

Biochemical characterization, localization, and tissue distribution of the longer form of mouse SIRT3

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Abstract: SIRT3 is a key mitochondrial protein deacetylase proposed to play key roles in regulating mitochondrial metabolism but there has been considerable debate about its actual size, the sequences required for activity, and its subcellular localization. A previously cloned mouse SIRT3 has high sequence similarity with the C-terminus of human SIRT3 but lacks an N-terminal mitochondrial targeting sequence and has no detectable deacetylation activity *in vitro*. Using 5' rapid amplification of cDNA ends, we cloned the entire sequence of mouse SIRT3, as well as rat and rabbit SIRT3. Importantly, we find that full-length SIRT3 protein localizes exclusively to the mitochondria, in contrast to reports of SIRT3 localization to the nucleus. We demonstrate that SIRT3 has no deacetylation activity *in vitro* unless the protein is truncated, consistent with human SIRT3. In addition, we determined the inhibition constants and mechanism of action for nicotinamide and a small molecule SIRT3 inhibitor against active mouse SIRT3 and show that the mechanisms are different for the two compounds with respect to peptide substrate and NAD⁺. Thus, identification and characterization of the actual SIRT3 sequence should help resolve the debate about the nature of mouse SIRT3 and identify new mechanisms to modulate enzymatic activity.

Keywords: sirtuin; NAD⁺-dependent deacetylation; inhibitor; mitochondrial function

Introduction

The silent information regulator 2 (Sir2 or sirtuin) proteins comprise a family of NAD⁺-dependent protein deacetylases and ADP-ribosyltransferases that are

highly conserved in both prokaryotes and eukaryotes.¹ Sirtuins are involved in many cellular processes including gene silencing,² cell cycle regulation,^{3,4} metabolism,^{5–8} apoptosis^{9–11} and the lifespan-extension

Abbreviations: AceCS2, acetyl-CoA synthetase 2; GDH, glutamate dehydrogenase; MPP, mitochondrial matrix processing peptidase; mSIRT3_L, a longer form of mouse SIRT3; NAM, nicotinamide.

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effects of calorie restriction.^{12,13} Mammals express seven sirtuins, SIRT1-7.¹⁴ SIRT1, 2, 3, and 5 possess deacetylase activity, whereas SIRT4 and 6 are ADP-ribosyltransferases. These proteins display a range of subcellular localization, including nuclear (SIRT1, 6, 7), cytosolic (SIRT2),¹⁵ and mitochondrial stores (SIRT3, 4, 5)¹⁶. These diverse locations in the cell reflect varying functions. SIRT1 has been studied most extensively and has been linked to glucose homeostasis,¹⁷⁻¹⁹ inflammation,²⁰ cancer,²¹⁻²³ and neuropathology.^{24,25} We have previously described small molecule activators of SIRT1 that improve insulin sensitivity, lower plasma glucose and increase mitochondria capacity in diet-induced obese and genetically obese mice.¹⁹

SIRT3 is a major mitochondrial deacetylase. Mitochondrial proteins show hyperacetylation in SIRT3 knockout mice, but not in SIRT4 or SIRT5 knockout mice.²⁶ Acetyl-CoA synthetase 2 (AceCS2), a mitochondrial enzyme that converts acetate into acetyl-CoA, was the first mitochondrial substrate of SIRT3 identified.^{6,27} Deacetylation of AceCS2 at lysine 642 by SIRT3 activates acetyl-CoA synthetase activity, providing increased acetyl-CoA to feed into the tricarboxylic acid cycle. Glutamate dehydrogenase (GDH), another mitochondrial protein involved in energy production, is deacetylated by SIRT3.²⁶ GDH can also be ADP-ribosylated by SIRT4 in turn to decrease its enzyme activity.²⁸ This indicates that SIRT3 could play an important role in cell metabolism. SIRT3 has also been shown to be involved in selective apoptosis pathways and cell growth control.²⁹ SIRT3 and SIRT4, but not SIRT5, have been implicated in the NAD⁺ salvage pathway that regulates the NAD⁺ level relating to cell survival.³⁰ In addition, variability in the hSIRT3 gene has been linked to human longevity.^{31,32}

Human SIRT3 (hSIRT3) has been shown to localize to mitochondria³³⁻³⁵ in a majority of the reported studies. One controversial report suggests that full length hSIRT3 is a nuclear protein under basal conditions but translocates into mitochondria during cellular stress including over-expression of hSIRT3.³⁶ A more recent report disagrees and shows full length hSIRT3 is localized only in mitochondria with either transient or stable over-expression in various human cell lines, and the hSIRT3 lacking the N-terminal 142 residues is present in the cytoplasm and nucleus at high expression levels.³⁵

The N-terminus of hSIRT3 contains a mitochondrial targeting sequence, an amphipathic α -helix rich in basic residues.³⁴ The full length hSIRT3 is enzymatically inactive. When the first 101 residues are cleaved by mitochondrial matrix processing peptidase (MPP) *in vitro*, the protein becomes activated. Both a full-length 44 kDa hSIRT3 and a smaller 28 kDa product were observed in whole-cell lysates of HEK293T cells transfected with FLAG tagged hSIRT3, whereas a 28-kDa product was the only form detected in the mitochondrial lysates. Only SIRT3 purified from fractionated

mitochondrial lysates showed NAD⁺-dependent protein deacetylase activity.³³ However, Scher *et al.*³⁶ reported that both the full length and 142-residue truncated hSIRT3 have similar deacetylation activity and substrate specificity.

Mouse SIRT3 (mSIRT3) was first cloned by Yang *et al.*³⁷ It was reported to be a 257 amino acid protein that aligned well with the C-terminal portion of hSIRT3 (residues 143-399) and localized in certain areas of the cytoplasm in discrete stacks. Mouse SIRT3 is expressed in many tissues, with high expression in brain, kidney, testes, heart and liver and is also expressed in brown adipose tissue.³⁸ The expression is up-regulated during caloric restriction and cold exposure. Although the reported mSIRT3 does not contain the N-terminal mitochondrial targeting sequence, Shi *et al.* showed that over-expressed mSIRT3 is localized to the inner mitochondrial membrane of NIH3T3 cells.³⁸ In a more recent report, when subcellular fractions prepared from mouse liver were probed with SIRT3-specific antibody, mSIRT3 is found to be a soluble mitochondrial protein, as with hSIRT3, under both basal and stress conditions.²⁶

Here we report the cloning of a longer form of mSIRT3 (mSIRT3_L), its tissue distribution, subcellular localization, the fragments that retain deacetylation activity, as well as its inhibition by nicotinamide and SRT1720, a small molecule SIRT3 inhibitor. Our rationale for attempting to clone a longer mSIRT3 was the lack of deacetylase activity for the recombinant short mSIRT3 with 257 amino acids and the equivalent hSIRT3-143-399. We also questioned the mitochondrial localization of mSIRT3 given the lack of an obvious mitochondrial targeting sequence. The long mSIRT3 that we have cloned matches the predicted mSIRT3 reported by Cooper and Spelbrink.³⁵

Results and Discussion

Cloning of the long mouse SIRT3

To identify cDNA that encodes the longer mouse SIRT3, we performed 5'-RACE PCR with mRNA from mouse liver. Two sequences were identified with an 8 base pair (bp) difference (TGTTACAG) at the 5' end of exon 2 of the mouse SIRT3 gene on chromosome 7 [Fig. 1(A)]. The sequence containing the 8 bp insertion encodes an open reading frame (ORF) for a protein of 257 residues (gene BC025878, protein AAH258219). This is the reported short form of mSIRT3 without the mitochondrial targeting sequence. Interestingly, the sequence lacking the 8 bp insertion encodes a much longer ORF for a protein of 334 residues (matching GenBank EST database AW258219). This longer ORF [Fig. 1(B)] is the same as predicted by Cooper and Spelbrink.³⁵ Both sequences had been reported by Yang *et al.*³⁷ previously. In their study, the longer ORF was not detected by reverse transcription-PCR (RT-

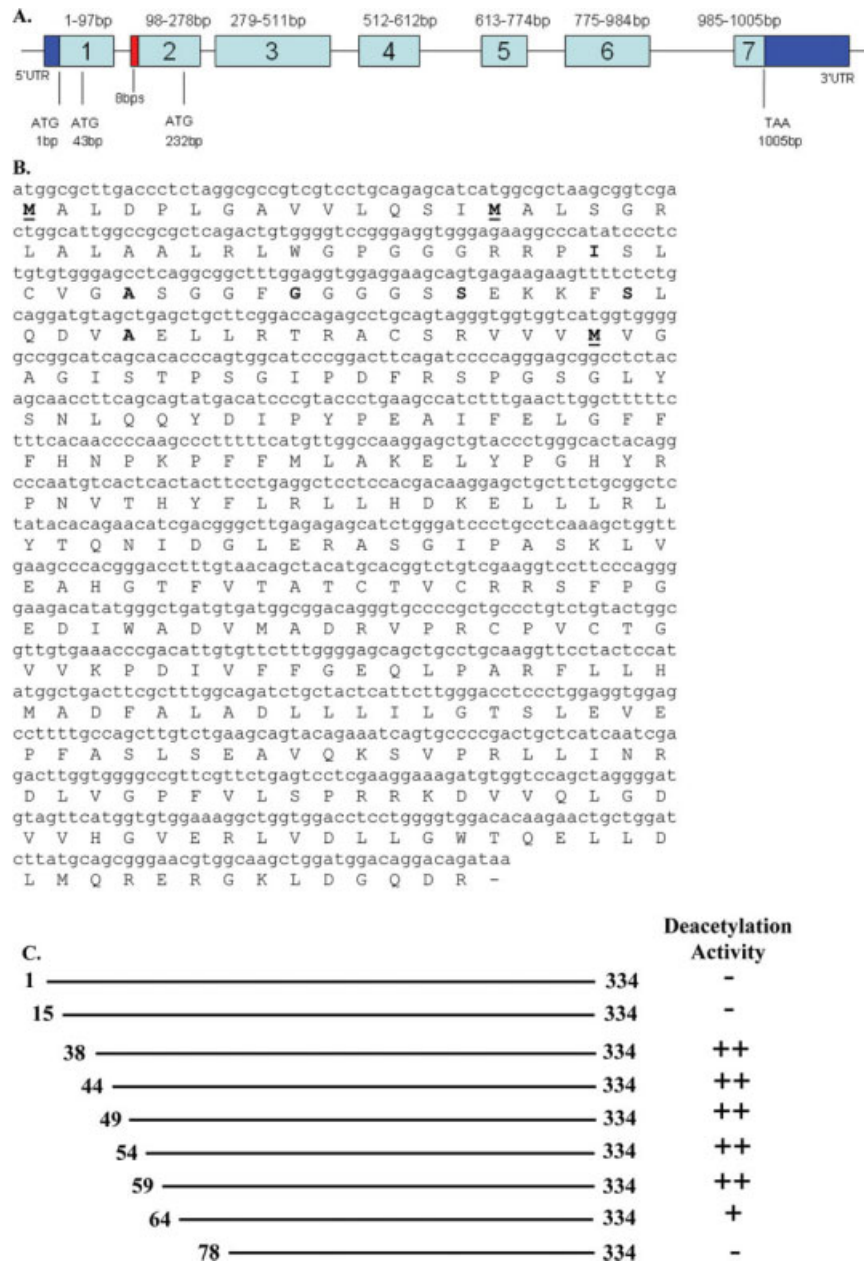


Figure 1. The cDNA and amino acid sequence of the longer mouse SIRT3 gene. (A) Schematic representation of the mouse SIRT3 gene. The exons are presented as rectangular boxes with the exon number labeled in each box and the base pair numbers of the coding sequences are labeled above each box. The 5' and 3' untranslated regions (UTR) are colored in dark blue. Coding sequences are colored in light blue. The location of the eight base pair insertion is colored in red. The three potential start codons and the stop codon are indicated. (B) The sequence of long mSIRT3. The three methionine residues that have been considered as the starting residue are in bold character and underlined. The other residues in bold are the starting residues for different N-terminal truncations that we generated and listed in Figure 1(C). (C) Schematic representation of all the mSIRT3 proteins that were generated for enzyme activity testing. The starting and ending residue numbers are labeled. The deacetylation activity for each protein is indicated as “-” no activity, “++” high activity, and “+” low activity.

PCR) in mouse tissues. The additional sequence was considered to have weak identity to human SIRT3 and its existence was dismissed. The two variants of mSIRT3 gene could be caused by alternative splicing. Alternative splicing at donor or acceptor sites located just a few nucleotides apart has been reported to be widespread in many species and results in subtle

changes in the transcripts that could cause changes in the encoded proteins.³⁹ We have not yet determined the frequencies of the two variants in mouse and what controls the splicing.

The longer mSIRT3 has three methionine residues at positions 1, 15, and 78 near the N-terminus. Residue Met78 is the first residue for the reported short

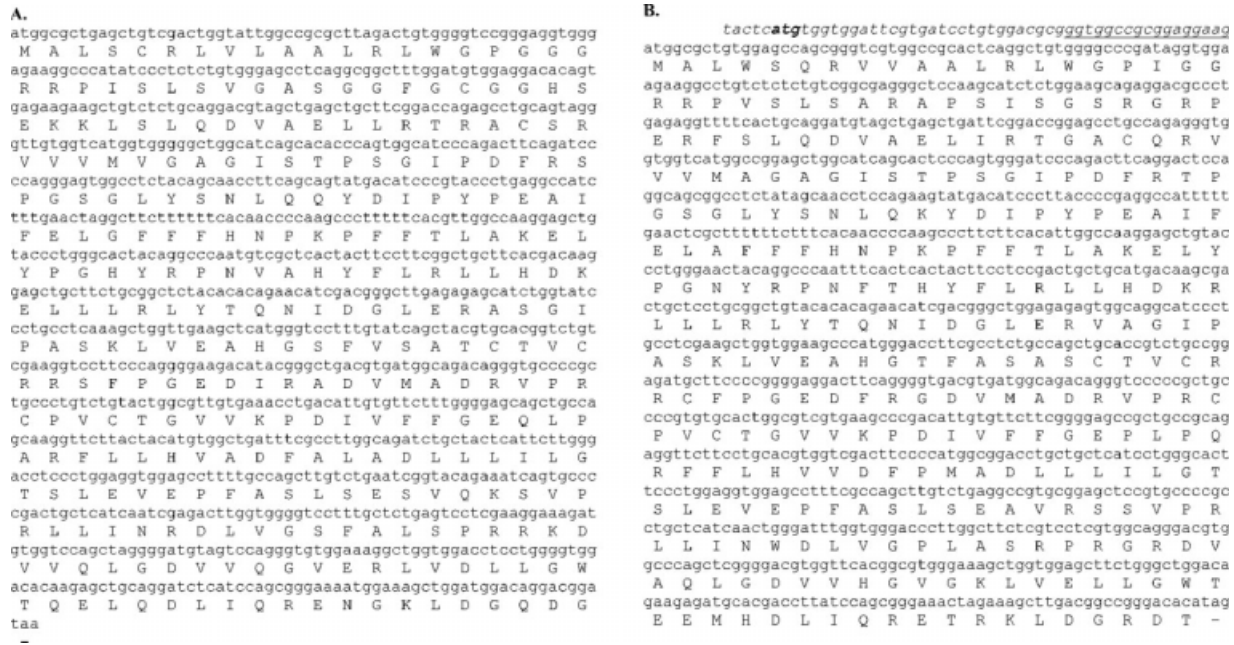


Figure 2. The cDNA and amino acid sequences of rat and rabbit SIRT3. (A) The sequence of rat SIRT3. (B) The sequence of rabbit SIRT3. The italic DNA sequence before the coding sequence was obtained by a BLAST search of the rabbit genome. The underlined portion was obtained from our 5'-RACE PCR. The potential start codon (ATG) before the coding codon is in bold.

mSIRT3 containing 257 amino acids. When we aligned the mSIRT3_L with human SIRT3 by TCOFFEE⁴⁰ (Fig. 3), it was Met15 (not Met1) of mSIRT3_L that aligned with Met1 of hSIRT3. Residues 15–30 of mSIRT3_L have high homology with the N-terminus of hSIRT3, whereas there is a 77-residue insertion in hSIRT3 that is absent in mSIRT3_L. The rest of the additional sequence in the mSIRT3_L (residues 31–77) is highly similar to hSIRT3. It is unclear which methionine, Met1 or Met15, is the starting residue for the mSIRT3_L (discussed in more detail below).

Cloning of SIRT3 from rat and rabbit

The SIRT3 of rat and rabbit have not been previously cloned. We performed RACE reactions as we did with cloning mSIRT3_L. The predicted rat SIRT3 (XM_215124) has 257 amino acids like the short mouse SIRT3. Based on the rat genome, rat has the sequence equivalent to mSIRT3_L-1-334. We performed 5'-RACE PCR multiple times and obtained two sequences with or without the 8 bps insertion (AGTATTAC) like the 5'-RACE PCR result for mSIRT3. The sequence without the 8 bps codes for 320 amino acid protein [Fig. 2(A)] and has high identity with mSIRT3-15-334 (Fig. 3).

The sequence we obtained for rabbit SIRT3 contains 319 amino acids [Fig. 2(B)] and aligned well with mouse SIRT3-15-334 (Fig. 3). A matching sequence was found by a BLAST search of the rabbit genome showing additional 5' sequence upstream of the coding sequence. We cloned 17 bps before the coding start codon ATG in our 5'-RACE product. There is another

ATG in the upstream 5' sequence [Fig. 2(B)] that is not in the current ORF (44 bps upstream). Considering the high similarity of rabbit SIRT3 with SIRT3 from human, mouse and rat, we believe that we have obtained the full-length rabbit SIRT3.

Mouse SIRT3 tissue distribution

We performed real-time RT-PCR against RNA from different mouse tissues with primers and a probe which is derived from the N-terminal region of mSIRT3_L that is not present in the short mSIRT3. The mRNA expression of mSIRT3_L is highest in kidney, brain and heart, followed by liver and testes, and is relatively low in lung, ovary and thymus (Fig. 4). We also performed RT-PCR using primers and a probe designed against the short mSIRT3 sequence and obtained a similar result (data not shown). Our results are consistent with previous studies of mSIRT3 mRNA expression.^{26,33,37,38}

Long mSIRT3 is a mitochondrial protein

The predicted longer form of mSIRT3 may contain N-terminal mitochondrial targeting sequences that are also present in human SIRT3. As noted by Cooper *et al.*, this longer mSIRT3 could begin at the most N-terminal methionine (Met1) or a second downstream methionine (Met15).³⁵ Therefore, we compared the localization of both of these proteins (mSIRT3_L-1-334 and mSIRT3_L-15-334) to that of the originally predicted mouse SIRT3 (mSIRT3_L-78-334). To determine the subcellular distribution of these proteins we

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humanSIRT3      -----MAFWGWRAAAALRLWGRVVERVEAGGGVGFQACGCRLVLGGRDDV
rabbitSIRT3     -----MALWSQRVVAALRLWGP-----
mouseSIRT3      MALDPLGAVVLQSIMALSGRLAALRLWGP-----
ratSIRT3        -----MALSCRLVLAALRLWGP-----
                ** : . *****

humanSIRT3      SAGLRGSHGARGEPLDPARPLQRPPRPEVPPRAFRRQPRAAAPSFFSSIKGRRRSISFSV
rabbitSIRT3     -----IGGRRFPVLSA-----
mouseSIRT3      -----GGGRRPISLCV-----
ratSIRT3        -----GGGRRPISLSV-----
                ****.:*:.

humanSIRT3      GASSVVGSGSSDKGLSLQDVAELIRARACQRVVVMVGAGISTPSGIPDFRSPGSGLYS
rabbitSIRT3     RAPSISGSRGRPE--RFSLQDVAELIRTGACQRVVVMVGAGISTPSGIPDFRTPGSGLYS
mouseSIRT3      GASGGFPGGGSSSEK-KFSLQDVAELLRTRACSRVVVMVGAGISTPSGIPDFRSPGSGLYS
ratSIRT3        GASGGFPGCGHSEK-KLSLQDVAELLRTRACSRVVVMVGAGISTPSGIPDFRSPGSGLYS
                *.. * * * .: :*****:*: *.*****.*****:*****

humanSIRT3      NLQQYDLPYPEAIFELPFFHNPKPFFTLAKELYPGNYKPNVTHYFLRLLHDKGLLRLY
rabbitSIRT3     NLQKYDIPYPEAIFELAFFHNPKPFFTLAKELYPGNYRPNVTHYFLRLLHDKRLLRLY
mouseSIRT3      NLQQYDIPYPEAIFELGFFHNPKPFFMLAKELYPGHYRPNVTHYFLRLLHDKELLRLY
ratSIRT3        NLQQYDIPYPEAIFELGFFHNPKPFFTLAKELYPGHYRPNVAHYFLRLLHDKELLRLY
                ***:*:***** ***** *****:*:*.***** *****

humanSIRT3      TQNIDGLERVSGIPASKLVEAHGTFASATCTVCRQRFPPGEDIRADVMADRVPRCPVCTGV
rabbitSIRT3     TQNIDGLERVAGIPASKLVEAHGTFASASCTVCRRCFPGEDFRGDVMADRVPRCPVCTGV
mouseSIRT3      TQNIDGLERASGIPASKLVEAHGTFVTATCTVCRRSFPGEDIWADVMADRVPRCPVCTGV
ratSIRT3        TQNIDGLERASGIPASKLVEAHGSFVSATCTVCRRSFPGEDIRADVMADRVPRCPVCTGV
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humanSIRT3      VKPDIVFFGEPLPQRFLLHVDFPMADLLLILGTSLEVEPFASLSEAVRSSVPRLLINRD
rabbitSIRT3     VKPDIVFFGEPLPQRFLLHVDFPMADLLLILGTSLEVEPFASLSEAVRSSVPRLLINWD
mouseSIRT3      VKPDIVFFGEQLPARFLLHMADFALADLLLILGTSLEVEPFASLSEAVQKSVPRLLINRD
ratSIRT3        VKPDIVFFGEQLPARFLLHVADFALADLLLILGTSLEVEPFASLSESVQKSVPRLLINRD
                ***** * * *:*:*.*****:*****:*:*.***** *

humanSIRT3      LVGPLAWHPRSRDVAQLGDVVHGVESLVELLWTEEMRDLVQRETGKLDGPDK
rabbitSIRT3     LVGPLASRPRGRDVAQLGDVVHGVGKLVELLWTEEMHDLIQRETRKLDGRDT
mouseSIRT3      LVGPFVLSPRRKDVVQLGDVVHGVERLVDLLGWTQELLDLMQRERGKLDGQDR
ratSIRT3        LVGSFALSPRRKDVVQLGDVVQGVERLVDLLGWTQELQDLIQRENGKLDGQD
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Figure 3. Sequence alignment of SIRT3 from human, mouse, rat, and rabbit. Conserved residues are labeled by “.”, functionally highly conserved residues are labeled by “:”, and less conserved residues are labeled by “.”. The arrow points to the putative MPP cleavage site. The three potential starting residues, Met1, Met15, and Met 78 of mSIRT3_L, are in bold.

transfected mouse embryonic fibroblasts with mSIRT3_L-1-334, 15-334, and 78-334 expressed with a C-terminal FLAG tag [Fig. 5(A)]. In immunofluores-

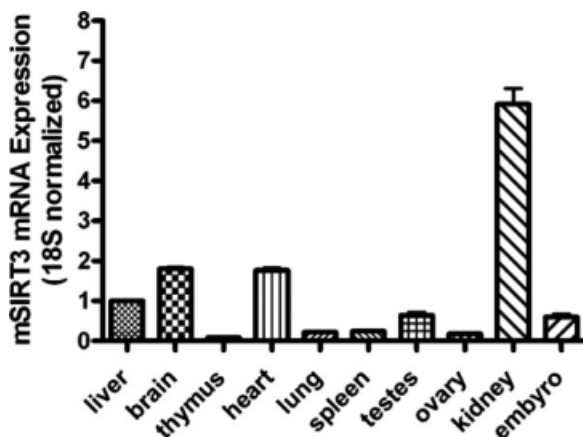


Figure 4. Tissue distribution of mSIRT3_L. The mRNA levels for mSIRT3_L were analyzed by real-time RT-PCR in several mouse tissues. The mSIRT3_L mRNA level in each tissue was normalized to 18S mRNA. The mRNA for mSIRT3_L in the liver was set to 1 and those in the other tissues were normalized to that in the liver.

cence assays, over-expressed mSIRT3 proteins were detected by anti-FLAG antibody (green) and mitochondrial compartments were labeled by staining for endogenous cytochrome c (red) [Fig. 5(B)]. Both mSIRT3_L-1-334 and 15-334 colocalized with cytochrome c in the mitochondria in all transfected cells, with no apparent alteration of mitochondrial morphology. In contrast, mSIRT3_L-78-334 was localized uniformly in the cytosol in 82% of cells and aggregated within the cytosol with partial cytochrome c overlap in 14% of transfected cells. mSIRT3_L-78-334 was also found in the nucleus when highly over-expressed.

Cell fractionation demonstrated that mSIRT3_L-1-334 and mSIRT3_L-15-334 are almost exclusively mitochondrial, whereas mSIRT3_L-78-334 is distributed amongst all fractions [Fig. 5(C)]. The mSIRT3_L-78-334 ran overlapping with another protein band that was present in all the fractions of the cell transfected with control vector. Our result is consistent with data obtained by Cooper *et al.*, where the authors demonstrate that the full-length hSIRT3 is exclusively mitochondrial but an N-terminal truncation mutant that corresponds to the original mouse start codon (hSIRT3-143-399) remains dispersed throughout the cell.³⁵ Taken together, our data suggest mitochondrial

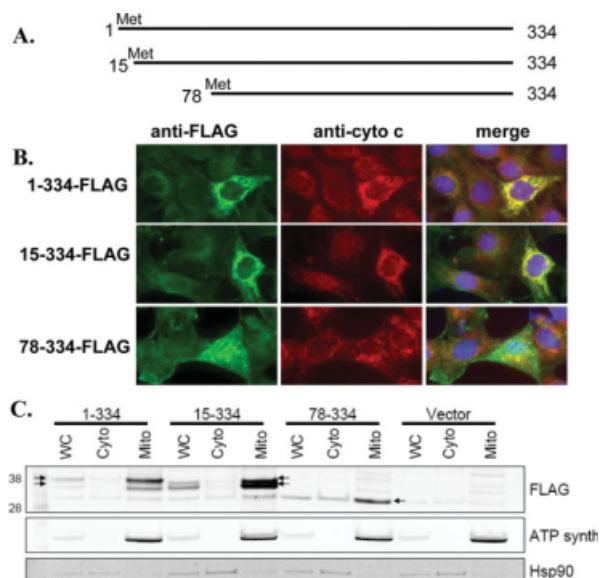


Figure 5. Long mSIRT3 localizes to mitochondria. (A) Schematic drawing of SIRT3 constructs. mSIRT3_L-1-334 begins at the most N-terminal Met, mSIRT3_L-15-334 starts at the second Met, and mSIRT3_L-78-334 represents the short mSIRT3 previously believed to be the full-length protein. (B) Mouse embryonic fibroblasts were transfected with C-terminal FLAG-tagged SIRT3 constructs and stained with both anti-FLAG (green) antibody to detect over-expressed mSIRT3 and anti-cytochrome c (red) to label mitochondria. Data are representative of three independent experiments. (C) Cell fractionation of MEFs transfected with either empty vector or plasmids shown in A. mSIRT3_L-1-334 and mSIRT3_L-15-334 are found predominantly in the mitochondrial fraction, whereas mSIRT3_L-78-334 is distributed throughout both cytosolic and mitochondrial fractions. Arrows point out the mSIRT3 protein bands that are discussed in the text.

localization of the longer mSIRT3 isoforms, whether they begin at the first (Met1) or second (Met15) methionine. In contrast, the originally predicted mouse protein (78-334) remains largely cytosolic and nuclear. The mitochondrial staining observed with mSIRT3_L-78-334 appears to alter mitochondrial morphology, similar to staining observed by Shi *et al.*³⁸ and noted as atypical by Cooper and Spelbrink.³⁵

Our data demonstrate that over-expressed mSIRT3_L, starting at either position 1 or 15, is a mitochondrial protein. To predict which methionine is the possible starting residue for mSIRT3_L, we analyzed the sequences with three programs that apply different approaches for predicting mitochondrial targeting sequences, MitoProt,⁴¹ Predotar,⁴² and TargetP.⁴³ The scores for mitochondrial targeting are highest for mSIRT3_L-15-334 in all predictions (Table I). The score for rat SIRT3-1-320 is higher than the predicted rat SIRT3-1-334 as well. In addition, we calculated the scores for human and rabbit SIRT3. Based on the

results from sequence alignment [Fig. 3(A)], the prediction of mitochondrial protein (Table I) and the homology among SIRT3 of mouse, human, rat and rabbit, mSIRT3_L-15-334 is most likely to be the full-length mouse SIRT3 protein that contains 320 amino acids.

Mouse SIRT3 deacetylase activity

The full length and several fragments of mSIRT3_L have been expressed and purified [Fig. 1(C)]. The presence of chaperone was required to obtain soluble protein. The proteins containing residues 1-334, 15-334, and 78-334 of mSIRT3_L do not have detectable deacetylation activity, whereas the other N-terminal truncated proteins except 64-334 displayed robust enzyme activity and have similar activity to hSIRT3-118-334, a very active hSIRT3 fragment. The 64-334 fragment of mSIRT3_L had lower deacetylation activity, indicating that truncating mSIRT3_L beyond residue 59 might adversely affect the proper folding of the protein. The proteins which include the mitochondrial targeting sequence (1-334 and 15-334) did not have detectable deacetylation activity similar to the full length hSIRT3.³⁴

MPP (matrix processing peptidase) is the putative mitochondrial enzyme that cleaves hSIRT3 between residue 101 and 102 to activate the enzymatic activity.³⁴ Based on the sequence alignment between mSIRT3_L and hSIRT3, mSIRT3_L-38-334 is equivalent to hSIRT3-102-399 (Fig. 3). The two arginine residues at -2 and -3 relative to the cleavage site of MPP and the hydrophobic residue at +1 position in hSIRT3 (MPP recognition site requirements) are conserved in mSIRT3_L, as well as in rat and rabbit SIRT3. Like hSIRT3, mSIRT3_L became active when the N-terminal residues were removed [Fig. 1(C)].

We observed [Fig. 5(C)] that there is another protein band under each of the protein bands of mSIRT3_L-1-334 and mSIRT3_L-15-334 that migrates to a similar location relative to each other but slightly higher than mSIRT3_L-78-334, the originally predicted mSIRT3 protein. This size difference suggests that the 78-334 protein is not the fragment produced through endogenous peptide cleavage. As mentioned above, the 78-334 fragment is not active in our *in vitro* deacetylation assay. This leaves open the possibility that mSIRT3_L-38-334, the fragment which corresponds to the human MPP processed SIRT3 (hSIRT3 102-399) and active in our *in vitro* assay, is more likely to be the endogenous active form of mouse SIRT3_L.

In our experiments, the human SIRT3-143-399 does not have deacetylation activity *in vitro*, although the predicted MPP cleaved form 102-399 is active (data not shown). These truncated proteins are similar in size, with 143-399 28.7 kD and 102-399 31.7 kD. Upon more careful inspection of the gel pictures in the report of Schwer *et al.*,³⁴ the mitochondrial hSIRT3 and MPP cleaved hSIRT3 were migrating slightly higher

Table I. Mitochondrial Targeting Scores for Long Mouse SIRT3 with Three Different Starting Methionine Residues (Met1, Met15, and Met78), as well as for SIRT3 of Human, Rat, and Rabbit

	Mouse SIRT3			Rat SIRT3		Human SIRT3	Rabbit SIRT3
	1-334 aa	15-334 aa	78-334 aa	1-334 ^a aa	1-320 aa	1-399 aa	1-319 aa
MitoProt II	0.6043	0.7847	0.0086	0.5418	0.7376	0.7455	0.941
Predotar	0	0.54	0	0.01	0.4	0.65	0.78
TargetP	0.12	0.814	0.18	0.109	0.819	0.555	0.938

Three different mitochondria targeting prediction programs were used.

Scores range from 0 to 1, with a higher score indicating higher probability for the protein to contain a mitochondria targeting sequence.

^a Indicates the rat SIRT3 that was predicted from the rat genome, but was not cloned.

than the 28.4 molecular weight marker. It could be that the 28-kDa hSIRT3 that was referred to be the active hSIRT3 is actually the 31.7-kDa hSIRT3-102-399.

Shi *et al.* have reported that mSIRT3 78-334 is a mitochondrial protein that can increase basal oxygen consumption of HIB1B cells by 1.8-fold when stably over-expressed.³⁸ Although we do see some amount of this protein in the mitochondria, over-expression of mSIRT3_L-78-334 appeared to distort mitochondrial structure and was localized to additional cellular compartments. Therefore over-expression of this form of mSIRT3 may not be reflective on the true activity of endogenous protein. We are in the process of immunoprecipitating over-expressed mSIRT3_L-1-334 and 15-334 to isolate sufficient amount of the processed form of mSIRT3 for N-terminal sequence identification.

SIRT3 inhibition studies

As part of the characterization of the purified mSIRT3_L-54-334 protein, we have tested two deacetylase inhibitors, nicotinamide (NAM) and SRT1720 as described in Material and Methods. NAM is a known inhibitor of the deacetylation activity of sirtuins,^{26,44-47} but the inhibition mechanism of NAM toward its substrates for SIRT3 has not been determined before. We also identified a small molecule SIRT3 inhibitor, SRT1720. Figure 6 shows the curves and the K_m , V_{max} , K_i and αK_i values calculated by the global fitting for each compound against NAD⁺ or AceCS2 peptide. The K_m and V_{max} values attained for NAD⁺ and AceCS2 peptide are within experimental error to previously obtained values without inhibitor used in the inhibition mechanism study.

SRT1720 inhibits mSIRT3_L-54-334 with αK_i of 0.34 μM against NAD⁺ [Fig. 6(A,E)]. The α value is significantly smaller than 1 indicating uncompetitive inhibition. However SRT1720 inhibits mSIRT3_L-54-334 in a competitive manner (α value $\gg 1$) against AceCS2 peptide with a K_i value of 0.56 μM [Fig. 6(B,E)]. This suggests that SRT1720 binding requires NAD⁺ to bind first to the enzyme and that SRT1720 competes for the same binding site of AceCS2 peptide or works as an allosteric competitive inhibitor to mSIRT3.

NAM has an α value much greater than one against NAD⁺ and AceCS2 peptide suggesting a competitive inhibition mechanism toward both substrates with K_i values of 2.84 μM [Fig. 6(C,E)] against NAD⁺ and 4.62 μM [Fig. 6(D,E)] against AceCS2 peptide. NAM has been reported as a non-competitive inhibitor toward both NAD⁺ and the peptide substrate and is part of the base-exchange reaction in SIRT1, SIRT2, yeast and bacterial sirtuins.⁴⁴⁻⁴⁸ However, it is likely that mSIRT3 has a different mechanism of inhibition for NAM. A more complete kinetic analysis of SIRT3 is ongoing to propose a reaction mechanism.

Under the current experimental conditions, the mechanism of inhibition is different for SRT1720 and NAM against NAD⁺ and AceCS2 peptide. These data suggest that SRT1720 and NAM bind to different sites on mSIRT3 and that mSIRT3 could be inhibited by different inhibitors via different sites and mechanisms.

It is still unclear what the biological and physiological roles of SIRT3 are and how inhibition of SIRT3 would affect glucose metabolism and insulin sensitivity. SRT1720 has previously been shown to activate SIRT1.¹⁹ However we have developed many other small molecule SIRT1 activators that are as effective as SRT1720 in controlling glucose homeostasis in ob/ob and diet-induced obesity mouse models but activate or have no effect on SIRT3. These data suggest that SRT1720, along with other SIRT1 activators, are potentially useful in treating type 2 diabetes. Developing SIRT3 selective modulators will help to address the functional roles of SIRT3 *in vivo*.

In summary, we have identified the correct form of mSIRT3. This longer form is more similar in length, localization, and activity to human SIRT3 than the shorter, originally cloned mouse ortholog. The identification of this protein increases our confidence of using mouse models to design small molecule modulators of human SIRT3 that could one day treat human disease.

Materials and Methods

Cloning of long mouse, rat, and rabbit SIRT3

To identify the 5' end of mSIRT3, 5'-RACE (rapid amplification of cDNA ends) polymerase chain reaction (PCR) was performed on the cDNA prepared

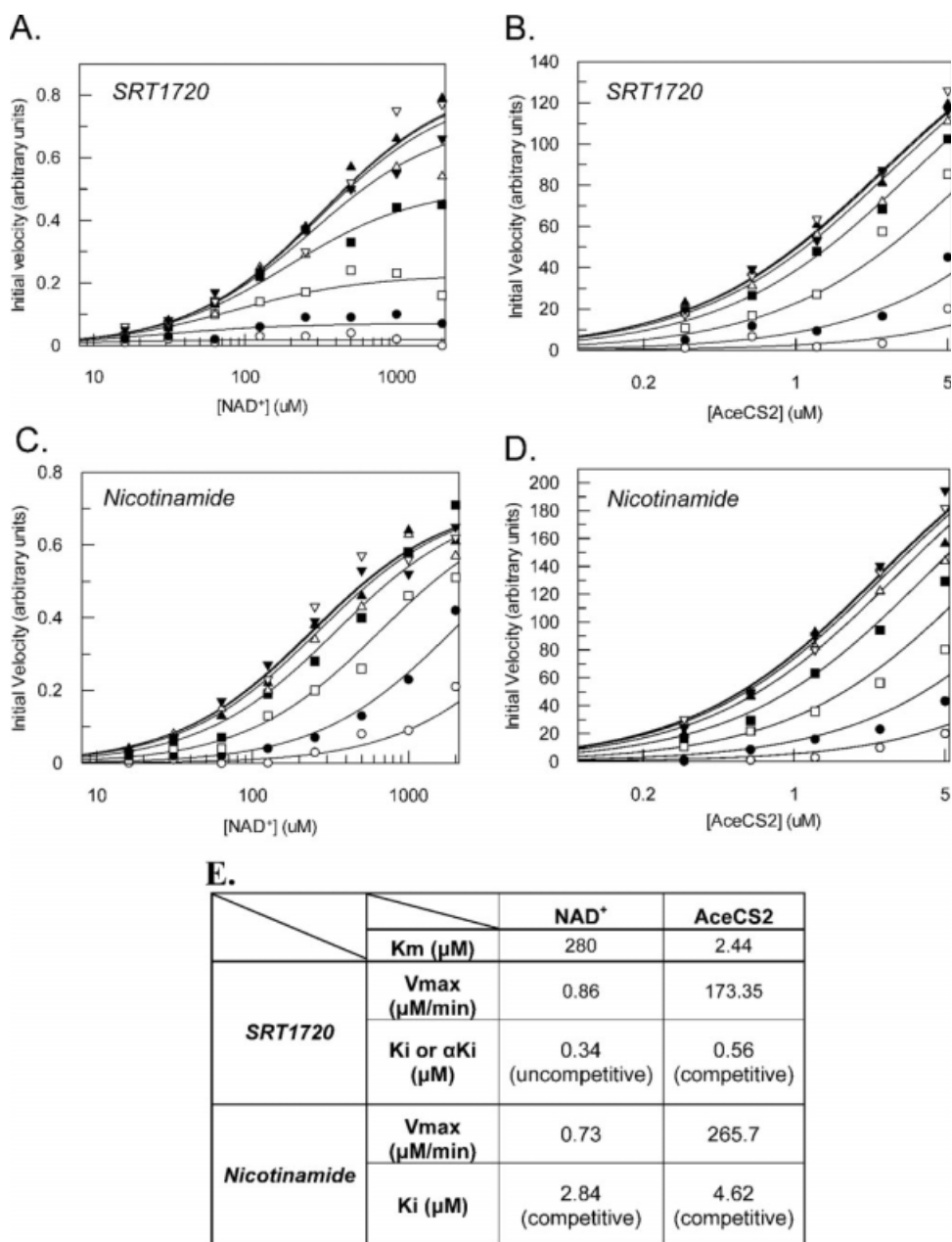


Figure 6. Inhibition of mSIRT3₅₄₋₃₃₄ by SRT1720 and nicotinamide. The global fitting curves of the observed rate of reaction with respect to varying substrate and inhibitor concentrations are displayed. (A) SRT1720 inhibition against NAD⁺. Eight concentrations of nicotinamide were used: ○ 15 μM, ● 3.75 μM, □ 0.94 μM, ■ 0.23 μM, △ 0.06 μM, ▲ 0.015 μM, ▽ 0.004 μM and ▼ 0.001 μM. (B) SRT1720 inhibition against AceCS2 peptide. The same eight concentrations of SRT1720 were used as in A. (C) Nicotinamide inhibition against NAD⁺. Eight concentrations of nicotinamide were used as follows: ○ 80 μM, ● 20 μM, □ 5 μM, ■ 1.25 μM, △ 0.31 μM, ▲ 0.08 μM, ▽ 0.02 μM and ▼ 0.005 μM. (D) Nicotinamide inhibition against AceCS2 peptide. Eight concentrations of nicotinamide were used: ○ 85 μM, ● 28.3 μM, □ 9.4 μM, ■ 3.2 μM, △ 1.05 μM, ▲ 0.35 μM, ▽ 0.12 μM and ▼ 0.04 μM. (E) The K_m and V_{max} values of the enzyme, V_{max} value in μM/min, K_i or αK_i values in μM, and the mechanism of inhibition for SRT1720 and nicotinamide are listed.

using the SMART RACE cDNA Amplification kit (Clontech) from RNA isolated from the liver of a female mouse (Institute of Cancer Research, CD-1) using TRI-reagent (Sigma). The mSIRT3 specific reverse primer was designed based on the gene with Genbank number of BC025878. The gene specific primer TTATCTGTCCTGTCCATCCA and universal primer mix were used for the PCR reaction. Nested

PCR was performed on the 5'-RACE PCR product with nested universal primers and mSIRT3 specific reverse primer: CCTGTCCGCATCACATCAGCCCAT. Specific DNA bands of mSIRT3 from the 5'-RACE products were generated and ligated into T-vector (Takara). Positive clones were obtained with two coding sequences. The novel isoform of mSIRT3 has 334 amino acids which contains 77 additional amino acids at the

N-terminus of previously reported mSIRT3 containing 257 amino acids.

The longer mSIRT3 was cloned into a modified pET21a vector with an N-terminal His₆ tag. Several N-terminal truncations of mouse SIRT3 were also generated in the same vector. The truncated proteins start at residue 15, 38, 44, 49, 54, 59, 64, or 78 and end at residue 334.

5'-RACE PCR was performed as above on cDNA derived from the liver and kidney mRNA purified from a Wistar male rat. Gene specific primers were designed according to the predicted rat SIRT3 sequence (XM_215124). The reverse primer used for 5'-RACE PCR was TCAGCTGTCTAGCGATGGGCATG and for nested PCR was TACGGGATGTCATACTGCTGAAGGTTGCT. The cloned rat SIRT3 contains 320 amino acids.

For cloning rabbit SIRT3, mRNA was purified from the liver of a male New Zealand white rabbit. The cDNA for 5'-RACE and 3'-RACE was prepared using the Clontech SMART RACE kit, and PCR was performed using a universal primer mix and gene specific primer. The gene specific reverse primer was CAAC-CAGCTTTGAGGCAGGGAT for 5'-RACE and the forward primer was TCACAACCCCAAGCCCTTTTCA for 3'-RACE. Both RACE reactions yielded product of interest. The cloned rabbit SIRT3 contains 319 amino acids.

For immunofluorescence and cell fractionation studies, the mSIRT3_L-1-334, 15-334, and 78-334 were cloned into the p3XFLAG-CMV-14 vector (Sigma) between KpnI and XbaI sites. Plasmid DNA was isolated using Endofree Plasmid Maxi kit (QIAGEN).

Reverse transcription-PCR amplification

Total RNA was purchased from Ambion and reverse transcribed in duplicate using a High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). Real Time PCR reactions were performed following conditions recommended by the manufacturer and analyzed using the 7300 Fast Real-Time PCR System (Applied Biosystems). Genes of interest were normalized to 18S ribosomal RNA (Integrated DNA Technologies). RT-PCR primers and probes were synthesized by Integrated DNA Technologies. The mouse SIRT3 probe was labeled at the 5' end with 6-FAMTM dye (6-carboxyfluorescein) and at the 3' end with BHQ-1 (Black Hole Quencher-1TM). The mouse 18S RNA probe was labeled at the 5' end with JOE (6-carboxy-4', 5'-dichloro-2', 7'-dimethoxyfluorescein) and at the 3' end with BHQ-1. The probe sequence was CTGCTT CGGACCAGAGCCTGCAGT from the N-terminal sequence unique to mSIRT3_L. The forward primer was GGCTTTGGAGGTGGAGGAA and reverse primer was GCCACTGGGTGTGCTGATG for mSIRT3 PCR. The probe sequence for 18S was TGCTGGCACCA-GACTTGCCCTC. The forward primer for 18S amplification was CGGCTACCACATCCAAGGAA and reverse primer was GAGCTGGAATTACCGCGGCT.

Immunofluorescence

Mouse Embryonic Fibroblasts (MEFs) (2×10^6) were transfected with 5 μ g plasmid DNA per reaction using Amaxa nucleofector kit V (Amaxa Biosystems). Cells were then plated in 2-well chamber slides (LabTekII) at a density of 2×10^5 cells per well. Twenty-four hours following transfection, cells were washed three times with PBS and fixed with 4% paraformaldehyde/PBS for 15 min at room temperature. Cells were washed as above, then permeabilized with 0.2% Triton X-100/PBS for 10 min on ice and incubated with 5%BSA/PBS blocking buffer for 1 h at room temperature. Cells were incubated for 1 h in 5% BSA/PBS with mouse anti-FLAG (1:100, Sigma) and rabbit anti-cytochrome c (1:100, Santa Cruz) primary antibodies. Cells were washed three times with 1% BSA/PBS and then incubated for 1 h in 5%BSA/PBS with FITC-conjugated anti-mouse (1:200, Jackson ImmunoRes) and TRITC-conjugated anti-rabbit (1:100, Jackson ImmunoRes) secondary antibodies. All antibody incubations were performed at room temperature. After three washes, mounting media with DAPI was added (VectaShield H-1200; Vector Bioloabs) and a coverslip was placed on each slide. Staining was visualized using a Zeiss Axiovert 200M fluorescence microscope and imaged with AxioVision 4.5 software. Quantification of subcellular localization of mSIRT3 constructs was based on scoring of 50 random transfected cells per slide for mitochondrial or cytosolic localization. Data are representative of three independent experiments.

Cell fractionation and western blotting

Mouse Embryonic Fibroblasts (MEFs) (4×10^6) were transfected with 5 μ g plasmid DNA per reaction using Amaxa nucleofector kit V (Amaxa Biosystems) and were plated in 10 cm dishes. Eighteen hours following transfection, mitochondria were isolated from whole cell lysate and cytosol using the Dounce homogenization protocol of the Pierce Mitochondria Isolation Kit (Pierce). Protein concentration of whole cell, cytosol, and mitochondrial fractions was measured by A280 and by the BCA Protein Assay Kit (Pierce). Each fraction (20 μ g) was run on a NuPAGE 4–12% SDS-PAGE gel (Invitrogen). Gels were transferred to nitrocellulose membranes (iBlot, Invitrogen) and membranes were blocked in Odyssey Blocking buffer (Li-cor) for 30 min. Membranes were incubated overnight with anti-FLAG (1:1000, Sigma), anti-ATP synthase subunit alpha (1:1000, Invitrogen), or anti-Hsp90 (1:500, Abcam) primary antibody in 0.1% Tween/Odyssey buffer. Blots were washed three times for 10 min with 0.1% Tween/PBS and incubated with IR Dye800 donkey anti-mouse secondary antibody (1:10,000, Rockland) for 1 h in 0.1% Tween/Odyssey buffer. Following three washes as above, blots were imaged using the Odyssey Infrared Imager (Li-cor).

Expression of mSIRT3 proteins

The full length and fragments of mSIRT_{3L} were expressed in BL21(DE3) Star cells (Novagen) with co-expression of chaperone groES-groEL-tig (pG-Tf2, Takara). A single colony was inoculated in LB media containing 20 µg/mL chloramphenicol and 100 µg/mL ampicillin for plasmid selection and 5 ng/mL tetracycline for chaperone induction at 37°C, 250 rpm until the OD₆₀₀ reached 0.3. The culture was then transferred to 18°C, 250 rpm until the OD₆₀₀ reached 0.6. IPTG was added to a final concentration of 0.3 mM and expression was continued at 18°C, 180 rpm overnight. Cells were collected by centrifugation and the pellet was resuspended in Lysis buffer (50 mM Tris-HCl, 300 mM NaCl, 10% glycerol, 10 mM MgCl₂, pH 8.8, 5 mL/gm of cell pellet) and sonicated to open the cells. Supernatant was separated from cell debris by centrifugation at 10,000 g for 40 min at 4°C and loaded onto a Ni-NTA column (Qiagen). The column was washed with 10 CV (column volume) of the Binding buffer (50 mM Tris-HCl, 10% glycerol, 300 mM NaCl, 10 mM MgCl₂, pH 8.0), 10 CV of the Wash buffer (50 mM Tris-HCl, 10% glycerol, 300 mM NaCl, 20 mM imidazole, 10 mM MgCl₂, pH 8.0) and eluted with the Elution buffer (50 mM Tris-HCl, 10% glycerol, 300 mM NaCl, 250 mM imidazole, 10 mM MgCl₂, pH 8.0). The eluted protein was dialyzed against the Binding buffer and then digested with TEV protease (Invitrogen) to remove the N-terminal His tag. The tag-free enzyme was isolated by a 2nd Ni-NTA column. The mSIRT_{3L}-54-334 was further purified to >95% purity by an S200 column (GE) judged by SDS-PAGE.

Enzyme assay

The partially purified full length and different fragments of mSIRT_{3L} were tested for deacetylation activity with the mass spectrometry based assay reported previously.¹⁹ Saturating amounts of peptide substrate (20 µM) and βNAD⁺ (3 mM) were added to the reaction with enzyme concentration starting at 4 µM and serially diluted in a 1:2 ratio. Reactions were incubated at 25°C and stopped at 0, 15, 30, 60, 90, 120, 150, 180 minute time points with 10% formic acid with 50 mM nicotinamide. The conversion of substrate to product was determined by mass spectrometry in conjunction with a RapidFire system (BioTrove).

Inhibition analyses

To identify the mechanism of inhibition of SRT1720, a small molecule SIRT3 inhibitor, and nicotinamide against mSIRT_{3L}-54-334, we used a single acetylated peptide derived from the SIRT3 substrate AceCS2 (EILVVKRLPKTRSG-K_{Ac}-VMRLLLRKIITSEAQ, K_{Ac} is acetylated lysine). For compound inhibition against acetylated AceCS2 peptide, eight concentrations of nicotinamide (15, 3.75, 0.94, 0.23, 0.06, 0.015, 0.004,

and 0.001 µM) or nicotinamide (85, 28.3, 9.4, 3.2, 1.05, 0.35, 0.12, and 0.04 µM) were used in the reaction. For each of the respective compound concentrations, the deacetylation rate was measured at five concentrations of acetylated peptide (5, 2.5, 1.25, 0.62, and 0.31 µM) with mSIRT_{3L}-54-334 (1.47 nM) and NAD⁺ (220 µM) kept constant. Substrate inhibition was detected at high concentrations forcing the use of lower amounts of acetylated peptide. The reactions were carried out at 25°C. The time course was determined from the linear portions of the AceCS2 K_m curve (ran at NAD⁺ K_m) and the reaction was stopped with 10% formic acid with 50 mM nicotinamide and the conversion of substrate to product was determined by mass spectrometry.

For compound inhibition against NAD⁺, eight concentrations of SRT1720 (15, 3.75, 0.94, 0.23, 0.06, 0.015, 0.004, 0.001 µM) or nicotinamide (80, 20, 5, 1.25, 0.31, 0.08, 0.02, and 0.005 µM) were used. For each of the respective compound concentrations, the deacetylation rate was measured at eight fixed concentrations of NAD⁺ (2000, 1000, 500, 250, 125, 63, 31, and 16 µM) with mSIRT_{3L}-54-334 (1.47 nM) and acetylated AceCS2 peptide (2 µM) kept constant at 25°C. The time course was determined from the linear portions of the NAD K_m curve (ran at AceCS2 K_m) and the reaction was stopped with 10% formic acid with 50 mM nicotinamide and the conversion of substrate to product was determined by mass spectrometry.

To determine the inhibition constants (K_i) and mechanism of action toward NAM and SRT1720, the initial velocities of several series of reactions were calculated by global non-linear regression. Briefly, varying concentrations of inhibitor were titrated against fixed concentrations of substrate (NAD⁺ or AceCS2). The observed rates of reaction with respect to substrates and inhibitors were then globally fit to the mixed noncompetitive inhibition equation [eq. (1)] using the software package GraFit 6.0.5 (Erithacus Software).

$$v = \frac{V_{\max} \cdot [S]}{K_m \left(1 + \frac{[I]}{K_i}\right) + [S] \left(1 + \frac{[I]}{\alpha K_i}\right)} \quad (1)$$

where, K_i and αK_i are the competitive and uncompetitive inhibition constants, respectively. The mechanisms of inhibition are determined from the alpha constant value. α = 1, indicates noncompetitive inhibition, α ≫ 1, competitive inhibition and α ≪ 1 uncompetitive inhibition.⁴⁹

GenBank deposition

The sequences for mouse, rat, and rabbit SIRT3 have been deposited at GenBank with accession numbers of EU886466, EU886468, and EU886467, respectively.

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