Mechanistic studies on the effects of nicotinamide on megakaryocytic polyploidization and the roles of NAD⁺ levels and SIRT inhibition

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(Received 19 June 2009; revised 20 August 2009; accepted 20 August 2009)

Objective. Megakaryocytic cells (Mks) undergo endomitosis and become polyploid. Mk ploidy correlates with platelet production. We previously showed that nicotinamide (NIC) greatly increases Mk ploidy in cultures of human mobilized peripheral blood CD34⁺ cells. This study aims to examine the generality of NIC effects, NIC’s impact on Mk ultrastructure, and potential mechanisms for the increased ploidy.

Materials and Methods. We used electron microscopy to examine Mk ultrastructure and flow cytometry to evaluate NIC effects on Mk differentiation and ploidy in mobilized peripheral blood CD34⁺ cell cultures under diverse megakaryopoietic conditions. Mk ploidy and NAD(H) content were evaluated for NIC and other NAD⁺ precursors. We tested additional inhibitors of the sirtuin (or SIRT) 1 and SIRT2 histone/protein deacetylases and, after treatment with NIC, evaluated changes in the acetylation of SIRT1/2 targets.

Results. NIC increased ploidy under diverse culture conditions and did not alter Mk ultrastructure; 6.25 mM NIC increased NAD⁺ levels fivefold. Quinolinic acid increased NAD⁺ similar to that for 1 mM NIC, but yielded a much smaller ploidy increase. Similar increases in Mk ploidy were obtained using NIC or the SIRT1/2 inhibitor cambinol, while the SIRT2 inhibitor AGK2 moderately increased ploidy. SIRT1/2 inhibition in cells treated with NIC was evidenced by increased acetylation of nucleosomes and p53. Greater p53 acetylation with NIC was associated with increased binding of p53 to its consensus DNA binding sequence.

Conclusion. NIC greatly increases Mk ploidy under a wide range of conditions without altering Mk morphology. Inhibition of SIRT1 and/or SIRT2 is primarily responsible for NIC effects on Mk maturation.

As precursors to platelets, megakaryocytic cells (Mks) are central to hemostasis. Mk differentiation includes several rounds of endomitosis to form polyploid cells and the formation of cytoplasmic extensions called proplatelets. However, Mk commitment and the mechanisms through which Mks differentiate and mature remain poorly under-
production in cultures of human mobilized peripheral blood (mPB) CD34<sup>+</sup> cells stimulated with Tpo [8]. Evaluating the conditions under which NIC increases Mk ploidy and identifying the mechanism(s) that underlie the effects of NIC would lead to a greater understanding of Mk differentiation and may allow further modulation of Mk maturation.

NIC has diverse functions in cells. NIC inhibits the activity of the silent information regulator 2 (Sir2) family of histone/protein deacetylases (sirtuins or SIRTs) [9–12]. SIRTs catalyze a unique NAD<sup>+</sup>-dependent deacetylation reaction and are important for a wide variety of biological processes, including transcriptional silencing, lifespan regulation, and regulation of apoptosis [13–15]. NIC is also a precursor of NAD<sup>+</sup> via the salvage pathway [16,17]. NAD<sup>+</sup> regulates a variety of intracellular activities, such as the DNA-binding specificity of p53 [18,19]. We have previously established that the p53-mediated Mk apoptotic program is intimately linked with terminal maturation and polyploidization, possibly through the downstream effects of p53 on MDM2 and BCL2 expression [20,21].

The aim of this study was to examine the generality of the NIC effects, NIC’s potential impact on Mk ultrastructure, and possible mechanisms for the profound effect of NIC on Mk ploidy.

Materials and methods

Unless noted, all reagents were obtained from Sigma-Aldrich (St Louis, MO, USA).

Human Mk culture

Cultures were initiated in T flasks with previously frozen mPB CD34-selected cells (AllCells, Berkeley, CA, USA or Fred Hutchinson Cancer Research Center, Seattle, WA, USA) and maintained at a concentration of 100,000 to 300,000 cells/mL as described [8]. Most cultures were supplemented with 100 ng/mL Tpo (Peprotech, Rocky Hill, NJ, USA or Genentech, South San Francisco, CA, USA) and treated with 6.25 mM NIC beginning at day 5 (Tpo + NIC). Selected cultures were also supplemented with 100 ng/mL stem cell factor (SCF; R&D Systems, Minneapolis, MN, USA) or were initiated in 12-well plates coated with fibronectin. For cultures containing Tpo plus 150 ng/mL stromal-derived factor–1α (SDF-1α; R&D Systems), NIC was added beginning at day 2. Cytokine cocktail cultures were supplemented with 1.5 ng/mL interleukin (IL)-3 (Peprotech), 10 ng/mL IL-6 (Peprotech), and 50 ng/mL SCF. On day 5, cells were treated with NIC and/or Tpo to generate four cultures: cocktail (IL-3, IL-6, SCF), cocktail + 6.25 mM NIC, cocktail + 100 ng/mL Tpo, or cocktail + Tpo + NIC. NAD<sup>+</sup> precursors nicotinic acid and quinolinic acid (QA) were added on day 5 to cells cultured with Tpo. The SIRT inhibitors cambinol (generously provided by A. Bedalov, Fred Hutchinson Cancer Research Center or purchased from Calbiochem, San Diego, CA, USA) and AGK2 (ChemDiv, San Diego, CA, USA) were added on day 5 to cells cultured with Tpo. The average cell and nuclear volumes were determined by analyzing whole cells and nuclei (released using 3% cetrimide), respectively, using a Multisizer 3 (Beckman Coulter, Fullerton, CA, USA).

Electron microscopy

Cells from Tpo only and Tpo + NIC cultures were removed on day 8 and transferred to fibronectin-coated 35-mm Petri dishes (BD Biosciences, San Jose, CA, USA) to induce cell adhesion. After 48 hours, cells were fixed for 4 hours (2.5% glutaraldehyde/2.5% paraformaldehyde in 0.1 M cacodylate buffer, pH 7.4) and processed by the NU Cell Imaging Facility. Briefly, after embedding the cells in Epoxy resin in an inverted capsular mold, ultra-thin sections were cut, stained with uranyl acetate and lead citrate, and examined with a JEOL-1220 transmission electron microscope at an accelerating voltage of 60 kV.

Flow cytometric detection of surface antigens, ploidy, and intracellular proteins

For CD41 expression, Mk apoptosis, and Mk ploidy, cells were prepared for flow cytometry and analyzed as described [8]. For intracellular detection of total and acetylated p53, cells were stained with fluorescein isothiocyanate–conjugated anti-CD41 antibody, fixed with 2% paraformaldehyde, permeabilized with 70% methanol, stained with phycocerythrin-conjugated-anti-p53 (Santa Cruz Biotechnology, Santa Cruz, CA, USA) and Alexa Fluor 647–conjugated-anti-AcK382p53 (BD Biosciences) antibody, treated with RNase, and stained with 7-aminactinomycin D. Expression of p53 and AcK382p53 proteins was quantified by subtracting the mean fluorescence intensity (MFI) of unstained CD41<sup>+</sup> cells from the MFI of the p53-labeled and AcK382p53-labeled CD41<sup>+</sup> cells. FACSARia flow cytometer and FACSDiva software (BD Biosciences) were used for data acquisition and analysis.

Measurement of intracellular NAD(H) levels

Three-hundred thousand cells were extracted in 0.6 mL of slightly basic extraction buffer containing 10 mM NIC and 0.05% (w/v) Triton X-100, as described by Wagner and Scott [22]. Standard solutions of NAD<sup>+</sup> and NADH were prepared in extraction buffer. To determine NAD(H) (sum of NAD<sup>+</sup> and NADH) content, the extracts were subjected to an enzymatic cycling reaction [23] performed in a 96-well plate using a reagent mixture containing 0.1 M bicine, 0.5 M ethanol, 4.17 mM ethylene diamine tetraacetic acid, 0.83 mg/mL bovine serum albumin, 0.42 mM MTT, and 1.66 mM phenazine ethosulfate. The reaction was started by adding 2 U yeast alcohol dehydrogenase. Color was developed in the dark for 30 minutes and detected at 570 nm. Each sample was run in duplicate. The NADH content was separately determined, after incubating extract samples at 60°C for 30 minutes to degrade NAD<sup>+</sup>.

Western blots

Nuclear lysates were prepared according to manufacturer’s protocol (NE-PER Kit; Pierce Biotechnology, Rockford, IL, USA). Thirty micrograms total protein per sample was denatured, separated by SDS-PAGE Ready-Gel (BioRad, Hercules, CA, USA), and transferred onto nitrocellulose membranes (BioRad). Membranes were blocked in nonfat dry milk for 1 hour and incubated overnight at 4°C with primary antibody for acetylated lysine residues (Cell Signaling, Danvers, MA, USA), nicotinamide mononucleotide adenyltransferase 1 (Nmnat1; Abcam, Cambridge, MA, USA), or AcK382p53 (Cell Signaling). After washing, membranes were incubated with horseradish peroxidase–conjugated goat anti-mouse- or anti-rabbit-IgG antibody (Cell Signaling). Bound antibodies were detected using...
chemiluminescence. The membranes were then stripped and reprobed with antibodies for nucleosomes (Chemicon, Billerica, MA, USA), actin (Abcam), or total p53 (Santa Cruz Biotechnology). Secondary antibodies were applied and detection was performed as described here. Densitometry was performed using ImageQuant 5.2 software (GE Healthcare, Piscataway, NJ, USA).

Electrophoretic mobility shift assays
Nuclear lysates were used for electrophoretic mobility shift assays (EMSAs) as described [20].

Results

NIC increases Mk maturation without altering normal Mk ultrastructure

The ultrastructure of Mks cultured with Tpo (Fig. 1A and B) was similar to that of Mks cultured with Tpo + NIC (Fig. 1C and D). α-Granules, dense granules, and mitochondria were visible in cells from both conditions. However, NIC-treated cells were larger and generally had a more complex nuclear structure, both of which are indicative of greater Mk maturation. NIC-treated cells also showed evidence of a demarcation membrane system (Fig. 1E). NIC-treated Mks had a greater number of short cytoplasmic projections and generally had a larger number of small cytoplasmic microparticles surrounding each cell. Proplatelets were present on some cells cultured with Tpo + NIC (Fig. 1F), and the morphology of the proplatelets was similar to that obtained by Cramer et al. [24]. Thus, NIC-treated Mks exhibit normal ultrastructure.

NIC increases ploidy under diverse Mk culture conditions

Because our previous cultures with NIC were carried out using Tpo as the only cytokine, we wanted to evaluate the generality of NIC effects. We first examined culture on fibronectin, as well as additional supplementation with SDF-1α or SCF. Culture on fibronectin delayed the increase in ploidy for Mks supplemented with Tpo (Fig. 2A). However, NIC increased the percentage of high-ploidy (≥ 8 N) Mks to a similar extent in the presence or absence of fibronectin. SDF-1α has previously been shown to increase Mk ploidy when added early in culture [25]. Adding 150 ng/mL SDF-1α resulted in a more rapid increase in ploidy compared to cultures with Tpo only, but the final ploidy was unchanged (Fig. 2B). Also, NIC increased ploidy to a similar extent in cultures with or without SDF-1α. SCF has been shown to synergize with Tpo to increase the number of Mks produced from umbilical cord blood or mPB CD34+ cells [26,27]. SCF increased Mk ploidy in cultures with or without NIC (Fig. 2C). Furthermore, NIC substantially increased Mk ploidy in cultures with or without SCF.

We also examined the ability of NIC to increase Mk ploidy in the absence of Tpo. mPB CD34+ cells were cultured with IL-6, IL-3, and SCF [28,29]. This cytokine combination yielded approximately 14% CD41+ cells by day 11 compared to 80% CD41+ cells in cultures with Tpo only (data not shown). On day 11, the percentage of high-ploidy Mks was approximately twofold lower in cocktail cultures compared to those with Tpo only, and reached only 9% high-ploidy Mks by day 11 (Fig. 2D). Addition of NIC to cocktail cultures beginning at day 5 increased the percentage of high-ploidy Mks at day 11 by more than twofold (Fig. 2D), indicating that NIC can act independently of Tpo. In contrast, addition of Tpo to cocktail cultures beginning at day 5 had a relatively small effect on the percentage of high-ploidy Mks. Adding both NIC and Tpo to cocktail cultures beginning at day 5 increased the percentage of high-ploidy Mks by a much greater extent (Fig. 2D), showing the synergy between Tpo and NIC. Together, these data demonstrate the versatility of NIC and its ability to increase the ploidy of Mks cultured under a variety of Mk-promoting conditions for donor samples that exhibit a wide range of Mk ploidy in cultures with Tpo only (Fig. 2A–D). Adding NIC to cultures with Tpo alone increased the percentage of high-ploidy cells by 3.9 (±0.5)-fold (n=29; p<10−15, using a paired t-test) for the donor samples used in this study.

NIC increases the level of intracellular NAD(H) in cultured Mks by up to fivefold

NIC may directly modulate Mk differentiation or it may act via an intermediary, such as NAD+, the level of which has been shown to increase greatly in diverse human cell types after NIC addition [18,30,31]. In order to explore this latter possibility, we first examined if NIC increases NAD+ levels in Mks. NAD(H) levels increased only slightly with Mk maturation in cultures with Tpo only (Fig. 3A). In contrast, NIC rapidly increased the NAD(H) content—by 60% for 1 mM NIC and threefold for 6.25 mM 4 hours after NIC addition. By day 9, the NAD(H) content was approximately threefold higher for 1 mM NIC and almost fivefold higher for 6.25 mM NIC compared to Tpo alone (Fig. 3A). For comparison, at day 9 the average volume of cells treated with 6.25 mM NIC was approximately 30% greater than cells cultured with Tpo alone and the average nuclear volume was 50% greater with NIC (data not shown). For all conditions, the level of NADH was much lower (approximately 10–20%) than that of NAD+ (data not shown). Thus, the increase in NAD(H) was predominantly due to an increase in NAD+. The increase in NAD(H) with NIC was dose-dependent (Fig. 3A), and the relative increase in NAD(H) for 1 mM vs 6.25 mM NIC compared to Tpo alone was similar to the relative increase in the fraction of high-ploidy Mks (Fig. 3B and C).

We also supplemented cultures with other NAD+ precursors. Nicotinic acid (NA), the other form of niacin, has been shown to increase NAD+ content in several cell types [32,33]. The NAD(H) content increased only slightly after addition of 3 mM or 6.25 mM NA to cells cultured with

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Tpo, and there was little if any effect of NA on CD41 expression or the percentage of high-ploidy Mks (data not shown). Thus, NA is not an effective NAD(H) precursor in cultured human Mks and does not increase Mk ploidy. NAD$^+$ can also be synthesized from tryptophan via QA [34], and QA increased intracellular NAD$^+$ levels in cultured glial cells.

Figure 1. Megakaryocytic cells (Mks) cultured with nicotinamide (NIC) exhibit normal ultrastructure. Transmission electron microscopy of Mks cultured with thrombopoietin (Tpo) only (A, B) and Tpo + NIC (C–F) at day 8. The images in (A–D) were acquired such that the magnification gave one cell per field of view. Because of the larger cell size with Tpo + NIC, it was necessary to use a lower magnification (2,500×) compared to that for Tpo only (4,000×). (E) An image of three cells exhibiting a demarcation membrane system (2,000×). (F) A proplatelet-bearing Mk (2,000×). n=nucleus, g=granules, dms=demarcation membrane system, pp=proplatelet. Scale bar: 2 μm.
QA increased NAD(H) levels in a dose-dependent manner (Fig. 3B). Despite the similar NAD(H) levels, Mks in cultures treated with 1 mM NIC exhibited much greater ploidy than those with 10 mM QA, which was just slightly higher than that of Mks in cultures with Tpo alone (Fig. 3C). QA is incorporated into NAD\(^+\) via the de novo synthesis pathway, which is distributed throughout the cell [34]. In contrast, NIC is incorporated into NAD\(^+\) via the salvage pathway, which in yeast cells is primarily localized to the nucleus [35,36]. Thus, the differential effects of NIC compared to QA could be due to differences in the location of NAD\(^+\) synthesis. Nicotinamide mononucleotide adenyltransferase 1 (Nmnat1) localizes exclusively to the nucleus [37] and is essential for NAD\(^+\) biosynthesis by catalyzing the formation of NAD\(^+\) from nicotinamide mononucleotide and ATP [38]. Cultured Mks treated with NIC expressed approximately twofold higher levels of Nmnat1 than cells treated with Tpo only (Fig. 3D), which is consistent with increased nuclear NAD(H) content in NIC-treated cells.

NIC increases Mk ploidy at least in part through SIRT inhibition

NIC has been extensively characterized as an inhibitor of SIRTs, which were originally identified as NAD\(^+\)-dependent class III histone deacetylases [39–41]. NIC is a potent inhibitor of SIRT1 [42], which deacetylates a wide range of histones and nonhistone proteins [41]. NIC also inhibits SIRT2, which deacetylates tubulin [11,43], histone H4 [44,45], and a growing
Figure 3. Effects of nicotinamide (NIC) and quinolinic acid (QA) on intracellular NAD(H) content, megakaryocytic cell (Mk) ploidy, and nicotinamide mononucleotide adenylyltransferase 1 (Nmnat1) levels. (A) Intracellular NAD(H) levels in cells cultured with thrombopoietin (Tpo) only, Tpo + 1 mM NIC, or Tpo + 6.25 mM NIC were measured using a cycling enzymatic assay on days 5 (4 hours after NIC addition), 7, and 9. Data shown are the mean ± standard error of mean (SEM) of n=7 (day 7) or n=5 (days 5 and 9) experiments. Based on a paired t-test, values of *p* < 0.005 (x), *p* < 0.05 (*) and *p* > 0.1 (+) are indicated in comparison to the Tpo only treatment. (B, C) Mobilized peripheral blood (mPB) CD34+ cells were cultured with 100 ng/mL Tpo. Beginning on day 5, cells were treated with either NIC or QA. (B) NAD(H) levels were measured on days 5 (4 hours after addition), 7, and 9. (C) The percentage of high-ploidy Mks was determined on days 5, 7, 9, and 11. (B, C) Data shown are the mean ± SEM of two experiments. Based on a paired t-test, values of *p* < 0.05 (*) and *p* < 0.1 (+) are indicated in comparison to the Tpo only treatment. (D) Western blot (representative of five biological experiments) of nuclear lysates prepared from primary Mks cultured with 100 ng/mL of Tpo only or supplemented with 6.25 mM NIC on day 5 were probed for nuclear NAD processing enzyme Nmnat1 and actin (loading control). Densitometry values for Nmnat1 or actin expression in cells grown with NIC were divided by the densitometry values for Nmnat1 or actin expression in cells grown with Tpo only. Ratios shown correspond to the mean ± SEM of protein expression in whole cell or nuclear lysates harvested from five biological experiments. Based on a paired t-test, values of *p* < 0.05 (*), and *p* < 0.1 (+) are indicated for the various time points in comparison to actin band density.
number of proteins [46]. We had previously concluded that inhibition of SIRT1 and SIRT2 was not responsible for NIC-mediated increases in Mk ploidy. This was based on our finding that the yeast Sir2p inhibitors sirtinol (inhibits mammalian SIRT1 [47–49]) and splitomicin (inhibits SIRT1 and SIRT2 [50,51]) did not affect Mk ploidy [8]. However, using these compounds in Mk cultures can be problematic due to splitomicin instability at pH 7.3 to 7.4 [45,52] and sirtinol toxicity in Mk cultures at doses below those reported to be effective in mammalian cells [8,47–49]. Therefore, we evaluated the effects of two recently described SIRT inhibitors.

Cambinol is a small molecule that inhibits both SIRT1 and SIRT2 [45]. When added on day 5 at 10 μM, cambinol increased the fraction of high-ploidy Mks to a similar extent as NIC (Fig. 4A). The Mk ploidy distributions (Fig. 4B) and mean ploidy values (4.2±0.2 for NIC and 3.9±0.2 for cambinol) were also similar for 10 μM cambinol and 6.25 mM NIC. Adding 3.125 mM NIC plus 5 μM cambinol slightly increased the fraction of high-ploidy cells compared to 6.25 mM NIC or 10 μM cambinol alone, but the difference was not statistically significant (Supplementary Figure E1A, online only, available at www.exphem.org). Synergistic addition of 3.125 mM NIC plus 5 μM cambinol had a greater effect on the fraction of Mks with ploidy ≥16 N (Supplementary Figure E1B, online only, available at www.exphem.org); the increase was statistically significant in comparison to 10 μM cambinol (p=0.03), but not to 6.25 mM NIC (p=0.14). No further increase in ploidy was obtained by adding 6.25 mM NIC plus 10 μM cambinol.

Figure 4. Sirtuin (SIRT) inhibitors increase megakaryocytic cell (Mk) ploidy. Mobilized peripheral blood (mPB) CD34+ cells were cultured with 100 ng/mL thrombopoietin (Tpo). On day 5, cultures were supplemented with (A, B) 10 μM cambinol (n=12) or (C, D) 10 μM AGK2 (n=11). For comparison, replicate cultures were either supplemented with 6.25 mM nicotinamide (NIC) on day 5 or maintained with Tpo alone. Flow cytometry was used to determine (A, C) the percentage of CD41+ cells with DNA content ≥8 N (high-ploidy Mks) and (B, D) the high-ploidy Mk distribution on day 11. Data shown represent the mean ± standard error of mean. Based on a paired t-test, values of p<0.0005 (x) and p<0.05 (*) are indicated for the various time points in comparison to the Tpo-only culture.
NIC increases Mk ploidy by inhibiting SIRT1 and/or SIRT2. In an effort to decouple the effects of SIRT1 vs SIRT2 inhibition, we evaluated AGK2, which has been shown to decrease SIRT2 activity in vitro without significantly affecting SIRT1 or SIRT3 activity at concentrations up to 10 μM [53]. When added at 10 μM, AGK2 moderately increased the percentage of high-ploidy Mks (Fig. 4C). At day 11, the AGK2-mediated increase in high-ploidy cells was about 30% as great as that for NIC. The Mk ploidy distribution (Fig. 4D) and mean ploidy value (4.3±0.2 for NIC; 3.3±0.1 for AGK2, and 3.0±0.1 for Tpo only) at day 11 for 10 μM AGK2 were intermediate between those for Tpo only and 6.25 mM NIC. It is unlikely that higher AGK2 concentrations would be more effective because the ploidy difference between 5 and 10 μM AGK2 was small (data not shown) and because 10 μM AGK2 inhibited cell growth to a similar extent as 6.25 mM NIC (Supplementary Figure E2F, online only, available at www.exphem.org). AGK2 (10 μM) had similar effects on the fraction of CD41+ cells (Supplementary Figure E2E, online only, available at www.exphem.org) as NIC, but AGK2 yielded a slower decline in cell viability (Supplementary Figure E2 G, online only, available at www.exphem.org) and a slower increase in the fraction of apoptotic Mks (Supplementary Figure E2H, online only, available at www.exphem.org).

NIC-mediated SIRT inhibition correlates with increased p53 binding to its consensus DNA binding sequence

EMSA experiments were performed to determine whether NIC-mediated p53 acetylation increased p53 DNA-binding activity. As reported previously [20], two low-mobility DNA complexes were formed during these experiments and their levels increased during megakaryopoiesis (Fig. 6A). Both of the low-mobility DNA complexes were
Figure 5. Nicotinamide (NIC) increases acetylation of p53. Mobilized peripheral blood (mPB) CD34⁺ cells were cultured with 100 ng/mL thrombopoietin (Tpo). On day 5 cultures were supplemented with 6.25 mM NIC. (A–D) Intracellular flow cytometry was used to detect expression of AcK382p53 and total p53 in (A, B) CD41⁺ and (C, D) CD41⁻ cells. Mean fluorescence intensity (MFI) was calculated by subtracting the MFI of an unstained sample from the MFI of a stained sample to correct for background fluorescence. Data shown represent the mean ± standard error of mean of two experiments. (A, C) MFI of cells stained with an antibody against AcK382p53 was divided by the MFI of cells stained with an antibody against total p53. This ratio is shown for cells maintained in Tpo with or without NIC. (B, D) MFI of cells grown with NIC was divided by the MFI of cells grown with Tpo only. This ratio was calculated for cells stained either with an antibody against AcK382p53 or against total p53. Based on a paired t-test, values of p < 0.1 (*) are indicated for the various time points. (E) Nuclear lysates of unselected cells were prepared at all time points shown and subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis. After electrophoresis, the proteins were transferred to polyvinylidene difluoride membranes for Western blots. After probing for AcK382p53 residues, blots were stripped and probed for total p53. Corresponding densitometry analysis is given. The blot shown is representative of two biological experiments.
used in subsequent densitometry analyses because the binding affinity of both regions varied during the course of CHRF cell Mk differentiation and both were specific to the p53 consensus binding sequence oligonucleotides [20]. Consistent with the greater DNA-binding activity reported for acetylated p53 [56,57], p53 DNA-binding activity was greater in cells cultured with NIC at every time point investigated (Fig. 6A and B). Although the differences were not statistically significant for the individual time points, nonparametric analysis of all of the EMSA data using the Kruskal-Wallis test confirmed that the consistently greater p53 DNA-binding activity observed for cells treated with NIC was significant ($p$ = 0.033). Results at days 7, 9, and 11 from another biological experiment conducted using the TransBinding p53 ELISA kit (Panomics, Freemont, CA, USA) also showed consistently greater p53 DNA-binding activity in the nuclear extracts of cells cultured with Tpo plus NIC compared to those cultured with Tpo alone (data not shown).

Discussion

We previously showed that NIC increases Mk ploidy and proplatelet production without significantly altering Mk-specific gene expression in cultures of mPB CD34$^+$ cells supplemented with Tpo [8]. Here, we show that NIC does not alter normal Mk ultrastructure (Fig. 1) and demonstrate the generality of NIC effects. NIC synergizes with Tpo to further enhance Mk maturation and is effective at increasing polyploidization under all of the Mk-promoting conditions we have investigated (Fig. 2). Therefore, NIC is likely to be effective under a wide range of other conditions that promote Mk maturation.

Increases in Mk ploidy may be due to direct modulation by NIC, but may also be mediated indirectly by elevated levels of NAD(H). We observed a fivefold increase in NAD(H) levels when 6.25 mM NIC was added to Mk cultures (Fig. 3). We also examined QA, which is incorporated into NAD$^+$ via the de novo synthesis pathway. QA at 10 mM increased the level of intracellular NAD(H) to a similar extent as 1 mM NIC, but had little effect on Mk ploidy. The de novo pathway is distributed throughout the cell [34], while the salvage pathway used for NIC incorporation has been shown to be primarily localized to the nucleus in yeast cells [35,36]. Greater levels of nuclear NAD-processing enzyme Nmnat1 in NIC-treated cells further support the hypothesis that NAD(H) synthesis in these cells is enhanced in the nucleus. Increased nuclear levels of NADH, induced by hypoxic conditions, cause a conformational change in hCtBP proteins that reduces their ability to bind with transcriptional corepressors [58], such as hdm2, and together inhibit p53 activation. Thus, higher levels of nuclear NADH result in increased p53 activity and greater activation of its downstream targets [59]. However, we note that cambinol would not be expected to increase NAD$^+$ synthesis and we did not observe

Figure 6. Nicotinamide (NIC) increases p53 DNA-binding activity. (A) Mobilized peripheral blood (mPB) CD34$^+$ cells were cultured using 100 ng/mL thrombopoietin (Tpo) with or without 6.25 mM NIC beginning on day 5. At selected time points nuclear extracts were prepared for electrophoretic mobility shift assay analysis. The gel shown is representative of three replicates from two biological experiments. Low mobility p53-DNA complexes (*) were formed using biotinylated p53 consensus binding sequence oligonucleotides. Nonspecific binding is indicated by a (+) sign. (B) Densitometry of low mobility p53-DNA complexes is shown as the mean ± standard error of mean of arbitrary density units for two replicates from one biological experiment.
any increase in NAD(H) levels when the megakaryocytic CHRF cell line was treated with cambinol (data not shown).

NIC has been extensively characterized as an inhibitor of SIRTs, which were identified as NAD+-dependent class III histone deacetylases [39–41]. SIRTs play important roles in transcriptional silencing, genetic stability, and cellular lifespan regulation [13,14,42]. Deacetylation by the nuclear protein SIRT1 [42] regulates the activity of proteins associated with DNA repair, apoptosis, and cell cycle regulation including NBS1, p53, FOXO3a, RelA/p65 (nuclear factor κB), and Ku70 [15,54,60–63]. In contrast, the tubulin deacetylase SIRT2 can be localized to either the nucleus or the cytoplasm [11,43]. Recent studies suggest that SIRT2 regulation of α-tubulin acetylation plays an important role in modulating the extension of cell processes. Knockdown of SIRT2 increased hippocampal neurite outgrowth [64] and the arborization complexity of primary oligodendrocyte precursors [65]. SIRT2-specific inhibitor AGK2 was 30% as effective as NIC in increasing Mk ploidy (Fig. 4C and D). In contrast, cambinol, which inhibits both SIRT1 and SIRT2, increased Mk ploidy almost as effectively as NIC (Fig. 4A and B). This suggests that NIC primarily increases Mk ploidy by inhibition of SIRT1 and/or SIRT2. Together with our previous finding that SIRT1 and SIRT2 mRNA levels increase during Mk maturation [8], our results suggest that SIRT1 and/or SIRT2 are involved in controlling the transition from endomitosis, and that inhibiting SIRT1/2 delays this process and increases Mk ploidy.

Functional inhibition of SIRT1 and/or SIRT2 in NIC-treated Ms was confirmed using two well-established targets of SIRT deacetylation: nucleosomes and p53 (Fig. 5). Histone deacetylation within nucleosomes generally leads to the repression of transcription because removal of acetyl groups from lysine residues leads to chromatin becoming more compact, such that specific regions of DNA are less accessible for transcription factor binding [66]. Increased acetylation of residue K382 on p53 in Mks treated with NIC (Fig. 5A, B, and E) was associated with increased binding of p53 to its consensus DNA binding sequence (Fig. 6). Acetylation of K382 on p53 can lead to cell cycle arrest and apoptosis [67,68], as well as activation and stabilization of p53, which may prevent p53 ubiquitination and degradation [69]. Although Mk viability and apoptosis did not substantially change upon treatment with NIC (Supplementary Figure E2C, D, G, H, online only, available at www.expem.org), p53 has widespread effects through its regulation of other cell cycle, DNA repair, and lifespan-related targets including CDC2, GADD45, p21, and 14-3-3s [70,71].

We previously demonstrated the importance of p53 in Mk polyploidization and apoptosis [20,21,72]. Gene expression analysis of mPB CD34+ cells cultured under Mk-promoting conditions revealed differential expression of several transcriptional targets of the p53 protein. Using RNAi to reduce p53 expression in CHRF cells, we showed that lower p53 activity leads to a greater fraction of polyploid cells, higher mean and maximal ploidy, accelerated DNA synthesis, and delayed apoptosis upon Mk differentiation [20]. Recent results in our laboratory indicate that posttranslational p53 modifications (phosphorylation and acetylation) take place in an Mk-specific manner and apparently play an important role in Mk maturation (data not shown). Consistent with this hypothesis are the findings reported here whereby NIC increased p53 acetylation and p53 DNA-binding activity. This suggests that the effect of NIC, as a SIRT1/2 inhibitor, is mediated at least in part by increased p53 acetylation. While it seems counterintuitive that increased p53 DNA-binding activity and p53 knockdown would both increase Mk ploidy, p53 has many different target genes, whose activity may be differentially regulated by SIRT1 and/or SIRT2. A genome-wide ChIP-on-chip analysis identified 1546 p53-binding sites after treatment with actinomycin D [73].

It is likely that mechanistic understanding of the modulation of p53 activity and its targets on Mk differentiation, as well as those of other SIRT target proteins, will lead to additional means for enhancing Mk polyploidization. One promising p53 target with increased transcription as a result of increased p53 acetylation is p21WAF1 [54]. Levels of p21WAF1 were approximately threefold higher in a Sir2-deficient cell line than with WT Sir2 [54]. Using quantitative reverse transcription polymerase chain reaction, we found that p21 expression was significantly reduced in p53 knockdown CHRF cells [20]. Expression of p53 and p21WAF1 was correlated with increasing nuclei size and DNA content in a human lung carcinoma cell line [74]. In addition, p21WAF1 expression increased in an erythroleukemia cell line induced to undergo polyploidization [75]. We are investigating p21 expression in Mks with inhibited or knocked-down SIRT1 and/or SIRT2 in comparison to primary human Mks cultured with Tpo alone to determine whether p21 expression correlates with increased Mk ploidy.

Acknowledgments

We are grateful to Genentech (San Francisco, CA, USA) for Tpo donation and to Dr. Antonio Bedalov (Fred Hutchinson Cancer Research Center; Seattle, WA, USA) for donation of cambinol and helpful discussions. We thank Aaron Kuhl for initial development of the NAD(H) assay. We thank Dr. Mohamed Eldibany of the NorthShore University (Evanston, IL, USA) Health System for help with analysis of electron microscopy images and Lennell Reynolds of the NU Cell Imaging facility for electron microscopy sample preparation. Supported by National Institutes of Health (NIH; Bethesda, MD, USA) grant R01HL48276, a grant from the NorthShore University Health System Pathology Department, and the Robert H. Lurie Comprehensive Cancer Center of Northwestern University (Chicago, IL, USA) Malkin Family Scholarship. S.P. was supported in part by NIH Biotechnology Predoctoral Training Grant T32 GM 008449. P.A.A. was
supported in part by funds through the Delaware Biotechnology Institute (Newark, DE, USA) and an Onassis Foundation fellowship.

Conflict of Interest Disclosure
No financial interest/relationships with financial interest relating to the topic of this article have been declared.

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Supplementary Figure E1. Nicotinamide (NIC) and cambinol synergize to slightly increase megakaryocytic cell (Mk) ploidy. (A, B) Mobilized peripheral blood (mPB) CD34⁺ cells were cultured with 100 ng/mL thrombopoietin (Tpo). On day 5, cultures were supplemented with 6.25 mM NIC, 10 μM cambinol, or 3.125 mM NIC plus 5 μM cambinol (n=4). (C, D) Cells from a subset of the experiments shown in (A, B) were also treated with 6.25 mM NIC plus 10 μM cambinol (n=2). Flow cytometry was used to determine (A, C) the percentage of CD41⁺ cells with DNA content ≥8 N (high-ploidy Mks) or (B, D) the high-ploidy Mk distribution on day 11. Data shown represent the mean ± standard error of mean.
Supplementary Figure E2. Effects of sirtuin (SIRT) inhibitors on CD41 expression, total-cell expansion, viability, and megakaryocytic cell (Mk) apoptosis. Mobilized peripheral blood (mPB) CD34+ cells were cultured with 100 ng/mL thrombopoietin (Tpo). On day 5 cultures were supplemented with (A–D) 10 μM cambinol or (E–H) 10 μM AGK2. For comparison, replicate cultures were either supplemented with 6.25 mM nicotinamide (NIC) on day 5 or maintained with Tpo alone. (A, E) Percentage of CD41+ cells in the viable population. (B, F) Total-cell fold-expansion. (C, G) The percentage of viable cells in culture. (D, H) The percentage of apoptotic Mks in the viable population. Data shown represent the mean ± standard error of mean for n = 12 (A–D) or n = 11 (E–H) experiments. Based on a paired t-test, values of p < 0.0005 (x) and p < 0.05 (*) are indicated for the various time points in comparison to the Tpo only culture.
Supplementary Figure E3. Nicotinamide (NIC) increases acetylation of nucleosomes. Whole cell lysates were loaded into gels for sodium dodecyl sulfate polyacrylamide gel electrophoresis. After electrophoresis, the proteins were transferred to nitrocellulose membranes for Western blots. After probing for acetylated lysine residues, the blots were stripped and probed for nucleosomes. Corresponding densitometry analysis is given. The blot shown is representative of four biological experiments.