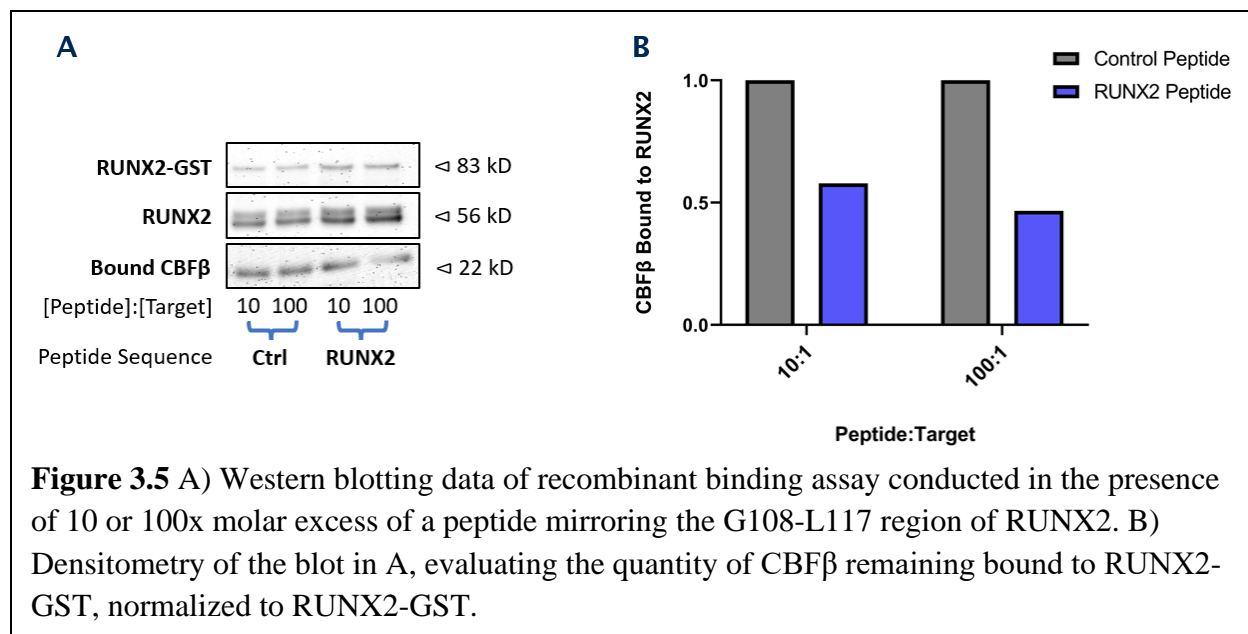
successful, with the above gels representing eluates from a 1L culture of NEB5 α cells. Purified CBF β produced much darker bands on the gel than did RUNX2-GST. Faint bands additional to proteins of interest were present in both purified CBF β and RUNX2.



Optimization of CBF β -RUNX2 recombinant pulldown assay. A pulldown assay was conducted to verify interaction of recombinant CBF β and RUNX2. Initial western blot was unreadable and successful interaction of recombinant CBF β and RUNX2 could not be concluded. Pulldown conditions and antibody concentrations were optimized to yield a cleaner blot, and verify conclusively that recombinant CBF β and RUNX2 do indeed interact. Optimized conditions were then used for further pulldown assays.



RUNX2 region G108-L117 and *CBFβ* region G61-G69 were selected for targeting via a synthetic peptide. We delved into the literature and compiled a list of all residues which appear to contribute to CBFβ-RUNX2 interaction, generating the above figure as guidance. Based on these data, we elected to use peptides mirroring the sequence of RUNX2 in region G108-L117 and the sequence of CBFβ in region G61-G69.



Recombinant RUNX2 and CBFβ interaction is inhibited by a peptide mirroring the sequence of RUNX2. When pulldown of recombinant CBFβ by recombinant RUNX2-GST was attempted in the presence of a peptide crafted after the sequence of RUNX2 (GNDENYSAEL), we saw a dose-dependent inhibition of CBFβ-RUNX2 interaction. This was evaluated by normalizing CBFβ to RUNX2-GST, then normalizing that to equal quantity of control peptide. We did not see inhibition of CBFβ-RUNX2 interaction when attempting pulldown in the presence of a peptide mirroring the sequence of CBFβ (ATGTNLSLQFF) (data not shown).

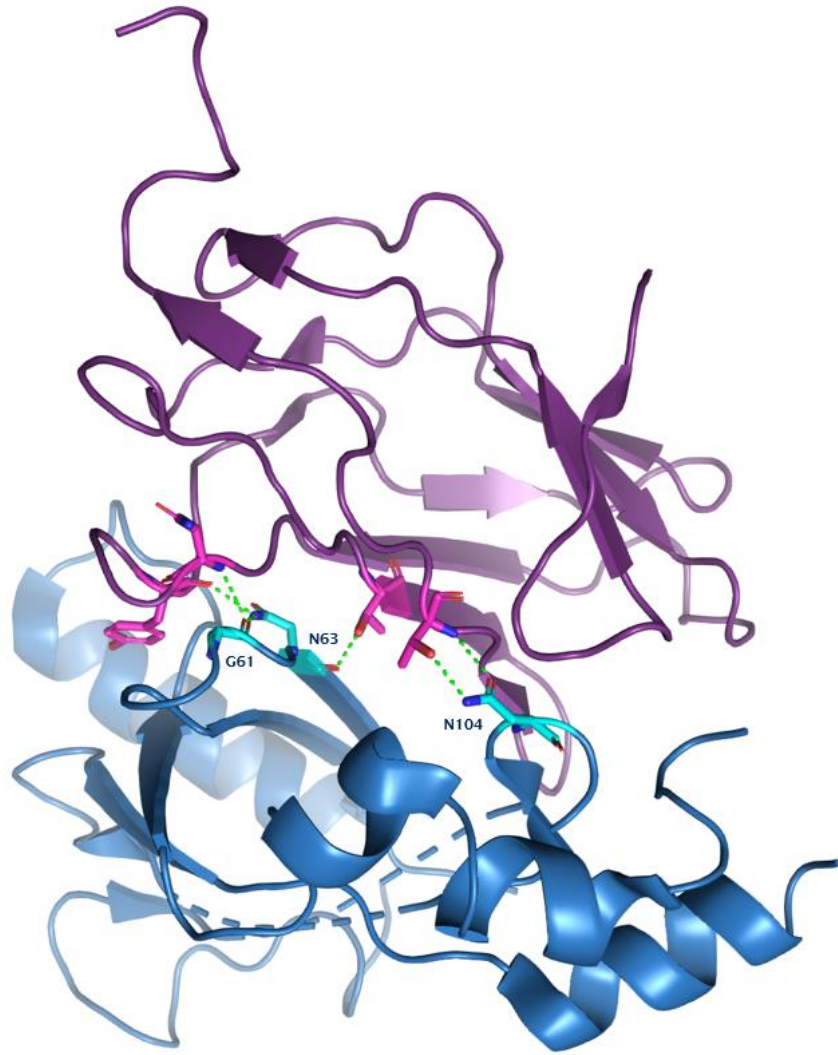
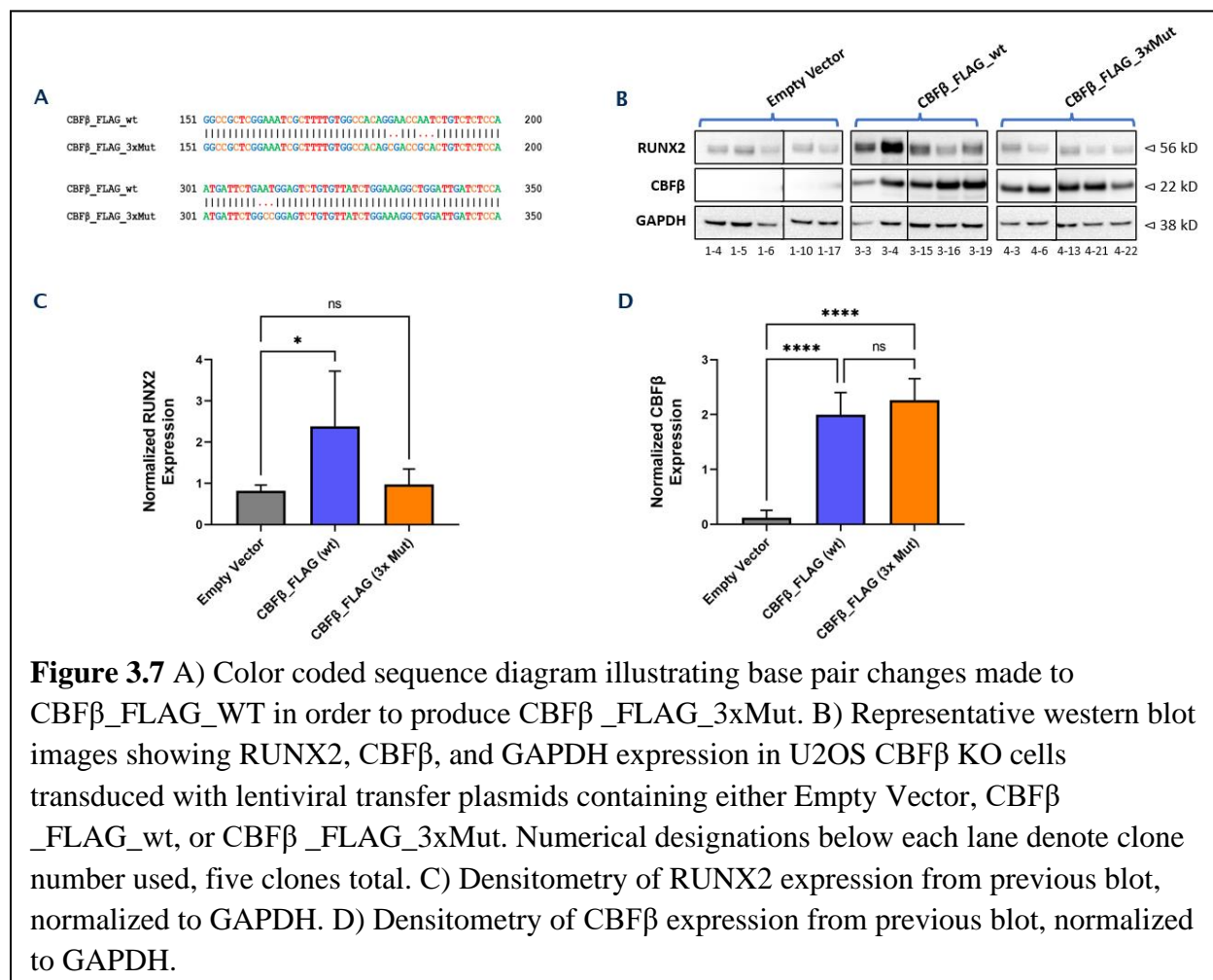
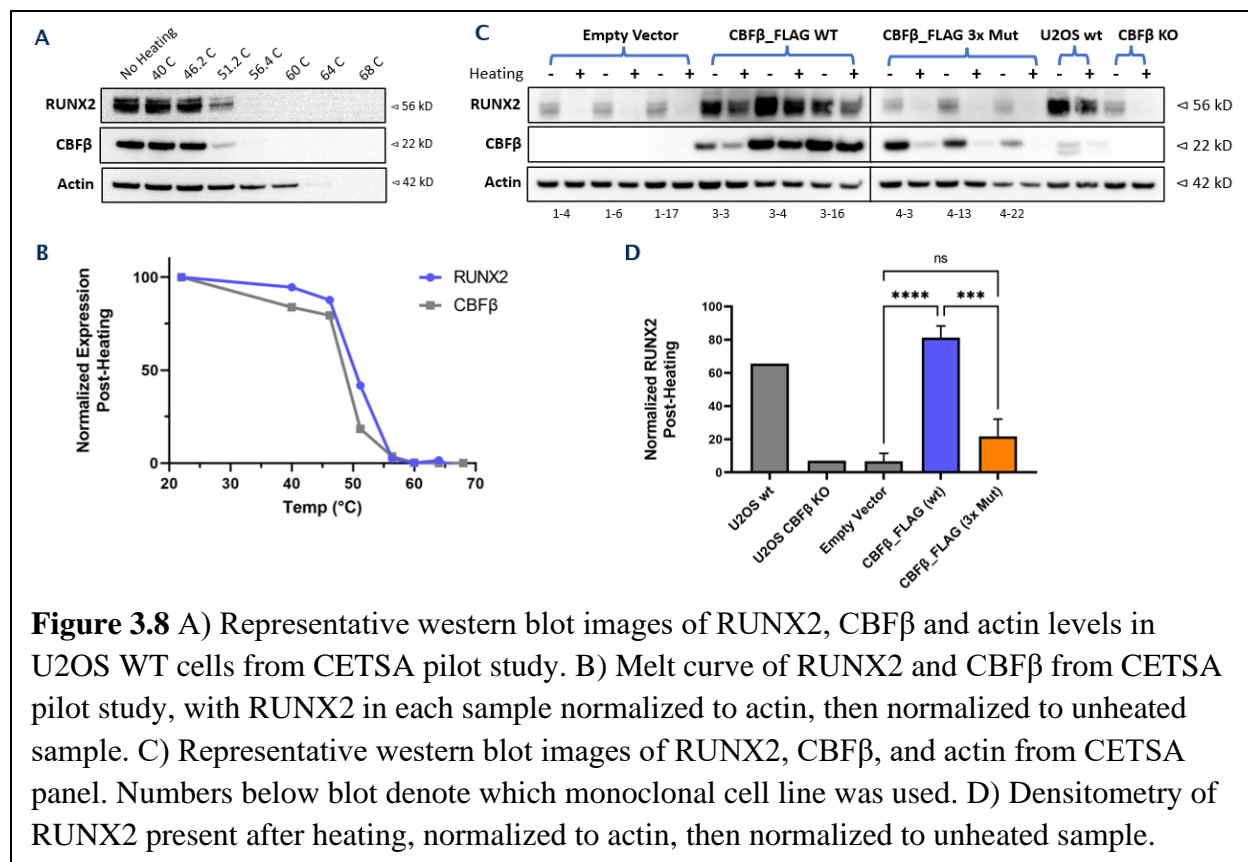


Figure 3.6 Rendering of PDB #6VGE showing RUNX2 (purple) interacting with CBFβ (blue). Residues of CBFβ targeted for mutation are visualized in teal, with residues on RUNX2 they interact with visualized in magenta. Hydrogen bonds between key residues of CBFβ and RUNX2 are highlighted in neon green.

Structure of CBFβ bound to RUNX2 provides visual clues of key amino acids. RUNX2 bound to CBFβ was rendered in Pymol referencing structure #6VGE, with color coding done to accentuate the two proteins as well as clearly display the key residues selected for mutation into alanine.



Stable expression of a mutant form of CBFβ is unable to rescue low RUNX2 protein expression. To investigate the significance of CBFβ-RUNX2 binding in the regulation of RUNX2 protein expression by CBFβ, we mutated previously validated residues glycine 61, asparagine 63, and an additional residue, asparagine 104, of CBFβ to alanine using site-directed mutagenesis (SDM) to generate CBFβ_FLAG_3xMut. Stable expression of CBFβ_FLAG_3xMut did not rescue low RUNX2 expression in the same manner as CBFβ_FLAG_wt. The expression level of CBFβ_FLAG_3xMut transgene did not differ significantly from that of CBFβ_FLAG_wt.



Mutations in CBF β shift the melting temperature of RUNX2. In order to investigate whether key mutations in CBF β had affected CBF β -RUNX2 interaction, we conducted a cellular electrothermal shift assay (CETSA) on our transduced cells. A melt curve was first conducted to determine the melting temperature (T_m) of RUNX2 in CBF β _FLAG_WT cells, and cell suspensions were subjected to a range of temperatures from 22°C to 68°C. Following normalization to actin and then unheated sample, the melting temperature of RUNX2 was found to be ~50°C. 49.5°C was selected for future heat treatments, and CETSA was conducted on five clones each of Empty Vector, CBF β _FLAG_WT, and CBF β _FLAG_3xMut cells as well as U2OS WT and U2OS CBF β KO cells. The T_m of RUNX2 in U2OS WT cells was similar to that of CBF β _FLAG_WT cells, while the T_m of RUNX2 in CBF β KO cells was similar to that of Empty Vector and CBF β _3xMut_FLAG cells. Most importantly, expression of CBF β _3xMut_FLAG

caused a statistically significant downward shift in RUNX2 Tm relative to cells expressing CBF β _FLAG_WT.

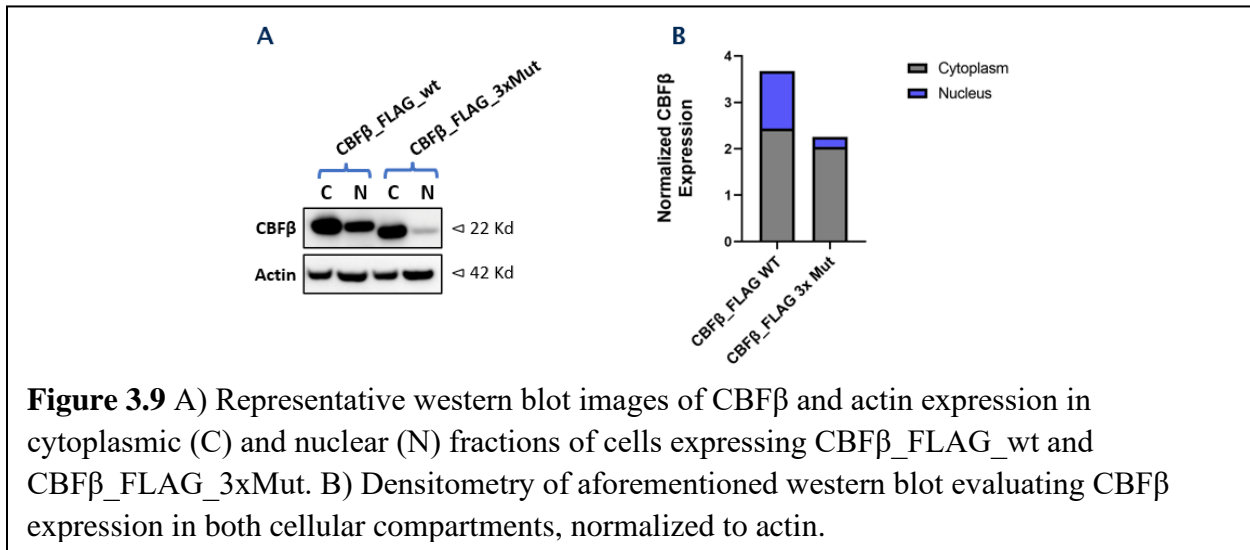


Figure 3.9 A) Representative western blot images of CBF β and actin expression in cytoplasmic (C) and nuclear (N) fractions of cells expressing CBF β _FLAG_wt and CBF β _FLAG_3xMut. B) Densitometry of aforementioned western blot evaluating CBF β expression in both cellular compartments, normalized to actin.

Mutations in CBF β inhibit nuclear shuttling of CBF β . In order to investigate the functional significance of point mutations in CBF β , we next subjected CBF β _FLAG_WT and CBF β _3xMut_FLAG-expressing cells to nuclear/cytoplasmic fractionation followed by western blotting for CBF β and actin. CBF β _FLAG_WT was present in both the cytoplasm and the nucleus, as expected, while CBF β _FLAG_3xMut was localized predominantly to the cytoplasm. Actin was present in both compartments and functioned as our housekeeping gene for this experiment.

Discussion

CBF β and RUNX2 were successfully cloned, expressed, and purified, verified via SDS-PAGE with Coomassie blue staining. Although recombinant CBF β produced a more intense band than RUNX2, this was likely due to the 8x concentration step imparted to CBF β during purification while RUNX2 remained stuck to affinity beads and was not concentrated. We did see a few additional bands in our purified CBF β , however further purification of recombinant CBF β was not possible due to aggregation issues when run on a Fast Protein Liquid Chromatography (FPLC) system. Additional bands were also present in recombinant RUNX2-GST, which likely correspond to free GST tag (26 kD) and *apo* RUNX2 (56 kD) produced by cleavage at HRV-3C protease site by endogenous proteases. Altogether, these impurities were faint in comparison to our protein of interest and therefore low in abundance (**Figure 3.2**).

We next performed and optimized a pulldown assay using our recombinantly expressed proteins, and confirmed that recombinant CBF β and RUNX2 do interact (**Figure 3.3**). Using this recombinant pulldown assay, we evaluated if targeting certain key residues of CBF β and RUNX2 would interrupt their interaction. The G61-G69 region of CBF β and corresponding G108-L117 face on RUNX2 were selected based on previous work highlighting these regions as important in CBF β -RUNX2 interaction (50,51) (**Figure 3.4**). We chose to synthesize a relatively short peptide targeting this region, as this would afford greater permeability into living cells when mixed with a cell penetrating peptide carrier (52–54). Repeat of pulldown assay in the presence of peptide GNDENYSAEL, crafted from RUNX2 region G108-L117, led to a dose-dependent inhibition of CBF β -RUNX2 interaction when normalized to RUNX2-GST (**Figure 3.5**). Conducting the pulldown assay in the presence of peptide GTNLSLQFF, crafted from CBF β region G61-G69, did not inhibit CBF β -RUNX2 interaction (data not shown). Nevertheless, as inhibition of CBF β -

RUNX2 interaction with peptide GNDENYSAEL was successful, and exhibited a dose-dependent effect, we felt confident in moving forward with further studies targeting residues G61 and N63 of CBF β .

With the likelihood of dual roles for CBF β occurring simultaneously, in distinct subcellular regions, and the protein interactions essential to each seemingly exclusionary of each other (43), studying each role individually proved difficult. In order to isolate and investigate the translational role of CBF β , we attempted to halt its transcriptional capabilities by generating cell lines expressing a mutant form of CBF β designed to be unable to bind with the RUNX proteins. With the residues key in CBF β -RUNX2 interaction now validated in our previous recombinant protein assay, we utilized SDM followed by lentiviral transduction to generate viral particles carrying CBF β _FLAG_3xMut and used these to infect U2OS CBF β KO cells. To be specific, CBF β _FLAG_3xMut was generated by mutating our previously validated residues, G61 and N63, to alanine. Additionally, we elected to also mutate N104 to alanine because of data suggesting it plays an important role in CBF β -RUNX2 interaction as well (50).

The side-chains of these three key residues, G61, N63, and N104, along with the hydrogen bonds they form with corresponding residues on RUNX2 are highlighted in **Figure 3.6**. Visualization with Pymol was helpful to make sure these residues do indeed lie on the RUNX2 interacting face of CBF β . We then evaluated RUNX2 expression in our cell lines expressing our triple mutant of CBF β , and found that CBF β _FLAG_3xMut transgene was unable to rescue low RUNX2 expression in CBF β KO cells (**Figure 3.7**). This observation was not due to hampered transgene expression, as CBF β levels were consistent between CBF β _FLAG_WT and CBF β _FLAG_3xMut.

We were somewhat surprised that CBF β _FLAG_3xMut was unable to rescue low RUNX2 expression in U2OS CBF β KO cells, as our previous data indicated that stabilization of RUNX2 via binding to CBF β and subsequent protection from degradation made a relatively minor contribution to overall RUNX2 levels. Previous research also suggested that CBF β binding and stabilization of RUNX proteins is not the dominant factor dictating RUNX protein levels (43), so this result was initially puzzling indeed. Previous research suggested that CBF β interaction with hnRNPK or RUNX proteins is mutually exclusive (43), so it's possible that these two binding partners of CBF β compete for the same residues, and our mutations had inadvertently altered the interaction of CBF β with both. Inhibiting the interaction of CBF β and hnRNPK would, in theory, lead to reduced expression of proteins under the purview of CBF β , as interaction with hnRNPK is the mechanism by which this translational role of CBF β is reported to occur (43). In order to fully unravel this, it is necessary to validate that 1) mutations in CBF β did inhibit binding with RUNX2 in living cells, 2) RUNX2 protein expression levels are strongly regulated by CBF β , and 3) mutations in CBF β alter its ability to facilitate translation of RUNX2 and other proteins. The first point will be addressed below while points two and three will be covered in a latter chapter through use of ribosome footprinting (55–57) and direct detection of biotinylated tags (DiDBiT) (58). Data generated for another project in our lab (not shown) seems to support the third point here, as CBF β _FLAG_3xMut cells appear to double at a slower rate than CBF β _FLAG_WT cells, which could be due to decreased overall protein synthesis resulting from the inability of CBF β to participate in protein translation.

While we had validated these residues as crucial to CBF β -RUNX2 interaction in our cell-free system, and used this data to generate CBF β _FLAG_3xMut, it was necessary to validate that we had indeed interrupted their binding in living cells. The simplest method to measure alterations

to a protein-protein interaction would have been to conduct a co-IP, which was our initial strategy. Unfortunately, RUNX2 proved incredibly challenging to co-IP, as its mass of 56 kD causes it to run parallel to the IgG heavy chain of any pulldown antibody used. Due to this, we were not able to get a clean blot for co-IP of RUNX2 as it was always obscured by background signal from the heavy chain of the pulldown antibody. Numerous workarounds to this were attempted, including usage of Protein A-HRP secondary antibodies, IgG light-chain specific HRP secondary antibodies, four different elution buffers, three different pulldown antibodies, and running SDS-PAGE in non-reducing conditions. To our chagrin, none of these were able to produce a clean blot, and stumped us as well as the technical support staff at four different reagent companies.

Due to these setbacks, we switched to CETSA, and used this assay to assess if CBF β _FLAG_3xMut had inhibited RUNX2 binding compared to CBF β _FLAG_WT. CETSA was first developed as a label-free method to evaluate target engagement in drug development, and is based on the principle that changes in the interactions of a particular protein will increase or decrease its melting temperature (T_m), thus shifting it from a soluble to an insoluble state (59). While this remains its most common usage (60), we hypothesized CETSA could also prove useful for our application. As cellular proteins have a wide range of melting temperatures (61) and their individual interactomes differ greatly between cell lines (62), it was necessary to first determine the T_m of RUNX2 in cells expressing CBF β _FLAG_WT. We conducted a melt curve using temperatures spanning 40°C to 68°C, and after normalizing to actin, we measured the T_m of RUNX2 to be approximately 49.5°C (**Figure 3.8A,B**). We then used this same temperature for future studies, and a decrease in RUNX2 protein remaining after heating, when compared to CBF β _FLAG_WT, was interpreted as a decrease in T_m and therefore a decrease in RUNX2 interaction with other proteins.

While RUNX2 protein is expressed at different levels in our cell lines, as shown previously, this did not confound our results as each cell line is normalized to its individual RUNX2 expression level. RUNX2 displayed a similar T_m between U2OS WT cells and CBF β _FLAG_WT cells, indicating that our stably expressed CBF β _FLAG_WT interacted with endogenous RUNX2 to the same extent as in U2OS WT cells (**Figure 3.8C,D**) and suggesting that our rescue experiment had successfully recapitulated normal CBF β -RUNX2 interaction. RUNX2 experienced a downward shift in T_m in both U2OS CBF β KO and Empty Vector cells, which was expected as RUNX2 in both lines had lost its interacting partner CBF β . Most prominently, RUNX2 experienced a statistically significant downward shift in T_m in CBF β _FLAG_3xMut cells, indicating that we had been successful in generating a CBF β mutant with inhibited RUNX2 binding.

CBF β lacks a nuclear localization signal (NLS) (42,63) and must be shuttled into the nucleus onboard the RUNX proteins (43,64); therefore inhibition of CBF β -RUNX protein binding should result in decreased nuclear accumulation of CBF β . We evaluated this by performing nuclear/cytoplasmic fractionation followed by western blotting, and were able to confirm these mutations interrupted nuclear shuttling of CBF β (**Figure 3.9**). Cells expressing CBF β _FLAG_WT demonstrated CBF β within both cytoplasmic and nuclear compartments, in line with previous studies in our hands and others (43), and confirming our affinity tag had not interrupted shuttling. In contrast to this, CBF β _FLAG_3xMut was predominantly localized to the cytoplasm, suggesting that our mutations had interrupted nuclear shuttling of CBF β . The nuclear fraction did still exhibit a faint band corresponding to CBF β , which may be explained by the actions of Crlz-1 (also known as SAS10 or UTP3) which interacts with CBF β (65) and has been postulated to play a role in CBF β nuclear shuttling (66,67) and OS development (68). While Crlz-1 may influence the cellular

distribution of CBF β , from our data it does not appear to be the predominant factor governing nuclear shuttling of CBF β .

Our choice of actin as a housekeeping gene for both nuclear and cytoplasmic fractions has been met with much skepticism by colleagues, as actin was long been thought to be exclusive to the cytoplasm. This was primarily due to a lack of reliable data detecting actin in the nucleus by immunofluorescence microscopy (69), however convincing evidence has since been put forth demonstrating actin in the nucleus of mammalian cells as well as numerous others (70–82). In previous experiments (not shown) we evaluated the purity of nuclear and cytoplasmic fractions generated with this method by blotting for nuclear-specific and cytoplasmic-specific markers Histone H3 and GAPDH, respectively, and validated that cross-contamination between fractions did not occur.

Our goal in generating CBF β _FLAG_3xMut was to inhibit the transcriptional activity of CBF β , and thereby de-couple the transcriptional and translational roles of CBF β . These data thus far suggest we have accomplished the former, as a mutant form of CBF β with reduced RUNX2 binding and subsequent nuclear shuttling is likely to be less effective as a transcriptional co-activator to the RUNX proteins. This assertion could be validated in future endeavors via qPCR of RUNX2 target genes implicated in the malignant phenotype of OS (15,16) such as vascular endothelial growth factor (VEGF) (11,12), alkaline phosphatase (ALP) (13), or matrix metalloproteinase 9 (MMP-9) (14). As mentioned previously, it is possible that these mutations have also affected the translational role of CBF β , and additional data generated by this project in a latter chapter will provide more detail into the influence of these specific residues as well as that of CBF β overall on the malignant phenotype of OS.

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Chapter 4

Profiling the interactome and translome of CBF β in osteosarcoma cells

Abstract

Previous chapters had generated a large assortment of data focusing on the potential for CBF β to regulate levels of RUNX2 in a post-transcriptional manner. In building on these previous data, we expanded our approach from investigating the influence of CBF β on solely RUNX2, to investigating the influence of CBF β on global phenomena in the cell. The assays performed in this chapter serve to answer two questions: 1) which proteins interact with CBF β ? And 2) which genes/proteins have expression regulated, at least in part, by CBF β ? Firstly, we performed affinity purification immunoprecipitation mass spectrometry (IP-MS) of CBF β _WT_FLAG cells, and generated a list of medium- and high-confidence binding partners of CBF β . We then performed gene ontology on these partners, and found multiple terms associated with protein translation to be highly enriched. Importantly, both members of the nascent polypeptide associated complex (NAC), a heterodimer crucial in protein expression, were identified as high confidence interactors. Additionally, we revealed proteins which specifically rely on residues G61, N63, and/or N104 of CBF β for their interaction. Secondly, we performed two somewhat orthogonal assays; Ribo-Seq (with parallel RNAseq) and Direct Detection of Biotinylated proteins (DiDBiT), to reveal a list of genes/proteins which may have their expression regulated by CBF β at transcription and/or translational level. We performed gene ontology on these targets, and observed enrichment of numerous pathways implicated in OS, and cancer at large, such as PD-L1, MAPK, and VEGF, among genes translationally regulated by CBF β . Notably, both Ribo-Seq and DiDBiT analyses yielded strong enrichment of genes associated with Rho GTPases, which are known to play key roles in tumor initiation and progression as well as proliferation and apoptosis.

Introduction

Osteosarcoma (OS) is the most common form of primary bone cancer in humans and canines, and most often presents early in life (1). The treatment outlook in OS hasn't changed in the last 30 years, in part due to the lack of identified driver mutations as well as OS being relatively uncommon when compared to other cancer types (2,3). In terms of location, OS lesions most frequently occur at the metaphysis, the portion of bone which contains the growth plate (4). The lack of identifiable driver mutations combined with the high degree of heterogeneity amongst OS patient tumors (5) has made development of targeted therapies challenging. Development of medicines which are able to overcome this heterogeneity could lead to dramatic successes in OS patients, but more study is needed to identify convergence points which could be targeted across large groups of patients. Protein translation has been proposed as such a convergence point (6), and this study will delve into this process. Core binding factor beta (CBF β) is a binding partner to the RUNX family of DNA-binding transcription factors, and formation of the CBF β -RUNX protein heterodimer allows RUNX proteins to better bind to DNA and facilitate transcription (7,8). RUNX2 is the master regulator of bone growth and differentiation (9), and high expression of it (10,11) as well as CBF β (12) are implicated in poor disease prognosis in OS. These factors led to the interaction of these proteins, and the roles of CBF β specifically, to be studied in detail in our lab (13).

CBF β has been proposed to participate in regulation of cap-dependent protein translation in breast cancer cells (14), although thus far this is the only cancer type where this noncanonical role of CBF β has been identified. Our study aims to interrogate whether this role also occurs in OS, and to what extent CBF β may impact OS malignancy. Previous chapters described efforts to explain the reduction in RUNX2 protein expression upon loss of CBF β , which may be due to a

post-transcriptional mechanism as loss of CBF β did not decrease RUNX2 mRNA expression. Although CBF β has been proposed to stabilize RUNX2, protecting it from proteasomal degradation (15,16), previous data in this dissertation suggest that stabilization of RUNX2 by CBF β may not fully explain this decrease in RUNX2 protein.

Building on our previous data, the current experiments aimed to delve deeper into the impact of CBF β on protein translation, and gene expression overall, in OS. The overall process of gene expression is very complex, with regulation at many steps, and assays such as western blotting or qPCR are limited in their ability to interrogate this process fully. While an increase in expression at the mRNA level is often referenced as impactful to a cell, there are numerous cases in which transcriptional and translational data diverge (17–20). In order to study the impact of CBF β on protein translation, and gene expression overall, we performed two somewhat orthogonal assays; Ribo-Seq (21–23) and DiDBiT (24). The quantity of a given protein is a convergence of two antagonistic processes: protein synthesis, and protein degradation. In this study, we are focused on the former, with Ribo-Seq revealing the translation efficiency of a given gene, and DiDBiT revealing the overall synthesis rate of a given protein.

Materials and Methods

Cell Lines and Culture Conditions

U2OS-derived cell lines were generated as previously described, and maintained in McCoy's 5A media supplemented with 10% FBS and 1% P/S.

FLAG Immunoprecipitation Mass Spectrometry

Sample Collection: Empty Vector, CBF β _WT_FLAG, and CBF β ebeta_3xMut_FLAG cells were grown onto 15 cm plates until reaching 70% confluence. Media was then removed, cells were washed with 15 mL PBS, which was then removed. 3 mL of Trypsin was then added and cells were incubated at 37°C until fully dissociated from plate. 7 mL of media was added to plate and suspended cells were transferred to a 15 mL conical tube. An additional 5 mL media wash was added to the plate to capture remaining cells, which were then put into the same 15 mL conical tube. Conical tubes were centrifuged 500g 5 min. and media was removed. Cell pellet was resuspended in 4 mL PBS and transferred to a 5 mL conical tube, and was then spun down 500g for 5 min. Following this, PBS wash was removed and cells were resuspended into 1 mL PBS, then transferred to a 1.5 mL conical tube. Cells were again spun down 500g for 5 min, PBS was removed, and 350 μ L of NET Lysis Buffer (50 mM Tris, 250 mM NaCl, 5mM EDTA, and 1% NP-40, in water, supplemented with 1x cOmplete protease inhibitor tablet and 1mM PMSF) was added per sample. Cells were lysed according to standard cell lysate collection procedure outlined in western blotting protocol in Chapter 2 and kept on ice moving forward. Protein concentration of each sample was then quantified via BCA, and samples were normalized to each other by addition of NET lysis buffer to yield samples all containing 350 μ g of protein. Isotype control

sample was created by combining equal protein amounts of CBF β _WT_FLAG and CBF β _3xMut_FLAG lysate together. All told, 20 samples were created, each containing 350 μ g of protein at 1.2 mg/mL.

Immunoprecipitation: Pulldown antibody (Sigma Aldrich M2 FLAG or SCBT α -GAPDH SC-47724) was added at 1:100 antibody μ g:lysate μ g and tubes were placed on a rotator overnight at 4°C. The next day, a 1,700 μ L slurry of magnetic protein G MagBeads (Genscript, Piscataway, NJ) was pipetted into a 1.5 mL conical tube, placed on a magnetic tube rack, and shipping diluent was removed. Beads were then resuspended in 1 mL NET lysis buffer, as prepared previously, followed by placement on magnetic rack and removal of supernatant. This wash was repeated 3 more times. Following this, the beads were resuspended in 1.5 mL of NET lysis buffer, and 65 μ L of washed bead slurry was added to each tube of lysate. The lysate + antibody + magnetic bead mixture was incubated on a rotator at room temperature for 1h. Following this, the tubes were placed on a magnetic separation rack and supernatant was removed, and beads were washed 3x as described previously. Supernatant was removed, and immunoprecipitated proteins were eluted by addition of 300 μ L of 500 μ g/mL FLAG peptide (MedChem Express, Monmouth Junction, NJ) in NET lysis buffer, and vortex mixed at 2,000 rpm for 10 min. Supernatant was removed and placed in a clean tube, and an additional elution was done with another 300 μ L of FLAG peptide in lysis buffer. Supernatant from second elution was combined with the elution from the first elution.

Protein Clean-Up: Samples were transferred to a clean 2 mL conical tube, and 600 μ L of MeOH (Thermo Fisher), 200 μ L of chloroform (Thermo Fisher), and 600 μ L MilliQ H₂O were added. Samples were vortexed for 5 minutes, then spun down 15,000g for 5 min. Aqueous and organic layers were removed with a gel loading pipette tip, leaving the interface in-between corresponding to protein. 1,200 μ L of MeOH was added to each tube, and vortex mixing and

centrifugation was repeated. Following this, supernatant was removed and pellet was allowed to air dry for 10 min.

Reduction, Alkylation, Digestion: Pellets were then resuspended in 200 μ L of 4M Urea and 50mM Ammonium Bicarbonate (Sigma Aldrich). 2.02 μ L of 500 mM TCEP (Sigma Aldrich) was added to each sample and samples were incubated at 55C with 2,000 rpm shaking for 30 min. 2.02 μ L of 1mM Iodoacetamide was added to each tube, and samples were vortex mixed 1,000 rpm for 20 min, at room temperature and sheltered from light. Following this, 275 μ L of 50mM Ambic, 1.25 μ L of ProteaseMAX, and 16 μ L of Trypsin (Promega Corporation Madison, WI) (200 μ g/mL in H₂O) were added to each tube. Tubes were incubated at 37°C with 1,600 rpm shaking for 4h. Following incubation, reaction was quenched by addition of 0.1% trifluoroacetic acid (TFA) (Sigma Aldrich) final concentration, and tubes were stored -80C until clean-up.

LC-MS/MS Sample Clean-Up: Peptides were incubated at room temperature until thawed. Samples were cleaned up using Sep-Pak C18 cartridges (Waters), which were primed in a series of wash steps. Using a vacuum manifold running at approx. 3 psi vacuum, cartridges were washed with 3 mL acetonitrile (Thermo Fisher), 3 mL of 0.5% acetic acid in water, 3 mL of 50% acetonitrile in water, and 3 mL of 0.1% TFA in water. Thawed peptide mixtures were vortex mixed for 2 minutes, then loaded into cartridges and pulled through sorbent at ~2 psi vacuum. Cartridges were then washed with 3 mL 0.1% TFA in water then 250 μ L of 0.5% acetic acid in water. Peptides were then eluted into a clean tube using 1 mL of 0.5% acetic acid, 80% acetonitrile in water, and evaporated to dryness in a Speed Vac.

LC-MS Parameters: For each sample, half of the total volume was loaded onto a disposable Evotip C18 trap column (Evosep Biosystems, Denmark) as per the manufacturer's instructions. Briefly, Evotips were wetted with 2-propanol, equilibrated with 0.1% formic acid,

and then loaded using centrifugal force at 1200g. Evotips were subsequently washed with 0.1% formic acid, and then 200 μ L of 0.1% formic acid was added to each tip to prevent drying. The tipped samples were subjected to nanoLC on a Evosep One instrument (Evosep Biosystems). Tips were eluted directly onto a PepSep analytical column, dimensions: 150 μ m \times 25cm C18 column (PepSep, Denmark) with 1.5 μ m particle size (100 Å pores) (Bruker Daltronics), and a ZDV spray emitter (Bruker Daltronics). Mobile phases A and B were water with 0.1% formic acid (v/v) and 80/20/0.1% ACN/water/formic acid (v/v/vol), respectively. The standard pre-set method of 60 samples-per-day was used, which is a 21 minute run.

Mass Spectrometry: Performed on a hybrid trapped ion mobility spectrometry-quadrupole time of flight mass spectrometer (timsTOF HT, (Bruker Daltonics, Bremen, Germany) with a modified nano-electrospray ion source (CaptiveSpray, Bruker Daltonics). In the experiments described here, the mass spectrometer was operated in Parallel Accumulation–Serial Fragmentation (PASEF) mode. Desolvated ions entered the vacuum region through the glass capillary and deflected into the TIMS tunnel which is electrically separated into two parts (dual TIMS). Here, the first region is operated as an ion accumulation trap that primarily stores all ions entering the mass spectrometer, while the second part performs trapped ion mobility analysis. The dual TIMS analyzer was operated at a fixed duty cycle close to 100% using equal accumulation and ramp times of 85 ms each.

Data-independent analysis (DIA) scheme consisted of one MS scan followed by MSMS scans taken with 36 precursor windows at width of 25Th per 1.09 sec cycle, over the mass range 300-1200 Dalton. The TIMS scans layer the doubly and triply charged peptides over a ion mobility $-1/k_0$ - range of 0.7-1.3 V^*sec/cm^2 . The collision energy was ramped linearly as a function of the mobility from 59 eV at $1/K_0=1.4$ to 20 eV at $1/K_0=0.6$.

Data Processing: Data was processed with Spectronaut 18.6 (Biognosys, Schlieren, Switzerland) using the directDIA workflow with the default settings. Briefly, trypsin/P Specific was set for the enzyme allowing two missed cleavages. Fixed modifications were set for Carbamidomethyl, and variable modification were set to Acetyl (Protein N-term) and Oxidation. For DIA search identification, PSM and Protein Group FDR was set at 0.01%. A minimum of 1 peptides per protein group were required for quantification. Proteins with an average log₂ratio >0.58 and in CBF β WT cells compared to EV, or CBF β WT cells compared to isotype control, with p and q values <0.05, were binned as medium confidence interactors of CBF β _WT_FLAG. Proteins possessing average log₂ratio >0.58 CBF β WT cells compared to both EV and isotype control, with p and q values <0.05, were binned as high confidence interactors of CBF β _WT_FLAG. High confidence interactors of CBF β _WT_FLAG possessing an average log₂ratio >0.58 in CBF β WT cells compared to CBF β _3xMut_FLAG cells were binned as high confidence interactors of CBF β reliant upon G61, N63, and/or N104 residues of CBF β for interaction. CBF β _FLAG_3xMut pulldown replicate 4 was eliminated from analysis as some sample was lost in processing.

Direct Detection of Biotinylated Tags (DiDBiT)

Sample Collection: DiDBiT was performed as previously described (24). Briefly, U2OS-derived cell lines were plated onto 15 cm dishes and grown to reach 70% confluence. Following this, media was removed, and cells were washed 2x with PBS. Cells were then starved of Methionine by 45 minute incubation in DMEM, high glucose, no glutamine, no methionine, no cystine (Gibco, Thermo Fisher). DMEM was then supplemented with 10% dialyzed FBS, 1% P/S, 1mM Sodium Pyruvate, 4mM L-Glutamine, and 200 μ M L-Cystine HCl (Thermo Fisher).

Following starvation, media was removed and cells were then incubated for 4h with aforementioned media supplemented with 1mM L-azidohomoalanine (Click Chemistry Tools, Vector Laboratories, Newark, CA). Click conjugation and sample clean-up were accomplished as previously described by Schiapparelli et al (24).

LC-MS/MS Analysis: Sample reconstitution, injection, and LC-MS method was the same as mentioned previously.

Data Processing: Surrogate peptide areas for peptides possessing the L-azidohomoalanine (AHA) (+523.2749) modification were summed from each protein and then summed together from two separate runs. The area for each peptide in CBF β _WT_FLAG cells was then divided by the areas in EV cells, and any protein with >20% higher signal between WT/EV was linked as a possible protein under the purview of CBF β . Proteins identified by at least 1 peptide possessing AHA modification, and quantifiable in both biological replicates, were included in analysis. Proteins which exhibited a 20% or greater increase in abundance of AHA modified peptides in CBF β _WT_FLAG cells compared to EV cells were binned as potential translational targets of CBF β . From this pool, proteins which also exhibited a 20% or greater decrease in abundance of AHA modified peptides in CBF β _3xMut_FLAG cells compared to CBF β _WT_FLAG were binned as potential targets of CBF β reliant upon residues G61, N63, and/or N104 of CBF β for their translation.

Ribosome Footprinting (Ribo-Seq)

Sample Collection: Ribosome footprinting was performed by TB-Seq, Inc. (South San Francisco, California) as previously described (21,25,26). Cells were grown on 15 cm plates (Corning) until reaching 70-80% confluence and then collected as follows. Six plates were used per cell line, per biological replicate. Media was removed and cells were washed with 5 mL of ice cold 1x Nuclease-Free PBS (Thermo Fisher) supplemented with 100 µg/mL Cycloheximide (Sigma Aldrich). Dishes were placed on ice in an RNase-free environment and wash was removed, then 3 more mL of Nuclease-Free PBS was added. Cells were collected via cell scraper and transferred into a 15 mL conical tube. An additional 2 mL of Nuclease Free PBS was added to plate and residual remaining cells were washed down, and added to the same 15 mL Falcon tube. Tubes were then spun down at 300g for 5 min. at 4°C, supernatant was removed, and pellet was flash frozen in liquid nitrogen before transferring to -80C storage prior to analysis. Pellets were stored on dry ice during transit to TB-Seq, who performed the Ribo-Seq analysis.

Ribosome Profiling: Cells were resuspended in ice cold lysis buffer (20 mM Tris HCl, pH 7.4, 100 mM NaCl, 5 mM MgCl₂ 1% Triton X100, 1 mM DTT, 20U/ml Turbo DNase I, 0.1 % NP40, 100 µg/ml cycloheximide) and the soluble cytoplasmic fraction was isolated by centrifugation at top speed in a microcentrifuge for 20 min at 4°C (21,23). Supernatants were collected and clarified lysates were digested with RNase I for 45 min at room temp. Digestion was stopped with SuperaseIN and monosomes purified by size exclusion chromatography on MicroSpin S-400 HR columns (GE Healthcare) as described (22). Size selection of footprints with length 25-33 nt was performed by electrophoresis on 15% TBE-urea gels. Illumina ready RIBO-seq libraries were prepared using a SMARTer smRNA-seq kit (TakaraBio, Kusatsu, Shiga, Japan). Library concentrations were measured by qubit fluorometer and their quality assessed on an

Agilent 2100 bioanalyzer. RIBO-seq libraries were sequenced on an Illumina Novaseq 6000 sequencer, single read, 1x50 cycles.

RNAseq: To obtain matched RNA-seq libraries total RNA was purified from an aliquot of cell lysate and rRNA was depleted from total RNA using a NEBNext rRNA depletion kit v2 (NEB) following manufacturer instructions. mRNA fragmentation was conducted for 20 min at 94 °C to generate RNA fragments of sizes similar to those of the ribosome footprints. SMARTer smRNA-seq kit (TakaraBio) was used to generate Illumina ready RNA-seq libraries. Library concentrations were measured by qubit fluorometer and their quality assessed on an Agilent 2100 bioanalyzer. RNA-seq libraries were sequenced on an Illumina Novaseq 6000 sequencer, single read, 1x50 cycles.

Data Analysis: Gene expression results for 19,811 protein-coding genes annotated in Ensembl 104 for the NCBI human hg38.p13 genome assembly are reported based on RIBO-seq and RNA-seq coverages. Based on the observed distribution of ribosome footprint lengths (see Section 4.3), we selected all RIBO-seq reads of lengths in the range 31-39 nucleotides as representing footprints of actively translating ribosomes. Shorter reads are thought to derive from ribosomes in an inactive state (ribosome in inactive “rotated conformation”, or ribosomes not charged with tRNA at the A_site) (27) whereas longer reads may represent stacked disomes in different states. In the case of RNA-seq reads, the vast majority of which have the maximum possible length 47 nt, reads of all lengths were included. Statistical analyses of differential gene transcription (RNA) and translation (RIBO) were performed with the quasi-likelihood F-test procedure implemented in the EdgeR package (28). EdgeR reports for each gene log₂-fold-change in expression between conditions, the p-values obtained testing each gene for differential coverage between conditions, and the False Discovery Rate (FDR) associated with each p-value threshold,

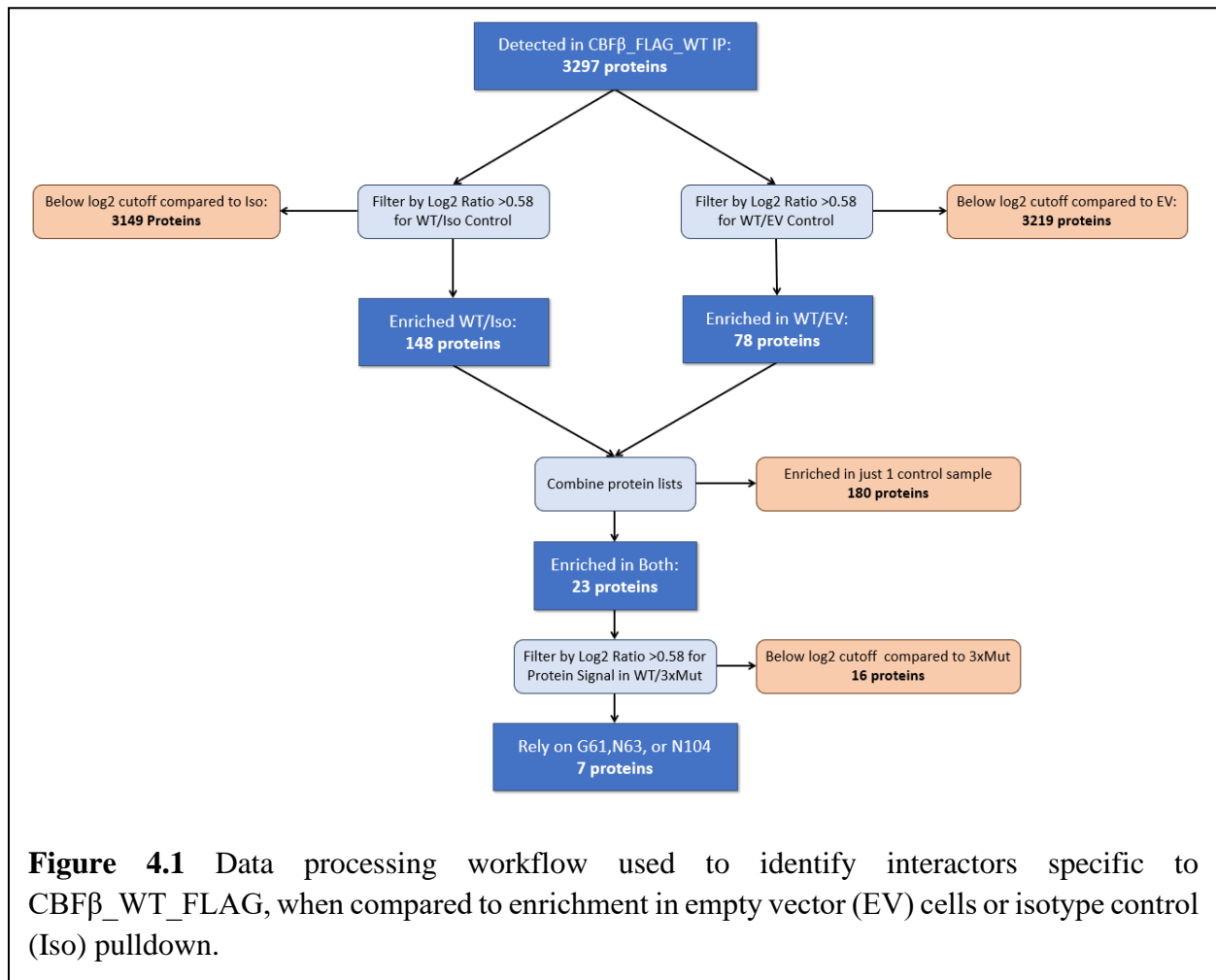
evaluated with the Benjamini Hochberg procedure. Changes in Translation Efficiency, i.e., changes in the ratio of levels of translation over levels of transcription, were evaluated with models and statistical analyses implemented in the DTEG procedure (29) (DTEG-TE) and in the anota2seq procedure (A2S) (30). Genes were selected for differential transcription (RNA-seq), translation (RIBO-seq), Translation Efficiency, or buffering.

Genes were selected by $p\text{-value} \leq 0.01$ in respective tests, as estimated by EdgeR (RNA and RIBO), DTEG (TE-DTEG) and anota2seq (TE-A2S and Buffering) differential-expression models. RNA is number of differentially transcribed genes based on RNA-seq coverage; RIBO is number of differentially translated genes based on RIBO-seq coverages; TE is number of genes with significant changes in Translation Efficiency. Buffering is number of genes with changes in transcription buffered by translational regulation.

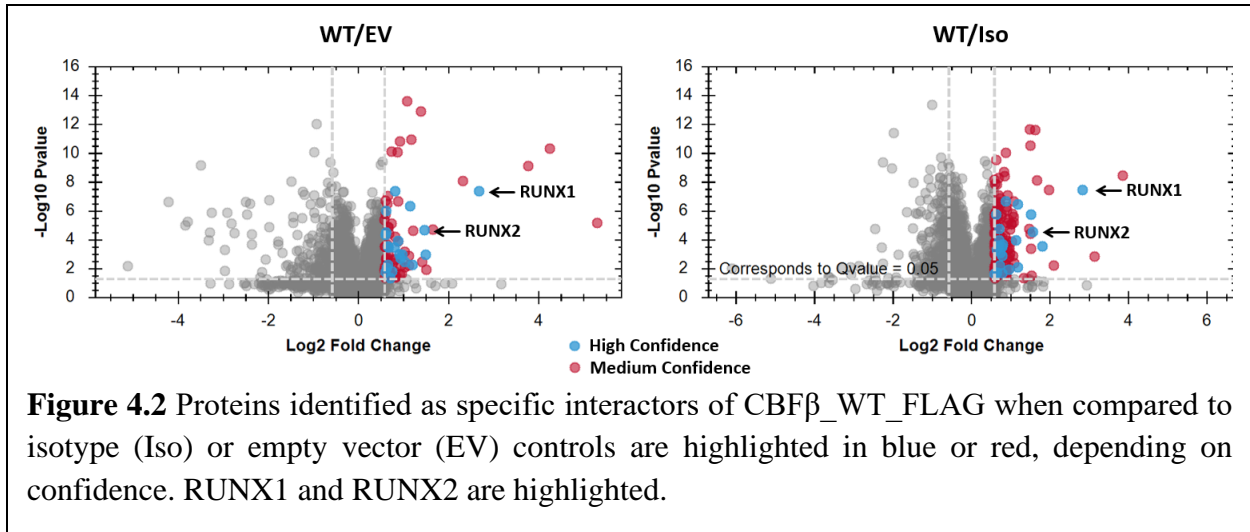
Gene Ontology and Pathway Analysis of Identified Proteins/Genes

Gene ontology, Reactome, and KEGG pathway analysis was performed using DAVID bioinformatics resource (31,32). Venn diagram figures were generated using DeepVenn (33). Gene ontology and pathway analysis figures were generated using SRPlot (34).

Results

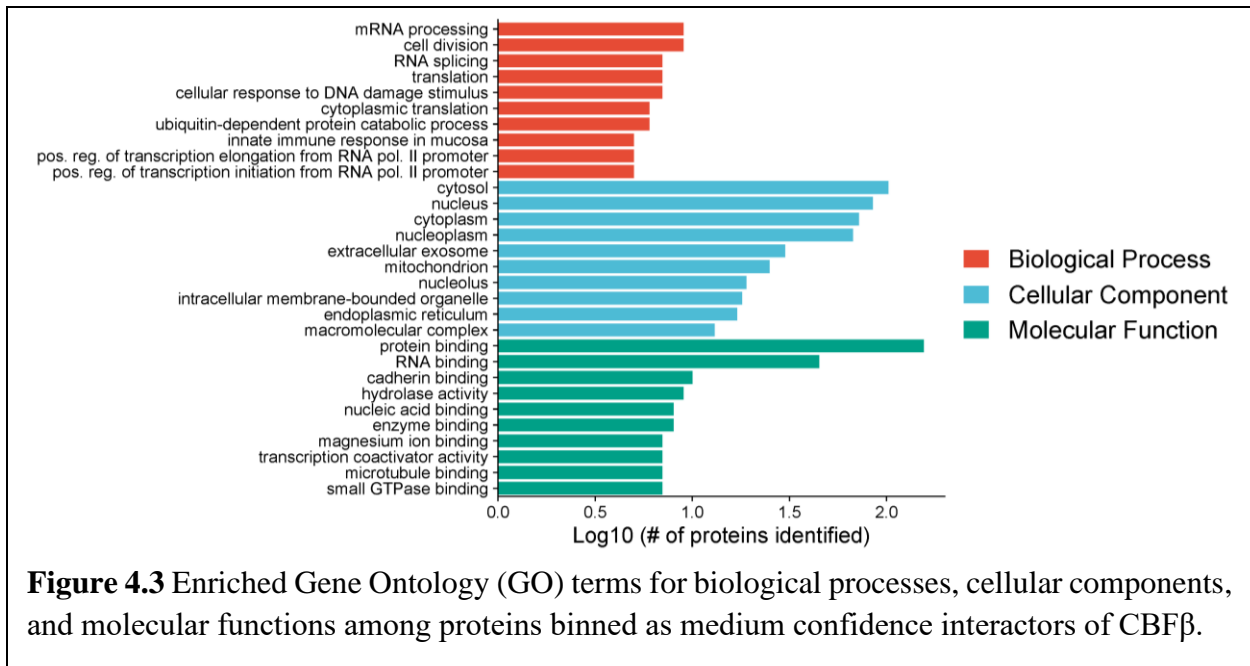


Proteins detected in IP-MS pulldowns were selectively filtered to identify specific interactors of CBFβ. Out of a total 3,297 proteins identified in pulldown of CBFβ_WT_FLAG, twenty-three (23) were enriched compared to both isotype and empty vector controls and were classified as high confidence interactors of CBFβ. 180 proteins were enriched compared to just one control, isotype or empty vector, and were binned as medium confidence interactors of CBFβ. Finally, of these twenty-three, seven were found to be significantly enriched in CBFβ_WT_FLAG compared to CBFβ_3xMut_FLAG.

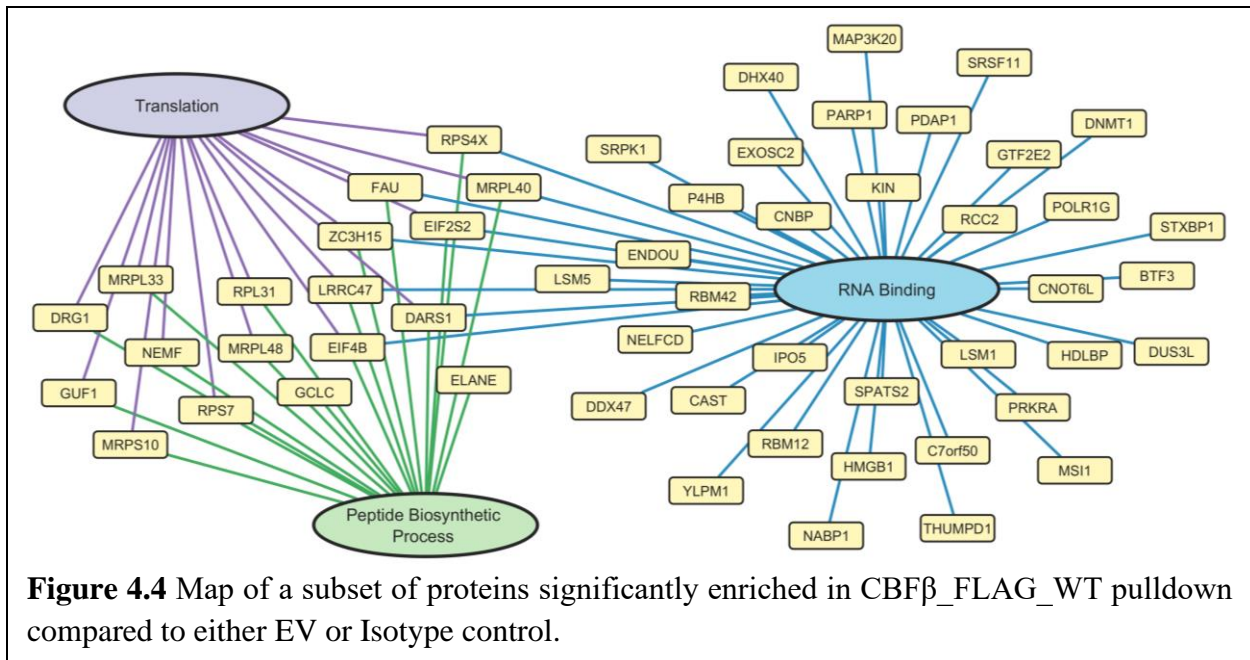


Volcano plots of proteins identified as medium or high confidence interactors of CBF β .

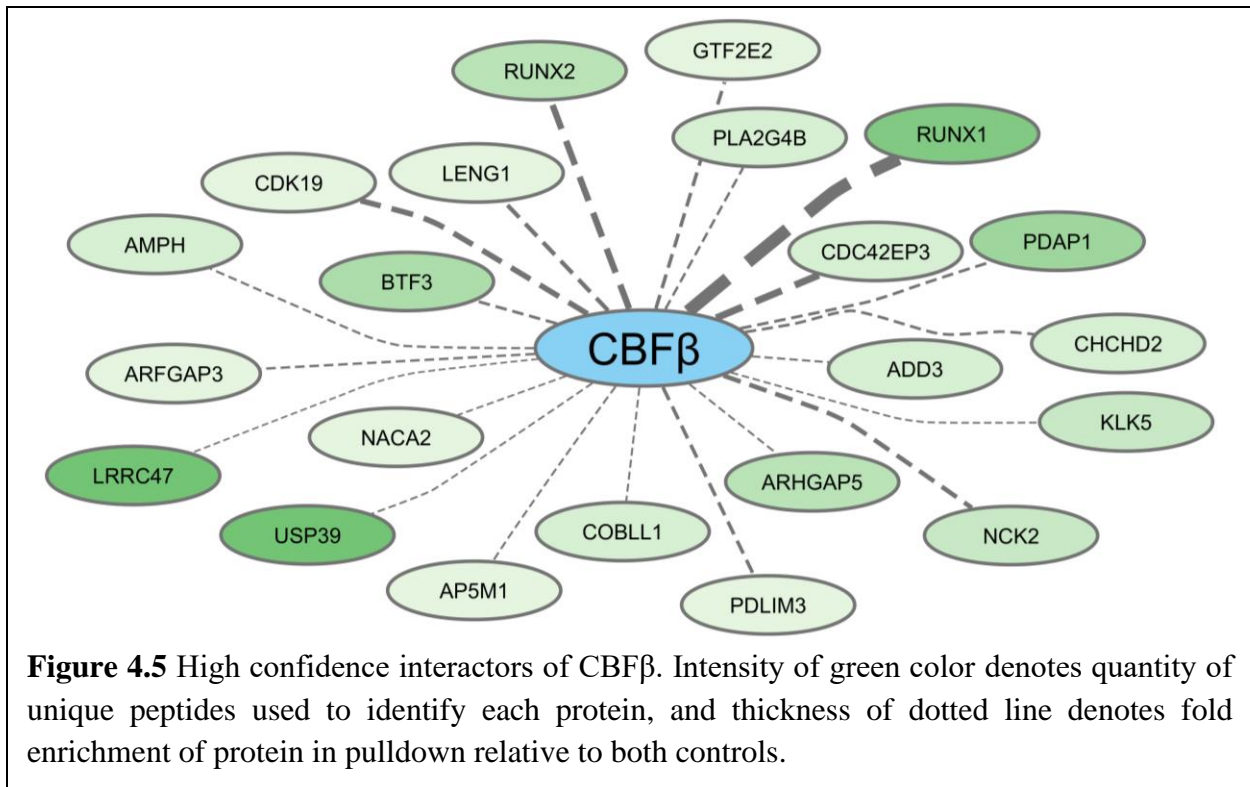
Both RUNX1 and RUNX2 showed up as high confidence interactors of CBF β . Each protein identified as a blue circle above was binned as a high confidence interactor of CBF β , while red circles were medium confidence interactors.



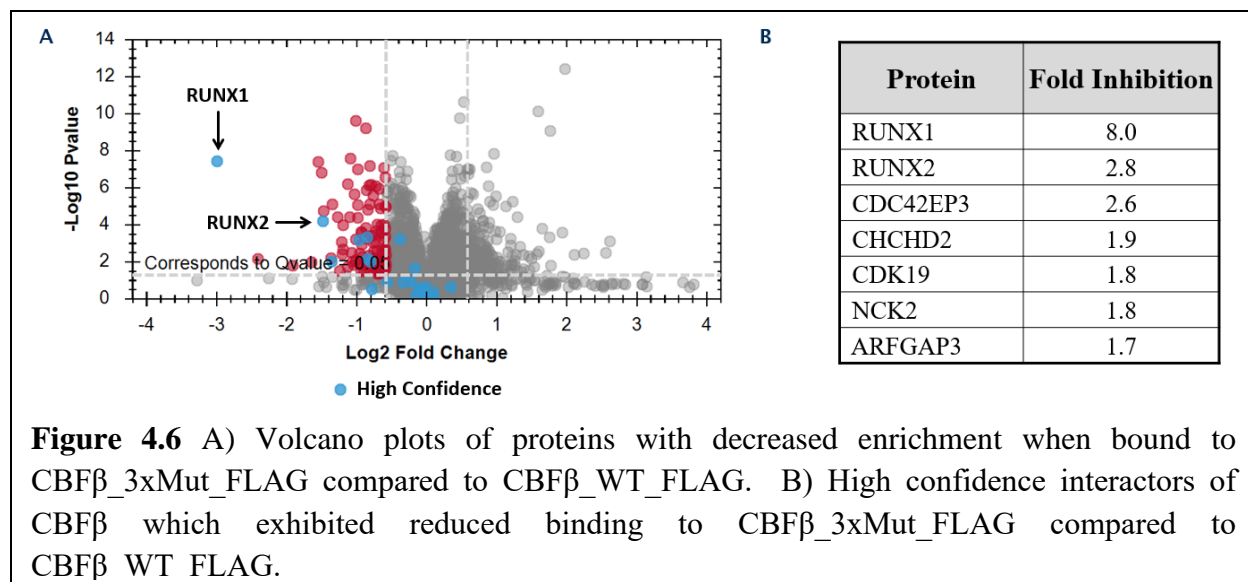
Enriched GO terms among medium confidence interactors of CBF β . The biological process terms translation and cytoplasmic translation came up highly ranked among proteins pulled down. Also highly ranked was mRNA processing and RNA splicing. A variety of cellular components were represented in the pulldown as well as numerous molecular functions.



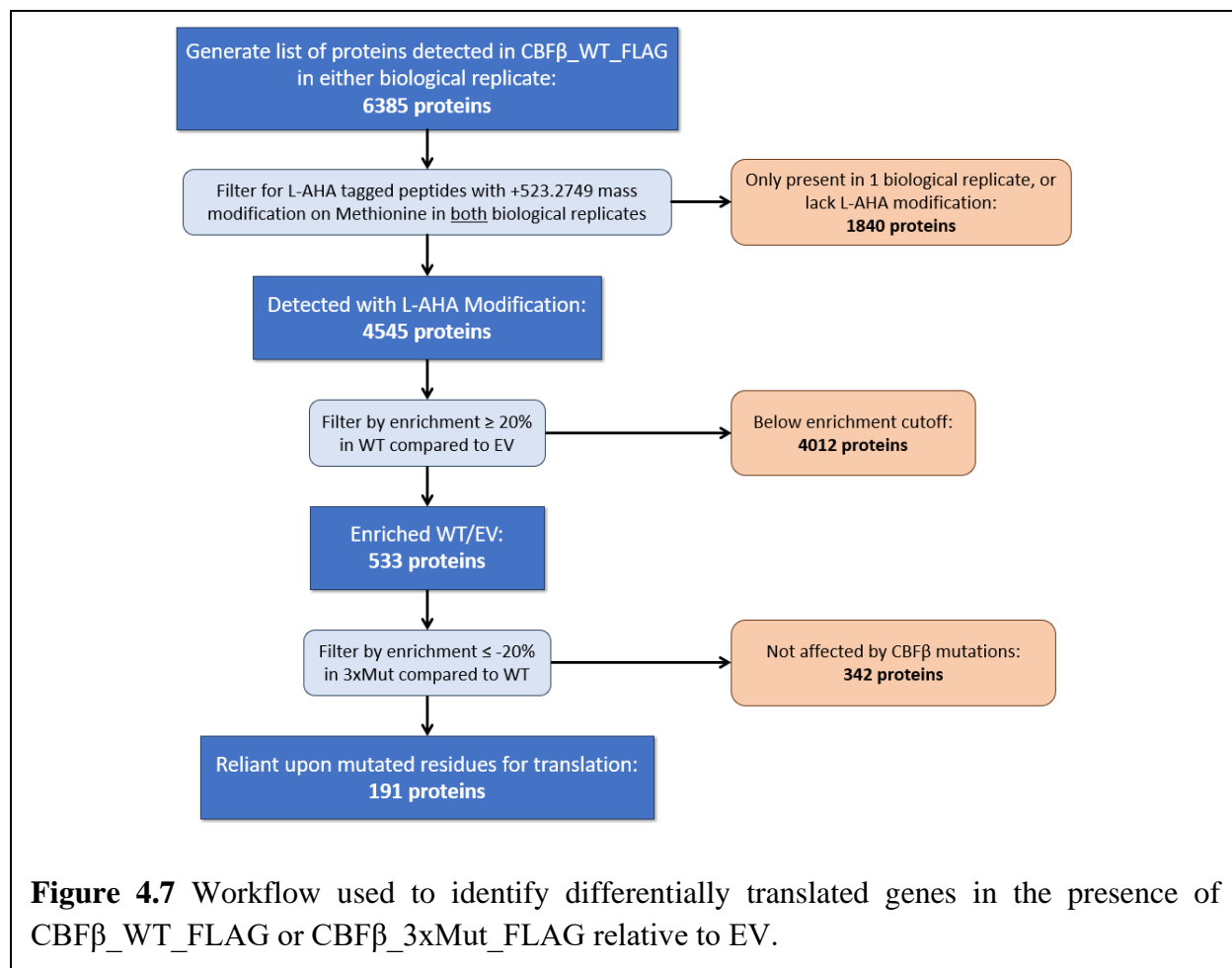
Network demonstrating specific proteins linked to GO terms surrounding protein translation. Each protein listed in graphic above was associated with at least one of the following GO terms: translation, peptide biosynthetic process, or RNA binding. Some proteins were associated with multiple terms. Of the total 224 total proteins identified, 54 were shown above.



Interaction network outlining high confidence interactors of CBF β . Both RUNX1 and RUNX2 showed up as high confidence interactors of CBF β , along with numerous others. RUNX1 experienced the highest fold enrichment in pulldown of CBF β _FLAG_WT compared to control, denoted by the thickest dotted line connecting it to CBF β .



High confidence interactors of CBF β with reduced binding to CBF β _3xMut_FLAG. In total, seven (7) proteins of the twenty-three (23) high confidence interactors identified displayed reduced enrichment when pulled down by CBF β _3xMut_FLAG. Of these, RUNX1 had the most drastic reduction in binding, with an eight-fold reduction in enrichment by CBF β _3xMut_FLAG.



Proteins detected in DiDBiT pulldowns were selectively filtered to identify differentially translated proteins. Out of a total 6,385 proteins identified, 533 were detected in both biological replicates with enrichment $\geq 20\%$ of L-AHA modified peptides in CBFβ_WT_FLAG cells compared to EV. Of these, 191 proteins exhibited a $\geq 20\%$ decrease in enrichment when expressed in CBFβ_3xMut_FLAG producing cells.

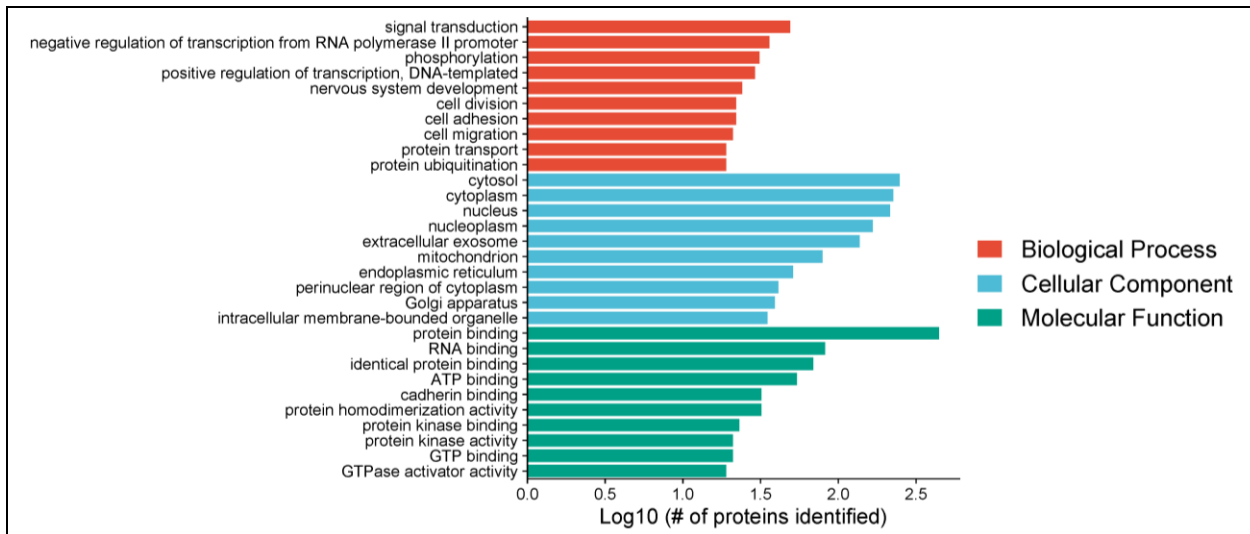
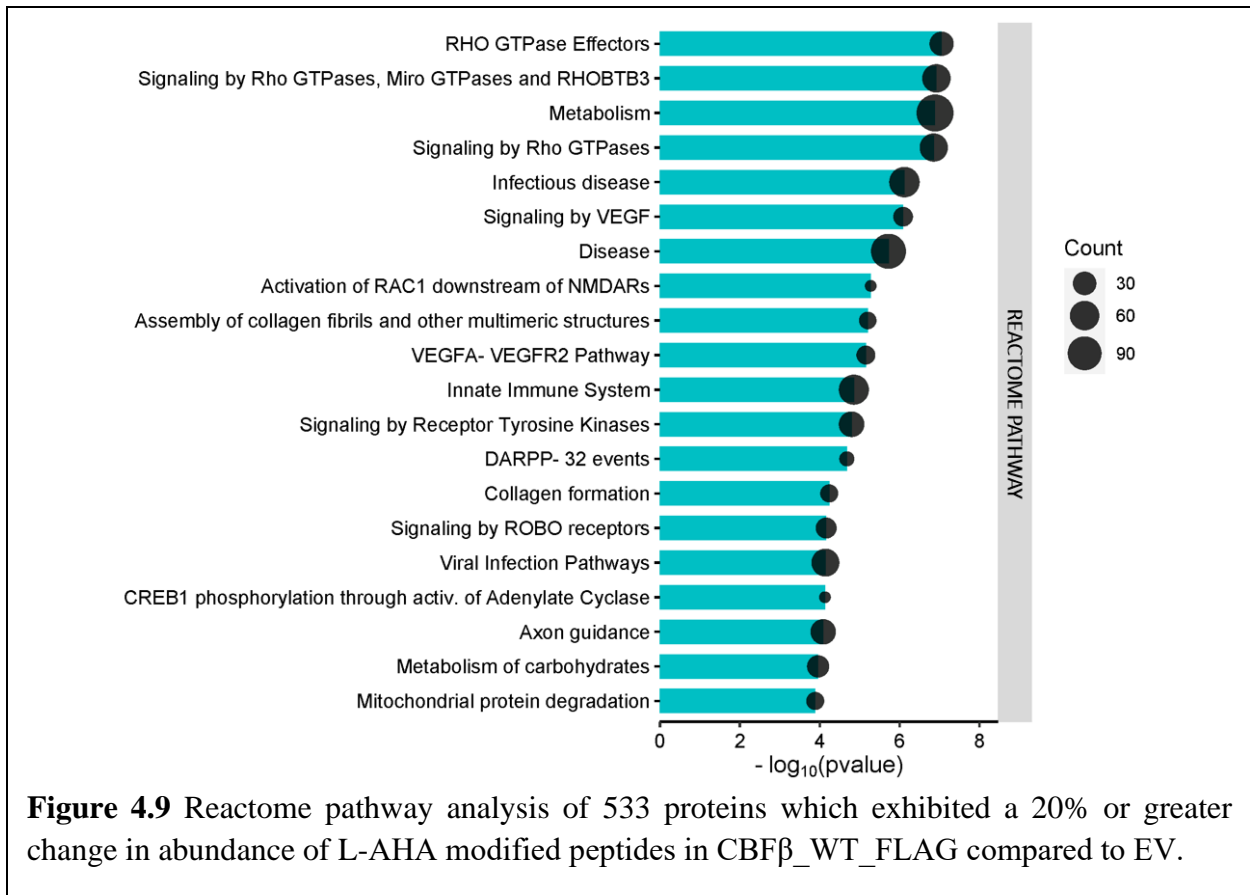
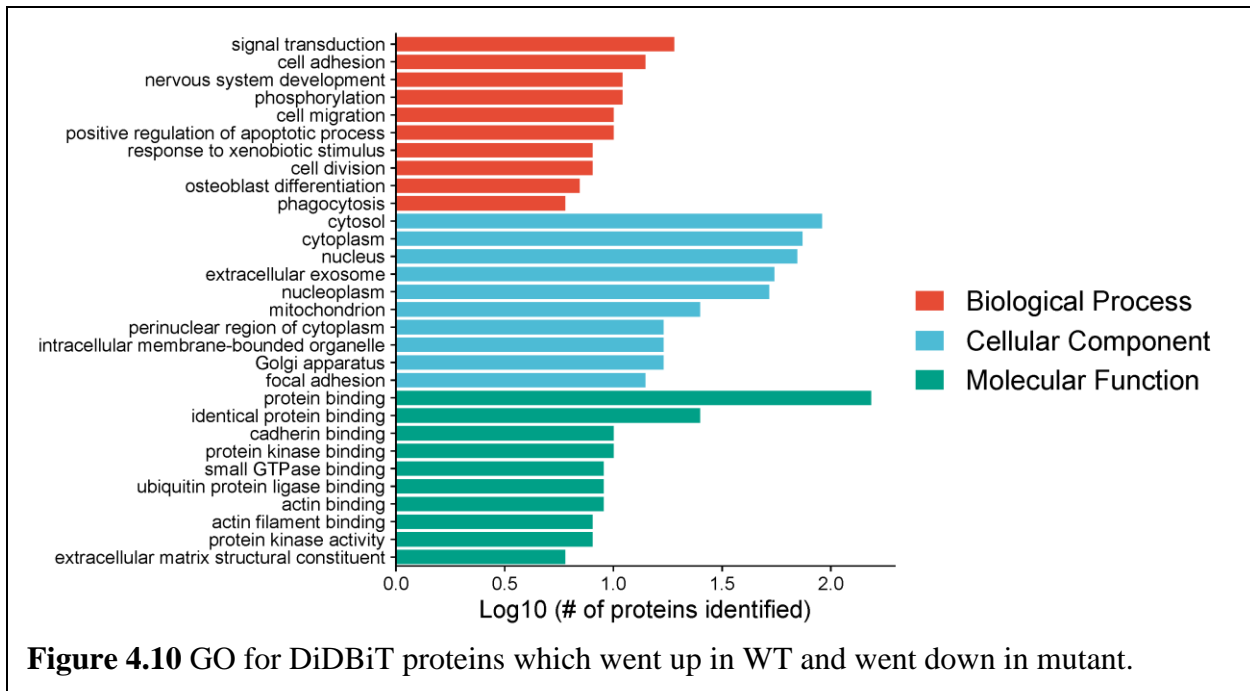


Figure 4.8 Enriched GO terms for biological processes, cellular components, and molecular functions among proteins which exhibited a 20% or greater change in abundance of L-AHA modified peptides in CBF β _WT_FLAG compared to EV.

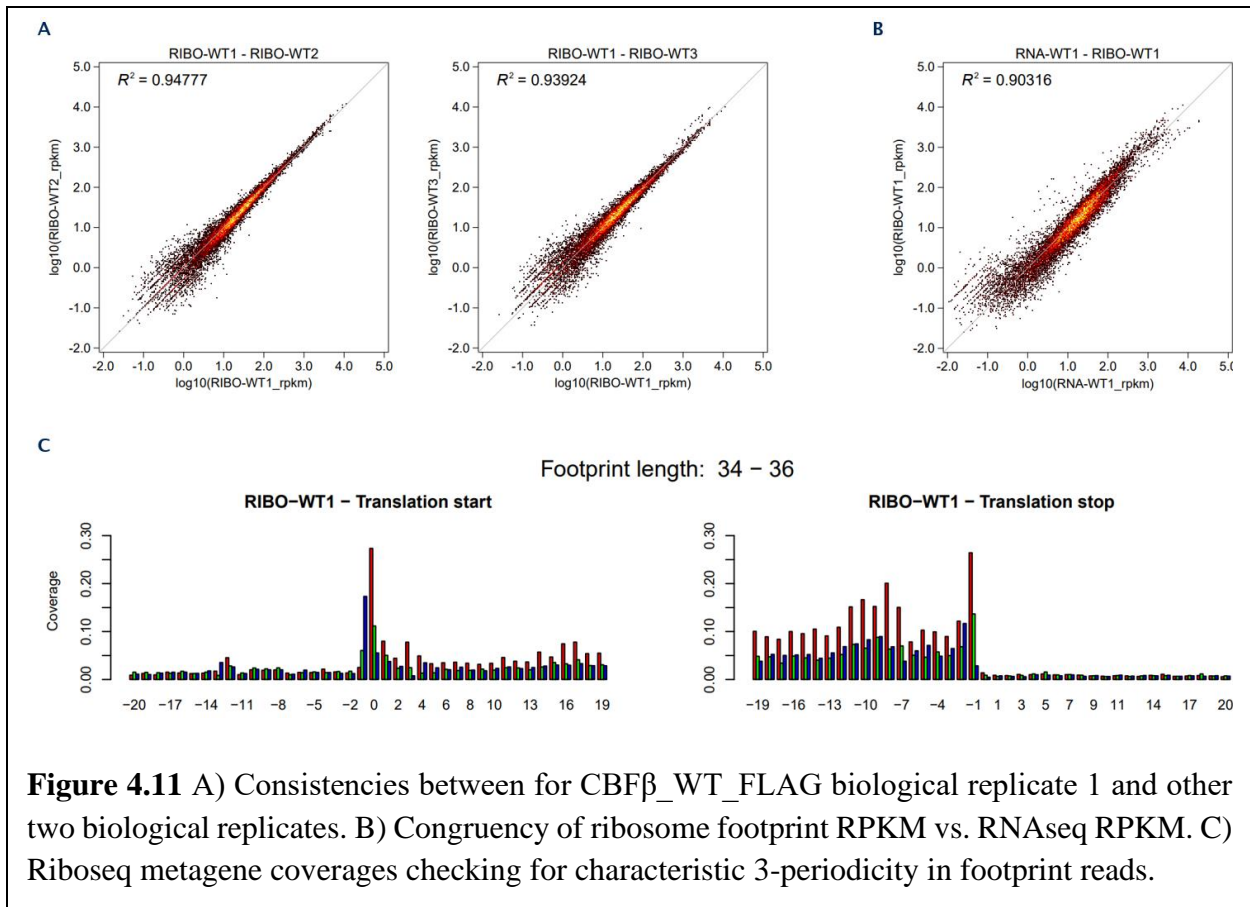
Enriched GO terms among proteins enriched in CBF β _WT_FLAG, as identified by DiDBiT. Biological process terms associated with transcription were highly ranked as well as those associated with cell division, adhesion, and migration. A variety of cellular components were represented as well as numerous molecular functions, including protein, RNA, ATP, and GTP binding.



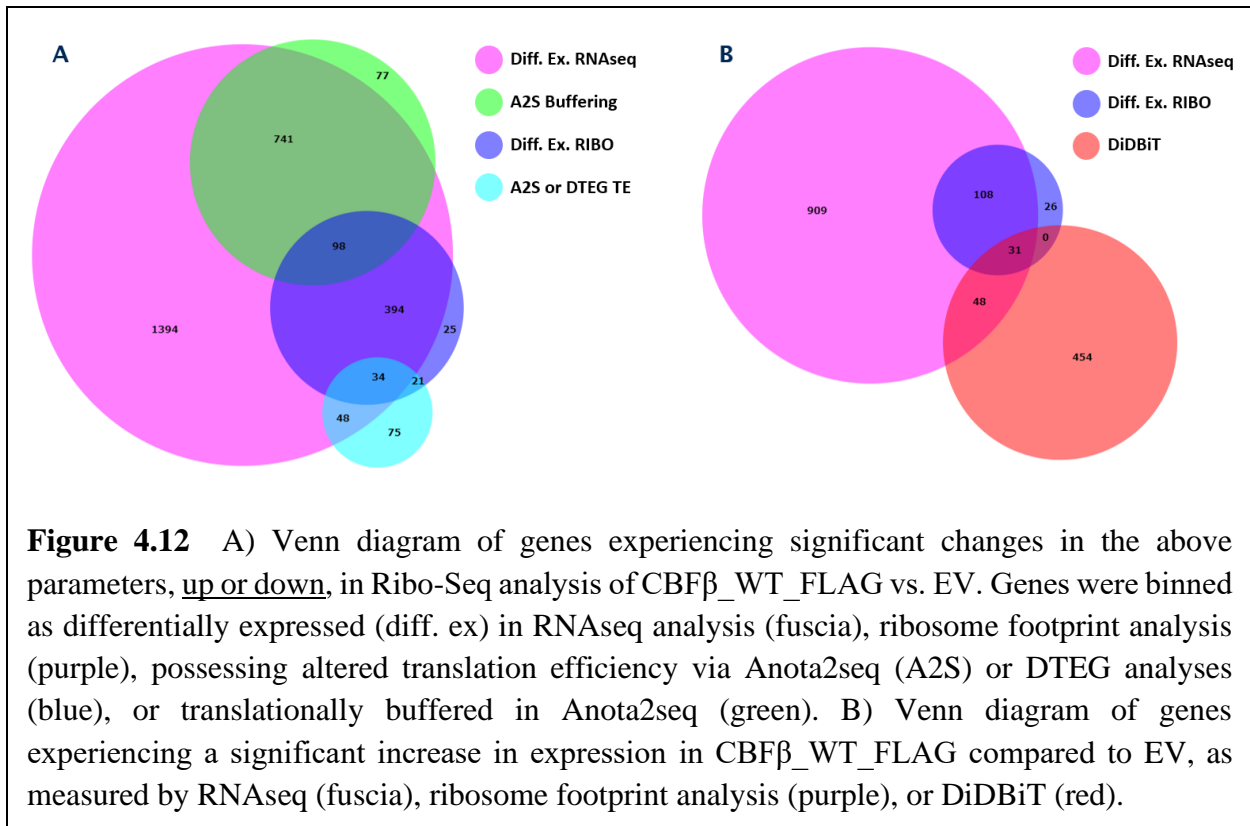
Enriched Reactome terms among proteins enriched in CBF β _WT_FLAG, as identified by DiDBiT. Rho GTPase-associated Reactome terms came up highly ranked among proteins identified in DiDBiT as well as those associated with VEGF.



Enriched GO terms among proteins enriched in CBF β _WT_FLAG and decreased in CBF β _3xMut_FLAG, as identified by DiDBiT. The biological process terms signal transduction, cell adhesion, and cell migration/division were highly ranked, as well as numerous cellular components. Binding to a variety of targets such as cadherin, protein kinases, and small GTPases were highly ranked in molecular function.



Quality control data for Ribo-Seq analysis. Representative images for the first biological replicate for CBF β _WT_FLAG are displayed above. Other biological replicates as well as those for EV cells were similar, with R squared values > 0.90 in all. Characteristic 3-periodicity of Ribo-Seq reads was present in WT1 as well as all other replicates.



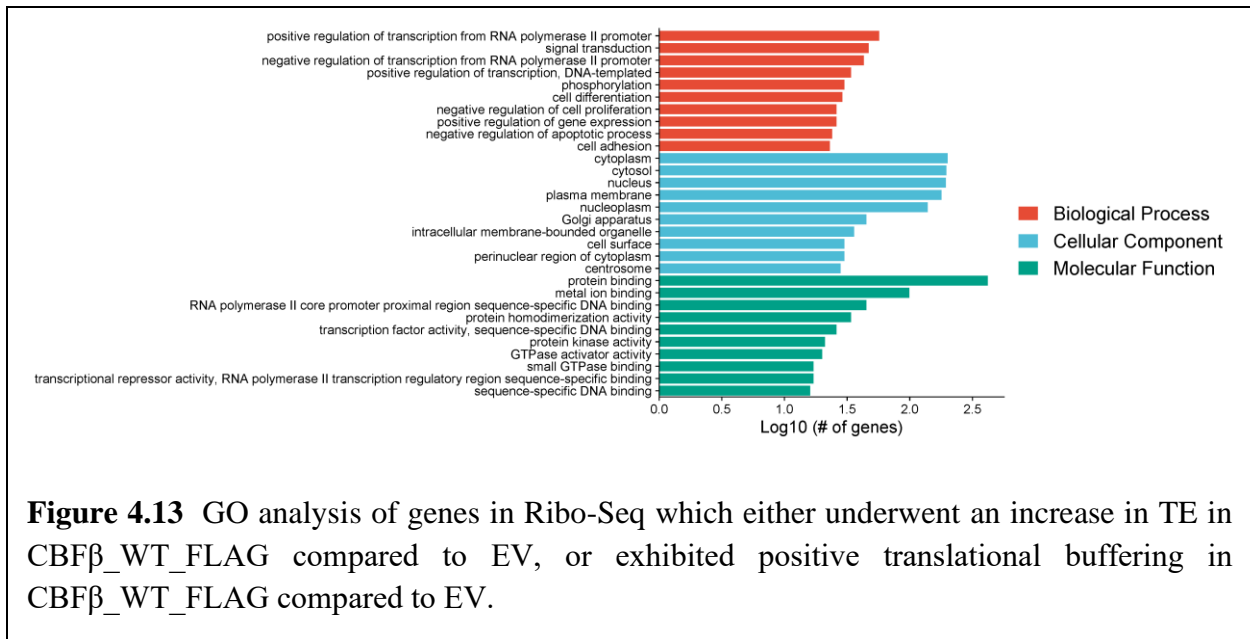
Grouping of genes identified in Ribo-Seq and comparison with those identified in DiDBiT.

A total of 2,715 genes were identified as differentially expressed at the RNA level between CBF β _WT_FLAG and EV cells. Of these, many were binned as buffered via A2S, differentially expressed in ribosome footprinting abundance, and altered in translational efficiency. Overlaps between the aforementioned groups are illustrated above. 79 genes identified as differentially expressed via DiDBiT were corroborated by RNAseq data, with ribosome footprints of 31 of these also significantly enriched in CBF β _WT_FLAG cells compared to EV.

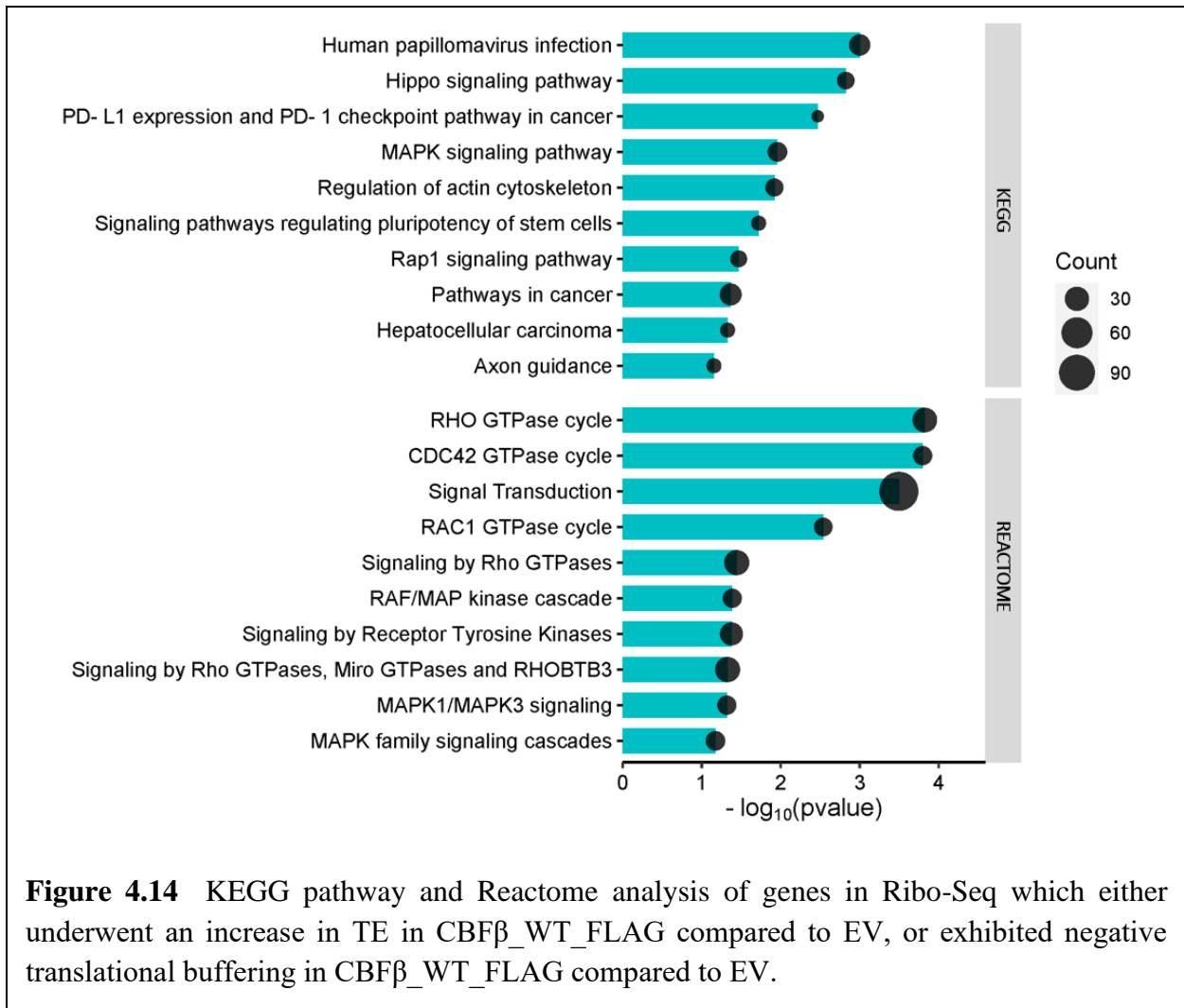
Table 4.1: Genes up-regulated in CBF β _WT_FLAG cells vs. EV across DiDBiT, RNAseq, and RIBO-Seq assays

| Gene | Name | HGNC ID |
|----------|---|------------|
| AHNAK | AHNAK nucleoprotein | HGNC:347 |
| ALDH2 | aldehyde dehydrogenase 2 family member | HGNC:404 |
| ARHGAP28 | Rho GTPase activating protein 28 | HGNC:25509 |
| CBFB | core-binding factor subunit beta | HGNC:1539 |
| CD70 | CD70 molecule | HGNC:11937 |
| CDC42EP3 | CDC42 effector protein 3 | HGNC:16943 |
| CDKN1A | cysteine rich DPF motif domain containing 1 | HGNC:33710 |
| COL6A3 | collagen type VI alpha 3 chain | HGNC:2213 |
| CPA4 | carboxypeptidase A4 | HGNC:15740 |
| ERBIN | erbB2 interacting protein | HGNC:15842 |
| EZR | ezrin | HGNC:12691 |
| FKBP9 | FKBP prolyl isomerase 9 | HGNC:3725 |
| GPHN | gephyrin | HGNC:15465 |
| HCFC1R1 | host cell factor C1 regulator 1 | HGNC:21198 |
| KRT86 | keratin 86 | HGNC:6463 |
| LAMB3 | laminin subunit beta 3 | HGNC:6490 |
| MIS18A | MIS18 kinetochore protein A | HGNC:1286 |
| NPC2 | NPC intracellular cholesterol transporter 2 | HGNC:14537 |
| PALLD | palladin, cytoskeletal associated protein | HGNC:17068 |
| PDCD2 | programmed cell death 2 | HGNC:8762 |
| PLXNB2 | plexin B2 | HGNC:9104 |
| PPME1 | protein phosphatase methylesterase 1 | HGNC:30178 |
| RHOBTB3 | Rho related BTB domain containing 3 | HGNC:18757 |
| SBF1 | SET binding factor 1 | HGNC:10542 |
| SERPINA1 | serpin family A member 1 | HGNC:8941 |
| SLC2A3 | solute carrier family 2 member 3 | HGNC:11007 |
| SYTL3 | synaptotagmin like 3 | HGNC:15587 |
| TBC1D10A | TBC1 domain family member 10A | HGNC:23609 |
| TBC1D2 | TBC1 domain family member 2 | HGNC:18026 |
| TBC1D22A | TBC1 domain family member 22A | HGNC:1309 |

Specific genes upregulated in CBF β _WT_FLAG cells vs. EV, as measured by DiDBiT, RNAseq, and RIBO-Seq. The above 31 genes were increased in expression across all three assays. Listed are gene ID, name, and HUGO Gene Nomenclature Committee (HGNC) identification.



GO analysis of genes with increased translation efficiency or positive translational buffering in the presence of CBFβ_WT_FLAG. Among genes identified, GO terms associated with transcription and gene expression overall ranked very highly. Also notable was genes involved in signal transduction as well as proliferation and cell adhesion. As in previous GO analyses, numerous cellular components as well as molecular functions were represented.



Reactome and KEGG pathway analyses of genes with increased translation efficiency or translational buffering in the presence of CBF β _WT_FLAG. Multiple signaling pathways, many implicated in cancer, were represented in Reactome pathway analysis of aforementioned genes. Rho GTPase actions as well as MAPK signaling ranked highly in KEGG pathway analysis.

Discussion

IP-MS of CBF β revealed 3,297 initial proteins detected, which were subjected to successive filtrations to reveal medium and high-confidence CBF β interactors (**Figure 4.1**). Encouragingly, both RUNX1 and RUNX2, known interactors of CBF β (15,35–38), were enriched relative to both controls (**Figure 4.2**). This corroborated our previous data on interaction of our transgene with endogenous targets, increasing our confidence in the legitimacy of novel interacting partners observed.

Among proteins identified as medium confidence interactors, a total of 224 proteins, gene ontology revealed high enrichment of biological process terms mRNA processing, translation, and cytoplasmic translation (**Figure 4.3**). This lined up well with our hypothesis, that CBF β may participate in protein translation. Medium confidence interactors were enriched in the cellular component term “nucleus”, which was expected, however GO term “cytoplasm” was also highly enriched. As the translational role of CBF β is postulated to occur in the cytoplasm, the enrichment of the cytoplasmic term bolsters these claims of a role for CBF β outside of the nucleus. It was interesting that “mRNA processing” came up highly enriched on the GO biological process terms, as well as “RNA binding” on molecular function terms. CBF β was suggested to interact with hnRNPk, a multifunctional protein involved in transcription, RNA splicing, and protein translation (39). While hnRNPk was not identified as an interactor in our screen, numerous other proteins implicated in mRNA processing were identified. Among the medium confidence interactors was eIF4B, a member of the eukaryotic translation initiation complex (40) which plays a key role in protein translation, and was previously reported to interact with CBF β (14). Cul5, an E3 ligase which has previously been reported to interact with CBF β (41), also showed up as a medium confidence interactor, further increasing our confidence in these data. Many of the

medium confidence interactors were associated with GO terms “translation” and “peptide biosynthetic process”, including eIF4B (**Figure 4.4**).

Further filtration for proteins significantly enriched when compared to both EV and Isotype controls yielded a list of twenty-three (23) proteins, which we binned as high confidence interactors of CBF β (**Figure 4.5**). As mentioned previously, RUNX1 and RUNX2 were among this group. RUNX1 experienced a higher enrichment in pulldown than RUNX2, as evidenced by the thicker line connecting it to CBF β . This was also encouraging, as previous studies reported CBF β -RUNX interaction to alter RUNX1 stability more drastically than RUNX2, possibly due to the CBF β -RUNX1 interaction being of a higher affinity than CBF β -RUNX2 (15). In addition to known interacting partners RUNX1 and RUNX2, twenty-one (21) proteins were also identified, none of which are mentioned in The Human Protein Atlas or String-db protein interaction databases. To our knowledge, this represents the first comprehensive study of CBF β interacting proteins in OS.

While the full relevance of these interactions remains to be understood, many of the proteins identified as high confidence interactors of CBF β are implicated in important cellular processes and disease. Among the list was ubiquitin specific peptidase 39 (USP39), a member of the deubiquitinase (DUB) family of enzymes. DUB's play key roles in maintaining protein homeostasis in the cell, and have been implicated in a wide array of diseases (42,43). USP39 is no exception, and has been reported to have an oncogenic role in osteosarcoma (44) as well as numerous other cancers (45–54). In fact, studies of USP39 in osteosarcoma were conducted in U2OS cells, the same parental line as the cells used in this project. This oncogenic function of USP39 is thought to take place through two mechanisms: 1) stabilization of beta-catenin via direct de-ubiquitination and 2) downregulation of TRIM26 by reduction of TRIM26 pre-mRNA

maturation (54). It's also possible that USP39 may keep pro-survival proteins such as Bcl-2, survivin, and Mcl-1 upregulated in these cells, further contributing to its oncogenic behavior. Also identified as a high confidence interactor of CBF β was NCK2, an adapter protein which associates with tyrosine-phosphorylated growth factor receptors. Elevated levels of NCK2 are implicated in melanoma proliferation, migration, and invasion (55) as well as invasion in breast cancer and prostate cancer cell lines (56). CDK19 was also identified, and upregulation of this protein correlates with unfavorable prognosis in hepatocellular carcinoma (HCC) (57).

Interestingly, NACA2 and BTF3, two members of the same complex, also came up on our screen. Together, NACA and BTF3 comprise the nascent polypeptide-associated complex (NAC), which is located at the ribosome exit tunnel (58). This complex functions as a chaperone, assisting in protein folding, and protects the nascent polypeptide sequences from proteolysis as the new protein is translated (59). Loss of this complex is embryonic lethal, and association with NAC is likely the first interaction undergone by the newly synthesized peptide (59). This complex is also implicated in controlling translation initiation (60) as well as oncogenic activity in colorectal cancer (61) and prostate cancer (62). Further study is required to elucidate how interaction with CBF β affects the activity of NAC, as this complex plays an indispensable role in nascent protein synthesis.

Analysis of CBF β _3xMut_FLAG-associated proteins in relation to those identified to interact with CBF β _WT_FLAG revealed seven (7) proteins which appear to be reliant upon residues G61, N63, or N104, or a combination, for their binding to CBF β . Among this list was RUNX1 and RUNX2, in agreement with previously shown CETSA data which suggested CBF β -RUNX2 interaction to be inhibited by these mutations. Also in this list were CDK19 and NCK2, as well as three other proteins: CDC42EP3, CHCHD2, and ARFGAP3.

Following identifying specific interactors of CBF β , we conducted DiDBiT (24) to elucidate which proteins may be increased in synthesis rate upon re-introduction of CBF β _WT_FLAG into CBF β KO cells. A total of 6,385 unique proteins were identified in CBF β _WT_FLAG cells, which were successively filtered for L-AHA methionine modification, presence in both biological replicates, and enrichment $\geq 20\%$ in CBF β _WT_FLAG cells relative to EV (**Figure 4.7**). This revealed a list of 533 proteins which are putative translational targets of CBF β . Further filtration in comparison of this list to proteins identified in CBF β _3xMut_FLAG cells revealed 191 proteins which may be reliant upon CBF β residues G61, N63, and N104 for their translation.

Gene ontology was applied to this list of 533 proteins, revealing high enrichment of biological process terms signal transduction, transcription, cell division, adhesion, and migration (**Figure 4.8**). Many of these pathways are de-regulated in cancer, and it's possible CBF β plays a larger role in them than previously thought. Numerous cellular components were identified, as well as molecular functions involving binding of RNA, ATP, and GTP.

Reactome pathway analysis was next applied to this list of 533 proteins, and revealed high enrichment of pathways involving Rho GTPase signaling as well as VEGF (**Figure 4.9**). Rho GTPase signaling has been implicated in OS (63) and VEGF is a well-known poor prognostic indicator in OS (64). Numerous proteins implicated in cancer were found on this list, notable inclusions being SYDE1 (65), BRD2 (66), NRP1(67), HMOX1 (68), and CD70 (69).

GO was next applied to the 191 proteins enriched in CBF β _WT_FLAG cells relative to EV as well as CBF β _3xMut_FLAG cells, which we are classifying as potentially dependent upon CBF β residues G61, N63, and/or N104 for their translation (**Figure 4.10**). Enriched biological process terms in this list included signal transduction and cell adhesion, as well as cell division

and osteoblast differentiation, all of potential interest in OS. A variety of molecular functions were represented, in particular numerous binding capabilities in the molecular function GO class.

It's important to note that DiDBiT is a measure of overall nascent protein synthesis rate, and looks purely at the rate of production of each protein. As such, a protein could theoretically be up-regulated at the mRNA level, experience no change in regulation at the translation level, and still show up as enriched in DiDBiT. This protein would therefore show up as up-regulated by CBF β _WT_FLAG on DiDBiT, although in terms of CBF β influencing protein translation, nothing significant may have occurred. Additionally, proteins which have longer half-lives would experience slower rates of translation (70) and may not be identified via DiDBiT, as insufficient quantities of these proteins may be produced in order to be detected via LC-MS/MS. In order to evaluate the role of CBF β in translational regulation more specifically, and generate a more comprehensive list of proteins without the inherent bias of DiDBiT, we performed Ribo-Seq, or Ribosome Footprinting analysis, on CBF β _WT_FLAG as well as EV cells.

Ribo-Seq consists of bulk RNA sequencing of a sample coupled with parallel processing and deep sequencing of individual portions of each mRNA strand protected by the presence of a ribosome (21). Comparing the quantity of ribosome reads ascribed to a gene to the quantity of mRNA present for that gene allows measurement of ribosomal quantity per mRNA strand, which allows us to infer the rate at which a particular mRNA is being translated.

Quality control checks of Ribo-Seq analysis (**Figure 4.11**) demonstrated that analysis was performed properly. This experiment was performed in three biological replicates per cell line, and demonstrated good consistency between replicates. Congruency, a measure of correlation between quantity of RNA produced and quantity of ribosome footprints detected, was high for replicates as well. In **Figure 4.11 B**, the genes which fall outside of the linear correlation for congruency are

those which are subject to alterations in translational efficiency (TE) or ribosomal buffering based on the presence of CBF β . Ribosome footprint reads passed QC check as well, with characteristic 3 periodicity displayed in all replicates shown in **Figure 4.11 C**. Although QC data for just the first biological replicate of CBF β _WT_FLAG cells was shown, QC graphs for all other samples were extremely similar.

Many genes were found to be altered in transcription quantity, translation quantity, translational efficiency, or ribosomal buffering in CBF β _WT_FLAG cells compared to EV (**Figure 4.12A**). There was considerable overlap between these categories, but inclusion in one category doesn't necessitate inclusion in another.

Transcription quantity (via RNAseq) measures the quantity of mRNA strands transcribed for a given gene. Translation quantity (via deep sequencing of ribosomal footprints) is a measure of quantity of a given protein being produced. The “bread and butter” of Ribo-Seq lies in comparing these two measurements, and generating two new metrics: translation efficiency (TE) and ribosomal buffering. TE is a measure of ribosome occupancy on a given gene, and is calculated by relating the quantity of ribosome footprint reads for that gene compared to the overall quantity of mRNA of that gene as measured by RNAseq. This is a measure of how efficiently the cell produces that protein from a given mRNA input. Higher efficiency means that mRNA is more actively translated. Ribosomal buffering occurs when a gene experiences a change in transcription, but the effect of this on protein quantity produced is blunted, or buffered, by a compensatory change in translation in the opposite direction. For example, if a gene is down-regulated at the mRNA level by a certain fold, the translation efficiency of this gene may also be increased, meaning the level of protein produced would not be reduced by the same fold. It's possible the overall quantity of protein produced remains the same, but without measuring buffering one may

presume there were no alterations in gene expression, when in fact radical changes in regulation had occurred.

As we are investigating the role of CBF β in translation, we are interested in the genes which experienced a change in ribosome footprints (Diff. Ex. RIBO), those altered in TE as measured via Anota2Seq (A2S) or DTEG algorithms (denoted A2S or DTEG TE), and those which experienced ribosomal buffering as defined by A2S (A2S buffering), as well as any overlaps between these groups. A gene may be included in just one of these categories, or multiple, and be of interest.

We next analyzed similarities between DiDBiT data and Ribo-Seq (**Figure 4.12B**). As mentioned previously, DiDBiT does not take rate of transcription into account, and focuses solely on the rate of nascent protein synthesis between CBF β _WT_FLAG and EV cells. For this reason, we avoided the A2S buffering category as well as A2S or DTEG TE and instead looked at overlap between overall production of mRNA (as measured by RNAseq), overall rate of protein synthesis (Diff. Ex. RIBO), and nascent proteins elevated in synthesis in CBF β _WT_FLAG cells via DiDBiT. Of the proteins identified as having differential synthesis quantity via Ribo-Seq or DiDBiT, 31 were found to be up-regulated in both experiments. These 31 proteins (**Table 1**) are certainly of high interest, and further study is needed to investigate the relative importance of a gene being identified only in DiDBiT, or Ribo-Seq, or across both.

RUNX2 was present in the list of differentially translated proteins as identified by DiDBiT, although it did not reach this cutoff in Ribo-Seq analysis. The RNAseq portion of Ribo-Seq analysis measured RUNX2 mRNA levels to be nearly identical between EV and CBF β _WT_FLAG cells, corroborating earlier data that this discrepancy in RUNX2 protein production must be via a post-transcriptional mechanism. In terms of ribosomal footprint reads,

RUNX2 RIBO reads per kilobase per million mapped reads (RPKM) was ~7.12 in EV and ~7.78 in CBF β _WT_FLAG cells, indicating elevated RUNX2 protein production in CBF β _WT_FLAG cells. Although this did not make it to the enrichment cutoff to be binned as differentially expressed in terms of ribosome footprint reads, it was encouraging to see a difference between the two cell lines in RUNX2 protein production which was not mirrored in transcription. Stabilization of RUNX2 via binding to CBF β may still play a role in the discrepancy of RUNX2 protein, but RUNX2 being identified as differentially synthesized via DiDBiT still suggests CBF β to influence overall production of RUNX2.

We next applied gene ontology to genes identified as positively influenced in translation in the presence of CBF β _WT_FLAG via Ribo-Seq (**Figure 4.13**). Included in this list were genes which underwent an increase in TE in CBF β _WT_FLAG cells compared to EV cells. Also included were genes that exhibited positive translation buffering in CBF β _WT_FLAG cells compared to EV. Positive buffering refers to genes which were down-regulated in transcription in CBF β _WT_FLAG cells relative to EV, but simultaneously exhibited a compensatory increase in translation efficiency. This does not necessarily lead to an increase in the overall quantity of protein produced, but does speak to up-regulation of expression of these genes at the translational level resulting from the presence of CBF β _WT_FLAG. Altogether, this list totaled 586 genes.

Enriched GO terms among these genes included biological process terms specific to transcription and gene expression. Interestingly, cell differentiation and cell adhesion were also mentioned, processes known to be implicated in cancer. As in previous GO studies, multiple cellular components were represented. Interestingly, the term “small GTPase binding”, enriched among DiDBiT candidates (**Figure 4.10**) was also enriched in Ribo-Seq analysis. Small GTPases as well as their downstream effectors have been implicated in a variety of cancers including OS

(71–74). Notably, multiple members and regulators of the RAS family, a group of proteins infamous in their association with cancer (75–77) were identified in this gene list.

Following this, these same genes were subjected to Reactome and KEGG pathway analysis (**Figure 4.14**). KEGG pathway analysis revealed enrichment of multiple pathways extensively implicated in cancer, such as PD-L1 (78), MAPK (79), and RAP1 (80), as well as enrichment of the “pathways in cancer” term itself. Furthermore, the KEGG term “hepatocellular carcinoma” (HCC) was also strongly enriched, which was interesting as proteins implicated in HCC were also among the list of high confidence interactors of CBF β identified via IP-MS (**Figure 4.5**). Reactome pathway analysis also demonstrated MAPK signaling cascade enrichment as well as multiple terms associated with GTPase cycles such as RHO, CDC42 and RAC1. Importantly, Rho GTPase activity was also highly ranked among genes identified via DiDBiT (**Figure 4.9**).

In conclusion, we have generated a comprehensive list of medium- and high-confidence interacting partners of CBF β in OS cells. This represents the first of its kind in OS. mRNA processing and translation terms were highly enriched in GO analysis of CBF β interactors, and the medium-confidence interacting partners list included proteins previously reported to interact with CBF β . Bonafide binding partners of CBF β , RUNX1 and RUNX2, appeared in our screen, giving us confidence in the validity of our assay. Numerous proteins identified in our assay as high-confidence interactors of CBF β have implications in cancer and OS specifically, and both members of the NAC complex, a heterodimer strongly implicated in protein translation, were among this list. Additionally, we identified seven proteins which may depend upon CBF β residues G61, N63, and/or N104 for their interaction. We also conducted DiDBiT as well as Ribo-Seq to identify genes which CBF β may translationally regulate, and pathway analyses of these genes revealed many enriched terms which are of interest in OS and cancer biology at large. Notably, DiDBiT and Ribo-

Seq both revealed Rho GTPase activity to be strongly enriched among their respective gene lists, suggesting CBF β may play a significant role in regulation of this pathway. As these experiments represent an enormous quantity of data with a wide array of future directions, more study is needed to understand the full significance of CBF β in OS, and the outcome of CBF β -mediated regulation of the most salient targets identified. Encouragingly, GO analysis of putative interacting partners, and regulatory targets, of CBF β revealed a multitude of terms strongly implicated in cancer. Based on these data, it appears highly likely that CBF β plays a role in protein translation in OS, and that proteins implicated in OS and cancer at large fall under its regulatory purview.

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Chapter 5

Conclusions and Future Directions

General Conclusions of the Dissertation Work

The studies within this dissertation entail efforts to describe a noncanonical role of CBF β in osteosarcoma (OS), and enhance our understanding of whether CBF β could present a novel therapeutic target in OS. While there has been considerable effort in researching new therapeutic approaches in OS, one aspect of OS which makes development of therapies difficult is the high degree of heterogeneity among patient tumors. This, coupled with OS having a very low prevalence, has caused OS therapeutic development to stagnate in comparison to other cancers. In fact, OS therapies have not appreciably changed in the last 30 years. Protein translation acts as a convergence point of multiple signaling pathways, and could provide a mechanism to overcome the heterogeneity of the OS patient population. This project focuses on studying the role of CBF β in protein translation in OS, and investigates it as a potential therapeutic target.

The data gathered in this project began with study of the interaction of RUNX2, a DNA binding transcription factor, and CBF β , a co-activator to RUNX2. RUNX2 expression, as well as that of its gene targets, is elevated in OS and correlated with poor prognosis, which brought this protein and its binding partner into our focus. Development of a CBF β knockout (KO) cell line and profiling via western blotting revealed a decrease in RUNX2 protein expression upon loss of CBF β , which was interesting and unexpected. The foundation of this research was built on another project studying a small molecule allosteric inhibitor of the CBF β -RUNX2 interaction, which was the reason for development of the CBF β KO cell line. With further profiling of this cell line, the mechanism of this reduction in RUNX2 expression became the focus of this work.

In chapter 2 we demonstrate that CBF β regulates RUNX2 expression via a post-transcriptional mechanism. Using western blotting, we confirmed the reduction in RUNX2 protein expression upon loss of CBF β , and via qPCR and RNA-seq we established that this discrepancy

did not occur at the mRNA level. This implied a post-transcriptional mechanism at play, and as RUNX2 is already being explored as a therapeutic target in OS, this new mechanism of CBF β regulation of RUNX2 expression could provide a novel therapeutic target in OS. Following this, using proteasome and translation inhibitors, we demonstrate that a reduction in RUNX2 stability due to loss of its binding partner CBF β does not fully explain this reduction in RUNX2 expression at the protein level. Around the same time, another study in breast cancer noted a similar reduction in RUNX1 expression upon loss of CBF β , and postulated this to result from CBF β playing a role in protein translation. To investigate whether this may also be occurring in OS, we used a Click-iT Alexa488 protein translation assay to measure global protein translation, and observed a reduction in global protein translation in U2OS cells upon loss of CBF β . This role was reported in breast cancer to occur via CBF β interaction with hnRNPk, and we used reciprocal co-immunoprecipitation (co-IP) to validate that this interaction occurs in OS cells as well. Finally, we used lentiviral transduction to re-introduce wild-type (WT) CBF β back into our CBF β KO cells, and verified that RUNX2 protein expression increased in the presence of CBF β . This confirmed that we were observing a CBF β -dependent effect, and not being misled by nonspecific CRISPR activity.

These results were very encouraging, but the dual roles of CBF β presented a complex mechanism to study. CBF β exerts a transcriptional role when present in the nucleus and bound to the RUNX proteins, yet this translational role reportedly occurs in the cytoplasm. Furthermore, CBF β cannot move freely from the cytoplasm to the nucleus by itself, and is instead shuttled into the nucleus onboard the RUNX proteins. In this manner, it is possible that interaction with RUNX proteins would reduce the cytoplasmic pool of CBF β , and negatively affect the performance of its

translational role. Additionally, knockout of one RUNX protein leads to compensatory increases in other RUNX proteins, making individual study of the translational role of CBF β difficult.

In chapter 3, we describe efforts made to design and validate a mutant form of CBF β , designed to display decreased affinity toward the RUNX proteins and instead remain in the cytoplasm. Using recombinant protein production combined with peptide inhibitors, we validated residues G61 and N63 of CBF β to be important in CBF β -RUNX2 interaction. We then generated a mutant of CBF β with alanines at these residues as well as N104, and re-introduced this mutant form into our CBF β KO cells to generate CBF β _FLAG_3xMut. We conducted cellular electrothermal shift assay (CETSA) and validated that our mutations had significantly interrupted CBF β -RUNX2 interaction, resulting in inhibited binding between RUNX2 and our mutant form of CBF β as compared to WT CBF β . Next, we conducted nuclear/cytoplasmic fractionation to investigate the functional consequence of our mutant form of CBF β on RUNX protein mediated nuclear shuttling. We observed a strong inhibition of shuttling of CBF β _FLAG_3xMut into the nucleus onboard RUNX proteins, validating that we had accomplished our goal of retaining CBF β in the cytoplasm in order to study its translational role.

At this point we had generated compelling data suggesting that CBF β may play a novel role in protein translation in OS, however our studies thus far had focused purely on the relationship between CBF β and RUNX2. In chapter 4 we extend our data collection beyond RUNX2 and performed three assays which look at all proteins within the cell. The assays in chapter 4 served to answer two questions: 1) what is the full list of proteins that CBF β interacts with in OS cells? And 2) what is the full list of proteins which may have their translation affected by the presence of CBF β ? We performed IP-MS to characterize the interactors of CBF β , and revealed a list of 23 proteins which we binned as high confidence interactors of CBF β in OS. Importantly,

the vast majority of these proteins have not been profiled in the literature at the time of writing this. Additionally, by performing IP-MS on our mutant form of CBF β , we revealed seven (7) proteins which are dependent upon CBF β residues G61, N63, and/or N104 for their binding to CBF β . Next, we performed direct detection of biotinylated peptides (DiDBiT) as well as Ribo-Seq, with the goal of elucidating which proteins may have their translation influenced by CBF β . These studies revealed hundreds of proteins which experience altered production in the presence of CBF β , with gene ontology (GE) of this list revealing strong enrichment of biological process terms cell division, cell adhesion, and cell migration. These processes are strongly implicated in cancer, and it's possible CBF β may influence these processes by altering production of proteins involved in them. Additionally, Reactome analysis of these proteins saw enrichment of Rho GTPase signaling and VEGF signaling terms, suggesting CBF β may play a role in regulating these processes. GO terms cell adhesion, migration, and division, were also enriched among proteins decreased in production in the presence of CBF β _FLAG_3xMut as compared to CBF β _WT_FLAG, suggesting these mutated residues may be important in the production of these proteins.

Future Directions

While the data from this project has significantly enhanced our understanding of CBF β in OS, there are numerous additional studies which could build on this work and dive deeper into whether this protein presents a viable therapeutic target. In **Chapter 2** we demonstrate that RUNX2 protein levels are affected by the presence of CBF β , and a decrease in RUNX2 stability upon loss of CBF β does not fully explain this occurrence. A limitation exists in the second portion of that statement, as the stability assays we employed run into confounding variables when used to study proteins with particularly long half-lives, such as RUNX2. As mentioned in the **Chapter 2 Discussion**, treatment with MG132 or Cycloheximide for these extended periods is likely to perturb normal cellular function, hampering accurate measure of the shift in RUNX2 stability upon loss of CBF β . As mentioned previously, these longer treatments may stimulate lysosomal degradation of RUNX2, which could be addressed by use of lysosomal protease inhibitors, although this would introduce additional variables to an already precarious experiment. For this reason, it may be better to switch approaches entirely.

To get a better picture of RUNX2 half-life, a useful assay involves growing cells in methionine-free media supplemented with L-azidohomoalanine (AHA), a methionine analog compatible with click chemistry conjugation. In short, cells would be incubated with AHA for ~4h, then switched to incubation in normal media for various durations. Lysate is then extracted, AHA is click conjugated to biotin, then RUNX2 is immunoprecipitated (IP) and quantified by western blot. Western blotting for biotin within the RUNX2 IP, and normalizing that to total RUNX2 signal, would yield a measure of quantity of labeled RUNX2 remaining over time. Usage of various post-AHA incubation durations in normal media would allow measure of the degradation of tagged RUNX2, without introducing toxicity into cells as, since AHA does not

interfere with protein translation or degradation. As mentioned in chapter 2, it's possible that lysosomal trafficking of RUNX2, potentially by SOX9, is confounding the stability data we generated, and an AHA pulse-chase experiment would avoid this issue.

AHA pulse chase would require significant optimization of RUNX2 IP, which we found intractable within our timeframe for this work, but is likely not impossible given adequate resources. Optimization of RUNX2 IP would also allow for treatment of cells with MG132, a proteasome inhibitor, followed by immunoblotting for ubiquitination of RUNX2, which would present clear data on altered degradation of RUNX2 based on CBF β status, and dovetail with stability data from AHA pulse-chase. While these would be useful, their overall benefit balanced against their cost would need to be evaluated first.

In chapter 2 we also re-introduced WT CBF β back into CBF β KO cells and observed a recovery of low RUNX2 expression. While we had verified the decrease in RUNX2 protein expression upon loss of CBF β occurs in a post-transcriptional manner, it would be useful to validate that the increase in RUNX2 protein expression upon re-introduction of CBF β also occurred via a post-transcriptional mechanism. This could be validated by way of qPCR of RUNX2 in each cell line normalized to HPRT, in the same manner as was originally tested in U2OS WT vs. U2OS CBF β KO cells in the beginning of chapter 2.

In chapter 3 we introduced key mutations into CBF β at residues G61, N63, and N104 (CBF β _FLAG_3xMut), with the goal of inhibiting binding to RUNX proteins, thereby preventing RUNX-mediated nuclear shuttling of CBF β from complicating study of its translational role. We demonstrated that our mutant exhibited decreased CBF β -RUNX2 binding and subsequent nuclear shuttling, as was the goal, however our mutant was unable to recover low RUNX2 expression seen in Empty Vector (EV) cells. Usage of the aforementioned AHA pulse chase assay on these 3xMut

cells would provide useful information as well, to test if RUNX2 stability differed between CBF β CBF β WT and 3xMut cells. RUNX2 expression may be unchanged relative to EV due to low stability, or perhaps these residues we mutated on CBF β also play a role in the translation of RUNX2 specifically. Usage of AHA pulse chase would assist in revealing which of these is the dominant factor at play. Additionally, usage of qPCR for RUNX2 expression in CBF β _3xMut_FLAG-expressing cells would also be informative.

In chapter 4 we conducted three large assays with the goal of answering two key questions: 1) what are all the proteins which CBF β interacts with in OS cells? And 2) what are all the proteins which CBF β may translationally regulate. To answer question 1, we performed IP-immunoprecipitation mass spectrometry, (IP-MS), and identified numerous new proteins which interacted with CBF β _WT_FLAG, the lion's share of which have not been reported anywhere in the literature. A necessary next step to further validate these proteins as specific interactors is to conduct IP followed by western blotting. Of course, it would not be feasible to purchase an antibody for every identified protein, as costs for that would be prohibitive, but blotting for certain proteins of interest for confirmation would be wise. Particularly interesting targets, such as NCK2, USP39, BTF3, and NACA2 could then be further validated via reciprocal co-IP, such as was performed towards the end of chapter 2 studying CBF β -hnRNPk interaction. Additionally, interaction between these proteins and CBF β could also be revealed by conducting cellular electrothermal shift assay (CETSA) on these proteins, in the event their stoichiometry in binding to CBF β is not high enough to be detected in a pulldown, or antibodies suitable to IP them are not available. These same experiments would be useful to validate the proteins we suggested to rely on CBF β residues G61, N63, and N104 for their interaction, as listed in **Figure 4.6**.

To answer question 2, we performed two orthogonal assays; direct detection of biotinylated peptides (DiDBiT) and Ribo-Seq. These are both complex assays that generated a myriad of data, and therefore a myriad of potential research directions. Building on DiDBiT data, further studies into the characteristics of CBF β _FLAG_3xMut cells compared to CBF β _WT_FLAG would prove very useful in unraveling the significance of these residues of CBF β on overall cancer cell behavior. For example, cell adhesion, migration, division, and osteoblast differentiation showed up as highly ranked biological processes among CBF β _WT_FLAG cells as compared to CBF β _FLAG_3xMut cells, suggesting our mutated residues of CBF β play a role in these processes. This could be further studied by use of migration/invasion or proliferation assays. These phenotypic experiments would bridge the gap between the specific molecular consequences revealed in this study and global effects which would influence cancer cell malignancy. This would also add more data which may further support CBF β as a therapeutic target in OS.

While our goal in generating CBF β _3xMut_FLAG was to decouple the transcriptional and translational roles of CBF β , the appearance of so many proteins which experienced diminished production in the presence of CBF β _3xMut_FLAG suggests we may have also affected the translational component of CBF β . In order to fully utilize CBF β _3xMut_FLAG as a tool in solely investigating the translational role of CBF β without the transcriptional role confounding data, Ribo-Seq could be performed on CBF β _3xMut_FLAG cells vs CBF β _WT_FLAG to measure how drastically these mutations did influence the role of CBF β in protein translation.

In terms of data from Ribo-Seq, while there was a small cohort of proteins which were upregulated in translational efficiency in the presence of CBF β , there were many more proteins which experienced a positive change in translational buffering upon loss of CBF β . That is to say, these proteins experienced a decrease in transcription, but this was coupled with a compensatory

increase in translational efficiency. Although this may not necessarily lead to a bulk increase in the quantity of these proteins, they did experience alterations in translational efficiency in the presence of CBF β , which is still important to analyze. Future studies which delve deeper into the effect of translational buffering in OS would be useful in giving these results greater context. Translational buffering has been implicated in cancer, although with it being a newly identified phenomenon, our knowledge on it and the overall impacts are more limited than that of more classical gene expression regulatory mechanisms.

The importance of CBF β in OS could also be measured via assays which compare CBF β KO cells to WT cells in terms of susceptibility to chemotherapies. While our lab has U2OS CBF β KO cells, the reliance of other parental cell lines on CBF β could be assessed by use of siRNA against CBF β combined with a chemotherapeutic. Functional assays such as migration/invasion would also be interesting to get a picture of the importance of CBF β in OS cell lines other than U2OS.