



SAMD9L (NM_152703.5) - cDNA + Protein - 2024-05-18

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MetSer LysGlnValS 6

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TGAAGAAGTA ACAGGATTAG TCCTGCAGGA ATTAAGTGA AAGGACCTTG 166
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alGluMetG1 yLeuProTrp GlyProAlaL euLeuIleLy sArgSerTyr 72

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AsnLysLeuA snSerLysSe rProGluSer AspAsnHisA spProGlyG1 89

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isThrLysLy sGluGluGlu AsnSerMetS erSerAsnI1 eAspTyrAsp 122

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ProArgGluI leArgAspI1 eLysGlnGlu GluSerIleL euMetLysG1 139

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uAsnValLeu AspGluValA laAsnAlaLy sHisLysLys LysGlyLysL 156

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euLysProGl uGlnLeuThr CysMetProT yrProPheAs pGlnPheHis 172

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AspSerHisA rgTyrIleGl uHisTyrThr LeuGlnProG luThrGlyAl 189

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luThrAlaTh rGluValAsp IleLysMetL ysPheSerAs nGluValPhe 222

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ArgPheAlaS erAlaCysMe tAsnSerArg ThrAsnGlyT hrIleHisPh 239

TGGAGTCAAG GACAAACCCC ATGGAGAAAT TGTGGGTGTG AAAATCACCA 766
eGlyValLys AspLysProH isGlyGluIl eValGlyVal LysIleThrS 256

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erLysAlaAl aPheIleAsp HisPheAsnV alMetIleLy sLysTyrPhe 272

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GluGluSerG luIleAsnGl uAlaLysLys CysIleArgG luProArgPh 289

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 eValGluVal LeuLeuGlnA snAsnThrPr oSerAspArg PheValIleG 306

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 luValAspTh rIleProLys HisSerIleC ysAsnAspLy sTyrPheTyr 322

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***SAMD9L* (NM_152703.5) - cDNA + Protein - 2024-05-18**

