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ORIGINAL ARTICLE Translocations at 8q24 juxtapose *MYC* with genes that harbor superenhancers resulting in overexpression and poor prognosis in myeloma patients

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Secondary *MYC* translocations in myeloma have been shown to be important in the pathogenesis and progression of disease. Here, we have used a DNA capture and massively parallel sequencing approach to identify the partner chromosomes in 104 presentation myeloma samples. 8q24 breakpoints were identified in 21 (20%) samples with partner loci including *IGH*, *IGK* and *IGL*, which juxtapose the immunoglobulin (Ig) enhancers next to *MYC* in 8/23 samples. The remaining samples had partner loci including *XBP1*, *FAM46C*, *CCND1* and *KRAS*, which are important in B-cell maturation or myeloma pathogenesis. Analysis of the region surrounding the breakpoints indicated the presence of superenhancers on the partner chromosomes and gene expression analysis showed increased expression of *MYC* in these samples. Patients with *MYC* translocations had a decreased progression-free and overall survival. We postulate that translocation breakpoints near *MYC* result in colocalization of the gene with superenhancers from loci, which are important in the development of the cell type in which they occur. In the case of myeloma these are the Ig loci and those important for plasma cell development and myeloma pathogenesis, resulting in increased expression of *MYC* and an aggressive disease phenotype.

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INTRODUCTION

Rearrangements at 8q24 have been reported in up to 47% of myeloma patients by a combination of fluorescence *in situ* hybridization (FISH), spectral karyotyping and classical cytogenetics.¹ In presenting myeloma this frequency is lower, with abnormalities of 8q reported in 15% of cases using mapping arrays and FISH.^{2,3} The gene of interest in this region is *MYC*, an oncogene that has a pivotal role in cell growth, proliferation, tumorigenesis and stem cells.⁴

The importance of *MYC* activation in myeloma has been shown through the use of the Vk**MYC* transgenic mouse model, where activation of *MYC* arises through AID-dependent somatic hypermutation during B-cell development, resulting in the onset of myeloma in these mice.⁵ *MYC* has also been shown to be activated in the transition from monoclonal gammopathy of undetermined significance to myeloma, implicating it in disease progression.⁶ Myeloma cells have been shown to have a dependency on MYC for survival, where inhibition of MYC by small hairpin RNA or small-molecule inhibitors results in cell death indicating that MYC is a promising therapeutic target.⁷

The mechanism of *MYC* activation is mainly through secondary translocations involving the immunoglobulin (Ig) loci (*IGH* > *IGL* > *IGL* > *IGK*), which juxtapose the strong B-cell enhancers present at these loci and *MYC*, resulting in overexpression of the oncogene.⁸ Unlike primary translocations in myeloma, which are often simple reciprocal exchanges of chromosomal material, the rearrangements that result from *MYC* translocations are often complex, involving many partner chromosomes.^{1,8} Interestingly, it has been reported that up to 74% of *MYC* rearrangements do not involve an Ig locus³ leading to the conclusion that other

mechanisms of activation may also be important in myeloma. Using FISH, partner chromosomes at 1p13, 1p21–22, 6p21, 6q12–15, 13q14 and 16q22 have been identified, but the specific loci involved have remained elusive.^{3,9–11}

The breakpoints on 8q24 have been mapped in a large number of myeloma cell lines and the majority are found within 1 Mb of *MYC*, but some can be greater than 3 Mb either telomeric or centromeric of the locus.^{8,12,13} The loci surrounding *MYC* are *POU5F1B* (centromeric) and *PVT1* (telomeric). *PVT1* is a non-coding RNA that has been shown to be the location of variant t(8;22) breakpoints in Burkitt's lymphoma,¹⁴ as well as generating fusion genes with *WWOX* and *NBEA* in myeloma patients with an 8q24 rearrangement.¹³

Here, we have used targeted capture followed by massively parallel sequencing to pull down the region surrounding *MYC* in a series of presenting myeloma cases in order to identify any translocations in this area and the mechanism of action involved.

MATERIALS AND METHODS

Cell selection

CD138-positive bone marrow plasma cells were selected to a purity >95% using magnetic assisted cell sorting (Miltenyi Biotech, Bisley, UK). Tumor DNA and RNA were extracted using the AllPrep kit (Qiagen, Manchester, UK). All patients were at presentation and had not received any treatment when the sample was taken.

FISH

Probes have been previously published with the addition of the *LPL* (8p22), CEP 8 and *MYC* (8q24.1-24.21) probes (Abbott, Maidenhead, UK).¹⁵⁻¹⁷

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MYC abnormalities were defined using the t(8;14) fusion probe, *LPL*/CEP 8/ *MYC* probes, IGH@ translocations (IGH@ break-apart probe followed by fusion probes for the common partner chromosomes). FISH results were interpreted alongside karyotype data, where available.

Targeted capture of the MYC locus

A targeted capture system was designed using the SureSelect system (Agilent, Santa Clara, CA, USA) that was based on tiling RNA baits across the *MVC*, *IGH*, *IGK* and *IGL* loci as previously described.¹⁸ The region captured surrounding *MYC* spanned from 127.5–129.8 Mb on chromosome 8, roughly 1 Mb on either side of *MYC*, which is located at 128.75 Mb. This region includes *POUSF1B* and *PVT1*, which are common sites of 8q24 translocations in myeloma.

DNA from 104 samples were assayed using 150 ng of DNA and a modified capture protocol with eight cycles of prehybridization PCR and 11 cycles of posthybridization PCR. Samples were barcoded using Illumina (San Diego, CA, USA) indexes and up to 27 samples were sequenced per lane on a HiSeq2000 generating 76-bp paired-end reads. After base calling and quality control metrics, the raw fastq reads were aligned to the reference human genome (build GRCh37) resulting in a median depth of 289 × per sample after de-duplication for the captured region.

Translocation breakpoints were identified in the sequencing data using DELLY.¹⁹ Breakpoints called using the bioinformatic approach were further filtered based on depth, unique mappability for 76 bp reads, number of supporting reads and whether or not they were detected in non-tumor samples. The coordinates of the breakpoints and superenhancers were compared with randomized data produced using Monte Carlo sampling using the Genomic Hyper Browser.²⁰

Statistical analyses were performed using SPSS version 19.0 (SPSS Inc., Chicago, IL, USA). Discrete data were expressed as frequencies and percentages and were assessed by Fisher's exact test or the χ^2 -test, as appropriate. Survival curves were plotted using the Kaplan-Meier method. Differences between curves were tested for statistical significance using the log-rank test. A multivariate Cox regression analysis was done to identify factors significantly associated with progression-free survival (PFS) and overall survival (OS), with statistical significance set at P < 0.05.

RESULTS

MYC rearrangements detected by FISH

MYC rearrangements were tested for using interphase FISH on $CD138^+$ selected plasma cells on patient samples from the UK MRC Myeloma IX trial using a combination of probes for *MYC* (8q24), *LPL* (8p22) and *CEP* 8. Interphase FISH was successful on 751 patients and was interpreted with data from FISH probes for the *IGH* locus and karyotyping. Table 1 shows the frequency of *MYC* abnormalities, split by translocation group in these samples. *MYC* abnormalities were detected in 26% of samples with the most frequent abnormality being gain of the locus, followed by a split probe signal. In 50% (32/63) of those with a split probe signal the *IGH/K/L* locus was identified as the partner. Of the 32 samples with an Ig partner 28 were with the *IGH* locus. In the remaining 50% the partner remained undetermined, but may involve the light chain loci in samples where cytogenetics or karyotyping was unsuccessful or uninformative. *MYC* rearrangements were not

associated with any particular primary translocation, but were associated with ISS II (P = 0.022) or ISS III (P = 0.0027). There was no negative impact on PFS or OS in patients with a *MYC* abnormality.

MYC rearrangements detected by sequence capture

The presence of unidentified *MYC* partner chromosomes led us to investigate possible partner chromosomes using an alternative technique. We had previously used a DNA capture technique followed by massively parallel sequencing to identify the translocation partner chromosomes to the *IGH/K/L* loci.¹⁸ In this assay we had also captured a region surrounding *MYC* and set out to identify potential partner chromosomes. A region of ~2 Mb was tiled around the *MYC* locus using RNA baits and used to capture DNA and associated translocations in this region.

We assayed 104 presentation myeloma samples for the presence of MYC translocations using the capture assay followed by massively parallel sequencing. The samples consisted of 55 samples from the Myeloma IX trial and an additional 49 samples from the UK Myeloma XI trial, for which FISH results are not available. The samples were categorized according to IGH translocation using a combination of FISH,^{15,21} gene expression²² and targeted capture of the IGH locus¹⁸ and consisted of a variety of samples with IGH translocations and 8q24 rearrangements as shown in Table 2; Supplementary Table 1. Only three samples analyzed by the capture technique had a split MYC locus, as determined by FISH or karyotyping. In these samples the translocation was detected using the capture and was found to be the IGH locus in one sample (673; verifying the FISH result) and in the other two samples (29 and 1310) was found to be chromosome 6 (also seen by karyotyping in one sample).

Breakpoints were identified and mapped in 21 samples and are shown in Figure 1. We identified breakpoints in 10 samples where either FISH or karyotyping failed to identify a translocation, as well as three samples where FISH did identify a translocation and also eight samples for which no FISH data were available. Interestingly,

Translocation	Assayed (%)	With 8q24 breakpoint (% of group)	P-value
t(4;14)	13(12.5)	0(0)	0.031
t(6;14)	8(7.7)	1(12.5)	NS
t(11;14)	28(26.9)	5(17.8)	NS
t(14;16)	17(16.3)	11(64.7)	< 0.001
t(14;20)	6(5.8)	1(16.6)	NS
HRD	31(29.8)	3(9.6)	0.037
Other ^a	1(0.9)	1(100)	NS
Total	104	21	

8q Status	Translocation (Tx) group							
	t(4;14)	t(11;14)	t(6;14)	t(14;16)	t(14;20)	HRD	HRD + Tx	None
Normal	49	74	4	17	6	294	20	90
Gain	7	17	2	8	4	46	10	17
Split	3	10	0	2	2	40	1	5
Del	5	0	0	0	0	4	1	14
Total abnormal (%)	23.4	26.7	33.3	37.0	50.0	23.2	37.5	28.5

Abbreviation: FISH, fluorescence in situ hybridization.

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there was an enrichment for t(14;16) and a depletion of t(4;14) and hyperdiploidy associated with an 8q24 breakpoint, Table 2, which was not seen with FISH alone.

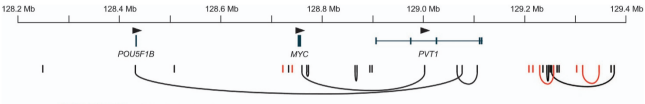
The positions of the breakpoints on 8q24 were spread out over 1.2 Mb. The majority of the breaks were situated telomeric of *PVT1* (51.5%) with 12% within *PVT1*, 15% between *MYC* and *PVT1* and 15% between *POU5F1B* and *MYC*. Of the three samples with breakpoints within *PVT1* two were found to have potential fusion genes, one with *FOXO3* and the other with *LINC00309*.

Nine samples had two breakpoints on 8q24, often with two different chromosome partners. In samples with two breakpoints, the sequencing reads indicate that *MYC* is only involved in one of the breakpoints (that is, only one side of the translocation was captured). The reason for this is not clear and will require long reads to clarify the final genomic rearrangements. Several samples had complex rearrangements resulting in multiple chromosomal

segments being joined together, often involving *MYC*, an Ig enhancer and the primary Ig partner oncogene (for example, sample 11/625 that has a t(6;14) and a t(6;8) or sample 12/0404 that has a t(11;14) and a t(8;11)). Details of breakpoints are shown in Table 3. It is possible that the Ig enhancer is influencing both the target oncogene of the primary translocation (for example, *CCND1* or *CCND3*) and *MYC* through the assembly of a complex t(8;11;14) or t(6;8;14) derivative chromosome.

Superenhancer colocalization with MYC

A total of 8/21 samples (38%) with *MYC* abnormalities had rearrangements with an Ig locus on the partner chromosome. Ig partner loci are recognized to upregulate expression of the target oncogene and have been shown to have clinical relevance as prognostic markers. Given that the mechanism of upregulation of



Legend: | IG translocation | Non-IG translocation

Figure 1. *MYC* locus breakpoints in myeloma. The locations of breakpoints are indicated by vertical lines corresponding in color to whether the partner chromosome belongs to an Ig loci (IGH@, IGK@ or IGL@; red) or a non-Ig locus (black). The genes and orientation are indicated according to their genomic location on chromosome 8. Arcs indicate the positions of two breakpoints found in one sample.

Sample	IGH translocation	1st Chr	1st breakpoint	Part of 1st chr positioned next to chromosome 8	1st Gene ^a	Distance (bp) to MM1.s enhancer from breakpoint	Distance (bp) to MM1.s superenhancer from breakpoint	2 nd Chr	2 nd breakpoint	2 nd Gene ^a
29	t(7;14)	chr22	36779252	Telomeric	МҮН9	0	0	chr8	129251475	PVT1-CCDC26
29	t(7;14)	chr6	7986298	Telomeric	TXNDC5	8862	8862	chr8	129375213	PVT1-CCDC26
176	None	chr1	118246012	Telomeric	FAM46C-GDAP2	16232	16232	chr8	129264632	PVT1-CCDC26
222	t(14;16)	chr2	64412927	Centromeric	LINC00309	3194	450109	chr8	129064694	PVT1
222	t(14;16)	chr2	64468628	Telomeric	LINC00309-LGALSL	12603	169177716	chr8	129104714	PVT1
471	None	chr22	23071965	Telomeric	IGL	5005	5005	chr8	128737937	POU5F1B-MYC
475	t(14;16)	chr12	25507375	Telomeric	KRAS-IFLTD1	20015	18435462	chr8	129368991	PVT1-CCDC26
478	t(11;14)	chr2	77937177	Telomeric	LRRTM4-SNAR-H	2872963	13042429	chr8	128896717 ^b	MYC-PVT1
592	t(14;16)	chr11	111195351	Telomeric	C110RF93-MIR4491	36912	36912	chr8	129247671	PVT1-CCDC26
592	t(14;16)	chr10	122670424	Centromeric	MIR5694	1102347	10042441	chr8	129248430	PVT1-CCDC26
673	t(14;16)	chr14 ^c	106119946	Telomeric	IGH	26346	26346	chr8	129209665	PVT1-CCDC26
730	None	chr22	41824373	Centromeric	TEF-TOB2	13346	3109392	chr8	129312555	PVT1-CCDC26
730	None	chr22	23280899	Centromeric	IGL	0	0	chr8	129345574	PVT1-CCDC26
984	t(14;16)	chr2	89151255	Centromeric	IGKC	95671	2922643	chr8	129228289	PVT1-CCDC26
984	t(14;16)	chr2	89130434	Centromeric	IGKC	34641	8060442	chr8	129254453	PVT1-CCDC26
1112	t(11;14)	chr11	69283791	Centromeric	CCND1	131027	2226257	chr8	128427581	POU5F1B-MYC
1112	t(11;14)	chrX	146700223	Centromeric	MIR510-FMR1-AS1	NA ^d	NA ^d	chr8	129075004	PVT1
1310	t(14;16)	chr6 ^e	108911604	Telomeric	FOXO3	0	0	chr8	128761005	MYC-PVT1
1310	t(14;16)	chr6 ^e	108908007	Centromeric	FOXO3	0	0	chr8	129000293	PVT1
11/088	t(14;16)	chr10	125858805	Telomeric	CHST15-OAT	0	0	chr8	129235977	PVT1-CCDC26
11/388	t(14;16)	chr22	23307780	Centromeric	IGL	5757	5757	chr8	128712616	POU5F1B-MYC
11/625	t(6;14)	chr6	41858885	Centromeric	USP49 (near CCND3)	377392	10585809	chr8	128244760	PCAT1-POU5F
11/741	t(14;16)	chr22	29210349	Telomeric	XBP1-ZNRF3	0	0	chr8	128772206	MYC-PVT1
11/741	t(14;16)	chr2	134989574	Centromeric	MIR3679-MGAT5	47048	22523506	chr8	128772558	MYC-PVT1
214	t(14;20)	chr22	23285257	Centromeric	IGL	0	0	chr8	129214819	PVT1-CCDC26
11/1212	t(14;16)	chr22	23258401	Centromeric	IGL	9496	9496	chr8	129301246	PVT1-CCDC26
11/1227	t(11;14)	chr14	105969653	Telomeric	MIR548AS-DAOA-AS1	24389	55470	chr8	128504864	POU5F1B-MYC
12/0213	t(11;14)	chr1	118302879	Telomeric	FAM46C-GDAP2	0	73103	chr8	128866931	MYC-PVT1
12/0365	t(14;16)	chr10	125858805	Telomeric	CHST15-OAT	Ő	0	chr8	129235977	PVT1-CCDC26
12/0365	t(14;16)	chr17	74521561	Centromeric	RHBDF2-CYGB	30899	645847	chr8	129262577	PVT1-CCDC26
12/0404	t(11;14)	chr11	69425933	Centromeric	MYEOV-CCND1	32171	2368440	chr8	128719531	POU5F1B-MYC

^aWhere >1 gene is named, the breakpoint is in the intergenic space between genes. ^bBreakpoint within inversion. ^cConfirmed by FISH. ^dComplex rearrangement containing 400 bp of chrX with translocations leading to chromosomes 14 (IGH@) and 8 (PVT1) as well as a t(8;11) and t(11;14). ^eConfirmed by karyotype underlined = derivative does not contain MYC.

Ig loci partner oncogenes is through colocalization of active enhancer elements with the oncogene, we examined the remaining samples with rearrangements for the presence of enhancer elements. We used data from two papers in which binding sites of BRD4 and MED1, which occur at transcriptionally active sites, enhancers and superenhancers in MM1.s cells had been annotated using only enhancer sites on the assembled derivative chromosome.^{23,24} As expected, the Ig loci breakpoints were typically close to or within an enhancer/superenhancer (minimum distance = 0, maximum distance = 26 kb) with the exception of *IGKC* breakpoints, which were ~292 kb from the nearest superenhancer (Table 3). This discrepancy is likely owing to MM1.s cells being a lambda light chain expresser that do not express a kappa light chain. If a cell line expressing a kappa light chain had been used in the analysis an enhancer may have been detected much closer.

In order to determine whether there was an enrichment of superenhancer sites in the vicinity of *MYC* breakpoints, we used the genomic locations of superenhancers from a variety of the cell types determined in Hnisz *et al.*²³ There was a statistically significant enrichment for breakpoints within 1 Mb of a superenhancer in the MM1.s myeloma cell line and in CD19⁺ B cells (P=0.0049 and 0.041, respectively). No enrichment was seen with enhancer locations in K562 cells, CD3⁺ T cells or

skeletal muscle myoblasts indicating a cell-specific enrichment for superenhancers in the B-cell lineage.

The partner loci to MYC are also mostly related to myeloma pathogenesis or B-cell biology (Table 3). In addition to the Ig loci, partner genes of MYC rearrangements included FAM46C, KRAS and *CCND1*. All of these genes are candidates in many genetic studies of myeloma. FAM46C is deleted in $\sim 20\%$ of samples and mutated in \sim 3% of samples.^{2,25} KRAS is mutated in 31% of samples and CCND1 is overexpressed in t(11;14) accounting for $\sim 15\%$ of samples.^{26,27} In addition, XBP1 is involved in plasma cell differentiation, the unfolded protein response and is mutated in a low percentage of relapsed refractory myeloma.^{26,28} The genes located next to breakpoints in the remaining samples may also be active in B-cell biology or myeloma pathogenesis as recurrent translocations are found near the CHST15 locus, which is known to be involved in B-cell signaling,^{29,30} and *FOXO3*, which is involved in B-cell development.³¹ Some rearrangements from these samples are shown in Figure 2 to illustrate the colocalization of superenhancers to MYC.

Impact on MYC expression

Previously published gene expression array data were available on 33 of the samples (Supplementary Table 2).² We categorized the

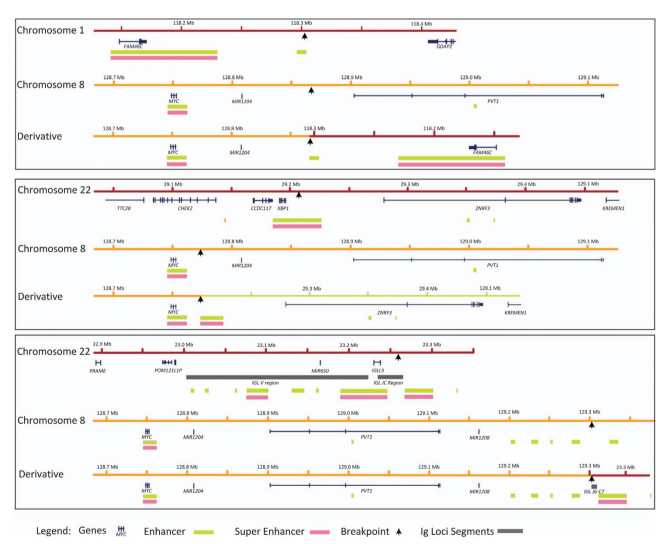


Figure 2. *MYC* rearrangements result in superenhancer colocalization on the derivative chromosome. Superenhancer elements are colocated near to MYC from a t(1;8) (top) or a t(8;22) (middle and bottom) where the partner chromosome gene (*FAM46C*, *XBP1* and *IGL*, respectively) has a known function in myeloma or B-cell biology.

samples according to the presence (n = 9) or absence (n = 24) of a MYC translocation, as detected by the capture assay, and used the MYC probe set (202431 s at) to examine any difference in expression between the groups. Those samples with any MYC translocation had higher expression of MYC compared with those without a translocation (median 703.2 vs 2313), but this did not reach significance with this number of samples (P = 0.065) Figure 3a. This was extended in the full Myeloma IX data set for which any translocation was detected by FISH or capture and for which expression array data were available. In this larger data set, samples with any translocation (n = 27) had a significantly higher expression of MYC (2003 vs 945.8, P = 0.009) compared with those with no translocation (n = 142), even allowing for those samples on which no capture had been performed (and may contain translocations not detected by FISH) (Figure 3b). The increase in expression of MYC is consistent with its deregulation being the central mechanism.

Impact on clinical outcome

A subset of patients (n = 55) from the capture panel had taken part in the UK MRC Myeloma IX trial, so we performed an analysis of the impact of *MYC* translocations on survival. *MYC* translocations had a significant impact on OS and PFS in univariate analysis (Table 3), which carried over into multivariate analysis, resulting in a significant decrease in OS and PFS for patients with a *MYC* translocation (Figure 4).

DISCUSSION

Here, we have investigated the partner loci involved in rearrangements with 8q24, namely *MYC*. We analyzed presenting cases of myeloma characterized by a range of different primary *IGH* translocation events and found evidence of a *MYC* translocation in 21% of samples, making it the most frequent translocation in presenting myeloma cases.

Although FISH analysis of 8q24 rearrangements found no significant association with primary translocation groups, we did find an enrichment for samples with a *MYC* breakpoint in samples that had a t(14;16) and a depletion of samples with a t(4;14) or hyperdiploidy when screened by the sequence capture technique.

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Other studies using FISH agree with our FISH-based results in that it has been reported that rearrangements at the *MYC* locus show a similar prevalence in hyperdiploid and non-hyperdiploid tumors¹ or show some trend to being associated with t(4;14).³ The discrepancy between our capture results and those determined by FISH may lie in the technique used to study the abnormality. One study used both FISH and mapping arrays to examine *MYC* breakpoints and found that 33% of breakpoints detected by array were not detected by routine FISH analysis.¹⁷ Therefore, a higher resolution assay, such as genome sequencing, may identify more breakpoints and explain some of the discrepant results. However, care must be taken regarding the frequency of sequence capture translocations within the primary translocation groups owing to the bias in sample selection and the relatively small number of samples studied.

It is notable that all groups find *MYC* rearrangements to be complex, often involving many chromosomes. We only capture the region surrounding *MYC* and are not able to detect any rearrangements downstream on the partner chromosome, which could result in inversions, insertions or duplications of DNA segments. Whole-genome sequencing of samples will provide more complex and detailed information regarding the final composition of the genome in these samples.

The translocations at 8q24 results in overexpression of MYC due to the colocalization of active superenhancers in the partner loci. The obvious examples of this belong to the known active enhancers in the B-cell lineage, the Ig loci, but in addition to these a series of previously unknown partners have also been identified. A similar study in B-cell lymphomas identified nonimmunoglobulin partners to MYC rearrangements and concluded that these are non-random processes that juxtapose MYC with genes involved in lymphomagenesis (namely BCL6, PAX5 and *IKAROS*).³² We have identified a different set of genes in this study and interestingly these genes are related to B-cell biology and myeloma pathogenesis. We hypothesize that in different diseases with MYC rearrangements there is a selection process, which results in active superenhancers from genes that are expressed in that cell type being placed near MYC. In all B-cell neoplasias these include the Ig loci, but in different B-cell subtypes the non-Ig loci differ to include those that are expressed in that cell type. In myeloma these non-lg partner genes include FAM46C, KRAS, XBP1

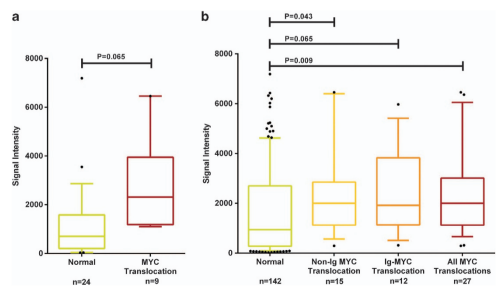


Figure 3. Expression of *MYC* in samples with a breakpoint is higher than in those without a breakpoint. (**a**) Expression data from 33 samples with sequence capture-determined translocations for those samples with no breakpoint at 8q24 (normal) and for those with an identified breakpoint (split *MYC*). (**b**) Expression data from 169 samples with any translocation detected by capture or FISH. Whisker plots show the 10–90 percentiles.

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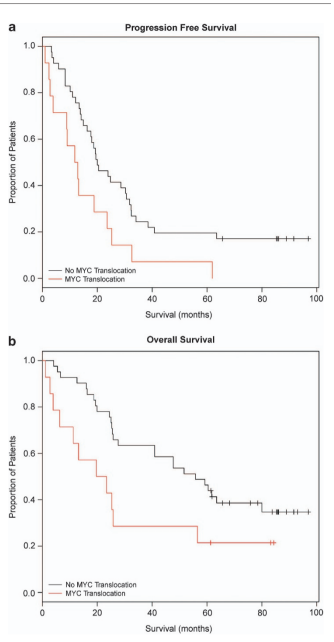


Figure 4. Progression-free survival (**a**) and Overall-free survival (**b**) in patients with a *MYC* breakpoint is significantly decreased compared with those without a breakpoint. PFS P = 0.032, OS P = 0.035. Data adjusted for confounding variables in Tables 4a, b and c.

and *CCND1*. The other partner genes identified here, which have no known function in myeloma or plasma cells, are likely also to be important in plasma cell development or myeloma disease as they have active superenhancers that are sequestered by *MYC*.

Some of the loci identified have active superenhancers up to 22 Mb away from the breakpoint. These distal superenhancers may be responsible for *MYC* overexpression, but as these results are based solely on data from the MM1.s cell line²⁴ it may be that in different myeloma cell types an enhancer closer to the breakpoint is active, and it is this as yet unidentified superenhancer that is located near *MYC* in this patient. This argument is exemplified by the lack of a superenhancer at the *IGK* locus in MM1.s and can be explained as this cell line expresses the lambda light chain and not the kappa light chain.³³ Similar studies involving a large cohort of myeloma cell lines would give further insight into the complex nature of superenhancers in this disease.

Table 4a. Clinical characteristics of MYC-translocated samples

Variable	Number of patients	P.	FS	OS		
		Months	P-value	Months	P-value	
MYC translocation	14	11.8	0.016	19.7	0.043	
No MYC translocation	41	20.0		55.8		
t(14;16)	9	9.0	0.006	11.2	0.095	
No t(14;16)	46	20.0		47.7		
Adverse IGH translocation	27	13.4	0.018	25.5	0.021	
No adverse IGH translocation	28	24.8		61.7		
ISS 1	7	61.9	0.013	NR	0.018	
ISS 2	15	23.9		60.4		
ISS 3	16	13.8		27.6		
Eligible for intensive pathway ^a	36	25.3	< 0.001	80.0	< 0.001	
Not eligible for intensive pathway ^a	19	11.8		23.4		

Abbreviations: OS, overall survival; PFS, progression-free survival. Univariate analysis—statistically significant. ^aPatients treated according to the intensive pathway received autologous stem cell transplantation, whereas patients not eligible entered the non-intensive pathway and were treated only with attenuated doses of chemotherapy.

Table 4b. Multivariate analysis of with an improved PFS	of variables	significantl	y associated
Variable	P-value	Hazard ratio	95% CI
Absence of adverse translocation	0.03	2.763	1.105-6.908
Absence of MYC translocation	0.035	3.094	1.081-8.856
Treatment on the intensive path	< 0.001	5.871	2.186-15.766

Table 4c. Multivariate analysis of with an improved OS	of variables	significantly	y associated
Variable	P-value	Hazard ratio	95% CI
Absence of adverse translocation	0.010	4.861	1.450–16.293
Absence of MYC translocation	0.032	4.077	1.125–14.785
Treatment on the intensive path	0.001	6.528	2.196-19.404

As the mechanism of action for *MYC* overexpression is through the juxtaposition of superenhancers specific to each disease, it is possible that this unifying mechanism can be therapeutically targeted. Given that it has been shown that superenhancers can be disrupted using BET-bromodomain inhibitors, such as JQ1,²⁴ it makes patients with *MYC* rearrangements good candidates for treatment with this class of drugs. We show here that patients identified with a *MYC* translocation have a poor PFS and OS compared with those with no rearrangement. If these patients could be identified in advance they may benefit from treatment with this class of targeted drugs.

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