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Jansen et al.

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(54) **METHOD FOR DIAGNOSING A PERSON HAVING MULTIPLE SCLEROSIS**

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(51) **Int. Cl.**

C12Q 1/68 (2006.01)

G01N 33/53 (2006.01)

C12P 19/34 (2006.01)

(52) **U.S. Cl.** **435/6**; 435/7.1; 435/91.2

(58) **Field of Classification Search** 435/6, 435/7.1, 91.2

See application file for complete search history.

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(57) **ABSTRACT**

Described is a method for diagnosing a person having multiple sclerosis (MS) or being at risk of developing MS, comprising the following steps:

providing a sample of a body fluid or tissue from said person, said sample containing at least one of the wild type SCF-Apoptosis-Response Gene- (wt-SARG-1-) protein and nucleic acids encoding wt-SARG-1, if taken from a person not having MS or a risk of acquiring MS,

detecting the presence of wt-SARG-1-protein or nucleic acids encoding wt-SARG-1 in said sample and

diagnosing MS or a risk of acquiring MS, if wt-SARG-1-protein or nucleic acids encoding wt-SARG-1 are not present in said sample.

11 Claims, 23 Drawing Sheets

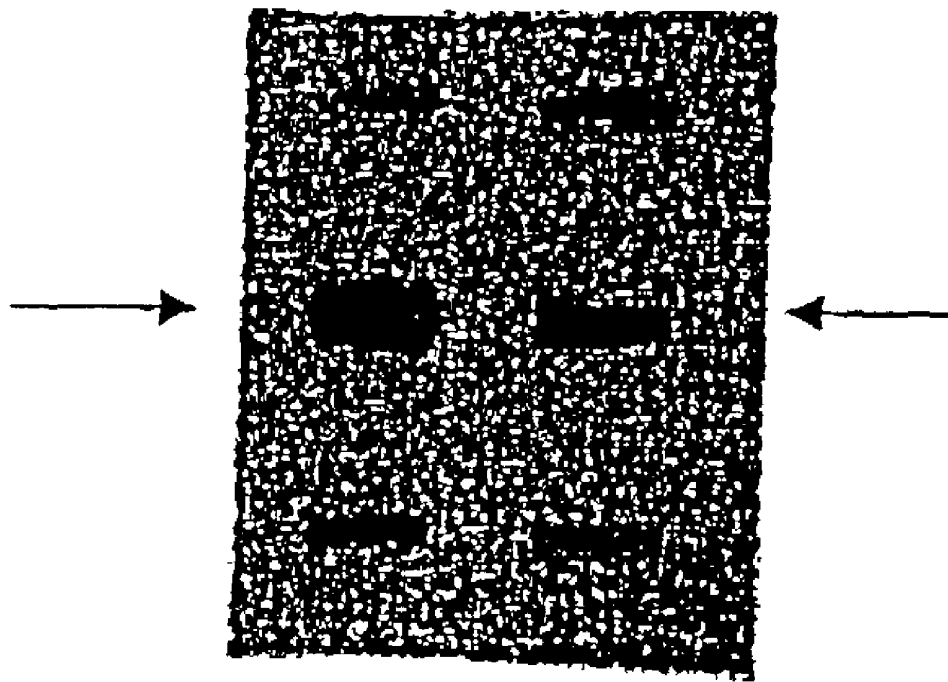
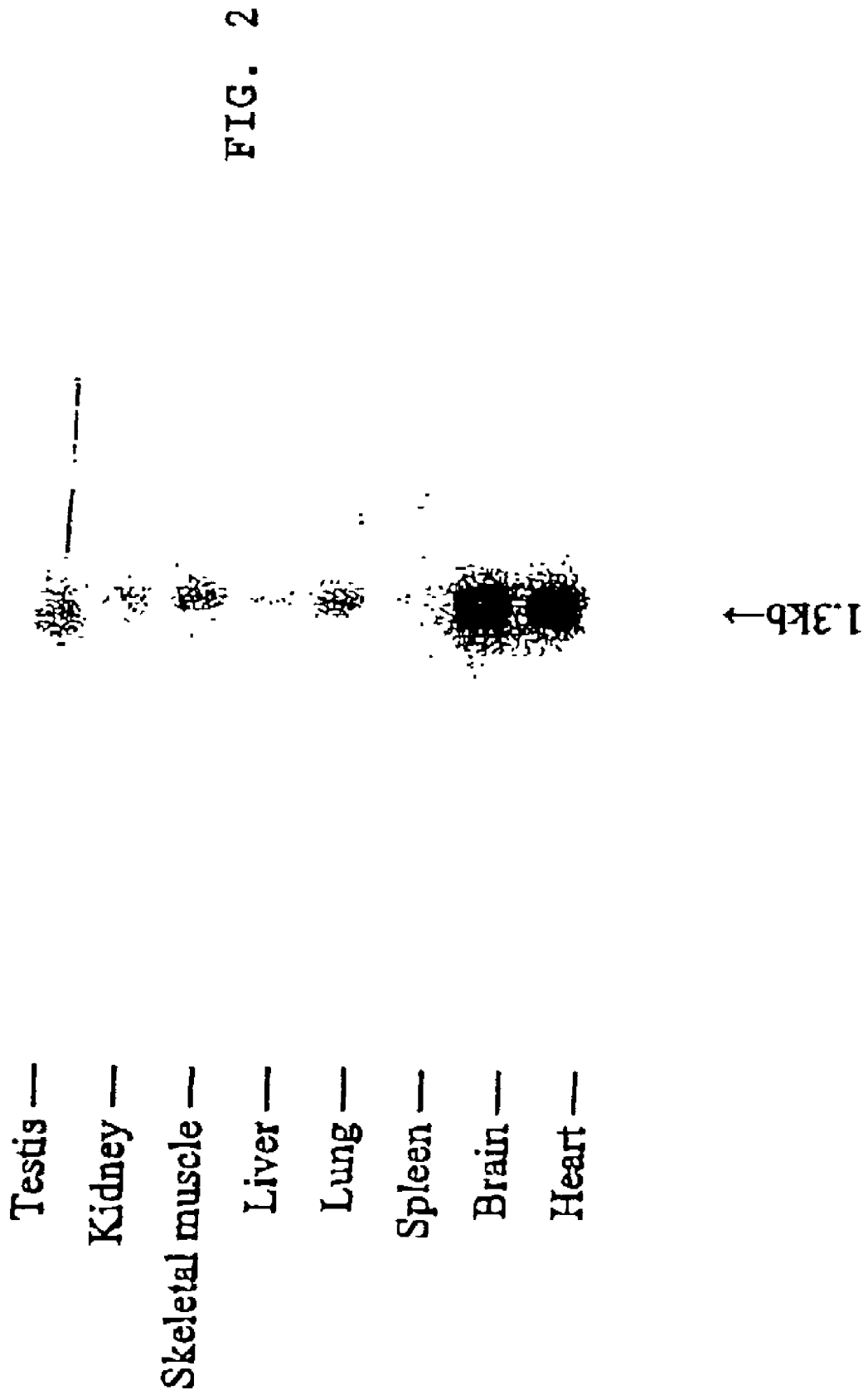


FIG. 1



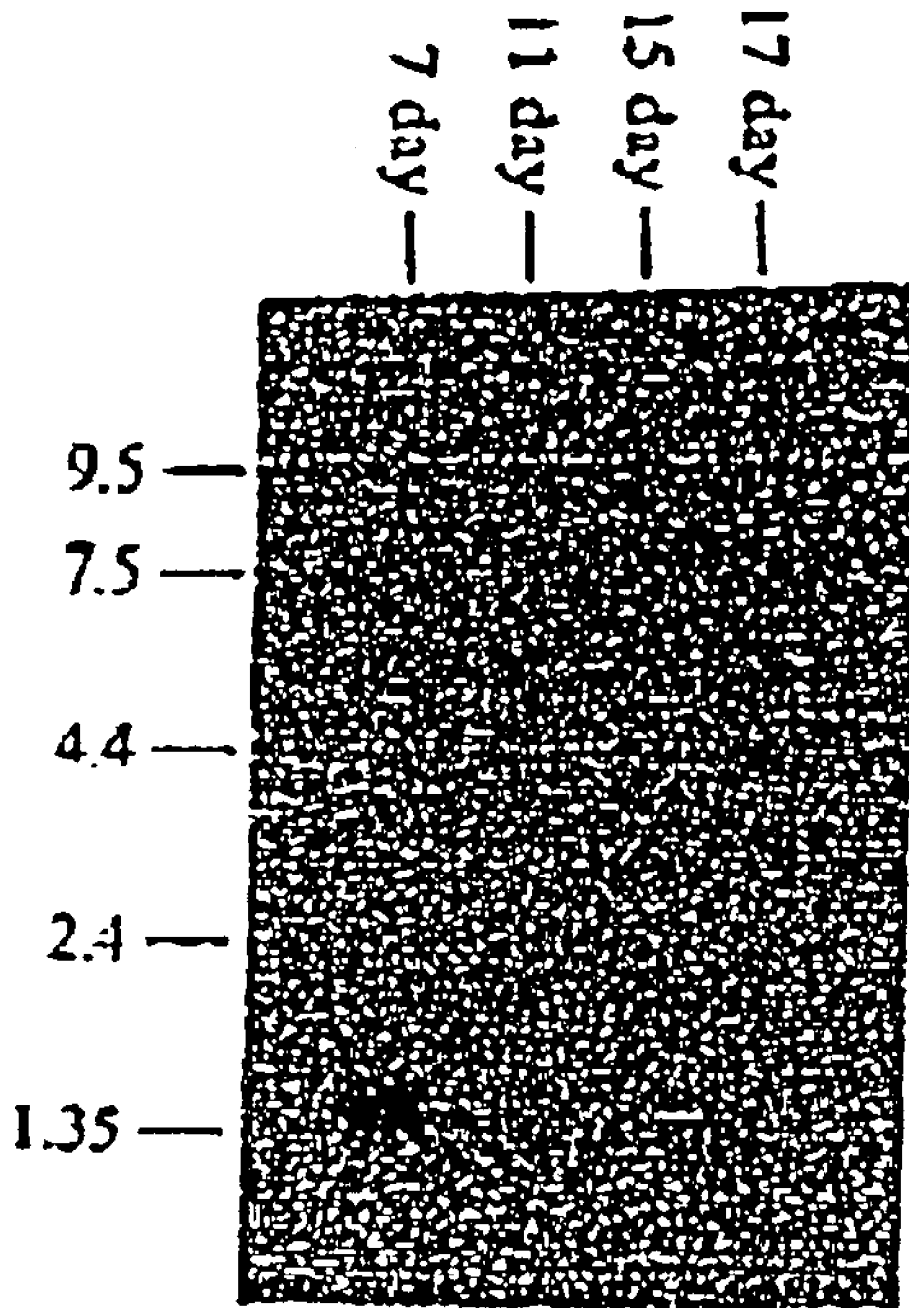


FIG. 3

FIG. 4

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>rSARG, 1037 bases
CCAGACTGGAAGCGAAGGCTGTGTTGCTGGGATGCCAGCTGCCGAGGGGC
TGCTTAAGCCTTGGCCCCACTACTTTCTGTTTCAGCCCCACTTCTGTGC
GTGTCTTACTCCATTACCCCCAGGGGCTGACATGGACCCAAATCCACGAG
CAGCCCTGGAGCGGCAGCAGCTGCGTCTCAGGGAGCGGCAGAAGTTCTTC
GAGGACATTTTACAGCCAGAGACAGAGTTTGTTCCTCCCTATCCCATCT
GCATCTCGAGTCACAAAGACCCCCATAGGTAGCATCTCCTCGATGGAAG
TGAATGTGGACACACTGGAGCAGGTGGAATTTATTGACCTTGGGATCAG
GATGGAGCAGATGTGTTCTTACCTTGTGAGGATTCTCCTCCAACCCCCA
GAGGTCTGGAGTGGATGACCACCCAGAGGAGCTGAGCCTGCTGGTACCCA
CGTCAGACAGGACCACATCCCGGACCTCCTCCTTGTCTCTGACTCCTCC
AACCTGCGCAGTCCAAATCCAAGTGATGGGGGAGGAGACACTCCCTTGGC
ACAGTCTGACGAGGAGGATGGGGACGGTGGAGGGGAGAACCTGGACCTT
GCAGTAGCAGAGGCCCTTACAAACTGAGCGATCTGGCTGTCTCCATG
GAGAGGAGACCTTAGGTCACCAGAGCACTCTGGAGAAGACCTGACACTT
TACTTACATCAGCACCAAAGGGAGGGAAAGGATGGTGGATGGTGTGCCTGA
GAGTTAGCCTCCCGCTTTACTGATAACGCTGTCTCTGCTGCCACGCCCC
ACAGTGTCTTCTTCTGAGGTAGGACTTCCAAGTGAGACTCTCGAAGGTGA
GGTGGGACAAGATGCCACTGTTTTCTTACTCCCTCCTGCCCCAAATGA
TCCTGTAGTCTCCACTAGTCTCCTAAGCCAGTGTCTCTGAGGGAAAGTT
CTGAGGAGTCCACTTTGCAGTTATCCTGCCTCTATAAGTCTTTCTGGG
AACGGATATGGTATAAATAATAAATAATACTGTACC
>mSARG, 1029 bases
CCAGACCGGAAGCGAGGCTGTGTTGCTGGGATCCAACGCCGGCGCTGCTC
GCTCCCAGCCCCCGCCGCGCTGTGCGGGAGCGCACCCAGGGAGCCAGC
GGGGCGGGGGCGCTGCAGGGGCTGACATGGACCCAAATCCGAGAGCAGCC
CTGGAGCGCAACAGCTGCGGCTCCGGGAGAGGCAGAAGTTCTTTGAGGA
CATTTTACAGCCAGAGACAGAGTTTGTCTTCCCTGTCCCATCTGCACC
TGGAGTCACAAAGACCCCCATAGGTAGCATCTCGTCTATGGAAGTGAAT
GTGGACACACTGGAGCAAGTGGAGTTTATTGATCTTGGGATCAGGATGG
AGCAGATGTGTTCTTGCCTTGTGAGGAGTCTCGCCAGCTCCCCAGATGT
CTGGAGTGGATGACCATCCAGAGGAGCTGAGCCTGCTGGTACCCACGTCT
GACAGGACCACATCCCGGACCTCCTCCTTGTCTCTGACTCCTCCAACCT
GCGCAGTCCAAATCCAAGTGATGGGGGAGGAGACACTCCTTGGCACAGT
CTGATGAGGAGGACGGGGATGACGGAGGGGACAGCCTGGACCCTGCAGC
TAGCAGTGGGCTCGTACAGACTGACCAGCCCGGCTGTCTCCATGGAAA
GGAGACCTTAGGCCAGCAGAGCCTGGAGAAGACCTGACACTTTCTTACT
TCAGCAACAAAGGGAGGGAAAGGATGGTGGATGGTGTGCTGAGAGTTAGC
TCCCTGTCTTACCGTAAAGCTATCCTGCTGCCACGCCCCACAGTGTCT
TTCTTCTGAGGTAGGACTTCCAAGTGAGACTTGGAGGTGAGGTGGGAC
AAGACGACAGTGTCTTCTTAGTCCCCTCCTGCCCCAGATGATCCTGTTG
TCTTCCACAGAGTCTCCTAAGCCAGTGTCTCTGAGGGGATGTTCTGAGGA
GTTCCACTTTCCAGTTATCCTGCCTCTATAAGTCTTTTGGGAACAGGAT
ATGGTATAAATAATAAATAATAATAATACC
>hSARG, 1082 bases
CCAGGCGGAGCCAGGGGCCACTGTTGGGATGCTGGCTGCAGTGGGGC
GCCCAAGCCAGTCCCCTCTGTCTTCTCTTTCGACTTTGCAGCTGTAC
TTGTTTTGCTCCTTACCCGAGGAGCTGACATGGACCCAAATCCTCGGG
CCGCCCTGGAGCGCCAGCAGCTCCGCTTCGGGAGCGGCAAAATTTCTTC
GAGGACATTTTACAGCCAGAGACAGAGTTTGTCTTCTCTGTCCCATCT
GCATCTCGAGTCGCAGAGACCCCCATAGGTAGTATCTCATCCATGGAAG
TGAATGTGGACACACTGGAGCAAGTAGAATTTATTGACCTTGGGGACCCG
GATGCAGCAGATGTGTTCTTGCCTTGGGAAGATCCTCCACCAACCCCCA
GTCTGTGGGGTGGACAACCATTGGAGGAGCTGAGCCTGCCGGTGCCTA
CATCAGACAGGACCACATCTAGGACCTCCTCCTCCTCCTCCGACTCC
TCCACCAACCTGCATAGCCAAATCCAAGTGATGATGGAGCAGATACGCC
CTTGGCACAGTCCGGATGAAGAGGAGGAAAGGGGTGATGCAGGGGCAGAGC
CTGGACCTGCAGCTAGCAGTGGGCCCTGCCTACAGACTGACCAGCTG
GCTATTCTCCATGAGACCACAGGCCAGCCAGAGCCTGTCCGGAGAAG
ACCAGACTCTTACTTGCAGTAGGCACCAGAGGTGGGAAGGATGGTGGGA
TTGTGTACCTTTAAGAAATTAACCCCTCCTGCTTTACTGCTAATTTTT
TCCTGCTGCAACCCTCCCACCAGTTTTTGGCTTACTCCTGAGATATGATT
```

FIG. 4 (Fortsetzung)

TGCAAATGAGGAGAGAGAAGATGAGGTTGGACAAGATGCCACTGCTTTTC
TTAGCACTCTTCCCTCCCCTAAACCATCCCGTAGTCTTCTAATACAGTCT
CTCAGACAAGTGTCTCTAGATGGATGTGAACTCCTTAACTCATCAAGTAA
GGTGGTACTCAAGCCATGCTGCCTCCTTACATCCTTTTTGGAACAGAGCA
CGGTATAATAATAAACTAATAATAATATGCC

FIG. 5

rSARG 1 **CCAGCTTGGAGCGTAGGCTGTGTGCTGGGTGCCAGCTGCCAAGGGCTGCTTAGCC**
mSARG 1 **CCAGCTTGGAGCGTAGGCTGTGTGCTGGGTGCCAGCTGCCAAGGGCTGCTTAGCC**
hSARG 1 **CCAGCTTGGAGCGTAGGCTGTGTGCTGGGTGCCAGCTGCCAAGGGCTGCTTAGCC**

rSARG 61 **TTGGCCCCACTACTTTCTGTTTCAGCCACACTTGGTGGGTCCTTACTCCGATACCC**
mSARG 56 **TTGGCCCCACTACTTTCTGTTTCAGCCACACTTGGTGGGTCCTTACTCCGATACCC**
hSARG 61 **TTGGCCCCACTACTTTCTGTTTCAGCCACACTTGGTGGGTCCTTACTCCGATACCC**

rSARG 120 **CCAGGGGCTGACATGGACCCAAATCCAGGACGCCCCGGAGCCGACGAGCTGCCATC**
mSARG 116 **CCAGGGGCTGACATGGACCCAAATCCAGGACGCCCCGGAGCCGACGAGCTGCCATC**
hSARG 120 **CCAGGGGCTGACATGGACCCAAATCCAGGACGCCCCGGAGCCGACGAGCTGCCATC**

rSARG 180 **ACGGAGCGGCGAGAGTTCTTCGAGGCAATTTTACAGCCGAGGACAGGATTTGCTTCCCC**
mSARG 175 **ACGGAGCGGCGAGAGTTCTTCGAGGCAATTTTACAGCCGAGGACAGGATTTGCTTCCCC**
hSARG 180 **ACGGAGCGGCGAGAGTTCTTCGAGGCAATTTTACAGCCGAGGACAGGATTTGCTTCCCC**

rSARG 240 **CTATCCCATCTGCATCTCGAGTCACAAAGACCCCCCATAGGTAGCATCTCTCTGATGGAA**
mSARG 235 **CTATCCCATCTGCATCTCGAGTCACAAAGACCCCCCATAGGTAGCATCTCTCTGATGGAA**
hSARG 240 **CTATCCCATCTGCATCTCGAGTCACAAAGACCCCCCATAGGTAGCATCTCTCTGATGGAA**

rSARG 300 **CTGAATGTGGACACACTGGAGCAAGTGGAGTTTATTGATCTTGGCCATCAGGATGGAGCA**
mSARG 295 **CTGAATGTGGACACACTGGAGCAAGTGGAGTTTATTGATCTTGGCCATCAGGATGGAGCA**
hSARG 300 **CTGAATGTGGACACACTGGAGCAAGTGGAGTTTATTGATCTTGGCCATCAGGATGGAGCA**

rSARG 360 **CATGTTCTTACCTTGTGAGGATTCCTCCACTCCCGAGCTTCTGGAGTGGATGAC**
mSARG 355 **CATGTTCTTACCTTGTGAGGATTCCTCCACTCCCGAGCTTCTGGAGTGGATGAC**
hSARG 360 **CATGTTCTTACCTTGTGAGGATTCCTCCACTCCCGAGCTTCTGGAGTGGATGAC**

rSARG 420 **CAACAGAGGAGCTGAGCCTGCTGGTACCACGTCAGACAGGACCAACTCCGGGACCTCC**
mSARG 415 **CAACAGAGGAGCTGAGCCTGCTGGTACCACGTCAGACAGGACCAACTCCGGGACCTCC**
hSARG 420 **CAACAGAGGAGCTGAGCCTGCTGGTACCACGTCAGACAGGACCAACTCCGGGACCTCC**

rSARG 480 **TCCTTGTCTCTCTTACTCCTCCATACCTGCCAGTCCAAATCCAGTGTATGGGGGA**
mSARG 475 **TCCTTGTCTCTCTTACTCCTCCATACCTGCCAGTCCAAATCCAGTGTATGGGGGA**
hSARG 480 **TCCTTGTCTCTCTTACTCCTCCATACCTGCCAGTCCAAATCCAGTGTATGGGGGA**

rSARG 534 **GGAGACTTCCCTTGGCAGGTCCTTGGAGGATGGGGAGCTGGAGGGGCGAGG**
mSARG 529 **GGAGACTTCCCTTGGCAGGTCCTTGGAGGATGGGGAGCTGGAGGGGCGAGG**
hSARG 540 **GGAGACTTCCCTTGGCAGGTCCTTGGAGGATGGGGAGCTGGAGGGGCGAGG**

rSARG 591 **CCGGACCTTGGAGCTAGCAGAGCCCTCTTACAACTGAGCGATTTGGCTGTTCTC**
mSARG 586 **CCGGACCTTGGAGCTAGCAGAGCCCTCTTACAACTGAGCGATTTGGCTGTTCTC**
hSARG 600 **CCGGAGCTTGGAGCTAGCAGAGCCCTCTTACAACTGAGCGATTTGGCTGTTCTC**

rSARG 647 **CATGGAGAGGAGACCTTAGGTCCTTCAGAGGACCTTGGAGAGACCTTGGAGAGGACCTTACT**
mSARG 643 **CATGGAGAGGAGACCTTAGGTCCTTCAGAGGACCTTGGAGAGACCTTGGAGAGGACCTTACT**
hSARG 660 **CATGGAGAGGAGACCTTAGGTCCTTCAGAGGACCTTGGAGAGACCTTGGAGAGGACCTTACT**

CODING

rSARG-coding

atggaccctccacgagcagccctggagcggcagcagctgcgtctcagggagcggcagaagtctctcagggac
atTTTAcagccagagacagagtttgttttccccctatcccatctgcctctcaggtcaaaagacccccataggt
agcatctcctcgatggaagtgaatgtggacacactggagcaggtggaatttatgaccttgccgatcaggatgga
gcagatgtgttcttaccttgtgaggattctcctccaaclcccagaggtctggagtggatgaccaccagaggag
ctgagcctgctggtaccacgtcagacaggaccacateccggacctcctccttgtcctctgactcctccaacctg
cgcagtccaaatccaagtgatgggggaggagacactcccttggcacagctctgacgaggaggatggggacggtgga
ggggcagaacctggaccttgcagctag
477bp

MDPNPRAALERQQLRLRERQKFFEDILOPETEFVFPPLSHLHLESQRPPIGSISMEVNVDTLEQVEFIDLADQDG
ADVFLPCEDSPTTQRSVDDHPEELSLLVPTSDRTTSRTSSLSDDSSNLRS PNPSSDGGGDTPLAQSDEEDGDDG
GAEPGPCS
158 amino acids

mSARG-coding

ATGGACCCAAATCCGAGAGCAGCCCTGGAGCGCCAACAGCTGCCGCTCCGGGAGAGGCAGAAGTCTTT
GAGGACATTTTACAGCCAGAGACAGAGTTTGTCTTCCCCTGTCCCATCTGCACCTGGAGTCACAAAGA
CCCCCATAGGTAGCATCTCGTCTATGGAAGTGAATGTGGACACACTGGAGCAAGTGGAGTTTATTGAT
CTTGCGGATCAGGATGGAGCAGATGTGTTCTTGCCTTGTGAGGAGTCTCGCCAGCTCCCCAGATGTCT
GGAGTGGATGACCATCCAGAGGAGCTGAGCCTGCTGGTACCCACGCTCTGACAGGACCACATCCCGGACC
TCTCCTTGTCTCTGACTCCTCCAACCTGCCGAGTCCAAATCCAAGTGATGGGGGAGGAGACACTCCC
TTGGCACAGTCTGATGAGGAGGACGGGGATGACGGAGGGGCAGACCTGGACCCTGCAGCTAG
477bp

MDPNPRAALERQQLRLRERQKFFEDILOPETEFVFPPLSHLHLESQRPPIGSISMEVNVDTLEQVEFIDLADQDG
ADVFLPCEESSPAPQMSGVDDHPEELSLLVPTSDRTTSRTSSLSDDSSNLRS PNPSSDGGGDTPLAQSDEEDGDDG
GAEPGPCS
158 amino acids

hSARG-coding

ATGGACCCAAATCCTCGGGCCGCCCTGGAGCGCCAGCAGCTCCGCCTTCGGGAGCGGCACAAAATCTTTCGAGGAC
ATTTACAGCCAGAGACAGAGTTTGTCTTCCCTCTGTCCCATCTGCATCTCGAGTCGAGAGACCCCATAGGT
AGTATCTCATCCATGGAAGTGAATGTGGACACACTGGAGCAAGTAGAACTTATTGACCTTGGGGACCCGGATGCA
GCAGATGTGTTCTTGCCTTGGGAAGATCCTCCACCAACCCCCAGTCTGCTGGGGTGGACAACCATTTGGAGGAG
CTGAGCCTGCGGGTGCTTACATCAGACAGGACCACATCTAGGACCTCCTCCTCCTCCTCCTCCGACTCCTCCACC
AACCTGCATAGCCCAATCCAAGTGATGATGGAGCAGATACGCCCTTGGCACAGTCCGATGAAGAGGAGGAAAGG
GGTGATGGAGGGCCAGACCCTGGAGCCTGCAGCTAG
486bp

MDPNPRAALERQQLRLRERQKFFEDILOPETEFVFPPLSHLHLESQRPPIGSISMEVNVDTLEQVELIDLGDGDA
ADVFLPCEDFPPTPQSSGVDDHLEELSLVPTSDRTTSRTSSSSSSDSSNLHSPNPSSDGGADTPLAQSDEEEER
GDGGAEPGACS
161 amino acids

FIG. 6

rSARG-coding	1	ATGGACCCCAATCCCGCGACAGCCCTGGAGCCGACGCACTGCGCTTCATGGAGCGGCG
mSARG-coding	1	ATGGACCCCAATCCCGAGCAGCCCTGGAGCCGCAACGCTGCGCTTCAGGAGAGGAG
hSARG-coding	1	ATGGACCCCAATCCCTGGGCGCCCTGGAGCGGCGAGCAGCTCCGCTTCAGGAGCGGCA
rSARG-coding	61	AGTTCTTCGAGGACATTTTACAGCCAGAGACAGAGTTTGTATTCCCCCTATCCCATCTG
mSARG-coding	61	AGTTCTTTTAGGACATTTTACAGCCAGAGACAGAGTTTGTCTTCCCCCTGTCCCATCTG
hSARG-coding	61	AAATTCTTCGAGGACATTTTACAGCCAGAGACAGAGTTTGTCTTTCCTGTGTCCTCATCTG
rSARG-coding	121	CATCTCGAGTCACAAAGACCCCCCATAGGTAGCATTCTCTCCATGGGAGTGAATGTGGAC
mSARG-coding	121	CACTTCGAGTCACAAAGACCCCCCATAGGTAGCATTCTCTCTATGGGAGTGAATGTGGAC
hSARG-coding	121	CATCTCGAGTCCCAAGACCCCCCATAGGTAGTTCCTCTCCATGGGAGTGAATGTGGAC
rSARG-coding	181	ACACTGGAGCAGGTGGAAATTTATGACCTTGCCTGATCAGGATGGAGCAGATGTCATTCTTA
mSARG-coding	181	ACACTGGAGCAAGTGGAGTTTATTTGATCTTTCCTGATCAGGATGGAGCAGATGTCATTCTT
hSARG-coding	181	ACACTGGAGCAAGTGGAGCTTATTTGACCTTGCCTGATCAGGATGGAGCAGATGTCATTCTT
rSARG-coding	241	CCTTGTGAGGATTCCTTCCAACTCCCCAGTCTCTGAGTGGATGCCCACCAGAGGAG
mSARG-coding	241	CCTTGTGAGGATTCCTTCCAACTCCCCAGTCTCTGAGTGGATGACCATCCAGAGGAG
hSARG-coding	241	CCTTGTGAGGATTCCTTCCAACTCCCCAGTCTCTGAGTGGATGACCATTTAGAGGAG
rSARG-coding	301	CTGAGCCTGCTGGTACCCACGTCAGACAGGACACATCCCGGACCTCCCTCTGCTCCCT
mSARG-coding	301	CTGAGCCTGCTGGTACCCACGTCAGACAGGACACATCCCGGACCTCCCTCTGCTCCCT
hSARG-coding	301	CTGAGCCTGCTGGTACCTTACATCAGACAGGACACATCCCGGACCTCCCTCTGCTCCCT
rSARG-coding	361	---SACTCCTCC---ACCTGGCGGTGCCAATCCAGTGAATGGGGAGGAGACACTCC
mSARG-coding	361	---SACTCCTCC---ACCTGGCGGTGCCAATCCAGTGAATGGGGAGGAGACACTCC
hSARG-coding	361	TCCSACTCCTCCCAACCTGCTTAGCCCAATCCAGTGAATGGGGAGGAGACTCC
rSARG-coding	415	TGGCACAGTC---TGAATAGGAGGATGGGGCAGTGGGGCAGGACCTGGACCTGG
mSARG-coding	415	TGGCACAGTC---TGAATAGGAGGATGGGGCAGTGGGGCAGGACCTGGACCTGG
hSARG-coding	421	TGGCACAGTCGGATGAATAGGAGGATGGGGCAGTGGGGCAGGACCTGGACCTGG
rSARG-coding	472	AGCTAG
mSARG-coding	472	AGCTAG
hSARG-coding	481	AGCTAG

FIG. 7

rSARG-coding	1	MDPNERAALERQQLRLRERQKFFEDILOPETEFVFFLSHLHLESORPPIGSISSMENVVD
mSARG-coding	1	MDPNERAALERQQLRLRERQKFFEDILOPETEFVFFLSHLHLESORPPIGSISSMENVVD
hSARG-coding	1	MDPNERAALERQQLRLRERQKFFEDILOPETEFVFFLSHLHLESORPPIGSISSMENVVD
rSARG-coding	61	NLEQVEFIDLADQDGADVFLCEDSPTFCRSGVDDHPEELSLLVPTSDRTTSRTSSLSS
mSARG-coding	61	NLEQVEFIDLADQDGADVFLCEESSEAFQMSGVDDHPEELSLLVPTSDRTTSRTSSLSS
hSARG-coding	61	NLEQVELIDLGEFADADVFLPCDEPPEPTQSSGVDDHPEELSLLVPTSDRTTSRTSESSS
rSARG-coding	121	DS-SNLRSPNFSDDGGGDTFLAQSDDEE--EDGEGAEPPGFCG
mSARG-coding	121	DS-SNLRSPNFSDDGGGDTFLAQSDDEE--EDGEGAEPPGFCG
hSARG-coding	121	SDSSINLRSPNFSDDGADFLAQSDDEEERED-EGAEPPGAGG

FIG. 8

FIG. 9A



FIG. 9B

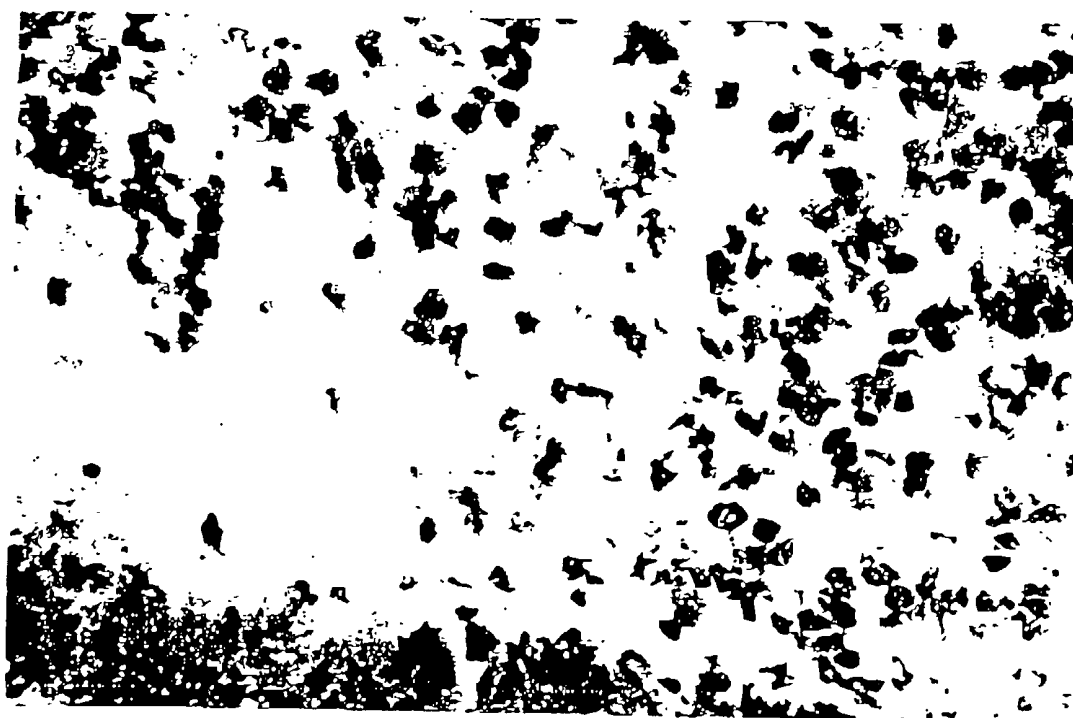


FIG. 9C

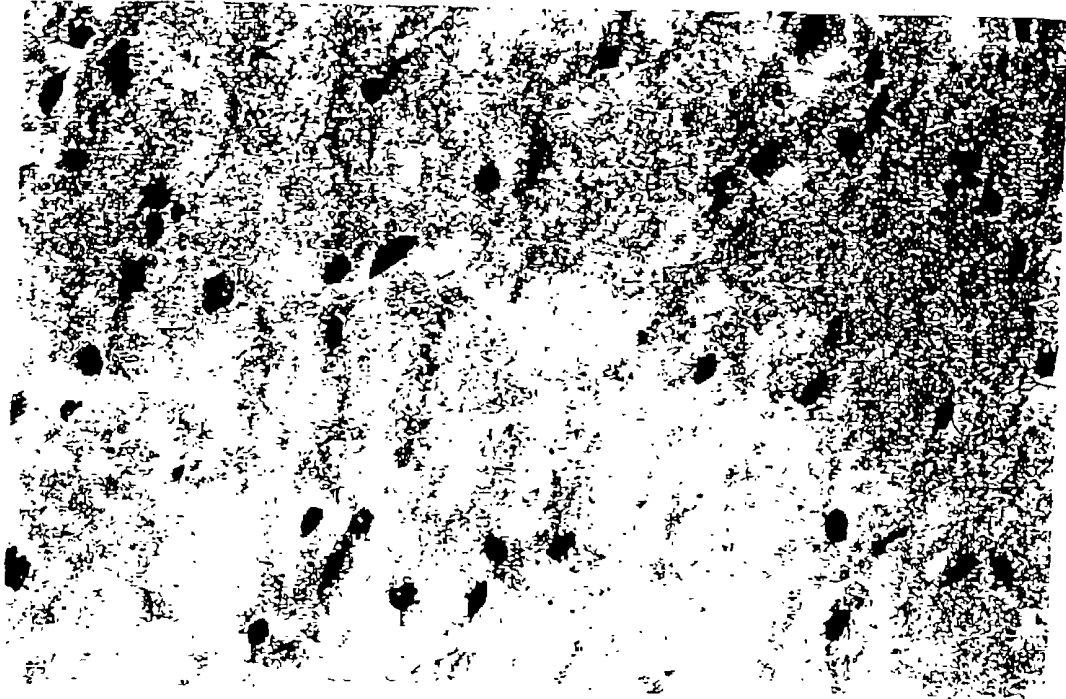


FIG. 9D

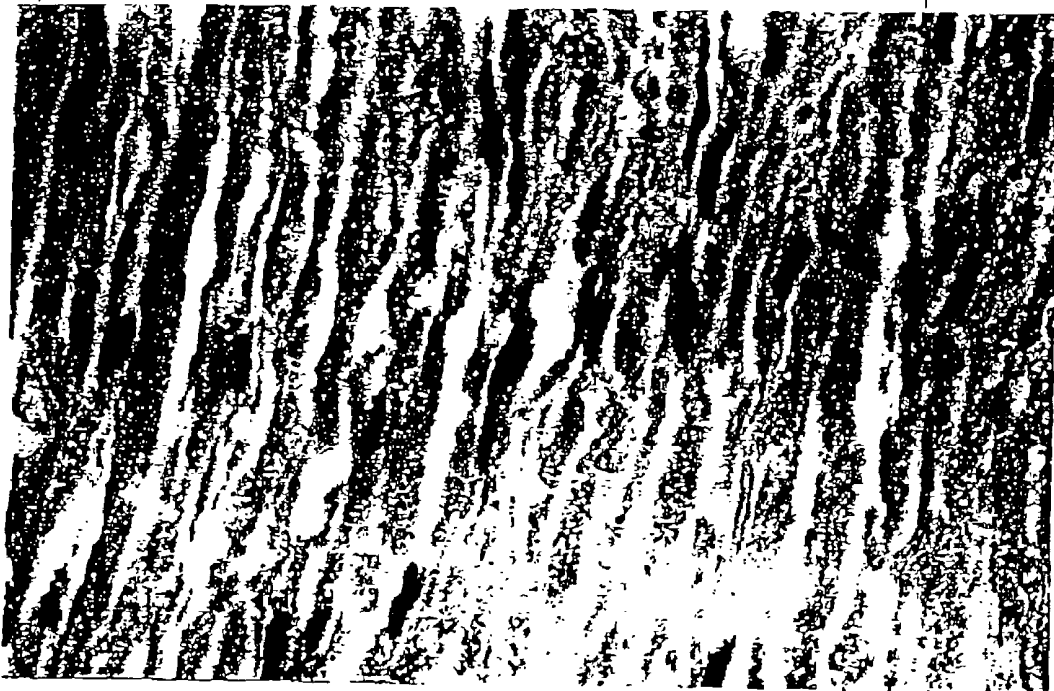


FIG. 9E

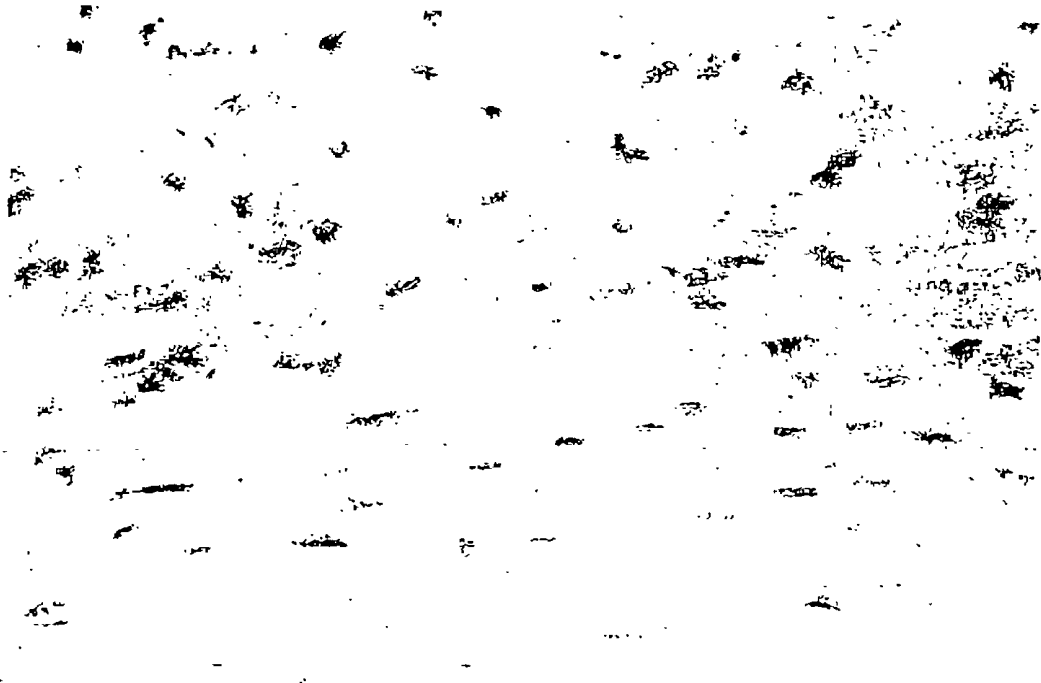
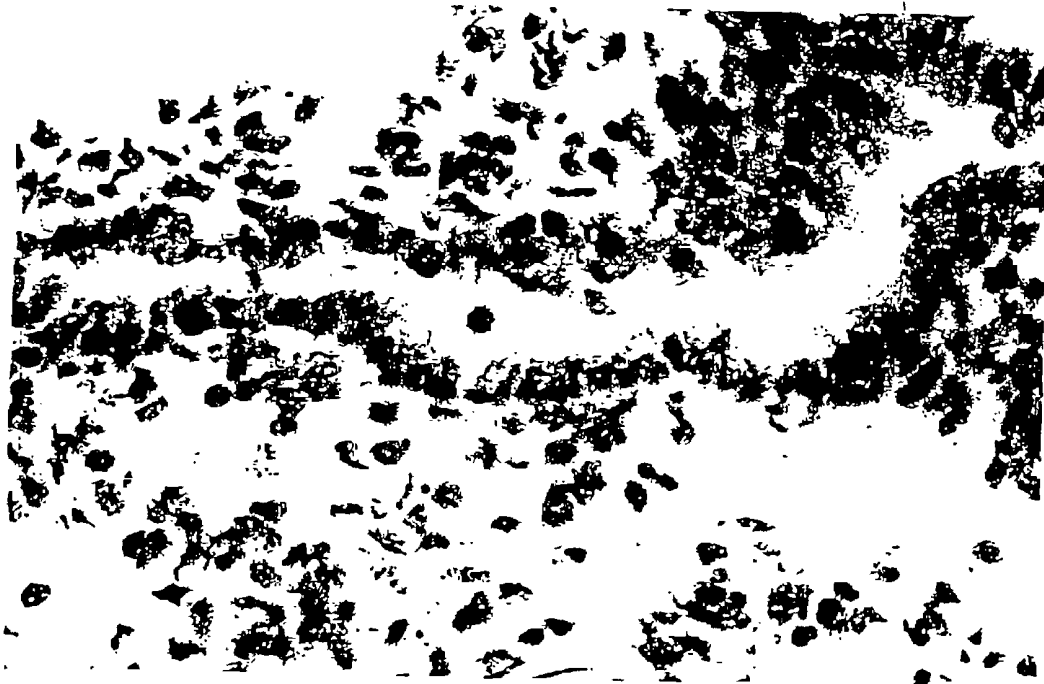


FIG. 9F



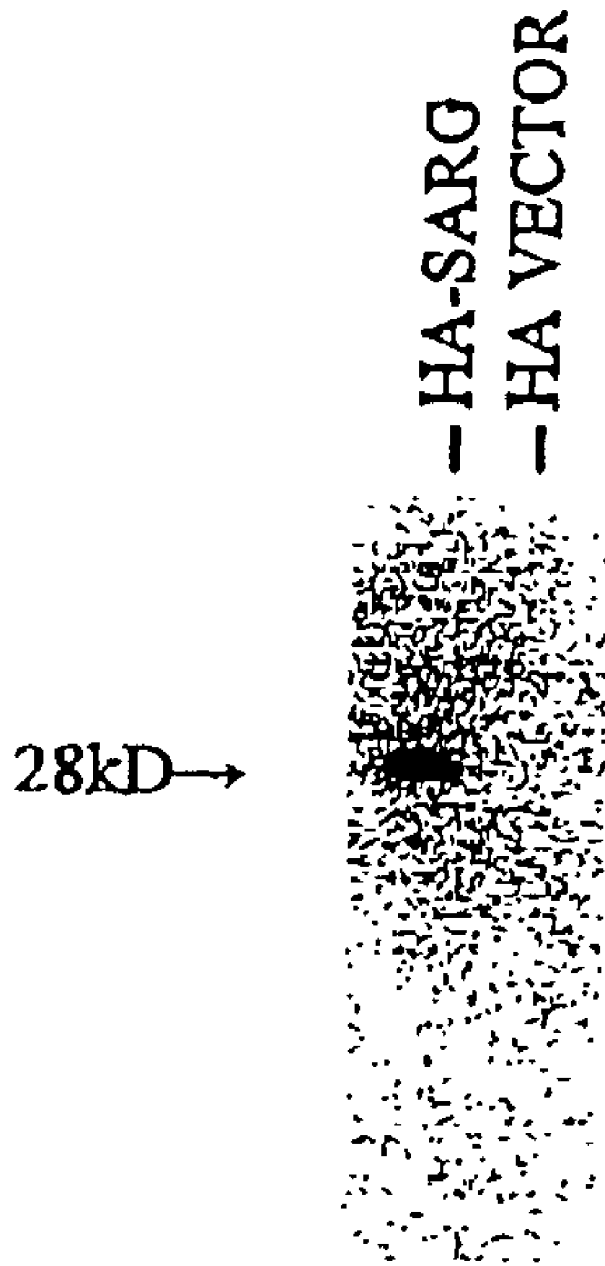


FIG. 10

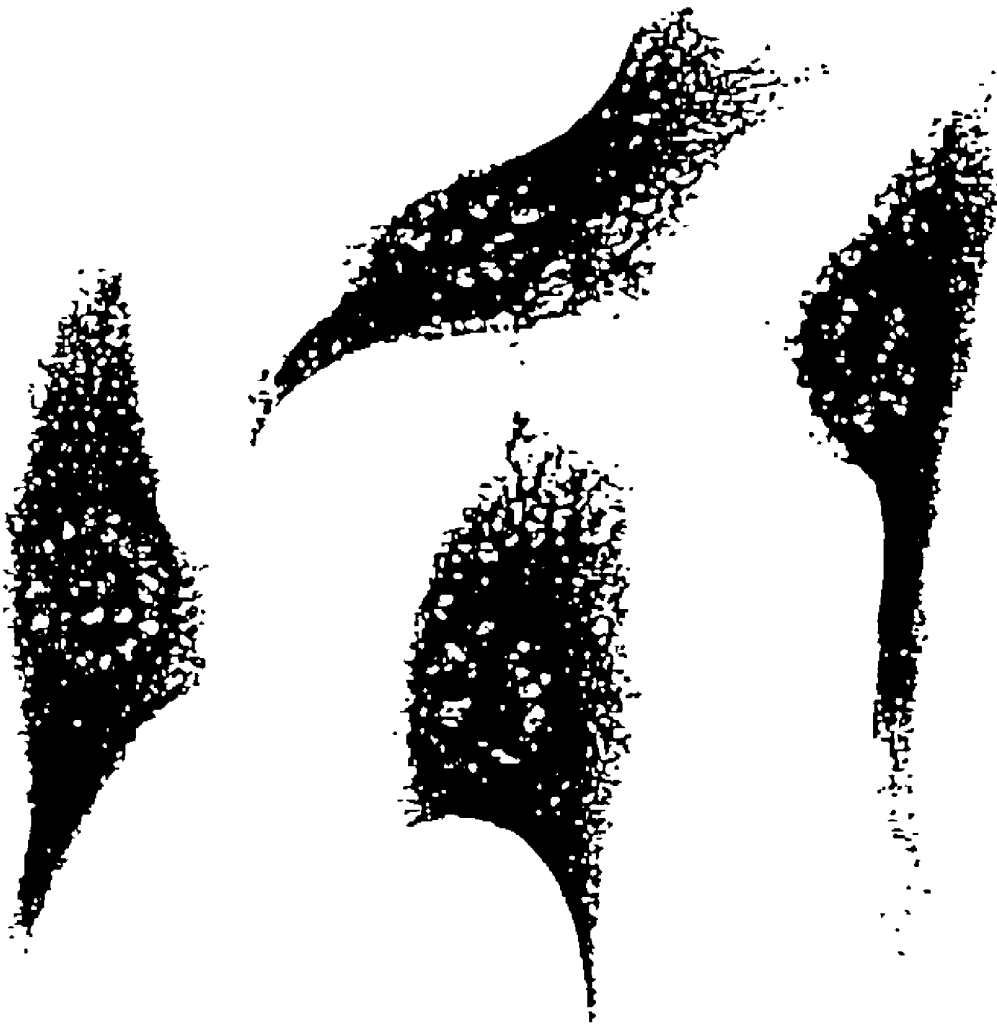


FIG. 11

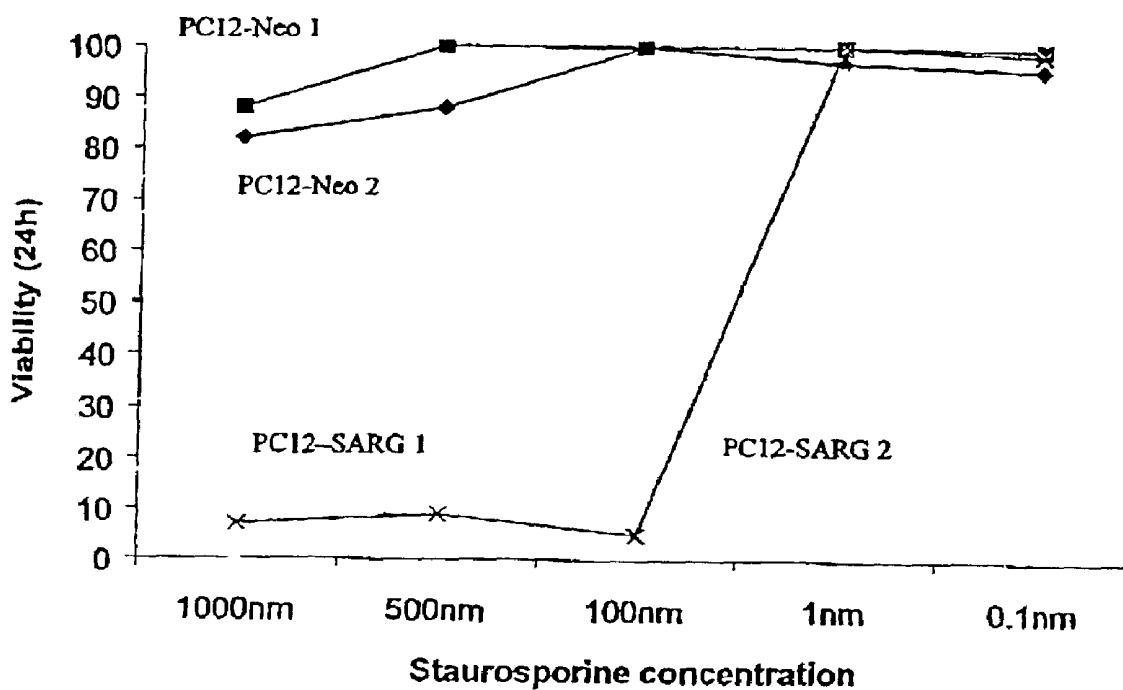


FIG. 12

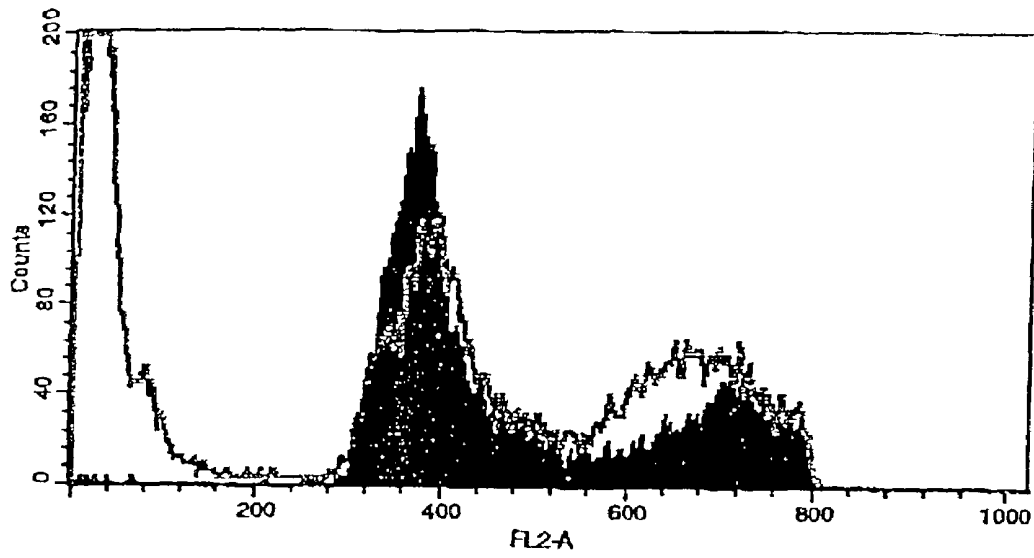
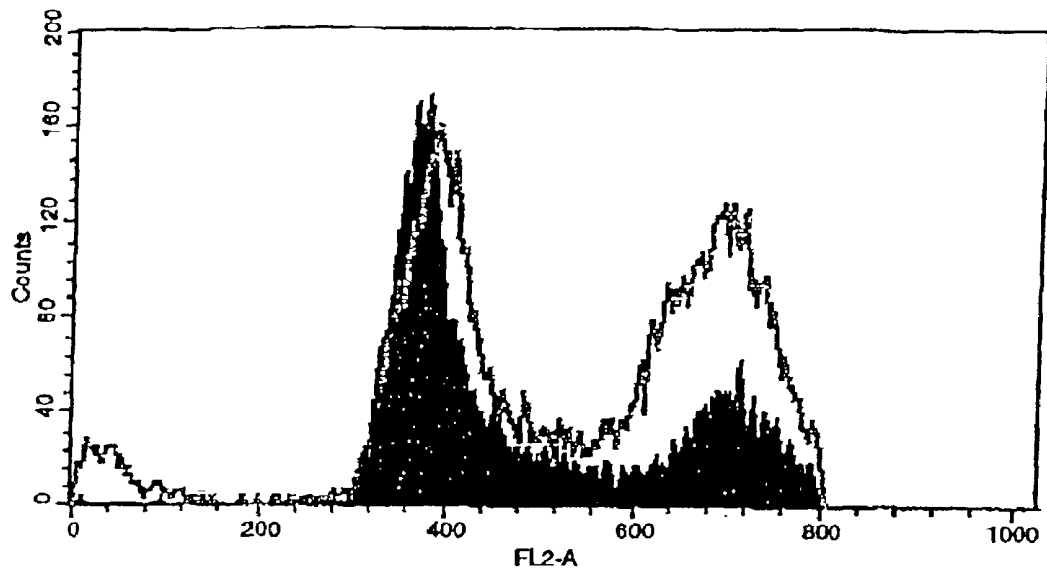


FIG. 13

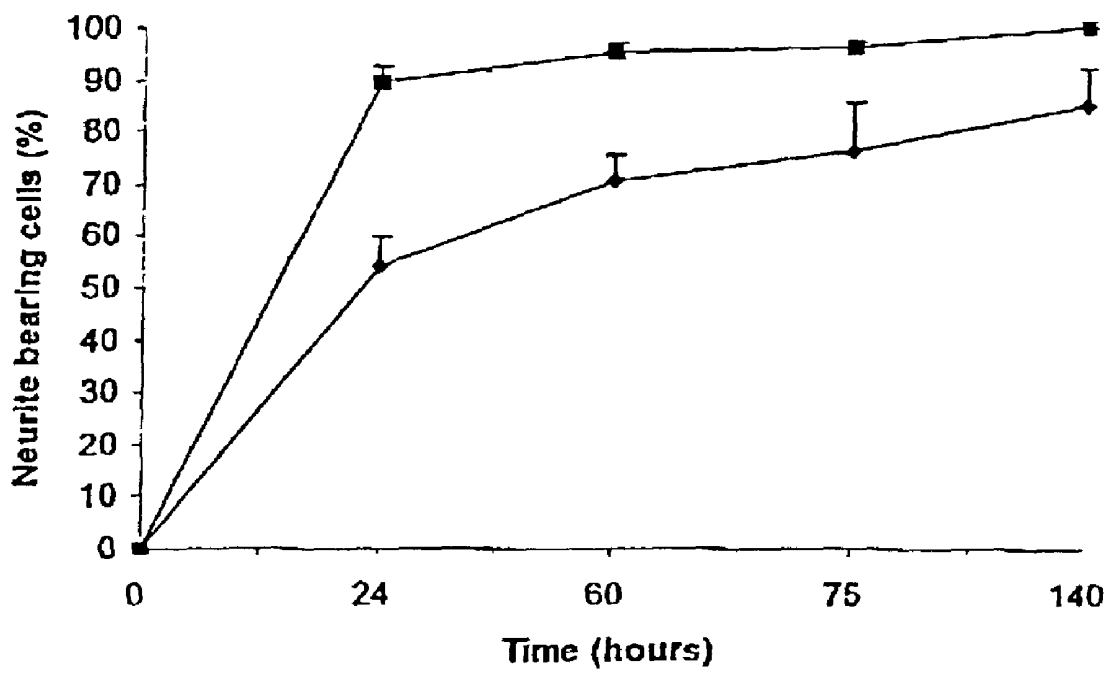
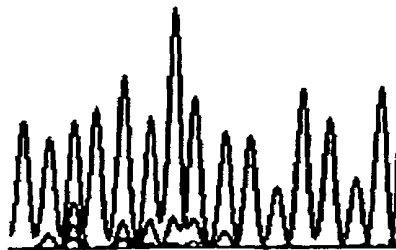


FIG. 14

t→c point mutation in a familial multiple sclerosis patient at nucleotide 67 of coding sequence. Substitution of phenylalanine (F) for leucine (L) at amino acid 23.

21 22 23 24 25
K F L E D

A A A T T C C T C G G G G A C



Wild type 1 MDPNPRAALERQQLRLRERQKFFEDILQPETEFVFPPLSHLHLESQRPPIGSISSEVNVD 60
Mutation 1 MDPNPRAALERQQLRLRERQKFLLEDILQPETEFVFPPLSHLHLESQRPPIGSISSEVNVD 60

Wild type 61 TLEQVELIDLGGPDAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRRTTSRTSSSSS 120
Mutation 61 TLEQVELIDLGGPDAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRRTTSRTSSSSS 119

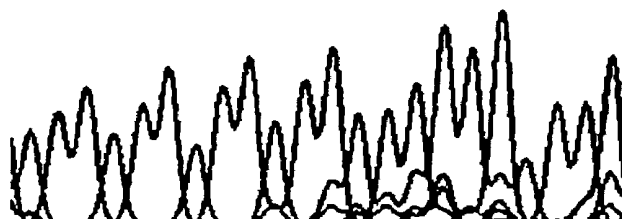
Wild type 121 SDSSTNLHSPNPSDDGADTPLAQSDDEEEERGDDGAEPGACS 161
Mutation 120 SDSSTNLHSPNPSDDGADTPLAQSDDEEEERGDDGAEPGACS 160

FIG. 15

c→t point mutation in a familial multiple sclerosis patient at nucleotide 359 of coding sequence. Substitution of Phenylalanine (F) for serine (S) at amino acid 120

115	116	117	118	119	120	121
T	S	S	S	S	F	S

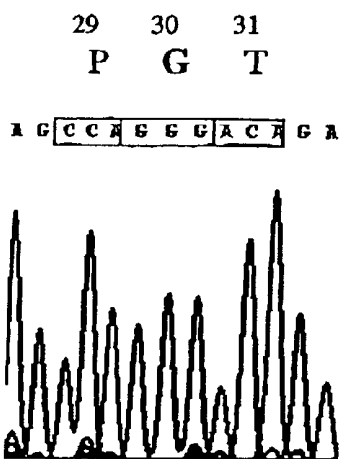
A C G T C C T C C C T C C T T T C T C C F



Wild type 1	MDPNPRAALERQQLRLRERQKFFEDILQPETEFVFPPLSHLHLESQRPPIGSISSMEVNVD	60
Mutation 1	M DENPRAALERQQLRLRERQKFFEDILQPETEFVFPPLSHLHLESQRPPIGSISSMEVNVD	60
Wild type 61	TLEQVELIDLGDPAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRTTSRTSSSSS	120
Mutation 61	TLEQVELIDLGDPAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRTTSRTSSSSF	119
Wild type 121	SDSSTNLHSPNPSDDGADTPLAQSDEEEERG DGG AEPGACS	161
Mutation 120	SDSSTNLHSPNPSDDGADTPLAQSDEESEERG DGG AEPGACS	160

FIG. 16

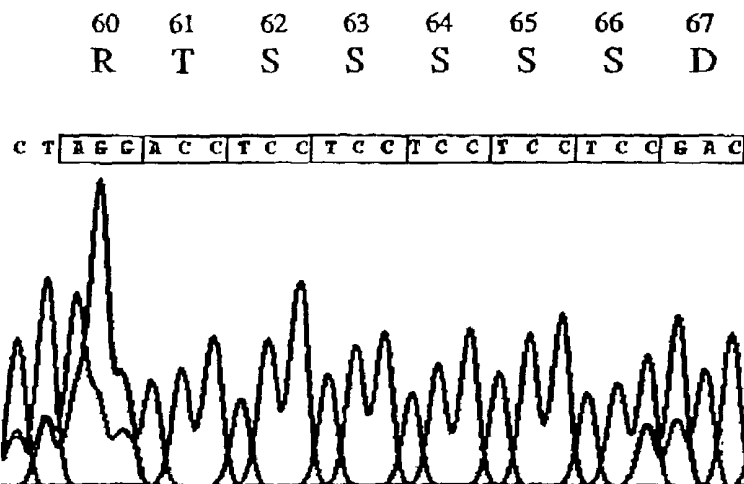
a→g point mutation in a familial multiple sclerosis patient at nucleotide 89 of coding sequence. Substitution of glycine (G) for glutamic acid (E) at amino acid 30.



Wild type 1	MDPNPRAALERQQQLRLRERQKFFEDILOPETEFVFFPLSHLHLESQRPPIGS ^I SSMEVNVD 60	
Mutattion 1	MDPNPRAALERQQQLRLRERQK ^F LEDILOPGTEFVFFPLSHLHLESQRPPIGS ^I SSMEVNVD 60	
Wild type 61	TLEQVELIDLGDPAADVFLPCEDPPPTPOS ^S GV ^D NHLEELSLPVPTSDR ^T TSRTSS ^S SSS 120	
Mutation 61	TLEQVELIDLGG ^P DAADVFLPCEDPPPTPOS ^S GV ^D NHLEELSLPVPTSDR ^T TSRTSS ^S SSS 119	
Wild type 121	SDSSTNLHSPNPSDDGADTPLAQSD ^E EE ^E ERG ^D GGAEPGACS 161	
Mutation 120	SDSSTNLHSPNPSDDGADTPLAQSD ^E EE ^E ERG ^D GGAEPGACS 160	

FIG. 17

Deletion of codon in familial multiple sclerosis patient. Loss of serine residue



Wild type 1	MDPNPRAALERQQLRLRERQKFFEDILQPETEFVFLSHLHLESQRPPIGSISSEVNVD	60
Mutation 1	MDPNPRAALERQQLRLRERQKFFEDILQPETEFVFLSHLHLESQRPPIGSISSEVNVD	60
Wild type 61	TLEQVELIDLGDPAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRRTSRTSSSSS	120
Mutation 61	TLEQVELIDLGDPAADVFLPCEDPPPTPQSSGVDNHLEELSLPVPTSDRRTSRT-SSSS	119
Wild type 121	SDSSTNLHSPNPSDDGADTPLAQSDDEEERGDGGAEPGACS	161
Mutation 120	SDSSTNLHSPNPSDDGADTPLAQSDDEEERGDGGAEPGACS	160

FIG. 18

FIG. 19

SARG INTRON/EXON STRUCTURE

Transcription start site initiator consensus YYCARR is underlined
Donor (GU) and acceptor (AG) splice sites are underlined in italics

Exons are in bold type

Coding exon sequences are in italics

EXON 1

CCAGGCCGGAGCCAGGGGCCCCACTGTTGGGATGCTGGCTGCAGTGGGGCGCCCCAAGCCCAGGT
CCCCTCTGTCTTCTCTTTCGACTTTGTCAGCTGTACTTGTTTTGTCTCCTCTACCCGCAGGAGCTGA
C

+1

ATGGACCCAAATCCTCGGGCCGCCCTGGAGCGCCAGCAGCTCCGCCCTTCGGGAGCGGCAAAAATT
CTTCGAGGACATTTTACAGCCAGAGACAGAGATTTGTCTTTCTCTGTGCCATCTGCATCTCGAGT
CGCAGAGAC

+139

INTRON 1

GTAGTCCCAAGTCTTGAGAAAGAGGGACTGGGGTAGGGTAGGGA
GGATGTCTGTGGGTCCGAAATCTGTGGCACTCTCTCCCCTCTGGTTTT
CTTGGCCCTCTATGCTTCTAACTTGGGACCTGACATGTAACCTCTCACTGT
CCTGGTGTGCAGCTTGGGTCTCTGACTTGGCCACTTCTTGATCCGCAG

EXON 2

+139

CCCCCATAGGTAGTATCTCATCCATGGAAGTGAATGTGGACACACTGGAGCAAGTAGAACTTATT
GACCTTGGGGACCCGGATGCAGCAGATGTGTTCTTGCCTTGCGAAGATCCTCCACCAACCCCCA
GTCGTCTG

+276

INTRON 2

GTATGCCCCCTCTGCTTTGGGGACTTCAGTGCCAGTCAGCCAGAGCCGGATGTCAGGCTCTGA
AACGAGGCTACAAGGCTGGGCTGGGGAAGTACACAAGTAAGGCCTGGAAG
TGGGTGTTTCTACCAATGAAACAGCTGCCTGTTCTGATTTTAGGGAAGTT
GACCCTGAGGGAGAAGTGGGTTACACATCTCTAATCCAAAATCTGGGAA
CGGTCAATCTCTTCTTTAATTTTACATTTGTTATATAAATAAATTAGTC
ACTATAATTAATAAATGTAATAATTGTAATTTTTATATTTGGCAACTTAA
GTAGTTTTAGTCAATTATAATGATATTAATATGTATTGAGTACTTTAGTAG
GTTCCAATACTGTACTAAAGTACTTTACATATATTATCTCAATCCTTACA

METHOD FOR DIAGNOSING A PERSON HAVING MULTIPLE SCLEROSIS

This application claims priority to U.S. Ser. No. 60/299, 765, filed Jun. 22, 2001, the entirety of which is hereby incorporated by reference.

The invention relates to a method for diagnosing a person having multiple sclerosis (MS) or being at risk of developing MS. Further, the invention relates to a method for diagnosing a person having cancer or being at risk of acquiring cancer.

Multiple sclerosis (MS) is a common demyelinating disease of the central nervous system (CNS) affecting up to 0.1% of the north European caucasian population and is considered an auto-immune syndrome directed against unidentified central nervous tissue antigens. The determination of susceptibility to MS development is complex and governed by both environmental and genetic factors (Ebers et al, 1995; Sawcer and Goodfellow, 1998; Sadovnick et al, 2000) with approximately 20% of patients having one or more affected relatives (Chataway et al, 1998). Although thought to be a polygenetic disease, candidate gene approaches have been adopted to isolate genes linked to MS (Weinshenker and Kantarci, 2000). Association with the Caucasian haplotype DRB*1501-DQA*0102-DQB1*0602 (Haines et al, 1998) and a point mutation in the protein tyrosine phosphatase receptor-type C (Jacobsen et al, 2000) have been linked to some cases. Recently, the importance of apoptosis in both T cell elimination and damage to neurons and oligodendrocytes in MS have been recognised (reviewed in Zipp, 2000).

However, to date no clear marker has been reported, although at least a portion of MS cases are clearly familial inherited. A mere recognition of MS or even providing a risk association would be very beneficial for early onset of therapy or preventive measures.

It is therefore an object of the present invention to provide an efficient diagnosis system for MS giving a clear indication and a clear correlation to this disease.

The subject matter of the invention is therefore a method for diagnosing a person having multiple sclerosis (MS) or being at risk of developing MS, characterised by the following steps:

providing a sample of a body fluid or a tissue from said person, said sample containing at least one of the wild type SCF-Apoptosis-Response Gene 1- (wt-SARG-1-) protein and nucleic acids encoding wt-SARG-1, if taken from a person not having MS or a risk of acquiring MS,

detecting the presence of wt-SARG-1-protein or nucleic acids encoding wt-SARG-1 in said sample and diagnosing MS or a risk of acquiring MS, if wt-SARG-1-protein or nucleic acids encoding wt-SARG-1 are not present in said sample.

Surprisingly, SARG-1 protein turned out to be a very specific marker for MS. Persons having either mutated SARG-1 protein or not expressing any SARG-1 protein due to mutations in SARG-1, have a clearly enhanced risk of MS. Investigations on the immuno-histochemical localisation of SARG-1 protein indicated that this protein is located in the grey and white matter of the CNS. The role of SARG-1 in apoptotic induction prompted a candidate gene approach to analyse the mutational status of SARG-1 in cases of familial MS. Indeed, DNA from 20 unrelated familial MS patients was examined by PCR amplification and DNA-sequencing of the SARG-1 locus and compared to SARG-1 sequences from healthy controls. It was found that

all control samples demonstrated wild-type SARG-1 genomic sequences whereas in DNA from MS patients only 6 from 20 DNA samples were even able to be amplified by PCR, i.e. 14 from 20 did not show any detectable SARG-1 signals. In 4 of the 6 other patients genetic alterations were seen. A T→C point mutation at nucleotide 67 resulting in a substitution of phenylalanine with leucine at amino acid 23 (numbering of amino acids and nucleotides according to FIGS. 4-8); a C→T point mutation at nucleotide 359 resulting in the substitution of phenylalanine for serine at amino acid 120 (FIG. 16), A→G point mutation a nucleotide 89 resulting in the substitution of glycine for glutamic acid at amino acid 30 (FIG. 17), deletion of a codon between amino acid 116 and 121 resulting in the loss of a serine residue (FIG. 18). Sequencing of only 20 control DNA samples revealed only wild-type sequence.

Surprisingly, it was also observed that changes in wild type SARG-1 or SARG-1 protein was seen in several cancer cells. In sequence analysis of over 30 cancer cells, a G residue is always present at position 280 resulting in a leucine residue at position 94 instead of a valine (A nucleotide) or methionine (T nucleotide). In a human melanoma cell line two mutations in SARG-1 are found: A→G point mutation at nucleotide 74 resulting in the substitution of aspartic acid for glycine and a C→T point mutation at nucleotide 289 resulting in the substitution of histidine for tyrosine at amino acid 97.

There is a number of restriction sites involved in these mutations: T→C at nucleotide 67 creates a number of restriction sites: Eco88I, XhoI, PaeR7I, Sfr274I, Ama781I, BcoI, BsoBI, Aval; C→T at nucleotide 359 creates an additional BseRI restriction-site.

Therefore, a further object of the present invention relates to a method for diagnosing a person having cancer or being at risk of acquiring cancer, characterised by the following steps:

providing a sample of a body fluid or tissue from said person, said sample containing at least one of the wt SARG-1 protein and nucleic acids encoding wt-SARG-1, if taken from a person not having cancer or being at risk of acquiring cancer, detecting the presence of wt-SARG-1 protein or nucleic acids encoding wt-SARG-1 in said sample and diagnosing cancer or a risk of acquiring cancer, if wt-SARG-1 protein or nucleic acids encoding wt-SARG-1 are not present in said sample.

The source of the sample is always dependent on the nature of the cancer or disease to be diagnosed. Especially preferred samples according to the present invention are derived from human blood, plasma, serum, lymph, nerve-cell containing tissue, cerebrospinal fluid, all biopsy-material, including tumor tissue, bone marrow, nervous tissue, skin, hair, tears, fetal material including amniocentesis material, uterine tissue, saliva, faeces, sperm, etc.

In principle, any method for detecting the presence of wt-SARG-1 protein or nucleic acid encoding wt-SARG-1 in the sample may be applied according to the present invention. Preferably methods are applied which allow also a characterisation of specific SARG-1 mutants if present, either by giving the information that a mutant is present or by analysing the nature of the mutant form in detail.

Especially detecting the presence of point mutations may be preferred within the present invention, i.e. non-wt-forms of SARG-1 differing from wt-SARG-1 or wt-SARG-1 protein in one nucleic acid residue, or one amino acid residue, respectively. The method according to the present invention may be designed to identify those point mutations, espe-

cially point mutations leading to the different amino acid sequence, e.g. exchange of one amino acid residue from the wild type SARG-1 protein.

Suitable methods for detecting the presence of wt-SARG-1 protein or nucleic acids encoding are known in the art, preferably nucleic acids encoding wt-SARG-1 are detected by nucleic acid amplification methods, especially polymerase chain reaction methods, single-strand conformation polymorphism (SSCP) analysis, restriction analysis, microarray technology, proteomics, etc. These methods have been shown as being fast, highly reliable and easily conductable on a high throughput basis. Those tests could be performed on standard tissue or body fluid samples, such as blood, hair or saliva.

On the other hand, preferred methods for detecting the presence of wt-SARG-1 protein encompass the application of a wt-SARG-1 protein antibody, especially a monoclonal antibody, e.g. in a ELISA-format. Such antibodies may be easily produced on an industrial scale with a high degree of standardisation potential.

The method according to the present invention is especially suited to be applied within a screening test format.

The SARG-1 intron/exon structure is given in FIG. 19. The transcription start site initiator consensus YYCARR is underlined. Donor (GU) and acceptor (AG) splice sites are underlined in italics; exons are in bold type. Coding exon sequences are in italic. SARG-1 is located on human chromosome 20q12-13.12.

The present invention also relates to a further aspect, to a nucleic acid molecule comprising a sequence according to Seq.ID.No. 1 (FIG. 4) encoding human wild-type SARG-1. Such nucleic acids may be used for diagnosis but also for therapeutic aspects by providing therapeutic molecules or gene sequences for gene therapy aspects, e.g. by antisense strategies, design of small molecule drugs.

The present invention also encompasses nucleic acid molecules comprising a sequence according to Seq.ID.No. 1, wherein one nucleic acid residue is exchanged by a different nucleic acid residue (e.g. T is replaced by C, G or A) wherein said exchange preferably results in a different SARG-1-protein amino acid sequence.

Especially preferred exchanges are selected from a T to C exchange at position 67 of Seq.ID.No. 1, an A to G exchange at position 74 of Seq.ID.No. 1, an A to G exchange at position 89 of Seq.ID.No. 1, a C to T exchange at position 289 of Seq.ID.No. 1 and a C to T exchange at position 359 of Seq.ID.No. 1. These exchanges relate to exchanges already observed in MS patients or cancer cells. Further exchanges resulting in a viable phenotype are also preferred.

Other preferred mutations in the nucleic acid molecule according to the present invention comprises a deletion in the coding region, preferably a deletion of one or more codons (e.g. 3 nucleic acids, or 6, 9, 12, etc.). One of these mutations leads to the deletion of a codon between amino acids 116 and 121, resulting in the loss of a serine residue (FIG. 19). Mutations leading to non-functional SARG-1 on SARG-1 protein may also be located in the controlling regions (5' or 3') and/or in the sequences, especially at critical positions for correct splicing.

When the nucleic acid molecule according to the present invention is used for a diagnostic purpose, it is not necessary to use the whole sequence. For use as a probe or performing a method according to the present invention a fragment of Seq.ID.No. 1, preferably having a length of at least 12, more preferred at least 15, especially at least 20, nucleic acid residues is suitable for performing various tests, especially diagnostic tests with these probes, e.g. as a probe to identify

or isolate nucleic acid samples or even chromosomal samples or as PCR primers, etc.

The nucleic acid molecules according to the present invention are not restricted to the coding sequence according to Seq.ID.No. 1, but also relate to the genomic counterparts including the whole-exon/intron-structure of this gene, especially also imitations in the non-coding-region resulting in non-wild type forms of the protein (or non-translated forms of the protein) are encompassed by the present invention.

The present invention also relates to a polypeptide being encoded by this nucleic acid molecule, e.g. comprising an amino acid sequence according to Seq.ID.No. 2. There is single potential N-gly-cosylation site with consensus Asn-X-Ser/Thr (amino acid residues 131-133)

```
MDPNPRAALERQQLRLRERQKFFEDILQPETEFVFPFLSHLHLESQRPPIG
SISSMENVDTLEQVELIDLGDPAADVFLPCEDPPPTPQSSGVNDHLEE
LSLPVPTSDRRTTSRTSSSSSDSSTNLHSPNPSDDGADTPLAQSDDEEER
GDGGAEPGACS
```

All threonine and serine residues may be O-glycosylated. Computer predictions indicate high likelihood of glycosylation of serine residues 91, 108, 113, 117, 118, 119, 120, 121, 123, 124 and 133 and the threonine residues 88, 107, 111, 112, 115 and 125.

Similarly, all threonine and serine residues may be phosphorylated. Computer predictions indicate high likelihood of phosphorylation of serine residues 54, 92, 108, 113, 116, 117, 118, 119, 120, 121, 123, 124, 129, 133 and 144 and the threonine residues 61, 88, 107, 112 and 139.

No signal sequences, characteristic domains or other 3-dimensional structures have been detected other than potential protein kinase recognition sites. Of course, also amino acid sequences also having an amino acid residue exchange or a deletion are also encompassed by the present invention.

Amino acid residue exchanges of the polypeptide according to the present invention are preferably selected from amino acid residues Phe23, Asp25, Glu30, His97 and Ser120, especially Phe23 to Leu 23, Asp25, to Gly25, Glu30 to Gly30, His97 to Tyr97 and Ser120 to Phe120 exchanges.

The present invention provides SARG-1 mutant forms as specific markers for (acute) myeloid leukaemia or other leukaemia subtypes as described hereinafter. Deletions of the SARG-1 gene may be partial or full to serve as marker. Diagnostic tests for screening for the presence or absence of such a marker are easily conceivable and reduced to practice by the skilled man in the art.

Preferred polypeptides according to the present invention are re-combinantly produced which exhibit structural differences compared to wt-SARG-1 protein, e.g. differential glycosylation, especially non-homogeneous glycosylation.

The present invention also relates to a method for making an antibody preparation comprising administering a polypeptide according to the present invention to an animal, allowing said animal to generate antibodies against said polypeptide, extracting antibody-containing body fluids or tissue from said animal and preparing an antibody preparation against said polypeptide from said body fluids or tissue. This method is especially applicable for making polyclonal antibodies.

For making monoclonal antibodies a method for making such an antibody preparation is preferred, comprising administering a polypeptide according to the present inven-

tion to an animal, allowing said animal to generate antibodies against said polypeptide, removing the spleen of said animal, preparing fusion cells of said spleen cells with suitable hybridoma generating cells, generating hybridoma cells producing monoclonal antibodies against said polypeptide, cloning and culturing said hybridoma cells, thereby expressing monoclonal antibodies, and preparation of said monoclonal antibodies. The skilled man in the art thereby relies on methods readily available for such purposes and e.g. described in "Antibodies: A laboratory manual" by Ed Harlow, Cold Spring Harbor Laboratory; David Lane, Imperial Cancer Research Fund Laboratories, 1988. Of course, also phage display peptides may also easily be generated.

The present also relates to a kit for performing the *in vitro* diagnosing method according to the present invention which comprises at least means for detecting the presence of wt-SARG-1 protein or nucleic acids encoding wt-SARG-1. The skilled man in the art can envisage the basis of the disclosure of the present application a wide number of suitable alternatives, e.g. anti-wt-SARG-1 protein antibodies, nucleic acid probes selectively binding to wt-SARG-1, nucleic acid primers defining a region being selective for a wt-SARG-1, a chip comprising said nucleic acid probes or said nucleic acid primers. Other preferred means or assays are assays in which proteins bind to SARG-1 such as antibodies or peptides including mutation specific antibodies, ELISAS, Western Blotting assays, flow cytometry assays and assays using immunohistochemical techniques including confocal microscopy.

A further aspect of the present invention relates to a transgenic non-human animal model of the present invention, especially an animal wherein the SARG-1 gene has been mutated or knocked out. A SARG-1 knock out mouse is especially preferred. Methods for providing such models, especially the mouse models, are readily available to the skilled man in the art. Such an animal model is extremely useful in studying genetic variations and mutations of SARG-1, especially with respect to its MS-related disorders.

The term "transgenic" is used herein to describe genetic material that has been or is about to be artificially inserted into the genome of a mammalian cell, particularly a mammalian cell of a living animal. The transgene is used to transform a cell, meaning that a permanent or transient genetic change, preferably a permanent genetic change, is induced in a cell following incorporation of exogenous DNA. A permanent genetic change is generally achieved by introduction of the DNA into the genome of the cell. Vectors for stable integration include plasmids, retroviruses and other animal viruses, YACs, and the like. Of interest are transgenic mammals, e.g. cows, pigs, goats, horses, etc., and particularly rodents, e.g. rats, mice, etc.

Transgenic animals comprise an exogenous nucleic acid sequence present as an extrachromosomal element or stably integrated in all or a portion of its cells, especially in germ cells. Unless otherwise indicated, it will be assumed that a transgenic animal comprises stable changes to the germline sequence. During the initial construction of the animal, "chimeras" or "chimeric animals" are generated, in which only a subset of cells have the altered genome. Chimeras are primarily used for breeding purposes in order to generate the desired transgenic animal. Animals having a heterozygous alteration are generated by breeding of chimeras. Male and female heterozygotes are typically bred to generate homozygous animals.

Transgenic animals fall into two groups, colloquially termed "knockouts" and "knockins". In the present invention, knockouts have a partial or complete loss of function

in one or both alleles of the endogenous SARG-1. Knockins have an introduced transgene with altered genetic sequence and function from the endogenous gene. The two may be combined, such that the naturally occurring gene is disabled, and an altered form introduced.

In a knockout, preferably the target gene expression is undetectable or insignificant. A knock-out of a SARG-1 means that function of the SARG-1 protein has been substantially decreased so that expression is not detectable or only present at insignificant levels or mutated according to the teachings according to the present invention to perform as suitable model for the situation in humans as described herein. This may be achieved by a variety of mechanisms, including introduction of a mutation or disruption of the coding sequence, e.g. insertion of one or more stop codons, insertion of a DNA fragment, etc., deletion of coding sequence, substitution of stop codons for coding sequence, etc. In some cases the exogenous transgene sequences are ultimately deleted from the genome, leaving a net change to the native sequence. Different approaches may be used to achieve the "knock-out". A chromosomal deletion of all or part of the native gene may be induced, including deletions of the non-coding regions, particularly the promoter region, 3' regulatory sequences, enhancers, or deletions of genes that activate expression of SARG-1. A functional knock-out may also be achieved by the introduction of an anti-sense construct that blocks expression of the native genes (for example, see Li and Cohen (1996) *Cell* 85:319-329). "Knock-outs" also include conditional knock-outs, for example where alteration of the target gene occurs upon exposure of the animal to a substance that promotes target gene alteration, introduction of an enzyme that promotes recombination at the target gene site (e.g. Cre in the Cre-lox system), or other method for directing the target gene alteration postnatally.

A "knock-in" of a target gene means an alteration in a host cell genome that results in altered expression or function of the native SARG-1. Increased (including ectopic) or decreased expression may be achieved by introduction of an additional copy of the target gene, or by operatively inserting a regulatory sequence that provides for enhanced expression of an endogenous copy of the target gene. These changes may be constitutive or conditional, i.e. dependent on the presence of an activator or represser.

The exogenous gene is usually either from a different species than the animal host, or is otherwise altered in its coding or non-coding sequence. The introduced gene may be a wild-type gene, naturally occurring polymorphism or mutation, or a genetically manipulated sequence, for example having deletions, substitutions or insertions in the coding or non-coding regions. The introduced sequence may encode wild-type human or animal SARG-1 protein or a mutation thereof, or may utilize the SARG-1 promoter operably linked to a reporter gene. Where the introduced gene is a coding sequence, it is usually operably linked to a promoter, which may be constitutive or inducible, and other regulatory sequences required for expression in the host animal. By "operably linked" is meant that a DNA sequence and a regulatory sequence(s) are connected in such a way as to permit gene expression when the appropriate molecules, e.g. transcriptional activator proteins, are bound to the regulatory sequence s).

Specific constructs of interest, include, but are not limited to anti-sense SARG-1, which will block native SARG-1 expression, expression of dominant negative SARG-1 mutations, and over-expression of a SARG-1. A detectable marker, such as lac Z may be introduced into the locus,

where upregulation of expression will result in an easily detected change in phenotype. Constructs utilizing the SARG-1 promoter region, in combination with a reporter gene or with the coding region are also of interest.

A series of small deletions and/or substitutions may be made in the SARG-1 to determine the role of different exons in DNA binding, transcriptional regulation, etc. By providing expression of SARG-1 protein in cells in which it is otherwise not normally produced, one can induce changes in cell behavior.

DNA constructs for homologous recombination will comprise at least a portion of the SARG-1 with the desired genetic modification, and will include regions of homology to the target locus. DNA constructs for random integration need not include regions of homology to mediate recombination. Conveniently, markers for positive and negative selection are included. Methods for generating cells having targeted gene modifications through homologous recombination are known in the art. For various techniques for transfecting mammalian cells, see Keown et al. (1990) *Methods in Enzymology* 185:527-537.

For embryonic stem (ES) cells, an ES cell line may be employed, or embryonic cells may be obtained freshly from a host, e.g. mouse, rat, guinea pig, etc. Such cells are grown on an appropriate fibroblast-feeder layer or grown in the presence of appropriate growth factors, such as leukemia inhibiting factor (LIF). When ES cells have been transformed, they may be used to produce transgenic animals. After transformation, the cells are plated onto a feeder layer in an appropriate medium. Cells containing the construct may be detected by employing a selective medium. After sufficient time for colonies to grow, they are picked and analyzed for the occurrence of homologous recombination or integration of the construct. Those colonies that are positive may then be used for embryo manipulation and blastocyst injection. Blastocysts are obtained from 4 to 6 week old superovulated females. The ES cells are trypsinized, and the modified cells are injected into the blastocoel of the blastocyst. After injection, the blastocysts are returned to each uterine horn of pseudopregnant females. Females are then allowed to go to term and the resulting litters screened for mutant cells having the construct. By providing for a different phenotype of the blastocyst and the ES cells, chimeric progeny can be readily detected.

The chimeric animals are screened for the presence of the modified gene and males and females having the modification are mated to produce homozygous progeny. If the gene alterations cause lethality at some point in development, tissues or organs can be maintained as allogeneic or congenic grafts or transplants, or in *in vitro* culture.

Binding partners of human SARG-1 include protein-O-mannosyltransferase 1 (POMT1), microtubule-associated protein 1 A (MAP1A), ATPase, Na⁺/K⁺ transporting beta 1 polypeptide (ATP1B1), SWI/SNF complex 60 kDa subunit (BAF60c) alpha-Actinin 2, exon 16, rab GDP dissociation inhibitor 1 (GDI1) and proteasome 26 S subunit, ATPase 3 (PSMC3).

It is a further aspect of the present invention to use SARG-1 binding proteins for modulating activity of SARG-1 proteins (including mutants) and vice versa. Moreover, SARG-1 (wild type) or SARG-1 binding proteins may be used for treating MS or cancer, preferably myeloproliferative disorders, polycythaemia, myelodysplasia and myeloid leukaemia, especially acute myeloid leukaemia, by administering an effective amount of SARG-1 or SARG-1 binding protein (or a complex thereof) to an MS or cancer

patient. Instead of wt-SARG-1 protein, also fragments of SARG-1 may be used which are suitable for advantageous treatment of such patient (e.g. fragments binding to the SARG-1 binding proteins). Minimum requirements for such fragments are easily found by the skilled man in the art especially using the mouse models described herein.

BRIEF DESCRIPTION OF THE DRAWINGS

The method is further described by the examples and drawing figures, without being restricted thereto.

FIG. 1 shows reverse northern blot demonstrating induction of the rat SARG-1 gene during apoptosis. Plasmid DNA was spotted onto nylon membranes and hybridised with radiolabelled cDNA isolated from apoptotic stem cells. Plasmids containing the 5' region of SARG-1 (arrowed) hybridise with cDNA only present in apoptotic cells. Control cDNA from self-renewing cells showed only background levels (data not shown).

FIG. 2 shows rat multiple tissue northern blot. The 342 bp SARG-1 fragment was radiolabelled and hybridised with RNA isolated from different rat tissues. A species of approximately 1.4 Kb is observed with an ubiquitous tissue distribution and highest expression in brain and heart.

FIG. 3 shows mouse embryonic tissue northern blot. The 342 bp SARG-1 fragment was radiolabelled and hybridised with RNA isolated from various stages of embryonic development. Clear regulation of a 1.4 Kb species is seen with highest levels after 7 days and re-expression of the gene product on day 17.

FIG. 4 shows full length rat mouse and human SARG-1 cDNA sequences.

FIG. 5 shows comparisons between full length rat mouse and human SARG-1 cDNA sequences.

FIG. 6 shows full length rat, mouse and human SARG-1 coding sequences with corresponding predicted amino acid sequence.

FIG. 7 shows comparison between the rat, mouse and human SARG-1 DNA coding sequences.

FIG. 8 shows comparison between the rat, mouse and human SARG-1 amino acid coding sequences.

FIG. 9 shows immunohistochemical analysis of mouse tissue with anti-SARG-1 peptide antibodies. A brown reaction product indicates SARG-1 expression. A: Mouse cerebral cortex (magnification $\times 40$), B: Cerebellum ($\times 40$), C: Spinal chord ($\times 40$), D: Peripheral nerve ($\times 60$), E: Heart ($\times 40$) and F: Lung tissue ($\times 40$).

FIG. 10 shows SARG-1 protein detected by western blotting of cellular extracts from MelJUSO melanoma cells transfected with pMH-SARG-1-HA. Anti-HA monoclonal antibody 3F10 detected a single species which migrates at 28 kD.

FIG. 11 shows immunohistochemical analysis of human SARG-1 protein in MELJUSO cells transfected with pMH-SARG-1-HA detected by the anti-HA monoclonal antibody 3F10 demonstrating cytoplasmic localisation of the protein.

FIG. 12 shows viability of PC12 cultures transfected with empty vector or pIRES2-EGFP-SARG-1 treated with 100 nM staurosporine.

FIG. 13 shows flow cytometrical analysis of PC12 cells transfected with empty vector or pIRES2-EGFP-SARG-1 treated with 100 nM staurosporine for 24 h. A sub G₀/G₁ peak characteristic for the induction of apoptosis is seen only in SARG-1 over-expressing cells.

FIG. 14 shows PC12 cells that overexpress SARG-1 protein (■) undergo accelerated NGF-mediated terminal differentiation when compared to cells transfected with empty vector (◆).

FIG. 15 shows genetic alteration in familial multiple sclerosis. A T→C point mutation at nucleotide 67 resulting in the substitution of leucine for phenylalanine amino acid 23.

FIG. 16 shows genetic alteration in familial multiple sclerosis. A C→T point mutation at nucleotide 359 resulting in the substitution of phenylalanine for serine at amino acid 120.

FIG. 17 shows genetic alteration in familial multiple sclerosis. An A→G point mutation at nucleotide 89 resulting in the substitution of glycine for glutamic acid at amino acid 30.

FIG. 18 shows genetic alteration in familial multiple sclerosis. Deletion of a codon between amino acids 116 and 121 resulting in the loss of a serine residue.

FIG. 19 shows the SARG-1 intron/exon structure.

EXAMPLES

Identification of a Novel SCF-Apoptosis-Response-Gene (SARG-1) Induced During Stem Cell Apoptosis

Withdrawal of growth factor from SCF dependent myelomonocytic progenitors results in rapid induction of cell cycle-independent apoptosis. Although over 95% of SCF-deprived cultures exclude vital dyes 12 hours after growth factor withdrawal, no proliferative response is seen on restimulation with SCF after this time point. A differential display screen was conducted to examine immediate-early (four hour) RA expression differences during the processes of myelomonocytic stem cell self-renewal and apoptosis induced by growth factor withdrawal. Amplification of approximately 1/3 of all cellular mRNA species by differential display PCR (Liang and Pardee 1992, 1995; Liang et al, 1993) with defined primer (Bauer et al, 1993) sets (Display systems) identified one fragment induced during apoptosis not present in self-renewing precursors. Following gel excision of this band and TA cloning into the pCR^{II} vector (Invitrogen) of reamplificants, specific expression induction was confirmed in a reverse northern procedure with representative dot blotted plasmid preparations hybridised with radiolabelled cDNA isolated prepared from independent cultures (FIG. 1). All positively identified differentially regulated clones were sequenced and found to contain an identical 342 base pair insert (excluding the downstream and upstream differential display primer pairs).

Molecular Weight of SARG-1 cDNA and Tissue Expression. Regulation During Embryonic Development

Radioactive labelling of the 342 bp fragment discovered above and probing of multiple tissue northern blots (Clontech) showed an approximately 1300 bp length for the mature mRNA in rat, mouse and human tissues. An ubiquitous low expression was seen with highest expression levels in brain and heart (FIG. 2). Northern blots of immobilised mRNA isolated from different stages of murine embryonic development probed with the radioactively labelled 342 bp fragment demonstrated regulation of SARG-1 mRNA during development with highest levels after 7 days and re-expression of the gene product on day 17 (FIG. 3).

Cloning of Full Length Rat, Murine and Human SARG-1.

Additional SARG-1 sequence information was obtained by a 5' rapid amplification of cDNA ends polymerase chain reaction (RACE) procedure from adapter ligated rat brain Marathon-ready cDNA (Clontech). A primer was constructed at the 3' proximal end of the 342 bp sequence and the PCR product cloned into the pCR[®] TA cloning vector (Invitrogen) and sequenced. The full length SARG-1 1062 bp gene transcript was sequenced and has a 479 bp open reading frame which encodes a 158 amino acid protein. Murine SARG-1 was isolated by screening a bacterial artificial chromosome bank derived from mouse strain 129SvJ with the full length rat SARG-1 cDNA. Homologous clones were isolated and a Xho I fragment sequenced. Mature murine SARG-1 RNA was then identified by PCR from mouse brain cDNA. Human SARG-1 was isolated from a phage bank of brain cDNA by homology to the full length rat sequence and sequenced. The full-human SARG-1 locus was amplified by PCR with primers spanning the SARG-1 cDNA from chromosomal DNA isolated from the peripheral blood mononuclear cells of healthy volunteers. The full length sequences of rat, mouse and human SARG-1 are shown in FIGS. 4, 5 and 6.

The Homologies Between Rat, Mouse and Human Sequences

Rat and mouse SARG-1 are 93% at the nucleic acid and 96% homologous at the predicted protein levels, respectively. Human SARG-1 displays 83% and 84% homologies to mouse and rat SARG-1 at the nucleic acid level and 84% and 86% homologies at the predicted protein levels, respectively (FIGS. 7 and 8), and is localised to the long arms of chromosome 20 (at 20q13.12 (Deloukas et al., 2001)), deletion of which are a common occurrence in a wide range of myeloproliferative disorders (Wattel et al, 1993 and Bench et al., 2000). Sequences contain a potential PEST region (Rechsteiner et al. 1996) (+), a single conserved potential N-glycosylation site (#) and three conserved PKC (\$) and casein kinase II (*) phosphorylation sites (see FIG. 5B).

Analysis of SARG-1 Distribution with Anti-SARG-1 Antibodies

SARG-1 antibodies were prepared by immunising chickens with a peptides (MDPNPAALERQQLR and DEEEERGDGQAEPGA) corresponding to the first 15 amino acids and to the c-terminal part of mouse, rat and human SARG-1 coupled to keyhole limpet hemocyanin. IgY was prepared from egg yolks and specific antibody prepared by affinity chromatography with column-immobilised peptide.

In immunohistochemical analysis of 5 mm acetone-fixed sections of mouse tissue, specific anti-SARG-1 IgY antibodies visualised with anti-IgY peroxidase conjugated second step antibodies, detected wide expression of the SARG-1 protein in nervous tissue. SARG-1 staining is seen throughout the cerebral cortex (FIG. 9A) and within the granular, Purkinje and molecular cell layers of the cerebellum (FIG. 9B). In the spinal chord, SARG-1 is expressed in both the gray matter neurophil containing nerve cell bodies, dendrites, glial cells and blood vessels and the white matter consisting largely of myelinated nerve tracts (FIG. 9C). SARG-1 expression is also colocalised to occasional peripheral nerve processes (FIG. 9D). SARG-1 is also ubiquitously expressed in cardiac muscle, (FIG. 9E), lung ciliated epithelia (FIG. 9F) and epithelial cells of the ileum and colon (data not shown). Staining of mouse embryonic tissue demonstrated strong staining of neural tissue, brain, heart, pla-

centa, uterus, the organ of corti, the dermis (stratum granulosum) and lining of the gut (data not shown).

Generation of SARG1 Over-dressing Cells

The MelJUSO melanoma cell line was maintained in Dulbecco's modified eagles medium (DMEM) supplemented with 10% fetal calf serum and an antibiotic-antimycotic mix containing 100 units/ml penicillin 100 µg/ml streptomycin, and 0.25 µg/ml amphotericin B (all from GibcoBRL) in a fully humidified air atmosphere containing 5% CO₂ at 37° C. The human SARG-1 coding sequence (hSARG-1₄₈₅) was cloned into the pMH expression vector (Roche) by standard molecular biology procedures (Sanbrook et al, 1989) under the control of a CMV promoter in frame with a c-terminal hemagglutinin (HA) peptide sequence. Semi-confluent cultures in 6 well plates were transfected with this vector (pMH-SARG-1-HA) or empty vector in the presence of Fugene (Roche) and clones isolated which displayed resistance to neomycin.

To examine the expression and function of SARG-1 in a neural cell culture system, standard molecular biology techniques (Sanbrook et al, 1989) were used to clone the rat SARG-1 coding sequence (rSARG-1₄₇₉) into the pRESII-EGFP eukaryotic expression vector (Clontech) under the control of a CMV promoter which bicistronically translates through an internal IRES sequence both SARG-1₄₇₇ and enhanced green fluorescent protein. The rat pheochromocytoma PC12 cell line was maintained in DMEM supplemented with 8% horse serum, 8% fetal calf serum and an antibiotic-antimycotic mix containing 100 units/ml penicillin, 100 µg/ml streptomycin, and 0.25 µg/ml amphotericin B (all from GibcoBRL) in a fully humidified air atmosphere containing 5% CO₂ at 37° C. Semi-confluent cultures in 6-well plates were transfected with pRES2-EGFP-SARG-1 or empty vector for 18 h in the presence of the uptake enhancing cationic lipid mix pFx1 (Invitrogen) at a 6:1 lipid to DNA ratio in serum-free opti-MEN (Gibco). Stable transfectant colonies were picked from separate wells after selection in 800 mg/ml geneticin (Gibco) and clones with similar EGFP fluorescence expanded from independent transfections.

Post Translational Modification of Mature Rat and Human Proteins

The molecular weights of rat, mouse and human SARG-1 protein predicted from the amino acid sequences are 17186.52, 17193.52 and 17492.76, respectively. IgY antibodies against SARG-1 detect a 28 kD species in western blotting expressed at low levels in native PC12 cells. Upon transfection of pRES2-EGFP-SARG-1 into these cells, expression levels of this species increase. The molecular weight of human SARG-1 protein was determined by western blotting of cellular extracts from MelJUSO melanoma cells transfected with pMH-SARG-1-HA. Anti-HA monoclonal antibody 3F10 (Roche) detected a single species which migrates at 28 kD (FIG. 10).

Subcellular Localisation of Transfected SARG-1 Protein

Immunohistochemical staining of SARG-1 transfected PC12 cells demonstrated an exclusive cytoplasmic localisation for the SARG-1 gene product. Immunohistochemical analysis with anti-HA monoclonal antibody 3F10 also demonstrated an exclusive cytoplasmic localisation for the SARG-1 protein (FIG. 11) primarily colocalising with expression of binding protein (bip) used as marker for the endoplasmic reticulum. Co-localisation of HA-SARG-1 with the golgi-specific antigen coat protein (b-cop) is minimal. In contrast, SARG-1 expression partly overlaps with expression of the lysosomal specific protein LAMP-2.

Role of SARG-1 in Apoptosis and Differentiation of Neural Cell Cultures

Treatment of PC12 cells with the protein kinase inhibitor staurosporine induces neurite outgrowth at concentrations of 100 nM (Hashimoto and Hagino, 1989) and apoptosis at concentrations above 1 mM (Fu et al, 1999). SARG-1 over-expressing PC12 cells undergo a rapid loss in viability on treatment with 100 nM staurosporine (Calbiochem) in comparison to vector control cultures (FIG. 12) which undergo neurite outgrowth. Cell death in SARG-1 over-expressing PC12 cultures is accompanied by cell shrinkage, development of a morphology characteristic of programmed cell death and loss of DNA from ethanol fixed cells in flow cytometrical cell cycle analyses (FIG. 13) characteristic of the apoptotic process (Fraker et al, 1995). No similar effects are seen with the staurosporine analogue X252a (Alexis), the phosphatidylinositol 3-kinase inhibitors LY-294002 (Alexis) or Wortmannin (Calbiochem), the MAP kinase kinase inhibitor PD 98059 (Alexis) or the protein kinase C inhibitor Bisindolylmaleimide 1 (Alexis). Over-expression of SARG-1 also enhances terminal differentiation of PC12 cells induced by nerve growth factor (FIG. 14).

Isolation of SARG-1 Binding Proteins

Yeast two hybrid screening are performable with the commercially available Matchmaker system from Clontech. The bait plasmid is constructed by cloning the full length hSARG-1 coding sequence into the pGBKT7 shuttle vector (Clontech) by PCR using the BamH1 and, EcoR1 restriction sites using standard methods (shuttle vector pGBKT7: The coding region of hSARG-1 is cloned as a c-terminal fusion to amino acids 1-147 of the GAL4 DNA binding domain (DNA-BD) containing a c-myc tag under the control of the yeast ADH1 promoter (P) by PCR amplification using the EcoR1 and BamH1 restriction sites and selected in yeast using the TRP1 nutritional marker). This construct is then used to lithium acetate transform the MAT α yeast strain AH109 which is auxotrophic for adenine (Ade), histidine (His) leucine (Leu) and tryptophan (Trp) and selected for pGBKT7 by growth on synthetic dropout (SD)-Trp medium. A human brain matchmaker library (Clontech) directionally constructed in pACT2 pretransformed in the MAT α Y187 (Ade⁻, His⁻ Leu⁻ and Trp⁻) yeast strain is purchased and at least 3x10⁶ clones mated with AH109 transformed hSARG-1-1 (shuttle vector pACT2: Human brain library inserts are directionally cloned in frame as c-terminal fusions to amino acids 768-881 of the GAL4 activation domain, the SV40 T-antigen nuclear localization sequence and a HA epitope tag under the control of the yeast ADH1 promoter (P) and selected in yeast using the LEU2 nutritional marker). Zygotes are isolated by Leu⁻ and Trp⁻ selection and protein interactions simultaneously by reporter gene activation of HIS3, ADE2, and MEL1 on -His, -Ade plates containing X- α -Gal (5-Bromo-4-chloro-3-indolyl-a-D-galactopyranoside), respectively. Positive colonies are restreaked and additionally tested for lacZ reporter activation with β -galactosidase. The inserts from positive colonies are amplified by PCR, repeats identified by restriction digestion and positive interactors identified by short run sequencing from the adapter end of the insert. The identity of potential interactors are ascertained by FASTA and BLAST searches against non-redundant and expressed sequence tag data bases at the national centre for biotechnology information (www.ncbi.nlm.nih.gov/BLAST/) and European bioinformatics institute (www.ebi.ac.uk/fasta33/) interfaces, respectively, to identify characterized proteins cloned in frame with the GAL4 activation domain. Single pACT2 plasmids

are then recovered by transformation of *E. coli* under ampicillin selection and positive interactions retested by cotransformation of AH109 with pGBKT7-SARG-1 and pACT2 containing the library insert selected by blue colony growth on -Ade, -His, -Leu, -Trp, X- α -gal plates. Bait and library genes are cloned into appropriate c-myc and HA epitope tagged eukaryotic expression vectors and interactions further confirmed by Western blotting of coimmunoprecipitated proteins either translated in vitro or from transiently cotransfected HEK 293 cells. The latter also allows colocalization experiments by confocal microscopical analysis following cloning and isolation of full length coding sequences. The cellular co-expression of interaction partners with SARG-1 is examined by either staining of sequential embryological slides generated with antibodies where available or by in situ hybridization of identified sequences. The identification of known binding partners enables the development of antisense or RNAi strategies to specifically reduce target expression or allows the use of specific functional inhibitors to further characterise the phenotypic changes seen in PC-12 apoptosis and differentiation. Binding partner identification also identifies the biochemical pathways which are influenced by SARG-1 expression which may also be specifically inhibited. Knockout models for identified binding partners may be crossed with the SARG-1 deficient mouse.

Pitfall Analysis

A relevant tissue is analysed where SARG-1 is highly expressed and plays a role in differentiation and apoptosis in a cell line model. Human gene banks are analysed to enable rapid identification of known RNA sequences. The SARG-1 sequence contains no cytoplasmic localization signals which could reduce transcriptional activation. Transfection of SARG-1 is not directly toxic to PC-12 or MelJuso cells and is therefore unlikely to be toxic in yeast. The matchmaker system uses multiple reporter genes with different promoter constructs to eliminate artefacts and has been used to isolate a number of binding partners (Corset et al, 2000; Galiegue et al, 1999; Ono et al, 2000; White et al, 2000). A eukaryotic cell line (HEK293) which allows efficient transient cotransfection of vector constructs is used to allow rapid screening of putative binding partners.

In a yeast two-hybrid analysis using the human SARG-1 protein as a bait to screen a human brain cDNA ebank the following binding partners were isolated:

- 1: Homo sapiens protein-O-mannosyltransferase 1 (POMT1), (LC2)mRNA: (gi: 12734916)
- 2: Homo sapiens microtubule-associated protein 1A (MAP1A), mRNA (XM_012387)
- 3: Homo sapiens ATPase, Na⁺/K⁺ transporting, beta 1 polypeptide (ATP1B1, mRNA (gi: 4502276)
- 4: Human SWI/SNF complex 60 KDa subunit (BAF60c) mRNA (gi: 1549246)
- 5: Homo sapiens ACTN2 gene for alpha-Actinin 2, exon 16 (gi: 6448557)
- 6: Homo sapiens rab GDP dissociation inhibitor 1 (GDI1), mRNA (gi: 4503970)
- 7: Homo sapiens proteasome (prosome, macropain) 26S subunit, AT-Pase, 3(PSMC3), mRNA (gi: 4506210)

Of these binding proteins, especially the ATPase beta 1 polypeptide is of specific interest for use as a pharmaceutical target with respect to the binding to SARG-1.

Generation of a Conditional SARG-1 Knockout Mouse

Effective homologous recombination between vector construct and chromosomal target sequences is normally achieved by a species specific region of homology of 7 kb

with at least 2 kb of homologous material adjacent to the drug resistance cassette (Johnson et al., 1995). The murine SARG-1 locus has been fully amplified by PCR (2420 bp) and sequencing demonstrated the presence of 3 introns.

For construction of the replacement vector, approximately 2.5 kb of additional non-coding chromosomal sequence information flanking the SARG-1 locus are obtained in the 129SvJ mouse. To this end, a radiolabelled SARG-1 cDNA sequence or a PCR product of the entire mSARG-1 locus is used to screen a 129SvJ mouse bacterial artificial chromosome (BAC) library (Incyte genomics) with an average insert size of 120 kb or a 129SvJ mouse genomic library constructed in Lambda FIX® II phage (Stratagene) with insert sizes of 9 to 23 kb, by standard methods. Following clone or phage isolation, genomic sequences flanking the coding region (approximately 2.5 kb) is characterized by either direct sequencing of isolated clones with primers internal to the SARG-1 coding region, sequencing of cloned restriction fragments bearing SARG-1 sequences or by inverse PCR. The sequence information obtained is used to design PCR primers to insert 129SvJ mSARG-1 genomic sequences into the conditional/hypomorphic pDELBOY-3X targeting vector (Rossi et al, 2001). This vector incorporates features that overcomes problems potentially associated with gene deletion. Artifactual phenotypes are generated in knockout mice due to the transcriptional activity of the neomycin cassette which can lead to disrupted regulation and splicing of the target locus and neighbouring genes (Pham et al, 1996; Olson et al, 1996). The neomycin cassette in the pDELBOY-3X vector is flanked by flip recombinase (frt) sites which permit efficient excision of the selection cassette in vitro and in vivo. Transient transfection of cells stably transfected with pDELBOY-3X with a vector expressing Flp recombinase leads to excision of the neomycin cassette.

Recently developed tools such as Flp recombinase-GFP fusion protein vectors, for example, allow enrichment of Flp-mediated recombination events in vitro by fluorescence activated cell sorting (Sabath et al., 2000). Conversely, crossing mice with the 129SvJ FLPer deleter mouse ubiquitously expressing enhanced FLP allows excision of the neomycin cassette in vivo. In addition, cloning of SARG-1 coding sequences between the loxP sites in the pDELBOY3X vector permits generation of a null allele either in vitro by transfection with cre recombinase expressing plasmids or in vivo by crossing mice generated with animals expressing cre recombinase in a tissue specific or. This strategy eliminates problems associated with heterozygote embryonic lethality and allows the precise tissue specific analysis of protein function. Knockout mice are generated by standard procedures (Papaioannou and Johnson, 2000; Gu et al, 1993). Briefly, embryonic stem cells (ES) are transfected by electroporation with linearized pDELBOY-3X-SARG-1 and homologous recombination events selected by neomycin/gancyclovir treatment and clones screened by PCR or southern blotting with a radiolabelled probe external to the targeting sequences. Blastocytes are isolated from 3.5 day pregnant mice, injected with ES clones and reimplanted in pseudopregnant mice. Following coat colour selection of chimeras, founder animals are mated with normal 129SvJ mice to produce a breeding line. Homozygotes are subsequently be obtained by inter-breeding. The phenotype of mice are screened by standard procedures. Analysis of histological sections prepared from embryonic and adult tissue, magnetic resonance imaging analysis combined with cell death assays such as TdT-mediated dUTP-X nick end labelling (TUNEL) provides direct evidence of the role of

SARG-1 in differentiation and apoptosis, respectively, during development. Cellular analysis includes hematopoietic stem cell characterisation (colony forming assays, flow cytometrical analysis for cluster of differentiation antigens) and apoptotic response measurements in neural culture systems.

Pitfall Analysis

As described above, the pDELBOY-3X vector eliminates artefacts produced by transcription from the neomycin cassette and overcomes the problems of heterozygote embryonic lethality. Screening is also aided by the availability of animals which have a single mutation in the c-kit gene. Identification of SARG-1 binding partners for which knockout mice exist allows breeding with SARG-1 deficient mice to further clarify phenotype.

Cellular Localization

Balb/c mice are mated and embryos isolated at 7, 11 (Theiler's stages 10-19), 14 and 17 days and snap frozen in liquid Nitrogen. 5 µm transverse and sagittal consecutive sections are made on a cryostat and frozen until required. SARG-1 expression is analysed by standard immunohistochemical analysis on sections with anti-SARG-1 IgY which detects acetone-fixed mSARG-1. Expression patterns are identified by reference to standard embryological texts (Kaufman, 1992) and the mouse atlas and gene expression database project (<http://genex.hgu.mrc.ac.uk/>). Expression patterns in 17 days embryos are additionally correlated with those seen in adult murine tissue. Following identification of SARG-1 binding partners, co-expression studies are performed either by immunohistochemistry if antibodies are available or in situ hybridisation.

Subcellular Localization and Biochemical Characterization of Epitope Tagged SARG-1

The cytoplasmic, vesicular subcellular localization of SARG-1 is determined by double staining of HEK293, MelJuso or PC-12 cells stably transfected with epitope-tagged SARG-1. The HA tag is stained with rat monoclonal anti HA mAb and visualised with a biotinylated monoclonal anti-isotype mAb followed by fluorochrome conjugated streptavidin (FITC, S, phycoerythrin or cy5) Organelle localized antigens are detected with antibodies to Bip/GRP78 (endoplasmic reticulum), β-cop (Golgi complex), Lamp-1 (lysosomes) and Ab-2 (mitochondria) either conjugated directly or by the use of second step antibodies and analysed on a Zeiss laser scan microscope. Cloned and native SARG-1 (predicted mass 17 Kd) migrates at 28 Kd in SDS-PAGE. Glycosylation of HA-tagged SARG-1 is initially investigated by immunoprecipitation on an anti-HA matrix and detection of glycosylated residues by periodate oxidation, incorporation of biotin hydrazide and detection with streptavidin conjugated alkaline phosphatase on western blots. N- and O-specific enzymatic deglycosylation reactions are performed with PNGase F and O-Glycosidase (Bio-Rad), respectively, and the molecular weight of SARG-1 monitored by western blotting. The SARG-1 sequence contains consensus protein kinase C and casein kinase II sites which are known to phosphorylate apoptotic regulators (Verma et al, 2001). The phenotypical differences seen between Trk activation by NGF which leads to PKC activation (Patapoutian et al., 2001) and treatment with the protein kinase inhibitor staurosporine in PC-12 possibly suggesting regulation of SARG-1 function by phosphorylation. To examine possible SARG-1 phosphorylation, immunoprecipitated SARG-1 is probed with polyclonal antibodies against phosphothreonine and phosphoserine in Western

blotting analysis and compared to phosphatase-treated protein. Mutant SARG-1 eukaryotic expression plasmids are then generated by site directed mutagenesis to delete potential casein kinase II phosphorylation sites and used to transfect PC-12 cells to monitor the effects of the protein kinase inhibitor staurosporine.

Analysis of Mutation in MS Samples

The immunohistochemical localization of SARG-1 protein to the grey and white matter of the CNS and the role of SARG-1 in apoptotic induction prompted a candidate gene approach to analyse the mutational status of SARG-1 in cases of familial MS. The DNA from twenty unrelated familial multiple sclerosis patients was examined by PCR amplification of the SARG-1 locus and DNA sequencing and compared to SARG-1 sequences from healthy controls. All control samples demonstrated wild type SARG-1 genomic-sequences. From MS patients, only 6 from 20 DNA samples were able to be amplified by PCR. Primer sets used spanned the whole coding region, intron 2/exon 3, and primer pairs specific for exons 1, 2 and 3. Control GAPDH primers were positive for all samples. In 6/20 samples that produced a PCR product, amplicants were TA cloned into the pCR®II vector (Invitrogen) and sequenced. In 4/6 patients, genetic alterations were seen. A T→C point mutation at nucleotide 67 resulting in the substitution of phenylalanine for leucine amino acid 23 (FIG. 16). A C→T point mutation at nucleotide 359 resulting in the substitution of phenylalanine for serine at amino acid 120 (FIG. 17). An A→G point mutation at nucleotide 89 resulting in the substitution of glycine for glutamic acid at amino acid 30 (FIG. 18). Deletion of a codon between amino acids 116 and 121 resulting in the loss of a serine residue (FIG. 19). Sequencing of over 20 control DNA samples revealed only wild type sequence.

Polymorphisms Other Mutations

In sequence analysis of over 30 cancer cell lines, a polymorphism in the human sequence at nucleotide 280 (coding sequence) results in either valine (g residue; shown), methionine (a) or leucine (t) residues at position 94 of the amino acid sequence. In a human melanoma cell line, two mutations in SARG-1 are found: an A→G point mutation at nucleotide 74 resulting in the substitution of aspartic acid for glycine and a C→T point mutation at nucleotide 289 resulting in the substitution of histidine for tyrosine at amino acid 97. Such SARG-1 mutations therefore are suitable markers for human melanomas.

TABLE

67	T → C	=	F → L	23
74	A → G	=	E → G	25
89	A → G	=	E → G	30
289	C → T	=	H → Y	97
359	C → T	=	S → F	120
Δ			S	116-121

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Pro Cys Glu Asp Pro Pro Pro Thr Pro Gln Ser Ser Gly Val Asp Asn
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 cca 1083

<210> SEQ ID NO 4
 <211> LENGTH: 1038
 <212> TYPE: DNA
 <213> ORGANISM: rat

<400> SEQUENCE: 4

ccagactgga agcgaaggct gtgttgctgg gatgccagct gccgaggggc tgcttaagcc 60
 ttggccccc ctactttctg ttccagcccc acttctgtgc gtgttact ccattacccc 120
 caggggctga catggaccca aatccacgag cagccctgga gcgccagcag ctgcgtctca 180
 gggagcggca gaagtcttc gaggacattt tacagccaga gacagagttt gttttcccc 240
 tatcccatct gcactctgag tcacaaagac ccccataggt tagcatctcc tcgatggaag 300

-continued

```

tgaatgtgga cacactggag cagggtggaat ttattgacct tgcggatcag gatggagcag 360
atgtgttctt accttgtgag gattctcctc caactcccca gaggtctgga gtggatgacc 420
accagagga gctgagcctg ctggtaccca cgtcagacag gaccacatcc cggacctcct 480
ccttgcctc tgactcctcc aacctgcgca gtccaaatcc aagtgatggg ggaggagaca 540
ctcccttggc acagtctgac gaggaggatg gggacggtgg aggggcagaa cctggacctt 600
gcagctagca gaggccctt acaaactgag cgatctggct gttctccatg gagaggagac 660
cttaggtcca ccagagcact ctggagaaga cctgacactt tacttacatc agcaccaaag 720
ggaggggaagg atggtggatg gtgtgcctga gagttagcct ccccgttta ctgataacgc 780
tgtctgctg ccacgcccc acagtgtctt cttctgaggt aggacttcca agtgagactc 840
tcgaaggta ggtgggacaa gatgccactg ttttcttact cccctcctgc ccccaaatga 900
tcctgtagtc tccactagt ctcctaagcc agtgtctctg agggaaagt ctgaggagtt 960
ccactttgca gttatcctgc ctctataagt ctttctggg aacaggatat ggtataaata 1020
ataaataata ctgtacca 1038
    
```

```

<210> SEQ ID NO 5
<211> LENGTH: 486
<212> TYPE: DNA
<213> ORGANISM: rat
    
```

<400> SEQUENCE: 5

```

atggacccaa atcctcgggc cgccctggag cgccagcagc tccgccttcg ggagcggcaa 60
aaattcttcg aggacattt acagccagag acagagtttg tcttctctct gtcccatctg 120
catctcgagt cgcagagacc ccccataggt agtatctcat ccatggaagt gaatgtggac 180
acactggagc aagtagaact tattgacctt ggggaccgag atgcagcaga tgtgttcttg 240
ccttgcaag atcctccacc aacccccag tcgtctggga tggacaacca tttggaggag 300
ctgagcctgc cgtgacctac atcagacagg accacatcta ggacctctc ctctccttc 360
tccgactcct ccaccaacct gcatagccca aatccaagt atgatggagc agatacgccc 420
ttggcacagt cggatgaaga ggaggaaagg ggtgatggag gggcagagcc tggagcctgc 480
agctag 486
    
```

```

<210> SEQ ID NO 6
<211> LENGTH: 158
<212> TYPE: PRT
<213> ORGANISM: rat
    
```

<400> SEQUENCE: 6

```

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1             5             10             15
Arg Glu Arg Gln Lys Phe Phe Glu Asp Ile Leu Gln Pro Glu Thr Glu
          20             25             30
Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
          35             40             45
Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
          50             55             60
Val Glu Phe Ile Asp Leu Ala Asp Gln Asp Gly Ala Asp Val Phe Leu
          65             70             75             80
Pro Cys Glu Asp Ser Pro Pro Thr Pro Gln Arg Ser Gly Val Asp Asp
          85             90             95
His Pro Glu Glu Leu Ser Leu Leu Val Pro Thr Ser Asp Arg Thr Thr
    
```

-continued

100				105				110							
Ser	Arg	Thr	Ser	Ser	Leu	Ser	Ser	Asp	Ser	Ser	Asn	Leu	Arg	Ser	Pro
	115						120					125			
Asn	Pro	Ser	Asp	Gly	Gly	Gly	Asp	Thr	Pro	Leu	Ala	Gln	Ser	Asp	Glu
	130						135					140			
Glu	Asp	Gly	Asp	Gly	Gly	Gly	Ala	Glu	Pro	Gly	Pro	Cys	Ser		
	145					150					155				

<210> SEQ ID NO 7
 <211> LENGTH: 1002
 <212> TYPE: DNA
 <213> ORGANISM: mouse

<400> SEQUENCE: 7

```

ggatccaacg ccggcgctgc tcgctccac gcccccgcg ccgcttctcg ggagcgcacc    60
cagggagcca gcggggcgcg ggcgctgcag gggctgacat ggacccaat ccgagagcag    120
ccctggagcg ccaacagctg cggctccggg agaggcagaa gttctttgag gacattttac    180
agccagagac agagtttgtc ttccccctgt cccatctgca cctggagtca caaagacccc    240
ccataggtag catctcgtct atggaagtga atgtggacac actggagcaa gtggagtta    300
ttgatcttgc ggatcaggat ggagcagatg tgttcttgc ttgtgaggag tcctcgccag    360
ctccccagat gtctggagtg gatgaccatc cagaggagct gagcctgctg gtaccacagt    420
ctgacaggac cacatcccgg acctcctcct tgcctctga ctccccaac ctgcgagtc    480
caaatccaag tgatggggga ggagacactc ccttggcaca gtctgatgag gaggacgggg    540
atgacggagg ggcagagcct ggaccctgca gctagcagtg ggcctctac agactgacca    600
gcccggctgt tctccatgga aaggagacct aggccagca gagcctggag aagacctgac    660
actttcctta cttcagcacc aaagggaggg aaggatggtg gatggtgtgc ctgagagtta    720
gctccccctg ctttaccgta acgctatcct gctgccacgc ccccacagtg cttttcttct    780
gaggtaggac ttccaagtga gacttgagag gtgaggtggg acaagacgca gctgctttct    840
tagtccccctc ctgccccag atgacctgtg tgtcttcac agagtctcct aagccagtgt    900
ctctgagggg atgttctgag gagttccact ttccagttat cctgcctcta taagttcttt    960
tggaacag ataggtata aataataat aataatatac ca                               1002

```

<210> SEQ ID NO 8
 <211> LENGTH: 477
 <212> TYPE: DNA
 <213> ORGANISM: mouse

<400> SEQUENCE: 8

```

atggacccaa atccgagagc agccctggag cgccaacagc tgcggctccg ggagaggcag    60
aagttctttg aggacatttt acagccagag acagagtttg tcttcccct gtccatctg    120
cacctggagt cacaaagacc ccccataggt agcatctcgt ctatggaagt gaatgtggac    180
aactggagc aagtggagtt tattgatctt gcggatcagg atggagcaga tgtgttcttg    240
ccttgtagg agtcctcgcc agctcccag atgtctggag tggatgacca tccagaggag    300
ctgagcctgc tggtaaccac gtctgacagg accacatccc ggacctctc cttgtcctct    360
gactcctcca acctgcgcag tccaaatcca agtgatgggg gaggagacac tcccttgcca    420
cagtctgatg aggaggacgg ggatgacgga ggggcagagc ctggaccctg cagctag    477

```

<210> SEQ ID NO 9

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<211> LENGTH: 158
 <212> TYPE: PRT
 <213> ORGANISM: mouse

<400> SEQUENCE: 9

```

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1           5           10           15
Arg Glu Arg Gln Lys Phe Phe Glu Asp Ile Leu Gln Pro Glu Thr Glu
           20           25           30
Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
           35           40           45
Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
           50           55           60
Val Glu Phe Ile Asp Leu Ala Asp Gln Asp Gly Ala Asp Val Phe Leu
           65           70           75           80
Pro Cys Glu Glu Ser Ser Pro Ala Pro Gln Met Ser Gly Val Asp Asp
           85           90           95
His Pro Glu Glu Leu Ser Leu Leu Val Pro Thr Ser Asp Arg Thr Thr
           100          105          110
Ser Arg Thr Ser Ser Leu Ser Ser Asp Ser Ser Asn Leu Arg Ser Pro
           115          120          125
Asn Pro Ser Asp Gly Gly Gly Asp Thr Pro Leu Ala Gln Ser Asp Glu
           130          135          140
Glu Asp Gly Asp Asp Gly Gly Ala Glu Pro Gly Pro Cys Ser
           145          150          155

```

<210> SEQ ID NO 10
 <211> LENGTH: 161
 <212> TYPE: PRT
 <213> ORGANISM: human

<400> SEQUENCE: 10

```

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1           5           10           15
Arg Glu Arg Gln Lys Phe Leu Glu Asp Ile Leu Gln Pro Glu Thr Glu
           20           25           30
Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
           35           40           45
Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
           50           55           60
Val Glu Leu Ile Asp Leu Gly Asp Pro Asp Ala Ala Asp Val Phe Leu
           65           70           75           80
Pro Cys Glu Asp Pro Pro Pro Thr Pro Gln Ser Ser Gly Val Asp Asn
           85           90           95
His Leu Glu Glu Leu Ser Leu Pro Val Pro Thr Ser Asp Arg Thr Thr
           100          105          110
Ser Arg Thr Ser Ser Ser Ser Ser Ser Asp Ser Ser Thr Asn Leu His
           115          120          125
Ser Pro Asn Pro Ser Asp Asp Gly Ala Asp Thr Pro Leu Ala Gln Ser
           130          135          140
Asp Glu Glu Glu Glu Arg Gly Asp Gly Gly Ala Glu Pro Gly Ala Cys
           145          150          155          160
Ser

```

<210> SEQ ID NO 11
 <211> LENGTH: 161

-continued

```

<212> TYPE: PRT
<213> ORGANISM: human

<400> SEQUENCE: 11

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1           5           10           15

Arg Glu Arg Gln Lys Phe Phe Glu Asp Ile Leu Gln Pro Glu Thr Glu
  20           25           30

Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
  35           40           45

Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
  50           55           60

Val Glu Leu Ile Asp Leu Gly Asp Pro Asp Ala Ala Asp Val Phe Leu
  65           70           75           80

Pro Cys Glu Asp Pro Pro Pro Thr Pro Gln Ser Ser Gly Val Asp Asn
  85           90           95

His Leu Glu Glu Leu Ser Leu Pro Val Pro Thr Ser Asp Arg Thr Thr
  100          105          110

Ser Arg Thr Ser Ser Ser Ser Phe Ser Asp Ser Ser Thr Asn Leu His
  115          120          125

Ser Pro Asn Pro Ser Asp Asp Gly Ala Asp Thr Pro Leu Ala Gln Ser
  130          135          140

Asp Glu Glu Glu Glu Arg Gly Asp Gly Gly Ala Glu Pro Gly Ala Cys
  145          150          155          160

Ser

```

```

<210> SEQ ID NO 12
<211> LENGTH: 161
<212> TYPE: PRT
<213> ORGANISM: human

<400> SEQUENCE: 12

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1           5           10           15

Arg Glu Arg Gln Lys Phe Phe Glu Asp Ile Leu Gln Pro Gly Thr Glu
  20           25           30

Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
  35           40           45

Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
  50           55           60

Val Glu Leu Ile Asp Leu Gly Asp Pro Asp Ala Ala Asp Val Phe Leu
  65           70           75           80

Pro Cys Glu Asp Pro Pro Pro Thr Pro Gln Ser Ser Gly Val Asp Asn
  85           90           95

His Leu Glu Glu Leu Ser Leu Pro Val Pro Thr Ser Asp Arg Thr Thr
  100          105          110

Ser Arg Thr Ser Ser Ser Ser Ser Ser Asp Ser Ser Thr Asn Leu His
  115          120          125

Ser Pro Asn Pro Ser Asp Asp Gly Ala Asp Thr Pro Leu Ala Gln Ser
  130          135          140

Asp Glu Glu Glu Glu Arg Gly Asp Gly Gly Ala Glu Pro Gly Ala Cys
  145          150          155          160

Ser

```

-continued

<210> SEQ ID NO 13
 <211> LENGTH: 160
 <212> TYPE: PRT
 <213> ORGANISM: human

<400> SEQUENCE: 13

```

Met Asp Pro Asn Pro Arg Ala Ala Leu Glu Arg Gln Gln Leu Arg Leu
  1           5           10           15

Arg Glu Arg Gln Lys Phe Phe Glu Asp Ile Leu Gln Pro Glu Thr Glu
  20           25           30

Phe Val Phe Pro Leu Ser His Leu His Leu Glu Ser Gln Arg Pro Pro
  35           40           45

Ile Gly Ser Ile Ser Ser Met Glu Val Asn Val Asp Thr Leu Glu Gln
  50           55           60

Val Glu Leu Ile Asp Leu Gly Asp Pro Asp Ala Ala Asp Val Phe Leu
  65           70           75           80

Pro Cys Glu Asp Pro Pro Pro Thr Pro Gln Ser Ser Gly Val Asp Asn
  85           90           95

His Leu Glu Glu Leu Ser Leu Pro Val Pro Thr Ser Asp Arg Thr Thr
  100          105          110

Ser Arg Thr Ser Ser Ser Ser Ser Asp Ser Ser Thr Asn Leu His Ser
  115          120          125

Pro Asn Pro Ser Asp Asp Gly Ala Asp Thr Pro Leu Ala Gln Ser Asp
  130          135          140

Glu Glu Glu Glu Arg Gly Asp Gly Gly Ala Glu Pro Gly Ala Cys Ser
  145          150          155          160
    
```

The invention claimed is:

1. A method for diagnosing in a patient multiple sclerosis (MS) or the increased risk for acquiring MS, wherein the method comprises the steps of
 - providing a sample of a body fluid or tissue from the patient; and
 - testing the sample to determine whether the sample contains a wild type SCF-Apoptosis-Response Gene-1 protein (SARG-1 protein) (SEQ ID NO. 2), whereby the absence of a wild type SARG-1 protein in the sample indicates that the patient has MS or an increased risk of acquiring MS.
2. The method according to claim 1, wherein the sample does not contain a SARG-1 protein.
3. The method according to claim 1, wherein the sample contains a mutant SARG-1 protein.
4. The method according to claim 1, wherein the testing is performed by contacting the sample with an antibody against the wild type SARG-1-protein.
5. A method for diagnosing in a patient multiple sclerosis (MS) or the increased risk for acquiring MS, wherein the method comprises the steps of
 - providing a sample of a body fluid or tissue from the patient; and
 - testing the sample to determine whether the sample contains nucleic acid (SEQ ID NO. 1) encoding a wild type SCF-Apoptosis-Response Gene-1 protein (SARG-1 protein) (SEQ ID NO. 2), whereby the absence of a nucleic acid encoding a wild type SARG-1 protein in

the sample indicates that the patient has MS or an increased risk of acquiring MS.

6. The method according to claim 5, wherein the sample does not contain a nucleic acid encoding a SARG-1 protein.
7. The method according to claim 5, wherein the sample contains a nucleic acid encoding a mutant SARG-1 protein.
8. The method according to claim 5, wherein the testing is performed by one or more techniques selected from the group consisting of a nucleic acid amplification, polymerase chain reaction (PCR), single-strand conformation polymorphism (SSCP) analysis, restriction analysis, microarray technology and proteomics.
9. The method according to claim 1, wherein the sample is derived from human blood, plasma, serum, lymph, nerve-cell containing tissue, cerebrospinal fluid, all biopsy-material, including tumor tissue, bone marrow, nervous tissue, skin, hair, tears, fetal material, amniocentesis material, uterine tissue, saliva, feces or sperm.
10. The method according to claim 5, wherein the sample is derived from human blood, plasma, serum, lymph, nerve-cell containing tissue, cerebrospinal fluid, all biopsy-material, including tumor tissue, bone marrow, nervous tissue, skin, hair, tears, fetal material, amniocentesis material, uterine tissue, saliva, feces or sperm.
11. The method according to claim 4, wherein the wt-SARG-1-protein antibody is a monoclonal antibody.

* * * * *

专利名称(译)	诊断患有多发性硬化症的人的方法		
公开(公告)号	US7208270	公开(公告)日	2007-04-24
申请号	US10/176372	申请日	2002-06-21
[标]申请(专利权)人(译)	严实BURKHARD 卢卡斯特里沃		
申请(专利权)人(译)	严实BURKHARD 卢卡斯特里沃		
当前申请(专利权)人(译)	严实, BURKHARD LUCAS, TREVOR		
[标]发明人	JANSEN BURKHARD LUCAS TREVOR		
发明人	JANSEN, BURKHARD LUCAS, TREVOR		
IPC分类号	C12Q1/68 C12P19/34 G01N33/53 A01K67/027 A61K38/00 A61K38/17 A61P25/00 A61P35/00 A61P35/02 C07K1/22 C07K14/47 C07K16/18 C12N15/09 C12P21/08 C12Q1/6883 G01N33/48 G01N33/564 G01N33/566 G01N33/574 G01N33/577 G01N33/68 G01N37/00		
CPC分类号	A61K38/1709 C07K1/22 C07K14/47 C07K14/4713 C07K16/18 G01N33/564 G01N33/57484 G01N33/6893 G01N33/6896 C12Q1/6883 A01K2217/05 C12Q2600/156 G01N2800/285 A61P25/00		
代理机构(译)	普洛思律师事务所		
审查员(译)	霍利克, 肯尼斯R.		
优先权	60/299765 2001-06-22 US		
其他公开文献	US20030113752A1		
外部链接	Espacenet USPTO		

摘要(译)

描述了一种诊断患有多发性硬化症 (MS) 或有发展MS风险的人的方法, 包括以下步骤: 从所述人提供体液或组织样品, 所述样品含有至少一种野生型SCF-凋亡 - 反应基因 - (wt-SARG-1-) 蛋白和编码wt-SARG-1的核酸, 如果取自没有MS或有获得MS的风险的人, 检测wt-SARG-1的存在 - 在所述样品中编码wt-SARG-1的蛋白质或核酸 如果wt-SARG-1-蛋白或编码wt-SARG-1的核酸不存在于所述样品中, 则诊断MS或获得MS的风险。

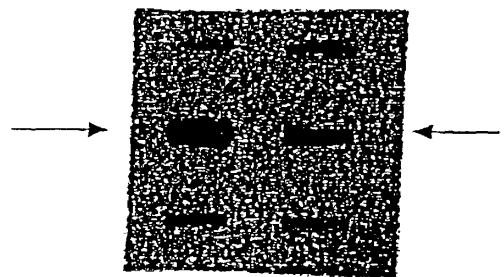


FIG. 1