Low NAD⁺ Levels Are Associated With a Decline of Spermatogenesis in Transgenic ANDY and Aging Mice

Mirella L. Meyer-Ficca¹,²*, Alexie E. Zwerdling², Corey A. Swanson¹, Abby G. Tucker², Sierra A. Lopez¹,², Miles K. Wandersee¹,², Gina M. Warner³,⁴, Katie L. Thompson³,⁴, Claudia C.S. Chini³,⁴, Haolin Chen⁵†, Eduardo N. Chini³,⁴ and Ralph G. Meyer¹,²*

¹ School of Veterinary Medicine, Utah State University, Logan, UT, United States, ² Department of Animal, Dairy, and Veterinary Sciences, College of Agriculture and Applied Sciences, Utah State University, Logan, UT, United States, ³ Signal Transduction and Molecular Nutrition Laboratory, Kogod Aging Center, Department of Anesthesiology and Perioperative Medicine, Mayo Clinic College of Medicine, Rochester, MN, United States, ⁴ Department of Anesthesiology and Perioperative Medicine Mayo Clinic, Jacksonville, FL, United States, ⁵ Department of Biochemistry and Molecular Biology, Johns Hopkins Bloomberg School of Public Health, Baltimore, MD, United States

Advanced paternal age has increasingly been recognized as a risk factor for male fertility and progeny health. While underlying causes are not well understood, aging is associated with a continuous decline of blood and tissue NAD⁺ levels, as well as a decline of testicular functions. The important basic question to what extent ageing-related NAD⁺ decline is functionally linked to decreased male fertility has been difficult to address due to the pleiotropic effects of aging, and the lack of a suitable animal model in which NAD⁺ levels can be lowered experimentally in chronologically young adult males. We therefore developed a transgenic mouse model of acquired niacin dependency (ANDY), in which NAD⁺ levels can be lowered using a niacin-deficient, chemically defined diet. Using ANDY mice, this report demonstrates for the first time that decreasing body-wide NAD⁺ levels in young adult mice, including in the testes, to levels that match or exceed the natural NAD⁺ decline observed in old mice, results in the disruption of spermatogenesis with small testis sizes and reduced sperm counts. ANDY mice are dependent on dietary vitamin B3 (niacin) for NAD⁺ synthesis, similar to humans. NAD⁺-deficiency the animals develop on a niacin-free diet is reversed by niacin supplementation. Providing niacin to NAD⁺-depleted ANDY mice fully rescued spermatogenesis and restored normal testis weight in the animals. The results suggest that NAD⁺ is important for proper spermatogenesis and that its declining levels during aging are functionally linked to declining spermatogenesis and male fertility. Functions of NAD⁺ in retinoic acid synthesis, which is an essential testicular signaling pathway regulating spermatogonial proliferation and differentiation, may offer a plausible mechanism for the hypospermatogenesis observed in NAD⁺-deficient mice.

Keywords: vitamin B3, niacin, nicotinamide, testis, aging, retinoic acid, spermatogonia, male fertility
INTRODUCTION

Associated with socioeconomic considerations, for example increased time needed for education and professional development, mean paternal age has increased over the past 44 years from 27.4 to 30.9 years (1). This is concerning because paternal age has been shown to negatively affect fertility, pregnancy rates and children’s health (2, 3). How exactly the aging process exerts its negative effects on male fertility is not clear, because of its pleiotropic effects on the body, including the testis (4, 5). Although underlying mechanisms are not yet well understood, one of the hallmarks of aging is a steady decline of cellular, tissue and plasma NAD⁺ concentrations, observed during chronological aging in humans, worms, flies, and mice (6–10). NAD⁺ and NADP⁺, and their reduced forms NADH and NADPH, are important coenzymes for most cellular redox reactions, and as such essential for maintaining cellular metabolism and respiration. In addition to its function as a redox cofactor, NAD⁺ is also consumed by enzymes involved in chromatin modification, gene regulation, and DNA repair, including poly(ADP-ribose) polymerases (PARP family of enzymes), as well as NAD-dependent protein deacetylases (sirtuins) and CD38 (11–13).

Unfortunately, the links between aging, low NAD⁺ levels and declining fertility are not well understood because systematic investigations have been hampered by basic metabolic differences present between laboratory rodents and humans in their ability to generate NAD⁺ from their diet.

In certain mammals, including humans, nicotinic acid (NA), nicotinamide (Nam) and Nam riboside (NamR), collectively referred to as niacin or vitamin B3, are the main nutritional precursors of NAD⁺ and its phosphorylated form, NADP⁺. Humans depend on dietary niacin as their main source of NAD⁺ and NADP⁺ precursors and can become niacin-deficient when their food lacks sufficient amounts of vitamin B3. Nicacin deficiency is characterized by very low levels of NAD⁺ and in its most extreme form, pellagra, can be debilitating and even deadly, which is now rare in western countries. However, milder forms of clinical niacin deficiency are commonly seen with increasing age, and in cancer patients, alcoholics and people without access to quality food (14, 15). While this may be clinically relevant on its own, it is unlikely that a lack of dietary vitamin B3 intake is at the root of age-related NAD⁺ decline. Instead, age-related increases in the activity of NAD⁺-consuming enzymes such as PARP1 and CD38, or potential mitochondrial dysfunction, or both, provide a more plausible explanation [(12, 16, 17), reviewed in (18)].

Physiological effects of low NAD⁺ status and their potential impact on male fertility have been difficult to study because of a lack of suitable animal models. Wild-type laboratory rodents are able to completely satisfy their NAD⁺ needs by metabolizing tryptophan (Trp) to NAD⁺ via the kynurenine (de novo synthesis) pathway and, unlike humans, do not depend on intake of dietary niacin. In order to address this problem and to investigate the impact of low NAD⁺ levels as a potential factor contributing to the decline of fertility in aging males, we therefore generated mice with tetracycline-inducible overexpression of a transgene encoding the enzyme human aminocarboxymuconate semialdehyde decarboxylase (hACMSD) to create a mouse model of human-like NAD⁺ metabolism (ANDY, acquired niacin dependency) (19). In this mouse, hACMSD overexpression diverts the central kynurenine pathway in the liver and kidney to produce acetyl-CoA instead of NAD⁺ which makes the animals dependent on dietary niacin intake as the main source of NAD+ synthesis, similar to humans (19) (Figure 1A). ANDY mice with hACMSD overexpression reproducibly become NAD⁺-deficient in various tissues over the course of 6 weeks on a defined diet that is devoid of niacin (ND diet), but not on a control diet that is chemically identical to ND but supplemented with 30 mg/kg nicotinic acid (CD diet). Previous data showed that ANDY mice had significantly lower NAD⁺ and NADP⁺ levels in blood, liver, and other tissues when they received a niacin-free ND diet and doxycycline (Dox, a water-soluble tetracycline) in their drinking water (19). If maintained at very low NAD⁺ levels, male ANDY mice sired smaller litters than control males (data not shown).

The goal of the current study has therefore been to investigate the impact of NAD⁺ deficiency on spermatogenesis in young adult ANDY mice to test the hypothesis that low NAD⁺ levels have a negative impact on male fertility, independent of chronological age.

MATERIALS AND METHODS

Animal Model and Induction of NAD Deficiency

Details of the generation of the transgenic animal model C57BL/6J-Gt(Rosa)26Sortm1(TTAa-M2)JaeCol1a1m6(tetO-hACMSD)MMF and the biochemical basis of NAD-dependency in these mice has been described previously (19). Briefly, administration of doxycycline, a water-soluble tetracycline, in the drinking water induces overexpression of the human aminocarboxymuconate semialdehyde decarboxylase (hACMSD) gene. Increased ACMSD activity renders these transgenic mice dependent on dietary niacin uptake in a manner similar to humans. In the absence of dietary niacin, these ANDY mice become measurably NAD⁺ deficient in blood and body tissues (19). Mice were bred and housed under standard conditions. Transgene expression was only induced in adult mice during the feeding trials. Breeding, postnatal and pubertal development occurred in the absence of doxycycline-induced transgene overexpression and on normal, niacin-containing chow diet. Animal studies and experimental procedures were approved by the Institutional Animal Care and Use Committees (IACUC protocol number 10056) of Utah State University and of Mayo Clinic, Rochester, Minnesota.

Defined Feeds and Feeding Trials

Standard chow diet was Teklad Rodent diet 8604 (24% crude protein, 63 mg/kg niacin, Envigo, Madison, WI, USA). Niacin-deficient diet (ND, TD.140376) and control diet (CD, TD.140375) were defined, purified diets compounded by Teklad laboratory animal diets (Teklad Custom Diets, Envigo) as modifications of AIN-93G standard chow (19). Both, ND and CD contained 10% alcohol-washed casein as a vitamin-free
NAD\(^+\) Requirements of the Testis

...niacin deficiency altered NAD metabolite profiles in ANDY mice. ANDY mice were kept on niacin-deficient (ND) for up to 12 weeks (ND-Dox_S) or longer (ND-Dox_L) or on control diet with Dox for ACMSD transgene induction (CD+Dox) or without Dox (CD+H2O). Old mice were 31 months old. (A) The NAD de novo synthesis pathway from tryptophan can provide all of the NAD in wild-type rodents in absence of all other dietary NAD precursors such as nicotinic acid (NA), nicotinate, nicotinamide (Nam), or nicotinamide ribonucleotide (nicotinamide riboside, NamR). Dox-mediated induction of a human ACMSD transgene overexpression diverts the central kynurenine pathway in the liver and kidney from NAD\(^+\) production towards acetyl-CoA formation, and ultimately makes these mice dependent on dietary niacin to maintain tissue NAD\(^+\) levels similar to humans. (B) Testicular NAD\(^+\) levels decline in ANDY mice on vitamin B3 (niacin) – free diet (ND). Data were generated using enzymatic cycling assays. (C) Metabolomic analyses confirms data in B, and indicates that a short-term period on ND diet (ND-Dox_S) results in a milder decline of NAD\(^+\) that is comparable to old mice. (D) Nam levels declined already after short-term dietary niacin-deficiency, while NamR (E) values appeared to increase again in mice with long-term niacin deficiency. Hypothetically, the latter may result from loss of spermatogonia, spermatocytes and spermatids that occurs at later stages of NAD\(^+\) decline. (F) Whole NA (nicotinate/nicotinic acid) did not change on the ND diet, while MNM (G) was significantly lowered in both short- and long-term ND fed mice compared to controls, but not old mice. Identical letters indicate group categories that are not significantly different from each other; different letters indicate statistically significant differences (One-way ANOVA with Tukey’s multiple comparison analysis, p<0.05 considered a significant difference; **p > 0.001).
testosterone was measured using standard RIA procedure with a
testosterone specific antibody (ICN Biomedicals, Costa Mesa,
CA) and ³H-T (NEN Life Science Products, Boston, MA).

Quantification of Metabolites in
Testicular Tissue
Testicular testosterone, nicotinamide adenine dinucleotide (NAD),
nicotinamide (Nam), nicotinamide riboside (NamR), nicotinic acid
(NA) and NAD mononucleotide (NMN) were quantified with
Ultrahigh Performance Liquid Chromatography-Tandem Mass
Spectroscopy (UPLC-MS/MS) on the Metabolon Platform
(Metabolon, Norristown, NC). Frozen testis samples were
prepared using the automated MicroLab STAR® system
(Hamilton), proteins were precipitated with methanol followed
by centrifugation. The resulting extract was analyzed by two
separate reverse phase (RP)/UPLC-MS/MS methods with positive
ion mode electrospray ionization (ESI), by RP/UPLC-MS/MS and
by HILIC/UPLC-MS/MS, both with negative ion mode ESI. UPLC-
MS/MS was performed on a Waters ACQUITY UPLC and a
Thermo Scientific Q-Exactive high resolution/accurate mass
spectrometer interfaced with a heated electrospray ionization
(HESI-II) source and Orbitrap mass analyzer operated at 35,000
mass resolution. The sample extract was dried, then reconstituted
in solvents compatible to each of the four spectroscopy methods. Each
reconstitution solvent contained a series of standards at fixed
concentrations to ensure injection and chromatographic
consistency. The MS analysis alternated between MS and data-
dependent MSn scans using dynamic exclusion. The scan range
varied slightly between methods but covered 70-1000 m/z. Raw
data was extracted, peak-identified and QC processed using
Metabolon’s hardware and software. Compounds were identified
by comparison to library entries of purified authenticated
standards, and peaks were quantified using area-under-the-curve.

Graphing and Statistical Analyses
GraphPad Prism software versions 7.04 & 9.2.0 (GraphPad
Software, San Diego, CA) were used for graphing and statistical
analyses (One-way ANOVA, Tukey’s multiple comparison,
Welch’s t-test, Pearson Correlation analysis;
p<0.05 was considered significant).

RESULTS

ANDY Mice on Niacin Free Diet Have
Significantly Reduced Testicular NAD⁺
In the absence of dietary niacin, blood NAD of ANDY mice
dropped steadily over time a time span of 6 weeks, and then
remained at significantly lower levels compared to those in
control animals (Supplementary Figure 1). Similar to blood,
testes of ANDY mice became niacin-deficient, i.e. had
significantly decreased NAD⁺ tissue concentrations. NAD⁺
levels in ANDY mice that were maintained on ND+Dox for 24
weeks dropped to about 1/3 of the NAD⁺ content measured in
ANDY mice fed niacin-containing CD diet or chow (Figure 1B),
as measured using a sensitive enzymatic cycling assay
(Supplementary Table 1). Similar changes were observed
using comparative metabolomics analyses of testes from
animals fed ND for 24 weeks (long-term, ND+Dox_L) or 12
weeks (short-term, ND+Dox_S). The metabolomic LC-MS/MS
quantification confirmed the significant lowering of testicular
NAD⁺ levels in the ND+Dox_L group (long-term on ND diet, i.e.
>12 weeks, one-way ANOVA, p-values from <0.0001 to 0.0052
with Tukey’s multiple comparison test, Figure 1C). Compared to
this group, NAD⁺ levels were higher in ANDY mice kept on ND
for 12 weeks (ND+Dox_S, p=0.0052), but still significantly lower
than the control groups (p-values from 0.0004 to 0.0055). NAD⁺
contents in controls CD+Dox and CD+H₂O were not
significantly different from each other, indicating that ACMSD
overexpression and doxycycline administration on their own did
not have any measurable effect on NAD⁺ levels in the testis.
Interestingly, NAD⁺ content in the testes of the ND+Dox_S
group was not significantly different from that of old mice at 31
months of age. Nam levels were low in both short- and long-term
ND groups (Figure 1D). Unexpectedly, NamR levels were
significantly higher in the ND+Dox_L group than ND+Dox_S
(p=0.0011), but not significantly different from the CD+Dox
control and the old mice (Figure 1E). NA values did not vary
between the different treatment groups (Figure 1F). Similar to
NAD⁺ and NamR, NMN was not significantly different between
mice in the ND+Dox_S group and old mice (Figure 1G).

Taken together, ACMSD overexpression in combination with
niacin-free feed significantly lowered testicular NAD⁺ levels
of ANDY mice, which is also reflected in an altered NAD⁺
metabolite profile. Moreover, the NAD⁺ levels created in
ANDY mice of the ND+Dox_S group were similar to those in
old mice at 31 months of age.

Declining Testicular Weight and Sperm
Counts in NAD⁺-Deficient ANDY mice
Sperm counts of mice that were kept on ND diet for ten weeks
decreased significantly compared to control animals, and were
similar to sperm numbers in old mice (Figure 2A). After two
additional weeks on deficient diet, sperm numbers declined
abruptly (Figure 2A). Along with falling sperm counts, testes
of mice in the ND+Dox group became significantly smaller than
testes of any other treatment group as soon as 10 weeks on this
diet, and continued to shrink until week 24 (Figure 2B). When
recovered on the CD diet for 9 weeks, testis weights returned to
normal values (Figures 2B, C). These results demonstrate that
decreasing NAD⁺ levels resulted in testicular shrinkage that was
reversed by niacin supplementation which restored NAD⁺ levels.

NAD⁺-Deficiency Causes a Reversible
Cessation of Spermatogenesis
Histological evaluation of the testicular shrinking process (Figure 3)
in testis from animals on the niacin-free ND diet revealed progressive
seminiferous epithelial defects compared to control animals on CD
+Dox (Figures 3A–D). Seminiferous tubules showed a lack of
ongoing spermatogenesis with severely decreased numbers of
spermatogonia and spermatocytes, as well as an abnormal spatio-
temporal organization (Figures 3C, D). Seminiferous epithelia of
animals kept on ND+Dox for 24 weeks, followed by recovery on niacin-containing CD+Dox for 9 weeks, were restored to full cell complements, consistent with the observed reversal of testicular shrinkage (Figures 2B, D). Sperminiferous tubes of older mice at 20 month-old appeared mostly normal, except for the appearance of sporadic abnormal sperminiferous tubes (asterisk in Figure 3F), while seminiferous tubes in testes of 31 month-old mice displayed marked and frequent disorganization of the seminiferous tubes (Figure 3G). After 24 weeks on ND+Dox diet, seminiferous tubules were lined mostly by Sertoli cells and some spermatogonia, and contained cells that appeared to be mostly residual round and some elongated spermatids (Figure 3H). Quantification of abnormal tubules in testis sections after 16 weeks on the indicated diets showed that NAD+ -deficient testes contained significantly more tubules with abnormal composition of the seminiferous epithelium than controls or mice that were first kept on ND+Dox diets for 24 weeks and then recovered on CD for 9 weeks (Figure 3I). Taken together, the histological results suggested that a lack of spermatogonial proliferation led to a paucity of promeiotic and meiotic germ cells, which together make up more than half of the testicular weight and size in a normal animal. The remarkable recovery of spermatogenesis and subsequent doubling of testicular volume to a normal state in animals recovered on CD diet further indicates that spermatogonial stem cells remained intact and capable of restoring full spermatogenesis once NAD+ levels returned to normal levels (Figures 2B, C and 3E).

**DISCUSSION**

The main results of this study are (i) that NAD+ deficiency can be produced in ANDY mouse testes on a niacin-free diet, and that...
the degree of this deficiency increases over time (Figure 1). To our knowledge, this is the first time this has been accomplished in a laboratory research animal. (ii) The degree of NAD+ decline that was achieved by keeping ANDY mice on niacin-free diet for 10-12 weeks was typical of an aging mouse (Figure 1). (iii) Low testicular NAD+ levels resulted in the attenuation of spermatogenesis and testicular atrophy due to impaired spermatogonial proliferation and differentiation (Figure 3). (iv) Recovery of mice on a niacin-containing control diet fully reversed testicular shrinkage and fully restored spermatogenesis (Figure 3E). Because NAD+ decline resulted in attenuation of spermatogenesis in ANDY mice, it may represent a link between low NAD metabolism as a hallmark of aging, and the decline of male fertility as males age.

Based on our data, low testosterone levels were not the determining factor for the observed hypo- and aspermatogenesis (Figure 4). However, the loss of mature germ cell stages in severely NAD+ depleted testes and overall seminiferous tubule histology was reminiscent of vitamin A-deficient males, where tubules appear to have only Sertoli cells and early stages of spermatogonia left in the tubular lumen (24). Vitamin A1 (retinol) is essential for spermatogenesis because it is the dietary precursor for RA synthesis (25) (Figure 4I). RA signaling is indispensable for spermatogonial proliferation and differentiation. If blocked by the inhibitor WIN 18,446 in adult rodents or humans, spermatogonial differentiation is disrupted and a vitamin A deficiency phenotype is created in the testis (24, 26–28). The rate-limiting step of RA synthesis is the oxidation of retinol to retinal by retinol dehydrogenase (RDH10), which is entirely dependent on the availability of NAD+ as a cofactor. In addition, the next step in RA synthesis is the conversion of retinal to retinoic acid, which is dependent on NADP+, whose levels are linked to cellular NAD+ stores (Figure 4D). This step, which is mediated by the aldehyde dehydrogenase (ALDHA) family of enzymes is also essential for testicular RA synthesis, and thus for the execution of spermatogenesis (27). Our finding that retinol appeared to accumulate to significantly higher levels in both NAD+-deficient and aging mice in a manner significantly inversely correlated with NAD+ levels (Figure 4E) therefore provides an intriguing clue that low NAD+ levels may block RA synthesis and thus cause the observed spermatogenic failure. However, additional investigations will be necessary to provide further confirmation of this hypothesis. Mechanisms underlying the aging process are still poorly understood, in part because effects of chronological aging are numerous and difficult to separate from environmental and intrinsic factors affecting a given individual over time. The NAD+ decline

![Figure 3](https://example.com/fig3.png)

**FIGURE 3** | Impact of testicular NAD+ levels and male age on the seminiferous epithelium. Hematoxylin/eosin-stained testicular sections of testes from young adult male fed (A) CD+Dox control diet, (B) ND+Dox diet for 8 weeks, (C) ND+Dox for 14 weeks; damaged tubules (D) ND+Dox for 24 weeks; normal tubules +/- absent (E) ND for 24 weeks followed by 9 weeks of recovery on niacin-containing CD diet; tubules mostly restored. (F) Control testis at 20 months of age. Asterisk marks a tubule with abnormal seminiferous epithelium. (G) Control testis of 31 month-old mouse. (H) After 24 weeks on ND diet, seminiferous tubules are lined mostly by Sertoli cells (SC, blue arrow heads) interspersed with spermatogonia (SG, red stars), as identified by histological morphology of the cells. Tubular lumen contain mostly cells resembling round spermatids (RS, black arrow) and occasionally elongated spermatids (ES)). (I) NAD+-deficient testis contain significantly more tubules with abnormal composition of the seminiferous epithelium in testis of mice that were on indicated diets for 16 weeks (CD+Dox, CD+water, ND+Dox) or ND+Dox that were subsequently recovered on niacin containing diet for 9 weeks. One hundred tubules were evaluated per testis section, ANOVA with Tukey’s multiple comparison, b is significantly different from a, p=0.0003 to 0.001. Scale bar: 250 mm in a.-f., 40 mm in (g) & h.
observed in aging animals and humans appears to be a consequence of the aging process, for example by means of failing mitochondrial activity, or through elevated consumption of NAD+ by PARP enzymes or elevated tissue activity of the NAD glycohydrolase CD38 (7, 16–18, 29–32). However, to what extent NAD+ decline itself may also be a driver of the aging process has remained an open question. The current study takes full advantage of the novel ANDY mouse model that allows for the first time that NAD+ levels in rodents can be lowered significantly, independent of the chronological age of the animal. The results of this initial investigation suggest that low and very low levels of NAD+ result in testicular decline in mice, similar to that observed in aging males. This finding suggests that NAD+ decline itself may promote aspects of the pathophysiology of aging.

NAD and NADP serve not only as an essential cofactor for enzymatic reaction in energy metabolism; they are also essential cofactors for several cellular mechanisms that protect the genome against DNA damaging insults, e.g. from reactive oxygen species (ROS). There is an age-related increases in ROS, the so-called “free radical theory of aging”, that is also evident in context of spermatogenesis and sperm quality (33–35). In fact, aging has been associated with reduce genetic quality in spermatogenic cells and sperm (36–38). Because NAD and NADP are required for both, maintaining a sufficient pool of the active antioxidant glutathione GSH, and for the enzymatic activity of PARP1, an important DNA repair factor, lower testicular NAD could potentially contribute to the aging-related accumulation of ROS and decline in sperm quality. Investigations are currently underway to address this important question.

A potential limitation of the present study is that the degree of testicular NAD+ decline produced in ANDY mice that were on ND for a long period of time (exceeding 12 weeks) may arguably be more severe than the NAD+ deficiency measured in the 31 month-old mice. On the other hand, while the spermatogenic defects observed in these old mice may be less severe, they were clearly detectable and may at least in part be caused by NAD+ deficiency. Furthermore, the dynamics of human testicular NAD+ decline with age may be different from mice, along with its importance for human male fertility, which will require further research. Additional investigations are currently underway to determine the role of NAD+ decline in the aging process in ANDY mice. In summary, this study is the first one to show that experimentally induced low testicular NAD+ levels result in reversible disruption of spermatogenesis, adding vitamin B3 to the list of vitamins that are essential for proper spermatogenesis in humans. The study also provides clues to the role of NAD+ decline in the age-related decline of testicular function and male fertility.
DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The animal study was reviewed and approved by Institutional Animal Care and Use Committees (IACUC) of Utah State University and of Mayo Clinic, Rochester, Minnesota.

AUTHOR CONTRIBUTIONS

MM-F and RM contributed conception and design of the study, experimentation, data analysis and preparation of the manuscript. AZ, CS, AT, SL, MW and HC performed experiments, collected and analysed data. GW, KT, CC, and EC contributed materials. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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