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# A functional link between NAD<sup>+</sup> homeostasis and N-terminal protein acetylation in *Saccharomyces cerevisiae*

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Edited by John M. Denu

Nicotinamide adenine dinucleotide (NAD<sup>+</sup>) is an essential metabolite participating in cellular redox chemistry and signaling, and the complex regulation of NAD<sup>+</sup> metabolism is not yet fully understood. To investigate this, we established a NAD<sup>+</sup>-intermediate specific reporter system to identify factors required for salvage of metabolically linked nicotinamide (NAM) and nicotinic acid (NA). Mutants lacking components of the NatB complex, *NAT3* and *MDM20*, appeared as hits in this screen. NatB is an N<sup>α</sup>-terminal acetyltransferase responsible for acetylation of the N terminus of specific Met-retained peptides. In NatB mutants, increased NA/NAM levels were concomitant with decreased NAD<sup>+</sup>. We identified the vacuolar pool of nicotinamide riboside (NR) as the source of this increased NA/NAM. This NR pool is increased by nitrogen starvation, suggesting NAD<sup>+</sup> and related metabolites may be trafficked to the vacuole for recycling. Supporting this, increased NA/NAM release in NatB mutants was abolished by deleting the autophagy protein *ATG14*. We next examined Tpm1 (tropomyosin), whose function is regulated by NatB-mediated acetylation, and Tpm1 overexpression (*TPM1-oe*) was shown to restore some NatB mutant defects. Interestingly, although *TPM1-oe* largely suppressed NA/NAM release in NatB mutants, it did not restore NAD<sup>+</sup> levels. We showed that decreased nicotinamide mononucleotide adenylyltransferase (Nma1/Nma2) levels probably caused the NAD<sup>+</sup> defects, and *NMA1-oe* was sufficient to restore NAD<sup>+</sup>. NatB-mediated N-terminal acetylation of Nma1 and Nma2 appears essential for maintaining NAD<sup>+</sup> levels. In summary, our results support a connection between NatB-mediated protein acetylation and NAD<sup>+</sup> homeostasis. Our findings may contribute to understanding the molecular basis and regulation of NAD<sup>+</sup> metabolism.

NAD<sup>+</sup> and its reduced form NADH are essential metabolites mediating redox reactions in cellular metabolism. NAD<sup>+</sup> is a substrate in protein modifications, such as sirtuin (Sir2 family proteins)-mediated protein deacetylation, and ADP-ribosylation by poly(ADP-ribose) polymerases. These protein modifi-

cations contribute to regulation of chromatin structure, DNA repair, circadian rhythm, metabolic responses, and life span (1–4). Aberrant NAD<sup>+</sup> metabolism is associated with a number of diseases, including diabetes, cancer, and neuron degeneration (2, 3, 5–8). Administration of NAD<sup>+</sup> precursors, such as nicotinamide mononucleotide (NMN),<sup>2</sup> nicotinamide (NAM), and nicotinamide riboside (NR), has been shown to ameliorate deficiencies related to aberrant NAD<sup>+</sup> metabolism in yeast, mouse, and human cells (3, 5–12). However, molecular mechanisms that underlie these beneficial effects remain unclear.

NAD<sup>+</sup> biosynthesis is highly conserved between yeast and vertebrates. In yeast, NAD<sup>+</sup> is synthesized *de novo* from tryptophan or salvaged from intermediates, such as NA, NAM, and NR (Fig. 1A). The NAD<sup>+</sup> levels maintained by these pathways converge at several different points and share cellular pools of ATP, phosphoribosyl pyrophosphate, and glutamine, while adding to total pools of ribose, AMP, phosphate, formate, alanine, glutamate, and others. NAD<sup>+</sup> is a NADP<sup>+</sup> precursor, which, like NAD<sup>+</sup>, is carefully balanced with its reduced form to maintain a favorable redox state. Some of these molecules contribute to other biosynthesis pathways or have signaling functions. Therefore, the cell must maintain these metabolites and their flux in a controlled manner. We do not fully understand all of the mechanisms by which the cell can sense and tune these metabolites, but some known NAD<sup>+</sup> homeostasis-regulatory mechanisms include transcriptional control, feedback inhibition, and enzyme or metabolite compartmentalization (1, 13–18). The complex and dynamic flexibility of NAD<sup>+</sup> precursors makes studying NAD<sup>+</sup> metabolism complicated. For example, NAM can both replenish NAD<sup>+</sup> pools and inhibit the activity of NAD<sup>+</sup>-consuming enzymes. In addition, little is known about the signaling pathways that regulate NAD<sup>+</sup> precursor homeostasis, in part due to the lack of sensitive and specific genetic screening systems to identify these NAD<sup>+</sup> homeostasis factors.

We have previously shown that NR-mediated NAD<sup>+</sup> synthesis plays important roles in the maintenance of NAD<sup>+</sup> pools and calorie restriction-induced life span (19). To further understand the regulation of NAD<sup>+</sup> homeostasis, we developed an NR-specific reporter-based bioassay to screen for mutants with

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This article contains Table S1.

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<sup>2</sup> The abbreviations used are: NMN, nicotinamide mononucleotide; NA, nicotinic acid; NAM, nicotinamide; QA, quinolinic acid; NR, nicotinamide riboside; NAT, N-terminal acetyltransferase; NMNAT, nicotinamide mononucleotide adenylyltransferase; rAPase, repressible acid phosphatase; oe, overexpression; EGFP, enhanced green fluorescent protein.

## NatB is a NAD<sup>+</sup> homeostasis factor

altered NR release. This screen system was based on our observations that yeast cells release and transport NR back into the cell (19), a phenomenon that is also conserved in human cells (20). Our studies have identified novel NAD<sup>+</sup> homeostasis factors, including the phosphate-responsive signaling (*PHO*) pathway (17), the SPS amino acid-sensing pathway (21), the conserved vacuolar NR transporter Fun26 (1, 17), and several NAD<sup>+</sup> metabolic enzymes (22).

In this study, we developed a reporter system targeting the NA/NAM branch of NAD<sup>+</sup> metabolism. The hypothesis is that cells defective in NA/NAM salvage (Fig. 1A) would show altered NA/NAM release. The *mdm20Δ* and *nat3Δ* mutants were among the top hits that showed increased NA/NAM release. Nat3 and Mdm20 are the catalytic and auxiliary subunits of NatB, respectively (23, 24). NatB belongs to a family of N-terminal acetyltransferases (NATs) that add an acetyl group from acetyl-CoA to the  $\alpha$ -amino group on the first amino acid of a protein. In yeast, there are four functional NATs, and each has distinct substrate specificity (23, 25, 26). About 20% of proteins are potential substrates of NatB in yeast and humans (27). N-terminal acetylation plays important roles in regulating protein stability, complex formation, and subcellular localization and has been implicated in various cellular processes and human diseases (27). Our studies are the first to link NatB to NAD<sup>+</sup> homeostasis. Here, we characterized the NatB mutants as well as the NatB downstream targets to provide a molecular basis underlying NatB-mediated regulation of NAD<sup>+</sup> homeostasis.

## Results

### A cell-based reporter assay to study NA/NAM homeostasis factors

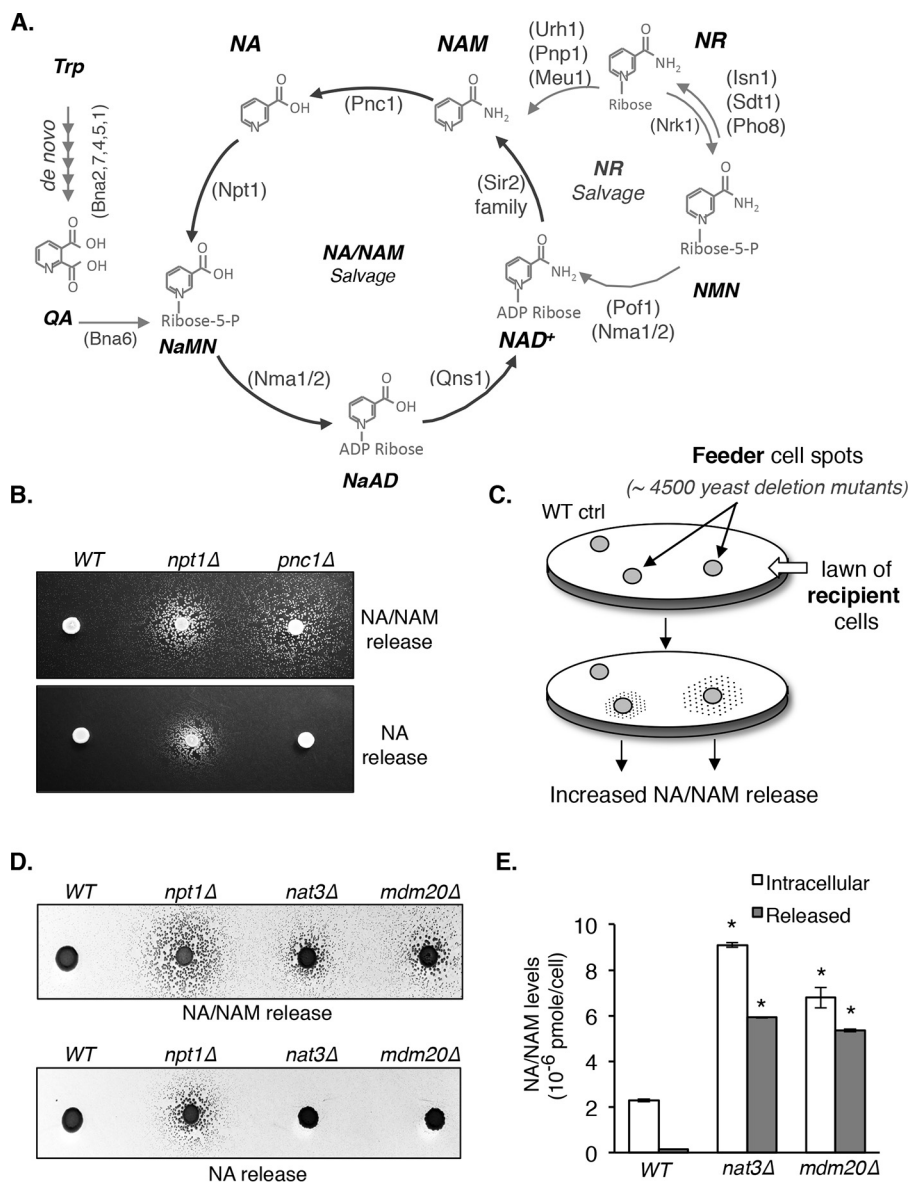
We first determined whether cells indeed release more NA and/or NAM when NA/NAM salvage is blocked. A cross-feeding assay is established using *bnaf6Δnrk1Δnrt1Δ* mutants as “recipient cells” (depend on NA or NAM for growth) and mutants of interest as “feeder cells.” In this system, recipient cells cannot grow on niacin-free SD (NA/NAM-free), but when feeder cells are placed in proximity, feeder cell–released NA or NAM supports recipient cell growth by “cross-feeding.” As a result, this assay determines relative levels of total NA and NAM released by feeder cells. As shown in Fig. 1B (top), recipient cells were spread onto niacin-free SD plates as a lawn. Next, *npt1Δ* (blocks NA salvage) and *pnc1Δ* (blocks NAM salvage) feeder cells were spotted on top of the lawn and allowed to grow. Recipient cells near the *npt1Δ* and *pnc1Δ* feeder spots appeared as satellite colonies after 3 days, and the number and size of the satellite colonies positively correlate with the amount of NA/NAM released from the feeders. To separate NAM release from NA/NAM release, *PNC1* was deleted in the recipient cell strain to block its ability to utilize NAM. Based on the biochemical function of Npt1 and Pnc1 (Fig. 1A), we expect that *npt1Δ* cells release more NA (and perhaps NAM) and *pnc1Δ* cells release more NAM. Indeed, as shown in Fig. 1B (bottom), *pnc1Δ* feeder cells failed to support the growth of the new recipient strain (*bnaf6Δnrk1Δnrt1Δpnc1Δ*, which cannot utilize NAM). In contrast, *npt1Δ* cells still supported the growth of *bnaf6Δnrk1Δnrt1Δpnc1Δ*. These results confirmed that mutants with altered NA/NAM homeostasis could be identified using this system.

### Components of the NatB complex are novel NA/NAM homeostasis factors

Next, we used the haploid yeast deletion collection as feeder cells to identify mutants with increased NA/NAM release (Fig. 1C). Mutants were assigned a score of 2–5 (WT = 1). A higher score represents more NA/NAM release. To eliminate false-positives, mutants with a score of  $\geq 3$  (81 mutants) were reexamined. A total of  $\sim 70$  mutants passed the secondary screen (Table S1). Known NAD<sup>+</sup> homeostasis factors were identified, including *npt1Δ* and *pnc1Δ*. The *mdm20Δ* and *nat3Δ* mutants were among the top hits. Nat3 and Mdm20 are subunits of NatB, a member of the NATs that adds an acetyl group from acetyl-CoA to the  $\alpha$ -amino group on the first amino acid of a protein (23, 24). Deletion of either NatB subunit results in stress sensitivity growth defects (23, 24, 28, 29). To verify that observed phenotypes are due to the featured mutations and not to secondary cryptic mutations in the deletion collection, we reconstructed all deletion mutants used in this study. We confirmed both *nat3Δ* and *mdm20Δ* indeed released more NA/NAM (Fig. 1D, top). Similar to *pnc1Δ* mutant (Fig. 1B), both *nat3Δ* and *mdm20Δ* released mostly NAM, not NA (Fig. 1D, bottom). Fig. 1E showed that increased NA/NAM release in *nat3Δ* and *mdm20Δ* cells correlates with increased intracellular NA/NAM levels determined by a quantitative liquid assay.

### NR metabolism contributes to increased NAM release in NatB mutants

To understand how NatB complex affects NAM/NAD<sup>+</sup> homeostasis, we first tested whether Pnc1 level or activity was reduced in NatB mutants because Pnc1 is a major NAM metabolic factor (nicotinamidase), and the *pnc1Δ* mutant showed strong NAM release (Fig. 1B). Although Pnc1 is not a predicted NatB substrate, Pnc1 may be indirectly regulated by NatB. Our results showed that Pnc1 activity (Fig. 2B) and protein expression (Fig. 2A) were only slightly decreased in *nat3Δ* cells. Interestingly, although *pnc1Δ* cells show WT level of NAD<sup>+</sup> under normal (NA-rich) growth conditions (19), the levels of NAD<sup>+</sup> were significantly reduced in NatB mutants (Fig. 2C). These studies suggest that additional factors probably contribute to the altered NAM/NAD<sup>+</sup> homeostasis observed in NatB mutants. Increased production of NR has been associated with increased activity of the Pi-sensing *PHO* signaling pathway (17). We therefore determined whether NatB mutants showed altered *PHO* activity by measuring the activity of *rATPase* (Pho5, a periplasmic phosphatase activated by *PHO* signaling). As shown in Fig. 2D (left), *PHO* activity was moderately increased in the *nat3Δ* mutant. As controls, deleting *PHO5* largely blocked the increase of *rATPase* activity. The residual *rATPase* activity observed in *nat3Δpho5Δ* cells probably came from Pho5-like phosphatases Pho11 and Pho12. Interestingly, observed *PHO* activation in *nat3Δ* cells appeared independent of the conventional *PHO* transcription factor Pho4-Pho2 complex, because deleting *PHO4* did not significantly reduce the *PHO* activation in *nat3Δ* cells (Fig. 2D, left). Consistent with this result, deleting *PHO4* did not prevent NA/NAM release in *nat3Δ* cells (Fig. 2D, right panel). Next, we determined whether and how NR contributes to increased NAM in NatB mutants.

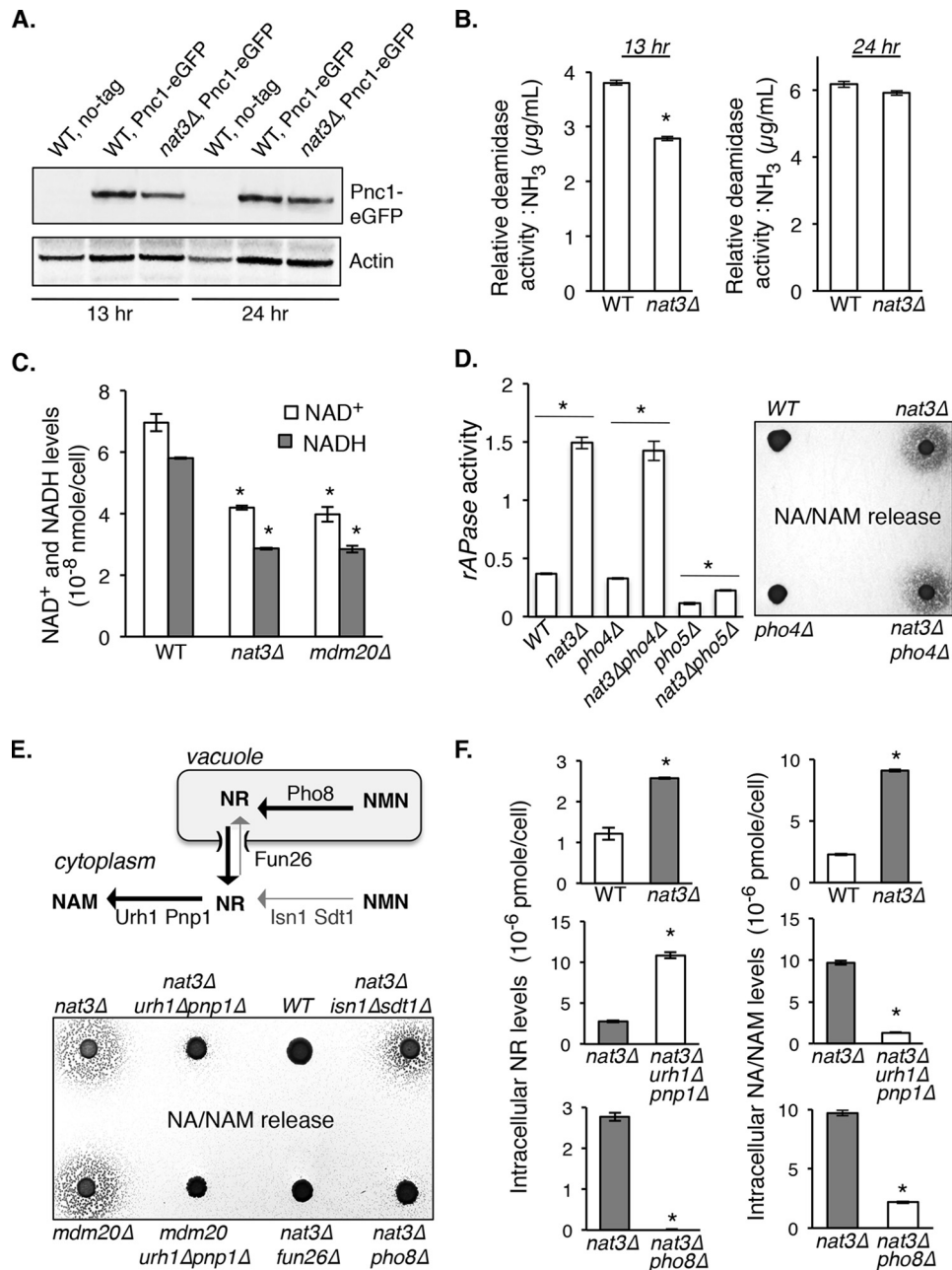


**Figure 1. Identification of yeast mutants with altered NA and NAM homeostasis.** *A*, a simplified model of the *Saccharomyces cerevisiae* NAD<sup>+</sup> synthesis pathway. NAD<sup>+</sup> can be synthesized *de novo* from Trp and by salvaging NAD<sup>+</sup> intermediates through the NA/NAM and NR cycles. *NaMN*, nicotinic acid mononucleotide. *NaAD*, deamido NAD<sup>+</sup>. Abbreviations of protein names are shown in parentheses. *B*, deletions of *NPT1* and *PNC1* confer increased NA/NAM and NAM release, respectively. NA/NAM-dependent recipient cells were plated onto niacin-free SD plates as a lawn. Next, *WT*, *npt1Δ*, and *pnc1Δ* feeder cells were spotted on top of the lawn and allowed to grow for 3–4 days at 30 °C. Recipient cells on the top panel (*bna6Δnrk1Δnrt1Δ*) can utilize both NA and NAM, whereas recipient cells on the bottom panel (*bna6Δnrk1Δnrt1Δpnc1Δ*) can only utilize NA. The number and size of the satellite colonies surrounding the feeder spots positively correlate with the amount of NA/NAM released from the feeders. *C*, overview of the genetic screen to identify mutants with increased NA/NAM release. Haploid single deletion mutants (feeder cells) were spotted onto a lawn of NA/NAM-dependent *bna6Δnrk1Δnrt1Δ* recipient cells, whose growth is dependent on NA/NAM released from the feeder cells in a dose-dependent manner. *D*, deletions of *NAT3* and *MDM20* confer increased NAM release. Various feeder strains were spotted onto a lawn of *bna6Δnrk1Δnrt1Δ* (top) and *bna6Δnrk1Δnrt1Δpnc1Δ* (bottom) recipient cells. For clarity, inverse images are shown. *E*, the *nat3Δ* and *mdm20Δ* mutants show significant increases in both released and intracellular NA/NAM levels. NA/NAM levels were determined in both growth media (released) and cell extracts (intracellular). Data shown are representative of three independent experiments. Feeding of collected lysate to recipient cells was conducted in triplicate. Error bars, S.D. The *p* values are calculated using Student's *t* test (\*, *p* < 0.05).

Intracellular NAM may come from NR via the reactions of cytoplasmic nucleosidases *Urh1* and *Pnp1* (Fig. 2E, top) (30). As shown in Fig. 2E (bottom), deleting *URH1* and *PNP1* largely diminished NA/NAM release in *NatB* mutants. Moreover, deleting the vacuolar phosphatase *PHO8* (Fig. 2E, top) (17) diminished NA/NAM release (Fig. 2E, bottom), whereas deleting the cytoplasmic nucleotidases *ISN1* and *SDT1* (Fig. 2E, top) had a lesser effect (Fig. 2E, bottom). These results suggest that increased NAM in *NatB* mutants probably results from vacuo-

lar NR, which is converted to NAM by cytoplasmic nucleosidases *Urh1* and *Pnp1*. Vacuolar NR may exit the vacuole through the *Fun26* transporter (human lysosomal hENT homolog) into the cytoplasm (17), where it is converted to NAM by *Urh1* and *Pnp1* (Fig. 2E, top). Supporting this, deleting *FUN26* indeed blocked NAM release in the *NatB* mutant (Fig. 2E, bottom). Moreover, *nat3Δ* cells carrying deletions of *URH1* and *PNP1* (which converts NR to NAM) showed increased intracellular NR levels (Fig. 2F, middle left) and reduced NA/NAM

## NatB is a NAD<sup>+</sup> homeostasis factor

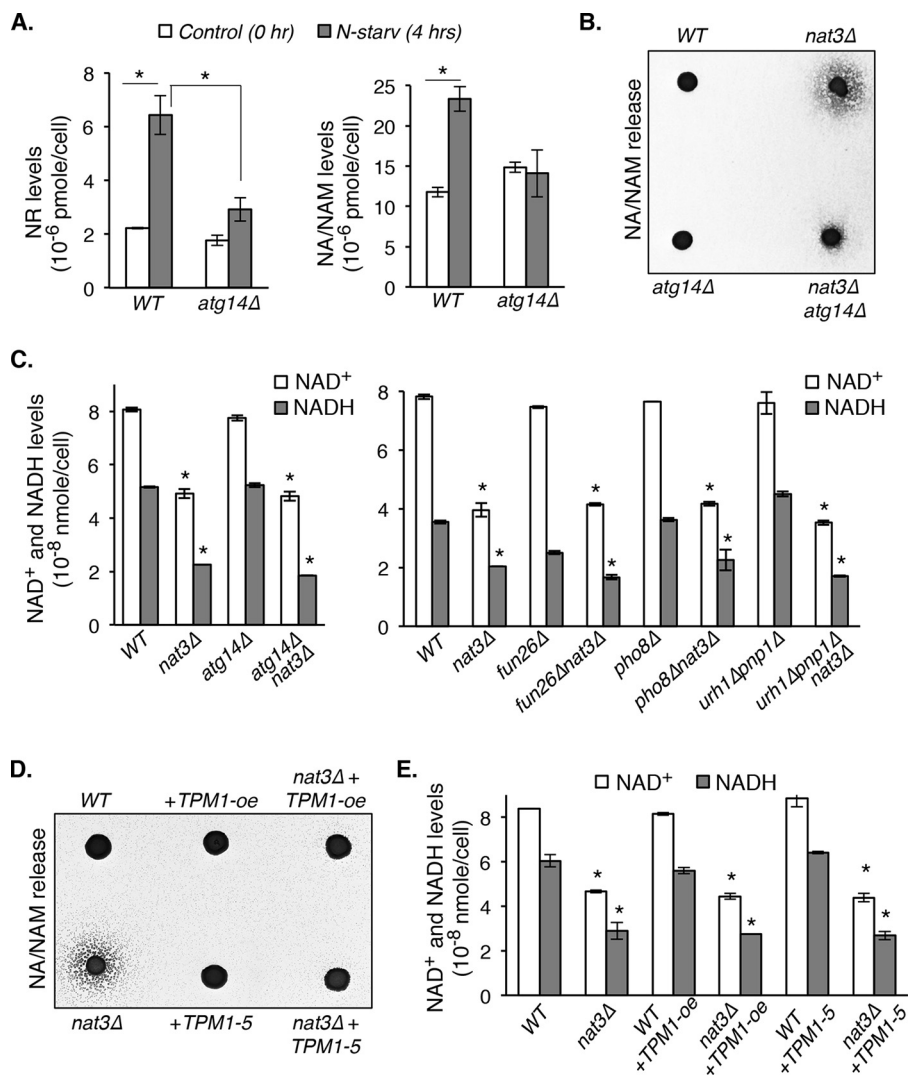


**Figure 2. Characterizations of NAD<sup>+</sup> homeostasis factors in NatB mutants.** *A*, Pnc1 protein levels in WT and *nat3Δ* cells. Pnc1 level is slightly decreased in *nat3Δ* mutant during late log phase (13 h). *B*, relative Pnc1 deamidase activities determined in WT and *nat3Δ* cell lysates. Pnc1 activity is slightly decreased in *nat3Δ* cells during late log phase (13 h). *C*, measurements of intracellular NAD<sup>+</sup> levels. Deleting *NAT3* significantly decreases NAD<sup>+</sup> levels. *D*, relative repressible acid phosphatase (*rAPase*) activities in WT and *nat3Δ* cells. Increased *rAPase* activity in *nat3Δ* cells indicates increased *PHO* signaling activity (left). Deleting *PHO4* does not abolish *rAPase* activity, whereas deleting *PHO5* largely diminishes the *rAPase* activity (left). Deleting *PHO4* does not block NA/NAM release in *nat3Δ* cells (right). Data shown are representative of three independent experiments. *E*, NR salvage contributes to increased NA/NAM production in NatB mutants. Intracellular compartmentalization of NR salvage factors (top). Deleting cytosolic nucleosidases *URH1* and *PNP1* (which convert NR to NAM) diminishes NA/NAM release in *nat3Δ* and *mdm20Δ* cells (bottom). Deleting vacuolar NR-producing factor *Pho8* also diminishes NA/NAM release, whereas deleting the cytosolic NR-producing factors *Isn1* and *Sdt1* has a lesser effect (bottom). Deleting vacuolar transporter *FUN26* also abolishes NA/NAM release in *nat3Δ* cells. This suggests that increased NAM in NatB mutants comes from vacuolar NR, which is converted to NAM by *Urh1* and *Pnp1* (top). *F*, intracellular NR (top left) and NA/NAM (top right) levels are increased in *nat3Δ* mutant. Deleting *URH1* and *PNP1* diminishes NA/NAM production (middle right) with a commitment increase of intracellular NR level (middle left). Deleting *PHO8* diminishes both NR and NA/NAM levels (bottom panels). For *B*, *C*, *D*, and *F*, the error bars denote S.D. values of triplicated samples. The *p* values are calculated using Student's *t* test (\*, *p* < 0.05).

levels (Fig. 2*F*, middle right). Deleting *PHO8* (which produces NR in the vacuole) blocked the increase of both NR and NA/NAM in *nat3Δ* cells (Fig. 2*F*, bottom panels). All of these results support vacuolar NR as the main source of increased NA/NAM in *nat3Δ* cells.

### Blocking NA/NAM production and release is not sufficient to restore NAD<sup>+</sup> levels in the NatB mutants

So far, our results suggest that in NatB mutants, increased NA/NAM release is probably due to increased NAM production, which originates from vacuolar NR. It remains unclear

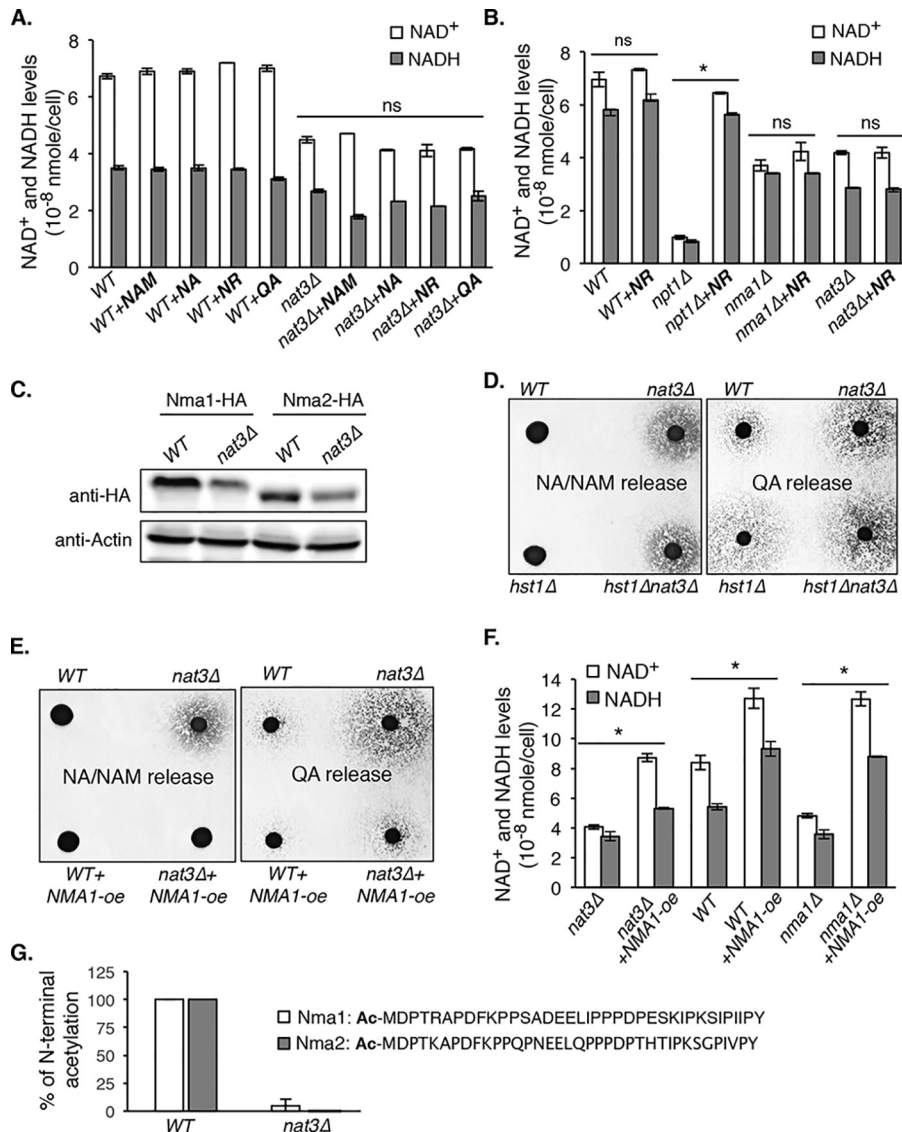


**Figure 3. Blocking NA/NAM production or release is not sufficient to restore NAD<sup>+</sup> levels in the NatB mutants.** A, nitrogen starvation (*N-starv*) increases NR (left) and NA/NAM (right) production, which are diminished by deleting *ATG14*, an autophagy-specific phosphatidylinositol 3-kinase subunit. B, increased NA/NAM levels in *nat3Δ* cells are diminished by deleting *ATG14*. C, blocking NA/NAM production by deleting *ATG14*, *FUN26*, *PHO8*, *URH1*, and *PNP1* is not sufficient to restore NAD<sup>+</sup> levels in *nat3Δ* cells. D, *TPM1* overexpression (*TPM1-oe*) and expression of a gain-of-function allele of *TPM1* (*TPM1-5*) diminish NA/NAM release in *nat3Δ* cells. E, *TPM1-oe* and expression of *TPM1-5* are not sufficient to restore NAD<sup>+</sup> levels in *nat3Δ* cells. Data shown are representative of three independent experiments. For A, C, and E, the error bars denote S.D. values of triplicated samples. The *p* values are calculated using Student's *t* test (\*, *p* < 0.05).

how NAD<sup>+</sup> and its intermediates are transported into the vacuole for turnover and storage. Because most cellular NAD<sup>+</sup> and NADH are bound to proteins, NAD<sup>+</sup> intermediates may enter the vacuole via vesicular trafficking and/or autophagy. Because autophagy is a critical regulator of organelle homeostasis and recycling of macromolecules (31, 32), we therefore asked whether inducing autophagy in yeast cells would indeed increase intracellular NR and NA/NAM levels. As shown in Fig. 3A, both NR (left) and NA/NAM (right) levels were increased by nitrogen starvation, a condition known to induce autophagy (31, 33, 34). These increases were largely dependent on *Atg14*, a key regulator of autophagic membrane tethering and fusion (31). These data suggest that autophagy may contribute to increased NA/NAM in NatB mutants. Supporting this, we showed that deleting *ATG14* largely blocked NA/NAM release in the *nat3Δ* mutant (Fig. 3B). Next, we tested whether deleting *ATG14* was sufficient to restore NAD<sup>+</sup> levels in the NatB

mutants. As shown in Fig. 3C (left), deleting *ATG14* did not rescue the low NAD<sup>+</sup> levels in *nat3Δ* cells. Similarly, deleting other factors contributing to NR and NAM production (Figs. 1A and 2E) in *nat3Δ* cells also failed to restore the NAD<sup>+</sup> level back to wildtype (Fig. 3C, right). Most reported NatB mutant phenotypes were attributed to defective N-terminal acetylation of tropomyosins (Tpm1 and Tpm2 in yeast) (23, 28, 29). Tropomyosins are actin filament-binding proteins that stabilize actin cables (29, 35–38). Overexpression of *TPM1* (*TPM1-oe*) and expression of a gain-of-function allele of *TPM1* (*TPM1-5*) were shown to restore actin cable formation in NatB mutants (23, 24, 28, 29). Interestingly, both *TPM1-oe* and *TPM1-5* largely blocked NA/NAM release in the *nat3Δ* mutant (Fig. 3D), suggesting a role for the actin cytoskeleton in NAM homeostasis regulation. However, *TPM1-oe* and *TPM1-5* still failed to restore the NAD<sup>+</sup> levels in *nat3Δ* cells (Fig. 3E). These studies indicate that the low NAD<sup>+</sup> phenotype of *nat3Δ* cells cannot

## NatB is a NAD<sup>+</sup> homeostasis factor



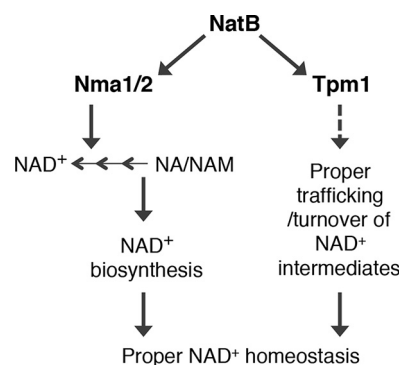
**Figure 4. NMNATs are responsible for the NAD<sup>+</sup> decrease in NatB mutants.** A, supplementing NAD<sup>+</sup> precursors (NA, NAM, NR, and QA) does not restore NAD<sup>+</sup> levels in the *nat3Δ* mutants. B, *nat3Δ* and *nma1Δ* mutants share similar low NAD<sup>+</sup> levels, which cannot be rescued by NR supplementation. As a control, NR is able to rescue the low NAD<sup>+</sup> level in *npt1Δ* cells. C, the levels of NMNATs proteins (Nma1 and Nma2) are decreased in *nat3Δ* cells. D, both *nat3Δ* and *hst1Δ* cells show increased QA release, which is indicative of increased *de novo* activities (right). Unlike *nat3Δ* cells, *hst1Δ* cells do not show increased NA/NAM release (left). These findings suggest that increased NA/NAM in *nat3Δ* cells is not simply due to increased *de novo* activity. The *npt1Δnrk1Δbna4Δ* mutant was used as the recipient cells for determining QA release (QA-dependent growth). E, *NMA1-oe* diminishes both NA/NAM and QA release in *nat3Δ* cells. F, *NMA1-oe* restores the NAD<sup>+</sup> levels in *nat3Δ* cells. Data shown are representative of three independent experiments. G, among specific Nma1 and Nma2 N-terminal peptides identified from WT cells, 100% are acetylated at the first methionine. Acetylations are mostly absent among specific Nma1 and Nma2 N-terminal peptides identified in *nat3Δ* mutant, indicating that Nma1 and Nma2 are N-terminally acetylated in a NAT3 (NatB)-dependent manner. For A, B, F, and G, the error bars denote S.D. values of duplicated samples. The *p* values are calculated using Student's *t* test (\*, *p* < 0.05).

be rescued simply by decreasing NAM production and/or release and that additional factors are probably involved.

### Nma1 and Nma2 are responsible for the low NAD<sup>+</sup> levels in NatB mutants

To gain further insight, we tested whether supplementing particular NAD<sup>+</sup> precursors to *nat3Δ* cells can restore the NAD<sup>+</sup> levels. Interestingly, all NAD<sup>+</sup> precursors tested (NA, NAM, NR, and QA) failed to restore NAD<sup>+</sup> levels (Fig. 4A). This suggests that *nat3Δ* cells may be defective in a step of NAD<sup>+</sup> biosynthesis that is common to the utilization of QA, NA, and NR. Nma1 and Nma2 are dual functional nicotinamide

mononucleotide adenylyltransferases (NMNATs), which utilize both NMN and nicotinic acid mononucleotide (*NaMN*) as substrates and are required for all three NAD<sup>+</sup> biosynthesis pathways (Fig. 1A). N-terminal acetyltransferases have specific targets that are largely determined by the sequence of the first two amino acids. NatB acetylates proteins with a Met-retained residue, followed by Asp, Glu, Gln, or Asn as the second residue (25, 26, 39, 40). Nma1 and Nma2 have a Met-Asp N terminus, but neither protein has been identified as a NatB target. Similar to *nat3Δ* mutant, *nma1Δ* cells showed decreased NAD<sup>+</sup> levels, which could not be rescued by supplementing NR (Fig. 4B). As



**Figure 5. A proposed model depicting factors contributing to altered NAD<sup>+</sup> homeostasis in NatB mutants.** We identified Nma1 and Nma2 as the acetylation targets of NatB. NatB regulates NAD<sup>+</sup> biosynthesis and NAD<sup>+</sup> intermediate trafficking via Nma1/Nma2 and Tpm1, respectively. Dashed line, the mechanism of this pathway remains largely unclear.

## Discussion

In this study, we characterized the NatB complex as a novel NAD<sup>+</sup> homeostasis factor. Mutants lacking components of the NatB complex, *NAT3* and *MDM20*, produce and release an excess amount of NA and NAM. Our studies showed that two pathways downstream of NatB contribute to NAD<sup>+</sup> homeostasis (Fig. 5). In one, NatB is important for proper NAD<sup>+</sup> biosynthesis by regulating Nma1/Nma2. NatB mutants have low NAD<sup>+</sup> levels (Fig. 2C), and all NAD<sup>+</sup> precursors examined failed to restore the NAD<sup>+</sup> levels (Fig. 4, A and B). This suggests that a NAD<sup>+</sup> biosynthesis factor(s) required for utilization of all NAD<sup>+</sup> precursors is defected in NatB mutants. Nma1 and Nma2 are such targets because they are the only factors required for all three NAD<sup>+</sup> biosynthesis pathways (*de novo*, NA/NAM salvage, and NR salvage) (Fig. 1A), and the N-terminal amino acid sequences are a match for NatB acetylation. In addition, *nma1Δ* and *nat3Δ* mutants showed similar NAD<sup>+</sup> utilization defects (Fig. 4B). Although Nma2 is present in both mutants, Nma2 is known to play a minor role in NAD<sup>+</sup> metabolism. Supporting this model, decreased Nma1 protein level was observed in the *nat3Δ* mutant (Fig. 4C), and overexpressing *NMA1* was sufficient to restore the NAD<sup>+</sup> levels (Fig. 4F). Our studies showed that Nma1 and Nma2 are indeed acetylated at the N terminus in a NatB-dependent manner (Fig. 4G). Further studies are required to understand the mechanisms of NatB-mediated regulation of Nma1/Nma2, but the decreased level of both proteins in *nat3Δ* suggests that the nonacetylated counterparts may be degraded at a higher rate by the N-end rule pathway. The N-end rule pathway recognizes N-terminal degradation signals and marks proteins for degradation by specific E3 ligases. This pathway can be broken up into three branches called the Arg/N-rule pathway, the Pro/N-rule pathway, and the Ac/N-rule pathway (42, 43). The Ac/N-rule branch recognizes N-acetylated proteins. However, in the absence of acetylation, the proteins may be stabilized or targeted by other N-end rule branches, which can lead to different rates of degradation (42, 44, 45). On the other hand, N-acetylated proteins may also bury their acetylated N termini within physically associated proteins (46–48). This provides two important functions. First, it shields the N terminus from Ac/N-rule E3 ligases. Second, it stabilizes the protein complex. The absence of a nor-

for the controls, NR efficiently restored the NAD<sup>+</sup> level in the *npt1Δ* mutant, which has functional NR salvage (Fig. 4B). The similarities between *nma1Δ* and *nat3Δ* mutants suggested that Nma1 and Nma2 activities are probably reduced in *nat3Δ* cells. Because N-terminal acetylation may affect the turnover of target proteins, we first determined whether reduced Nma1/Nma2 activities were due to decreased Nma1/Nma2 protein levels. As shown in Fig. 4C, Nma1 and Nma2 protein levels were indeed decreased in *nat3Δ* cells (Fig. 4C). In addition to Nma1 and Nma2, NAD<sup>+</sup> homeostasis factors Bna2, Bna5, and Hst1 are also potential NatB targets. Bna2 and Bna5 are biosynthesis enzymes in the *de novo* pathway (Fig. 1A), and Hst1 represses the expression of *BNA* genes (13). Because blocking the *de novo* pathway does not affect NAD<sup>+</sup> levels under standard growth conditions (NA-rich media), the low NAD<sup>+</sup> phenotype of *nat3Δ* mutant is unlikely to be due to reduced *de novo* activities. To determine *de novo* activities in *nat3Δ* cells, we developed a cell-based readout for QA production. In this assay, more QA release indicates increased *de novo* activities, and vice versa. As expected, *hst1Δ* released more QA (Fig. 4D, right) due to derepression of *BNA* genes, and deletions of *BNA* genes completely blocked QA release (data not shown). Similar to *hst1Δ* mutant, *nat3Δ* mutant also released more QA, and double deletions of *NAT3* and *HST1* did not further increase QA release (Fig. 4D). In comparison, *hst1Δ* cells did not release more NA/NAM (Fig. 4D, left), indicating that increased NA/NAM in the *nat3Δ* mutant is not simply due to increased *de novo* activities. This is in line with our findings that in *nat3Δ* cells, increased NA/NAM arises from NR (Fig. 2E). Overall, these results demonstrate that the low NAD<sup>+</sup> phenotype of *nat3Δ* cells is mainly due to reduced protein levels of Nma1 and Nma2. To confirm this, we examined whether overexpression of *NMA1* is sufficient to reduce NA/NAM release. As shown in Fig. 4E (left), *NMA1-oe* completely blocked NA/NAM release in *nat3Δ* cells. Moreover, *NMA1-oe* also reduced QA release in *nat3Δ* cells (Fig. 4E, right). The reduction in QA and NA/NAM release is probably due to more efficient NA/NAM and QA assimilation by *NMA1-oe*. Supporting this, *NMA1-oe* was sufficient to restore the NAD<sup>+</sup> levels in *nat3Δ* cells back to WT levels (Fig. 4F). Interestingly, *NMA1-oe* also increased NAD<sup>+</sup> levels in WT cells. Next, we examined whether Nma1 and Nma2 are N-terminal acetylated by NatB. Immunoprecipitated Nma1-HA protein complexes obtained from WT and *nat3Δ* cells were treated with chymotrypsin and then analyzed by tandem mass spectrometry. Both Nma1 and Nma2 N-terminal peptides were identified in WT and *nat3Δ* cells. As shown in Fig. 4G, among identified specific Nma1 and Nma2 N-terminal peptides from WT cells, all of them (66 for Nma1, 25 for Nma2) were acetylated on the first methionine. Conversely, of the identified Nma1 and Nma2 N-terminal peptides from *nat3Δ* cells (20 for Nma1, 10 for Nma2), only one Nma1 peptide showed acetylation (Fig. 4G). These results indicate that Nma1 and Nma2 are indeed N-terminally acetylated in a NatB-dependent manner.



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mally present acetylated N terminus can drastically decrease the affinity of a protein to complex with its partners (24, 47), leaving it more dissociated and prone to targeted degradation (42). Therefore, it is possible that in addition to protein stabilization, N-terminal acetylation may also be important for Nma1 and Nma2 protein-protein interactions. Current research suggests that Nma1 and Nma2 may function as tetramers (49–51).

In the other pathway, NatB- and Tpm1-mediated proper vesicular trafficking appears to specifically affect NA/NAM homeostasis (Fig. 5). *TPM1* encodes one of two tropomyosin proteins in yeast, which bind to actin filaments and stabilize actin cables. Acetylation of *TPM1* by NatB greatly improves its affinity for actin filaments (24). During cell division and growth, actin cables provide tracks for organelle segregation and endocytic trafficking (35, 38, 52, 53). Interestingly, although overexpression of *TPM1* and expression of a dominant *TPM1-5* completely blocked NA/NAM release (Fig. 3D), they failed to restore the NAD<sup>+</sup> levels in the NatB mutants. This is in line with the model that Nma1 and Nma2 abnormalities are responsible for observed NAD<sup>+</sup> defects in NatB mutants. *NMA1* overexpression, however, was sufficient to block NA/NAM release (Fig. 4E) and restore NAD<sup>+</sup> level in NatB mutants (Fig. 4F). These results show that the two NatB-controlled NAD<sup>+</sup> homeostasis pathways are interconnected. How may *TPM1* and actin cytoskeleton affect NAD<sup>+</sup> homeostasis? It is possible that NAD<sup>+</sup> and related precursors (free and protein-bound) are trafficked by means of vesicles, which would require Tpm1 and actin cables to mediate their directed movement (35, 37). Deleting *TPM1* alone abolishes the appearance of actin cables, whereas deleting *TPM2* does not (36). Neither *tpm1Δ* nor *tpm2Δ* single mutant (*tpm1Δtpm2Δ* double mutant is lethal) appeared in our screen for high NA/NAM release mutants. It is probably because Tpm1 and Tpm2 have some functional redundancy (36, 54), and they can substitute for each other when one is absent. During asymmetrical cell division, targeted and increased vesicular transport to daughter may concomitantly release NA/NAM and other NAD<sup>+</sup> intermediates. It is possible that unacetylated Tpm1 fails to efficiently assist the formation and delivery of endocytic vesicles, leading to increased NA/NAM release observed in the NatB mutants.

Our studies showed that increased NA/NAM in NatB mutants came from the NR pool (Fig. 2E). Increased NA/NAM release in NatB mutants was abolished by deleting cytosolic nucleosidases (*URH1/PNP1*), which convert NR to NAM. Increased NR mostly originates from the vacuolar pool because deleting vacuolar NR-producing phosphatase (*PHO8*) abolished NA/NAM increase, whereas deleting cytosolic NR-producing nucleotidases (*ISN1/SDT1*) had a lesser effect (Fig. 2E, bottom). Vacuolar NR may exit the vacuole through Fun26 transporter (human lysosomal hENT homolog) and enter the cytoplasm (17), where it is converted to NAM by nucleosidases. Indeed, deleting *FUN26* blocked NAM release in the NatB mutant (Fig. 2E, bottom). It remains unclear how NAD<sup>+</sup> and its intermediates are transported into the vacuole. Our studies suggest that autophagy may also play a role. Autophagy is a critical regulator of organelle homeostasis and recycling of macromolecules (55). It has been reported that NatB mutants failed to properly segregate mitochondria during cell division

(mitochondrial inheritance defects) (23, 24, 28), which may lead to increased mitochondrial turnover by autophagy. Supporting this, many mutants we identified in our screen affect mitochondrial function (Table S1). In addition, we showed that intracellular levels of both NA/NAM and NR were increased by starvation conditions known to induce autophagy (Fig. 3A). Finally, deleting *ATG14*, a key regulator of autophagic membrane tethering and fusion (56), blocked NAM release in the *nat3Δ* mutant (Fig. 3B). Although *TPM1-oe* shared similar phenotypes with *ATG14* deletion, it is unclear whether Tpm1-mediated vesicular trafficking and autophagy function in the same pathway to regulate NA/NAM homeostasis. Further evidence is required to detail the functional link between these factors.

It remains unclear why NatB mutants release more NA/NAM and not NR. Our studies show that the NR pools in NatB mutants are quickly converted to NAM by cytoplasmic Urh1 and Pnp1 (Fig. 2E). In line with this, the NatB mutants were previously identified in a different screen as “low-NR” mutants (17). Although Urh1 and Pnp1 are not predicted direct targets of NatB (23, 25, 26), their activities and localization could be indirectly affected by NatB mutations. Some NatB mutant phenotypes may also be due to aberrant energy homeostasis (57). Supporting this, the Glc7 phosphatase, which inhibits Snf1 (yeast AMPK) activity, is acetylated by NatB, and Snf1 activation was observed in *nat3Δ* cells (57). In addition, Snf1 activation was associated with Pnp1 activity in starvation-induced ribonucleotide salvage (58). Another NAM metabolic factor that may contribute to increased NA/NAM release is Pnc1. Although Pnc1 levels and activities are only slightly decreased in NatB mutants (Fig. 2, A and B), its localization may be altered in NatB mutants. Pnc1 has been shown to localize in peroxisomes in normal conditions. In response to nutrient starvation and stress, Pnc1 expression is induced and becomes ubiquitous (15, 59). In addition, Pnc1 has been shown to be released extracellularly (60), and it is most likely retained in the periplasmic space (61). It is possible that during asymmetrical cell division, increased vesicular transport to daughter may concomitantly increase NAM release. Pnc1 may be released to convert NAM to NA to support NAD<sup>+</sup> biosynthesis and/or eliminate NAM toxicity (62–64). In mammals, eNAMPT is also released extracellularly to help metabolize NAM (65); however, yeast does not have NAMPT.

In summary, our studies have uncovered novel NAD<sup>+</sup> homeostasis factors. It would be informative to further investigate the regulation of NAD<sup>+</sup> homeostasis by NatB in future studies. These studies may also provide insights into the regulation of NAD<sup>+</sup> metabolism and the molecular basis of disorders associated with aberrant NAD<sup>+</sup> metabolism.

## Experimental procedures

### Yeast strains, growth media, and plasmids

Yeast strain BY4742 *MATα his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0* acquired from Open Biosystems (66) was used for this study. Standard media, such as yeast extract-peptone-dextrose (YPD), synthetic minimal (SD), and synthetic complete (SC) media, were made as described (67). Niacin-free SD and SC were made by using niacin-free yeast nitrogen base (Sunrise Science Prod-

ucts). Niacin (vitamin B3) comprises both NA and NAM. All gene deletions were generated by replacing wildtype genes with reusable loxP-*Kan<sup>r</sup>*-loxP cassettes as described (68). Multiple deletions were carried out by removing the *Kan<sup>r</sup>* marker using a galactose-inducible Cre recombinase (68). *PNC1*, *NMA1*, and *NMA2* were tagged by the HA or EGFP epitope tag in the genome using the pFA6a-3HA-HIS3MX or pFA6a-link-yoEGFP-CaURA3 plasmid as described previously (69, 70). The *NMA1* and *TPM1* overexpression constructs, pADH1-*NMA1* and pADH1-*TPM1*, were made in the integrative pPP81 (*LEU2*) vector as described (71). The resulting constructs were verified by DNA sequencing. *TPM1-5* in pRS316 was a gift from Dr. B. Plevoda. *TPM1-5* encodes an extended 7 amino acids at the N terminus of *TPM1* (23).

### NA/NAM cross-feeding spot assays

The *bnA6Δnrk1Δnrt1Δ* mutant (which utilizes both NA and NAM) and *bnA6Δnrk1Δnrt1Δpnc1Δ* mutant (which only utilizes NA) were used as “recipient cells,” and yeast strains of interest were used as “feeder cells.” First, recipient cells were plated onto niacin-free SD (NA/NAM-free) as a lawn (~10<sup>4</sup> cells/cm<sup>2</sup>). Next, 2 μl of each feeder cell strain (~10<sup>4</sup> cells; cell suspension was made in sterile water at A<sub>600</sub> of 2) was spotted onto the lawn of recipient cells. Plates were then incubated at 30 °C for 3 days. The extent of the recipient cell growth indicates the levels of NA/NAM released by feeder cells.

### Genetic screen using the yeast deletion collection

The haploid yeast deletion collection (~4500 strains) established in the BY4742 background was acquired from Open Biosystems (72). To screen for mutants with increased NA/NAM release, 2 μl of each strain was directly taken from the frozen stock and then spotted onto niacin-free SD plates spread with the *bnA6Δnrk1Δnrt1Δ* recipient cells at a density of ~10<sup>4</sup> cells/cm<sup>2</sup>. After incubation at 30 °C for 3 days, we scored the cross-feeding activity (which indicates the level of NA/NAM release) of each mutant by comparing the diameter of the cross-feeding zones with that of the wildtype. Mutants were assigned a score of 2–5 (WT = 1). A higher score represents more NA/NAM release. To eliminate false-positives, mutants with a score of ≥3 (81 mutants) were reexamined. Cells from the frozen stock were first recovered in fresh media for 2 days. 10<sup>4</sup> cells of each strain were then spotted and scored as described above. Both the *bnA6Δnrk1Δnrt1Δ* mutant (which utilizes both NA and NAM) and *bnA6Δnrk1Δnrt1Δpnc1Δ* mutant (which only utilizes NA) were used as “recipient cells.” A total of 63 mutants passed the secondary screen, and interestingly, most mutants release more NAM. Known NAD<sup>+</sup> homeostasis factors were identified, including *npt1Δ* and *pnc1Δ* (Table S1).

### Measurement(s) of NAD<sup>+</sup>, NADH, NR, NA, and NAM

Total intracellular levels of NAD<sup>+</sup> and NADH were determined using enzymatic cycling reactions as described (71). NR levels were determined by a liquid-based cross-feeding bioassay (17, 19). Relative NA/NAM levels were determined similarly as NR with some modifications. To prepare cell extracts for intracellular NA/NAM determination, ~2.5 × 10<sup>9</sup> (~250 A<sub>600</sub> units) of cells (donors of interest) grown to late log phase

(~16-h growth from an A<sub>600</sub> of 0.1) were lysed by bead-beating (Biospec Products) in 800 μl of ice-cold 50 mM ammonium acetate solution. After filter sterilization, 50–200 μl of clear extract was used to supplement 8-ml cultures of recipient cells (the NA/NAM-dependent *bnA6Δnrk1Δnrt1Δ* mutant) with starting A<sub>600</sub> of 0.05 in niacin-free SC. To determine extracellular NA/NAM levels, supernatant of donor cell culture was collected and filter-sterilized, and then 4 ml was added to 4 ml of recipient cell culture with total starting A<sub>600</sub> of 0.05 in 2× niacin-free SC. A control culture of recipient cells in niacin-free SC without supplementation was included in all experiments. After incubation at 30 °C for 24 h, growth of the recipient cells (A<sub>600</sub>) was measured and normalized to the cell number of each donor strain. A<sub>600</sub> readings were then converted to NA/NAM concentrations using the standard curve established as described previously (19).

### Deamidase activity assay

About 300 A<sub>600</sub> unit cells grown overnight in YPD were harvested, and cell lysate was obtained by beads beating in breaking buffer (10 mM Tris-HCl, pH 7.5, 150 mM NaCl, Roche protease inhibitors). Cell lysate containing 70 μg of total cellular proteins was added to 300 μl of deamidase reaction mix (10 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl<sub>2</sub>) (15, 73) using 8 mM NAD<sup>+</sup>, NMN, NmR, or Nam as substrates, followed by incubation at 30 °C for 45 min. 100 μl of the deamidase reaction mix was then added to 1 ml of ammonia assay mix (3.4 mM α-ketoglutarate, 0.23 mM NADPH, 50 mM phosphate buffer, pH 7.4, 10 units of glutamate dehydrogenase) followed by reaction at room temperature for 15 min (15, 73). The amount of ammonia was calculated by the decrease in absorbance at 340 nm using standard curve derived from the ammonia standard solutions (Sigma).

### Repressible acid phosphatase (rAPase) activity assay

The *rAPase* liquid assay was performed as described (74) on cells grown to late log phase in SDC. In brief, ~2.5 × 10<sup>7</sup> cells were harvested, washed, and resuspended in 150 μl of water. 600 μl of substrate solution (5.6 mg/ml *p*-nitrophenylphosphate in 0.1 M sodium acetate, pH 4) was added to cell suspension, and the mixture was incubated at 30 °C for 15 min. The reaction was stopped by adding 600 μl of ice-cold 10% trichloroacetic acid. 600 μl of this final mixture was then added to 600 μl of saturated Na<sub>2</sub>CO<sub>3</sub> to allow color (neon yellow) development. Cells were pelleted to acquire the supernatant for A<sub>420</sub> determination. The *rAPase* activities were determined by normalizing A<sub>420</sub> readings to total cell number (A<sub>600</sub>).

### Nitrogen starvation

We followed previous nitrogen starvation regimens that were shown to induce autophagy in yeast cells with some modifications (31, 33, 34). 8 ml of cells grown in SC overnight were harvested by centrifugation, washed twice before inoculating into 250 ml of SC – niacin at A<sub>600</sub> of 0.2. SC – niacin was used to reduce the interference of extracellular niacin (NA/NAM) while determining intracellular NA/NAM. After 6 h at 30 °C, 2 × 100 ml of cells were collected and washed twice in water. One tube of washed cells were subject to lysate collection (con-

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trol, 0 h) for NA/NAM and NR measurements. The other tube of washed cells were inoculated into 250 ml of SC –niacin –nitrogen (without ammonium sulfate and amino acids) and allowed to grow for 4 h (N-starvation, 4 h) before lysate collection for NA/NAM and NR measurements. SC –niacin and SC –niacin –nitrogen were made with special yeast nitrogen base from Sunrise Science Products.

### Protein extraction and Western blot analysis

A total of 100 ml of cells were grown in SC or SD to mid-logarithmic phase ( $A_{600}$  of 1–2), concentrated by centrifugation. Total protein extract from cell lysate was obtained by bead-beating in lysis buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1% Nonidet P-40, and Roche cComplete Mini protease inhibitor mixture). 30 or 70  $\mu$ g of total protein was loaded in each lane. After electrophoresis, the protein was transferred to nitrocellulose membrane (Whatman). The membranes were then washed and blotted with either anti-HA antibody (Pierce) or anti-actin antibody (Abcam). Protein was visualized using anti-mouse (GE Healthcare) or rabbit (Pierce) antibody conjugate to the horseradish peroxidase and the ECL reagents (GE Healthcare). The chemiluminescent image was analyzed using the Alpha Innotech imaging system and software provided by the manufacturer.

### Immunoprecipitation and analysis of N-terminal acetylation of Nma1 and Nma2

About 500 ml of WT and *nat3* $\Delta$  cells carrying HA-tagged Nma1 grown to  $A_{600}$  of 1–2 were collected by centrifugation and washed twice in PBS. Cell lysate was obtained by bead-beating in immunoprecipitation lysis buffer (25 mM Tris-HCl, 150 mM NaCl, 1% Nonidet P-40, 3 mM EDTA, 5% glycerol, and protease inhibitor mixture). Lysate was cleared at  $100,000 \times g$  for 1.5 h. Immunoprecipitation was carried out using 1.8 ml of lysate at 2.8 mg/ml with 75  $\mu$ l of Pierce<sup>TM</sup> anti-HA magnetic beads. The magnetic beads were washed five times with end-over-end rotation: wash 1 (20 min) in lysis buffer, washes 2–4 (15 min) in TBS with 0.5% Tween 20, and wash 5 with PBS. The magnetic beads were then washed three more times in 300  $\mu$ l of 50 mM ammonium bicarbonate with end-over-end rotation for 20 min. The magnetic beads were then brought up in 300  $\mu$ l of 50 mM ammonium bicarbonate for on-bead digestion, and 2.5  $\mu$ l of 250 ng/ $\mu$ l Pierce<sup>TM</sup> chymotrypsin protease (catalog no. 90056) was added. Digest was left at room temperature overnight with end-over-end rotation. The next day, the supernatant was separated from the magnetic beads and collected using a magnetic stand. The magnetic beads were washed with 100  $\mu$ l of 50 mM ammonium bicarbonate for 20 min, and the supernatant was pooled. Samples were immediately sent to the University of California Davis proteomics core for tandem mass spectrometry analysis (LC-MS/MS). Digested peptides were analyzed by LC-MS/MS on a Thermo Scientific Q Exactive Plus Orbitrap mass spectrometer in conjunction with Proxeon Easy-nLC II HPLC (Thermo Scientific) and Proxeon nanospray source. The digested peptides were loaded into a 100- $\mu$ m  $\times$  25 mm Magic C18 100- $\text{Å}$  5U reverse phase trap, where they were desalted online before being separated using a 75  $\mu$ m  $\times$  150 mm Magic C18 200- $\text{Å}$  3U reverse phase column. Peptides were

eluted using a 60-min gradient with a flow rate of 300 nl/min. An MS survey scan was obtained for the  $m/z$  range 350–1600, and MS/MS spectra were acquired using a top 15 method, where the top 15 ions in the MS spectra were subjected to high-energy collisional dissociation. An isolation mass window of 1.6  $m/z$  was used for the precursor ion selection, and a normalized collision energy of 27% was used for fragmentation. A 15-s duration was used for the dynamic exclusion. The total number of identified Nma1 and Nma1-specific peptide sequences (peptide spectrum matches) were determined using Byonic software (75) with 1% maximum spectra decoy false-discovery rate. The Skyline software (41) was used to calculate the peak area intensity of the acetylated N-terminal peptides identified by Byonic. The percentages of N-terminal acetylated Nma1 and Nma2 peptides shown in Fig. 4G were calculated from two independent experiments using the Scaffold software with peptide threshold set at 1% false-discovery rate.

**Author contributions**—T.C. designed and performed the experiments. C.J.T.R. performed experiments shown in Figs. 2 (A and B) and 4 (D and E). M.S. and B.S.P. performed the mass spectrometry studies and data analysis shown in Fig. 4G. S.J.L. helped conceive the experiments, analyzed the data, and co-wrote the manuscript with T.C. All authors approved the final version of the manuscript.

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