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Citation	Journal Of Clinical Pathology, 2007, v. 60 n. 6, p. 664-669
Issued Date	2007
URL	http://hdl.handle.net/10722/57537
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ORIGINAL ARTICLE

Epigenetic dysregulation of the death-associated protein kinase/p14/HDM2/p53/Apaf-1 apoptosis pathway in multiple myeloma

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Aim: To study the role of gene promoter hypermethylation of the putative tumour suppressor genes involved in the death-associated protein (DAP) kinase/p14/HDM2/p53/Apaf-1 apoptosis pathway in multiple myeloma (MM).

Method: DNAs from 55 primary MM marrow samples and myeloma cell lines were analysed for aberrant promoter methylation of *DAP kinase*, *p14* and *Apaf-1* genes by methylation-specific polymerase chain reaction (MSP).

Result: In the methylated positive control, the sensitivity of M-MSP for DAP kinase was 1×10^{-3} . Aberrant hypermethylation of DAP kinase was found in 29/55 (52.7%) primary MM samples, whereas hypermethylation of p14 or Apaf-1 was undetectable in any of the samples tested. 5-Azacytidine treatment of two myeloma cell lines, WL2 and HS-Sultan, led to de-methylation and re-expression of DAP kinase, thereby confirming gene silencing associated with promoter hypermethylation. Hypermethylation of DAP kinase the overall survival.

Conclusion: Of the putative tumour suppressor genes in the DAP kinase/p14/HDM2/p53/Apaf-1 apoptosis pathway, only *DAP kinase* is frequently methylated in MM, which is associated with gene silencing and might be of prognostic significance. *p14* and *Apaf-1* were not methylated in MM.

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Accepted 22 May 2006

Multiple myeloma (MM) is a disease with a low proliferative rate. Plasma cells actively synthesising DNA are typically <1%, and increase only in advanced disease.^{1 2} However, neoplastic plasma cells are protected from apoptosis by interacting with the marrow microenvironment.^{1 2}

Death-associated protein (DAP) kinase is a proapoptotic calcium/calmodulin-regulated serine/threonine kinase with a multidomain structure. It participates in a wide array of apoptotic mechanisms initiated by interferon- α , tumour necrosis factor- α , activated Fas and detachment from the extracellular matrix.³ DAP kinase counteracts oncogene-induced transformation by activating p53 in a p19ARF-dependent manner, thereby providing an intrinsic p53-dependent apoptotic checkpoint that is turned on by oncogenes at the initial stages of transformation.³

 $p14^{ARF}$ is one of the two genes encoded by the INK4A/ARF locus at chromosome 9p21.⁴ Alternative first exons 1 α and 1 β , under the control of different promoters, specify the 5' ends of $p16^{INK4A}$ and $p14^{ARF}$, respectively. These alternative exons are spliced to the same splice acceptor site in exon 2, leading to translation in alternative reading frames. p14 physically associates with HDM2, which possesses E3 ubiquitin ligase activity. HDM2 is a negative regulator of p53, mediating its degradation in the proteosome. p14^{ARF} binding to HMD2 leads to inhibition of the E3 ubiquitin ligase activity, and localisation of HDM2 to the nucleus, precluding its interaction with p53.⁴ Therefore, by stabilising p53, p14^{ARF} acts as a tumour suppressor.

In normal cells, aberrant activation of a cellular oncogene may trigger an intrinsic tumour suppression mechanism, leading to apoptosis and thus eradication of the transformed cell.⁵ The DAP kinase/p14/HDM2/p53 pathway is one such mechanism, leading to the downstream activation of Apaf-1 and formation of the apoptosome.⁶ In MM, p53 is functional in the majority of cases, and mutation or deletion is rarely demonstrated.^{7 8} Therefore, other tumour suppressor components of this pathway may be potentially affected in the pathogenesis of MM, in which oncogene activation is a frequent event.^{1 2}

DNA methylation, catalysed by DNA methyltransferase, involves the addition of a methyl group to the carbon 5 position of the cytosine ring in the CpG dinucleotide, converting it to methylcytosine.^{9 10} In many cancers, the CpG islands of selected genes are aberrantly methylated (hypermethylated), resulting in transcriptional repression. This may serve as an alternative mechanism of gene inactivation.

We have demonstrated frequent epigenetic dysregulation of cell cycle control and Jak/STAT signalling in MM.^{11–14} However, as study of isolated genes or random assortment of genes precludes interpretation of the role of hypermethylation of putative tumour suppressor genes in a specific pathway, we investigated the role of aberrant gene promoter hypermethylation in disrupting the DAP kinase/p14/HDM2/p53/Apaf-1 pathway in MM. *p*53 was not investigated, as it does not harbour a CpG island in the promoter region (accession number: J04238).

MATERIALS AND METHODS

Patients, diagnosis and treatment

A total of 34 male and 21 female patients with MM, with a median age of 57 (25–87) years, were studied. The diagnosis of MM was based on standard criteria.¹⁵ Complete staging examination included bone marrow examination, skeletal survey, serum and urine protein electrophoresis, and serum immunoglobulin (IgG, IgA and IgM) levels. Monoclonal

Abbreviations: 5-AC, 5-azacytidine; DAP, death-associated protein; HSCT, haematopoietic stem cell transplantation; MM, multiple myeloma; MSP, methylation-specific polymerase chain reaction; OS, overall survival

Gene	Forward primer (5'–3')	Reverse primer (5'-3')	Tm/cycles	Reference
DAP kinase				
M-MSP	GGATAGTCGGAT	CCCTCCCAAA	63°C/35	Chan <i>et al</i> ¹⁹
	CGAGTTAACGTC	CGCCGA		(2002)
U-MSP	GGAGGATAGTTG	CAAATCCCTCCC		
	GATTGAGTTAATGTT	AAACACCAA		
P14				
M-MSP	GTGTTAAAG	AAAACCCTC	65°C/35	Esteller <i>et al</i> ¹⁸
	GGCGGCGTAGC	ACTCGCGACGA		(2000)
U-MSP	TTTTGGTGT	CACAAAAAC		
	TAAAGGGTGGTGTAGT	CCTCACTCACAACAA		
Apaf-1				
M-MSP*	TATTGCGATATTGTT	GAAACGTAACTAAA	64°C/35	Genebank
	TTAAATTCGA	CCTCAAAAACG		
U-MSP**	TATTGTGATATTGTT	CAAAACATAACTAA		AB070 829
	TTAAATTTGA	ACCTCAAAAACAC		

Primers for Apaf-1: M-MSP*, nucleotides 785–809; U-MSP**, nucleotides 938–963 in Genbank accession number: AB070829.

proteins were identified by electrophoresis, immunoelectrophoresis and immunofixation. There were 6 (10.9%) patients with Durie-Salmon stage I, 14 (25.5%) with stage II and 35 (63.6%) with stage III. Patients with stage I or asymptomatic disease did not receive chemotherapy until disease progression. Patients below 65 years of age were induced with three to six courses of vincristine, adriamycin and dexamethasone for initial myeloma cytoreduction. Those with a human leucocyte antigen-identical donor and below 50 years of age received myeloablative allogeneic haematopoietic stem cell transplantation (HSCT; n = 8), and those between 50 and 65 years of age (n = 10) received non-myeloablative HSCT. Patients without a suitable donor received autologous HSCT up to the age of 65 years (n = 11). Patients >65 years of age were given melphalan and prednisolone until they reached maximal response or the plateau phase. Patients progressing from the plateau phase received melphalan, cyclophosphamide, thalidomide, or vincristine, adriamycin and dexamethasone. Patients with local symptoms due to disease received local radiotherapy. The study was approved by the institutional review board of Queen Mary Hospital (Hong Kong, Hong Kong) and informed consent obtained.

Methylation-specific polymerase chain reaction

DNA was extracted from 55 myeloma marrow samples at diagnosis and from myeloma cell lines (WL2 and HS-Sultan) by the standard method. Methylation-specific polymerase chain reaction (MSP) for aberrant gene promoter methylation was performed as described previously.^{16 17} Treatment of DNA with bisulphite for conversion of unmethylated cytosine to uracil (but not affecting methylated cytosine) was performed with a commercially available kit (CpGenome DNA modification kit, Intergen, New York, New York, USA). The genes p14, DAP kinase and Apaf-1 were tested. Primers for the methylated (M-MSP) and unmethylated (U-MSP) promoters are shown in table 1.18 19 DNA from 19 normal bone marrow donors was used as negative control, while methylated control DNA (CpGenome Universal Methylated DNA, Intergen) was used as positive control in all experiments. MSP was performed in a thermal cycler (9700, ABI Biosystems, Foster City, California, USA) with the following cycling conditions: 95°C for 4 min, 35 cycles of 95°C for 45 s, specific annealing temperature for 30 s, 72°C for 30 s and a final extension of 10 min at 72°C. The MSP mixture

contained 50 ng of bisulphite-treated DNA, 0.2 mM dNTPs, 2 mM MgCl₂, 10 pmol of each primer, $1 \times$ PCR buffer and 2.5 U of AmpliTaq Gold DNA Polymerase (ABI Biosystems) in a final volume of 50 µl. In all, 10 µl of PCR products were loaded onto 6% non-denaturing polyacrylamide gels, electrophoresed and visualised under ultraviolet light after staining with ethidium bromide.

Sensitivity of MSP

One microgram of DNA of the methylated positive control (Intergen) was diluted serially (10-fold) in normal marrow DNA, modified by bisulphite and then amplified by MSP.

5-Azacytidine treatment of the WL2 and HS-Sultan cell lines

The human myeloma cell lines WL2 and HS-Sultan were cultured in Rosewell Park Memorial Institute medium supplemented with 10% fetal calf serum. For treatment with 5azacytidine (5-AC), cells from myeloma cell lines, WL2 (kindly donated by Dr Andrew Zannettino, Myeloma and Mesenchymal Research Laboratory, Division of Haematology, Institute of Medical and Veterinary Science, Adelaide, Australia) and HS-Sultan (purchased from ATCC), were seeded in six-well plates at a density of 10⁶ cells/ml, and cultured with 5 and 0.5 μ M of 5-AC (Sigma, St Louis, Missouri, USA) for 5 days, with fresh medium containing 5-AC replenished every 2 days. Cells on days 0 and 5 of 5-AC treatment were harvested.

Reverse transcription-PCR

One microgram of total cellular RNA was reverse transcribed with M-MLV reverse transcriptase (Life Technologies, Rockville, Maryland, USA). cDNA was amplified by PCR with *DAP kinase*specific primers (forward: 5'-GATAGAAATGTCCCCAAA-CCTCG-3' and reverse: 5'-TCTTCTTGGATCCTTGACCAGAA-3', which spanned nucleotides 781–1124, Genbank accession number: X76104). RNA that was not reverse transcribed served as control. Reverse transcription-PCR for glyceraldehyde 3phosphate dehydrogenase (forward: 5'-GAAGGTGAAGGTC-GGAGTC-3' and reverse: 5'-GAAGATGGTGATGGGATTC-3') was performed to confirm RNA integrity. PCR conditions for *DAP kinase* were 35 cycles of 95°C for 30 s, 58°C for 60 s and 72°C for 60 s, and those for glyceraldehyde 3-phosphate dehydrogenase 30 cycles of 95°C for 30 s, 60°C for 30 s and



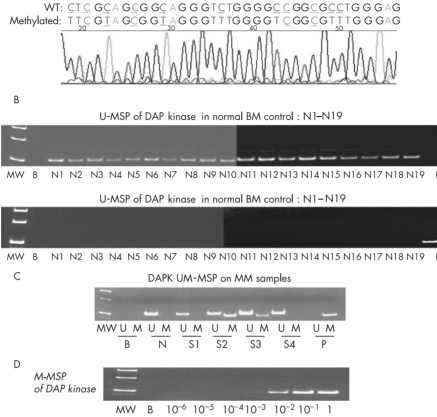


Figure 1 Methylation-specific polymerase chain reaction (MSP) of death-associated protein (*DAP*) kinase. (A) Sequence analysis of the methylated MSP (M-MSP) product showed the expected changes after bisulphite treatment. Wild-type (WT) cytosine (C) residues that remained unchanged in methylated CpG were coloured blue and underlined, while those that were changed to thymidine (T) were coloured red and underlined. (B) Unmethylated MSP (U-MSP) and M-MSP showing the expected results, with all 19 normal controls (N1–N19) amplifying in U-MSP but not in M-MSP, and the methylated positive control (P) amplifying only in M-MSP but not in U-MSP. MW, molecular weight marker; B, reagent blank; P, positive control of methylated DNA; N1–N19, normal marrow DNA. (C) For the MM samples, samples 2 and 3 (S2, S3) showed DAP kinase hypermethylation. B, reagent blank; P, positive control of methylated DNA; S1–4, MM marrow DNA. (D) M-MSP of the serially diluted methylated positive control DNA, showing a sensitivity of 1 in 10³. BM, bone marrow; MM, multiple myeloma.

72°C for 30 s. PCR products were electrophoresed on 6% nondenaturing polyacrylamide gels, stained with ethidium bromide and visualised under UV illumination. The identity of the PCR products was confirmed by DNA sequencing.

Statistical analysis

Association between *DAP kinase* hypermethylation and other clinical parameters (age, sex, Durie–Salmon stage, paraprotein subtypes) was studied by the χ^2 test (for categorical variables) or Student's t test (for continuous variables). Overall survival (OS) was measured from the date of diagnosis to the date of last follow-up or death. Survival was estimated by the Kaplan–Meier method and compared by the log-rank test. All p values were two-sided.

RESULTS

Methylation-specific polymerase chain reaction

None of the 19 normal control marrows showed aberrant methylation of *p14*, *DAP kinase* and *Apaf-1* (fig 1B, 2A,B). The positive and negative controls showed the expected MSP results (normal DNA, U-MSP positive/M-MSP negative; methylated DNA: U-MSP negative/M-MSP positive). Sequencing of the MSP products showed the expected nucleotide changes after bisulphite treatment (fig 1A, 2A,B). For the primary MM samples, *DAP kinase* was methylated in 29 cases (52.7%; fig 1C),

whereas *p14* and *Apaf-1* were not methylated in any of the samples (fig 2A,B).

Sensitivity of DAP kinase M-MSP

For the positive control, the sensitivity of M-MSP was 1×10^{-3} for *DAP kinase* (fig 1D).

5-AC treatment of WL2 myeloma cell line

DAP kinase was completely methylated and consequently not expressed in WL2 and HS-Sultan also. 5-AC treatment of both cell lines resulted in partial demethylation of *DAP kinase*, as evidenced by positive U-MSP amplification and gene expression on day 5 (fig 3).

Statistical analysis

DAP kinase hypermethylation was not associated with age (p = 0.16), sex (p = 0.59), Durie–Salmon stage (p = 0.76) and paraprotein subtypes (p = 0.37; table 2). However, *DAP kinase* hypermethylation was significantly associated with an inferior median OS (methylated 34 months vs unmethylated 80 months, p = 0.013; fig 4).

DISCUSSION

We showed that *DAP kinase* was frequently methylated in MM samples, whereas *p14* and *Apaf-1* were not. Therefore, abroga-

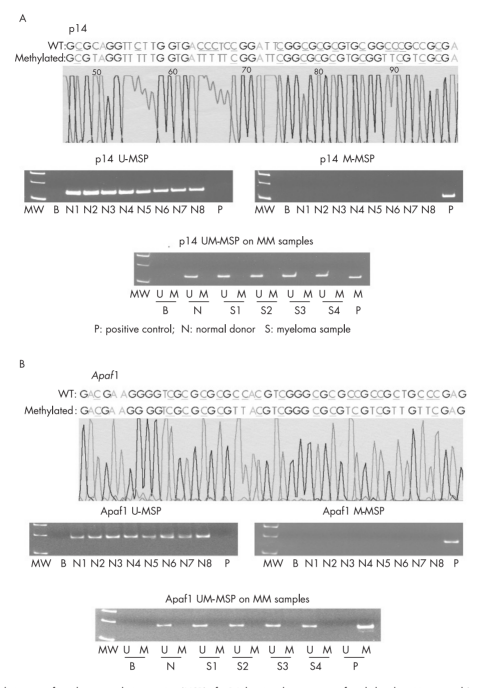


Figure 2 (A) Methylation-specific polymerase chain reaction (MSP) of *p14* showing the sequence of methylated positive control (upper panel: wild-type (WT) cytosine (C) residues that remained unchanged in methylated CpG are coloured blue and underlined, while those that were changed to thymidine (T) are coloured red and underlined), absence of *p14* hypermethylation in normal control DNA (middle panel: presence of unmethylated MSP (U-MSP) but absence of methylated MSP (M-MSP) amplification in normal control DNA) and primary myeloma marrow samples (lower panel: presence of U-MSP) but absence of M-MSP amplification in normal control DNA). (B) MSP of *Apaf-1* showing the sequence of methylated positive control (upper panel: WT cytosine (C) residues that remained unchanged in methylated CpG are coloured blue and underlined, while those that were changed to thymidine (T) are coloured red and underlined), absence of p14 hypermethylated CpG are coloured blue and underlined, while those that were changed to thymidine (T) are coloured red and underlined), absence of p14 hypermethylated CpG are coloured blue and underlined, while those that were changed to thymidine (T) are coloured red and underlined), absence of p14 hypermethylated CpG are coloured blue and underlined. while those that were changed to thymidine (T) are coloured red and underlined), absence of p14 hypermethylation in normal control DNA (middle panel: presence of U-MSP but absence of M-MSP amplification in normal control DNA) and primary myeloma marrow samples (lower panel: presence of U-MSP but absence of M-MSP amplification in normal control DNA). B, reagent blank; M, M-MSP; MM, multiple myeloma; MW, molecular weight; N1–N8, normal marrow DNA; S1–S4, myeloma marrow samples; U, U-MSP

tion of the putative tumour suppressor genes involved in the *DAP kinase*-related apoptotic pathway in some cases might collaborate with other antiapoptotic signals from the microenvironment to enhance the survival of neoplastic plasma cells.^{1 2} Moreover, as concomitant abrogation of multiple tumour suppressors of this apoptosis pathway does not confer additional survival benefit, our findings suggested that hypermethylation of tumour suppressor components of this apoptosis pathway appears to be mutually exclusive.

Despite the sensitivity of *DAP kinase* MSP being only 1 in 10^3 , the methylation status of the neoplastic plasma cells should preferably be studied by CD138-sorted plasma cells.

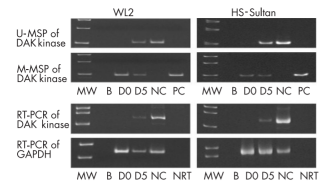


Figure 3 5-Azacytidine (5-AC) treatment of WL2 and HS-Sultan. Deathassociated protein (*DAP*) kinase was unmethylated in the normal marrow control (NC), which was associated with expression of *DAP kinase* transcript. On day 0 (D 0), *DAP kinase* was completely methylated in WL2 and HS-Sultan, with a corresponding absence of *DAP kinase* transcript. At 5 days after treatment (D 5), partial de-methylation of *DAP kinase* occurred, leading to positive amplification in unmethylated methyl-specific polymerase chain reaction (U-MSP), and expression of the *DAP kinase* transcript. B, blank; MW, molecular weight marker; NC, normal marrow control; NRT, non-reagent control; PC, positive control of methylated DNA.

DAP					
kinase	Methylated	Unmethylated	p Value		
Number	29	26			
Median	60.0	54.5	0.16		
age					
Sex			0.59		
Male	19	15			
Female	10	11			
g subtype			0.37		
lgG	19	15			
lgA	7	4			
lgD	0	1			
Light chain	3	6			
Durie–Salmon			0.76		
stage					
≼∥	11	9			
III	18	17			

In the myeloma cell lines WL2 and HS-Sultan, *DAP kinase* hypermethylation was associated with the absence of transcript, which was re-expressed after 5-AC treatment. As 5-AC is a DNA methyltransferase inhibitor, and thus a demethylating agent, re-expression of the transcript after 5-AC treatment confirmed gene silencing-associated gene promoter hypermethylation.

In this study, *DAP kinase* hypermethylation was associated with an inferior OS but not with Durie–Salmon stage. The latter could be due to the retrospective nature of the study, the small number of patients and the heterogeneity of treatment. For instance, 29 (52.7%) of our patients have undergone bone marrow transplantation. Therefore, the prognostic value of *DAP kinase* methylation needs verification in future prospective clinical trials with a larger number of patients receiving uniform treatment.

The oncogenic role of *DAP kinase/p14/Apaf-1* has been demonstrated in previous studies. DAP kinase expression was shown to suppress the metastatic potential of tumour cells in

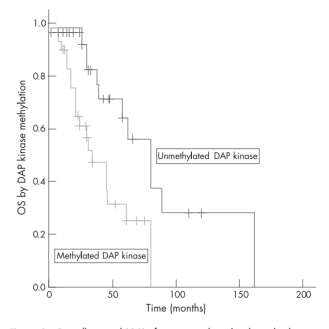


Figure 4 Overall survival (OS) of patients with and without deathassociated protein (*DAP*) *kinase* hypermethylation. *DAP kinase* hypermethylation was associated with an inferior OS.

Take-home messages

- Death-associated protein (DAP) kinase, a tumour suppressor gene regulating p53-mediated apoptosis, is potentially important in myeloma pathogenesis because of its frequent methylation.
- DAP kinase hypermethylation might be of prognostic significance.

animal models.²¹ Modest frequency of DAP kinase hypermethylation (10-20%) has been identified in lung, colon, and head and neck cancers.²² In contrast, frequent (66–72%) DAP kinase hypermethylation was demonstrated in B-cell lymphoma and MM.^{23 24} Moreover, DAP kinase has been shown to be methylated in patients with monoclonal gammopathy of undetermined significance at a frequency comparable with that with MM.25 p14 hypermethylation was found in 10-30% of renal, gastric and colorectal cancers.²² In haematological malignancies, p14 hypermethylation was present in 40% of chronic myeloid leukaemia cases in acceleration,²⁶ but in only 8% of acute lymphoblastic leukaemia cases,²⁷ and in none of the therapy-related myelodysplastic syndrome/acute myeloid leukaemia cases.28 In MM, one previous study showed p14 hypermethylation in 9% of the patients.²⁹ The infrequent *p14* hypermethylation was consistent with results from another study, which showed p14 expression in MM.³⁰ Furthermore, as the *p14/p16* locus was not found to be deleted in MM, ³¹ *p14* does not appear to be targeted in MM. Finally, while hypermethylation of Apaf-1 has been detected in melanoma,³² and recently in leukaemia,²⁶ this is the first report demonstrating the absence of Apaf-1 hypermethylation in MM.

In conclusion, of the putative tumour suppressor genes in the DAP kinase/p14/HDM2/p53/Apaf-1 apoptosis pathway, only *DAP kinase* is frequently methylated in MM, which is associated with gene silencing. *p14* and *Apaf-1* were not methylated in

Hypermethylation of p14, DAPK and Apaf-1 in multiple myeloma

MM. The high sensitivity of DAP kinase MSP might be a useful marker for monitoring of minimal residual disease.

ACKNOWLEDGEMENTS

We thank Dr R Robert Orlowski (Department of Medicine, Division of Hematology/Oncology, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina, USA) for his critical review of the manuscript.

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Competing interests: None declared.

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