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| Citation         | Roig, J. 2002. “Nercc1, a Mammalian NIMA-Family Kinase, Binds the Ran GTPase and Regulates Mitotic Progression.” Genes & Development 16 (13): 1640–58. https://doi.org/10.1101/gad.972202. |
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Nercc1, a mammalian NIMA-family kinase, binds the Ran GTPase and regulates mitotic progression

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The protein kinase NIMA is an indispensable pleiotropic regulator of mitotic progression in Aspergillus. Although several mammalian NIMA-like kinases (Neks) are known, none appears to have the broad importance for mitotic regulation attributed to NIMA. Nercc1 is a new NIMA-like kinase that regulates chromosome alignment and segregation in mitosis. Its NIMA-like catalytic domain is followed by a noncatalytic tail containing seven repeats homologous to those of the Ran GEF, RCC1, a Ser/Thr/Pro-rich segment, and a coiled-coil domain. Nercc1 binds to another NIMA-like kinase, Nek6, and also binds specifically to the Ran GTPase through both its catalytic and its RCC1-like domains, preferring RanGDP in vivo. Nercc1 exists as a homooligomer and can autoactivate in vitro by autophosphorylation. Nercc1 is a cytoplasmic protein that is activated during mitosis and is avidly phosphorylated by active p34Cdc2.

Microinjection of anti-Nercc1 antibodies in prophase results in spindle abnormalities and/or chromosomal misalignment. In Ptk2 cells the outcome is prometaphase arrest or aberrant chromosome segregation and aneuploidy, whereas in CFPAC-1 cells prolonged arrest in prometaphase is the usual response. Nercc1 and its partner Nek6 represent a new signaling pathway that regulates mitotic progression.

[Key Words: NIMA; NEK; Ran; RCC1; cdc2/MPF; mitosis]

Received December 20, 2001; revised version accepted May 16, 2002.

Cell division is timed and controlled in part through the interplay of a specialized set of protein kinases and phosphatases, the best known of which are the cyclin-dependent kinases (CDK). Recent studies indicate, however, that several other families of protein kinases also play important roles at different stages of this intricate process [Nigg 2001]. Polo-like kinases [Glover et al. 1998], Aurora kinases [Bischoff and Plowman 1999], and NIMA-like kinases [Neks] [Fry and Nigg 1997; Kandli et al. 2000] have been implicated in such processes as centrosome separation and chromosome condensation in prophase, nuclear envelope breakdown and spindle assembly in prometaphase, as well as in exit from mitosis and cytokinesis.

Neks are named for the Aspergillus nidulans protein kinase encoded by the nimA gene [Osmani and Ye 1996]. Early data suggested that NIMA cooperates with p34Cdc2/cyclin B during the onset of mitosis, perhaps by enabling nuclear entry of cyclin B/p34Cdc2 [Wu et al. 1998]. Moreover, both Cdc2 and NIMA must be inactivated for mitotic exit. Temperature-sensitive mutations of the nimA gene [Osmani et al. 1991a] or expression of the noncatalytic domain of NIMA [Lu and Means 1994] arrest Aspergillus cells in G2 [thus the name NIMA, never in mitosis] without interfering with p34Cdc2 activation. Conversely, overexpression of NIMA causes chromatin condensation and abnormal spindle formation without activating p34Cdc2 [Osmani et al. 1988a; O’Connell et al. 1994]. G2 arrest of nimA mutants can be bypassed by mutation of different anaphase-promoting complex (APC) subunits. Double nimA + APC mutants can enter mitosis when shifted to restrictive temperature, although mitotic cells show aberrant nuclear envelopes and spindle organization, pointing to an involvement of the NIMA protein kinase in mitotic processes beyond the control of the G2/M transition [Osmani et al. 1988b, 1991b; Lies et al. 1998].

NIMA protein levels are maximal during mitosis, and NIMA protein kinase activity seems to parallel NIMA protein content [Osmani et al. 1991b; Ye et al. 1995]. NIMA is hyperphosphorylated in vivo during mitosis and can be phosphorylated in vitro by p34Cdc2 [Ye et al. 1995]. Such in vitro phosphorylation alters NIMA protein kinase activity modestly; however, once phosphorylated in mitosis, NIMA is rapidly degraded, and this degradation is necessary for mitotic exit [O’Connell et al. 1992; Pu and Osmani 1995].
The ability of recombinant NIMA to induce chromatin condensation in fission yeast (O’Connell et al. 1994) and vertebrate cells (accompanied in the latter by nuclear membrane breakdown, O’Connell et al. 1994, Lu and Hunter 1995) as in Aspergillus, suggests that a protein kinase with similar specificity participates in cell cycle control in higher metazoa. At least eight mammalian NIMA-related kinases, or Neks (Nigg 2001), have been identified; however, none has emerged as a bona fide functional homolog of NIMA, that is, as necessary for mitotic progression, or able to induce chromatin condensation if overexpressed. The Neks are most closely related to NIMA in their catalytic domain sequences, but each diverges substantially from NIMA in its non-catalytic C-terminal tail, including the Neurospora crassa NIMA-related kinase that has the capacity to complement the nima mutation (Pu et al. 1995). Nek2, the mammalian homolog most similar in overall amino acid sequence to NIMA, is involved in the regulation of centrosomal structure and function (Mayor et al. 1999), but does not appear to be involved in other aspects of mitosis. The functions of other Neks are largely unknown, although recently Nek6, and the closely related Nek7, recent mammalian additions to the family (Kandli et al. 2000), were shown to phosphorylate the protein kinase p70 S6 kinase on Thr412 within a hydrophobic motif, a phosphorylation that, together with the PDK1-mediated phosphorylation of Thr252 in the activation loop, mediates activation of the p70 S6 kinase (Belham et al. 2001).

Herein we describe a new member of the NIMA-like family of protein kinases, which we designate Nercc1 kinase. This enzyme was identified by its tight binding to overexpressed recombinant Nek6. Nercc1 kinase is activated during mitosis, binds specifically to the Ran GTPase, and is a substrate for Cdc2 phosphorylation. Overexpression of both active and inactive variants of the Nercc1 kinase is toxic to cells, inhibiting cell division and correct segregation of the chromosomes, resulting in either prometaphase arrest or aneuploidy. Nercc1 kinase appears to play one or more central roles in the control of mitotic progression, possibly regulated by p34Cdc2 and the Ran GTPase.

**Results**

**Cloning of Nercc1, a novel protein kinase in the NIMA family**

Immunofluorescence purification of a Flag–Nek6 polypeptide overexpressed in HEK293 cells results in the recovery of an associated 120-kD polypeptide. Incubation of the Flag–Nek6 immunoprecipitate with Mg\(^{2+}\) plus [γ-\(^{32}\)P]ATP yields \(^{32}\)P incorporation into both Nek6 and p120 to a similar extent, suggesting that p120 is a substrate for Nek6, a protein kinase itself, or both [Fig. 1a]. Tryptic digests of the 120-kD band were analyzed by electrospray ionization mass spectrometry. Spectra corresponding to multiple peptide sequences were identified on each of three successive ORFs predicted by GENESCAN (Burge and Karlin 1997) on the human BAC clone 201F1 [AC007055]. The sum of the molecular masses of the three polypeptides predicted by these ORFs was close to 120 kD, suggesting that the exon–intron boundaries had been determined incorrectly by GENESCAN. Further analysis, using GENEMARK (Borodovsky and McIninch 1993) or GENESCAN, yielded predictions containing all three original ORFs (AAD31938, AAD31939, and AAD31940) in polypeptides of ~100 kD (107 kD and 91 kD, respectively).

Using the predicted sequences and supporting ESTs, a cDNA containing the complete coding region for a protein of 979 residues containing all the peptides identified by MS was cloned by PCR (see Supplementary Fig. 1a; the nucleotide sequence along with a translation of the coding region is shown in Supplementary Fig. 1b; Supplemental Material available online at http://www.genesdev.org). The predicted protein product (Fig. 1b) has a calculated molecular mass of 107,034 D, a theoretical pI of 5.50, and contains all 29 peptides detected in the tryptic digest of the 120-kD protein band. The polypeptide has a typical eukaryotic protein kinase domain situated near the N terminus [residues 52–308] that shows all the features of a functional serine/threonine protein kinase. The catalytic domain is most similar to the NIMA-related family of protein kinases (~39%–44% identity and 56%–66% similarity with vertebrate Neks, 33% identity and 49% similarity with NIMA). Immediately following the catalytic domain is a nuclear localization signal [NLS] composed of two classical nuclear localization motifs (residues 347–726; Fig. 1c), followed by a segment containing seven consecutive glycine residues (752–760), encompassed within a PEST region [734–779], the polypeptide is likely to act as a flexible hinge. An acidic serine/threonine/proline-rich segment [761–830] follows, which includes two motifs that conform to the SH3-domain-binding sequence PXXP [X_{32}PXPPXXP; X_{60} and X_{81}KXXPXXP; X_{88}], and seven SP and TP sites [four overlapping the PXXP motifs], immediately succeeding this region is a predicted coiled-coil domain [891–940], followed by the protein C terminus. We designate this polypeptide as Nercc1 kinase, based on the similarity of the kinase domain to the NIMA/Nek kinases, and the presence of an RCC1-like domain.

Although there is no significant identity in the primary sequences of the C-terminal noncatalytic segments of NIMA and the Nercc1 kinase, these two segments do share several related features, namely, a nuclear localization signal immediately following the catalytic domain, a proline-rich segment containing multiple SP and TP sites (some of which, in the case of NIMA, are probably phosphorylated during mitosis and appear to be important for regulation; Fry and Nigg 1995; Osmani and
Ye 1996), and a coiled-coil domain. In addition, several PEST regions (involved in control of protein stability) are found in both protein kinases. A striking difference between Nercc1 protein kinase, NIMA, and the Neks characterized thus far in higher eukaryotes is the presence in the Nercc1 kinase of a domain homologous to the RCC1 protein. RCC1 is a guanine-nucleotide-exchange factor for the small G-protein Ran, and is composed of seven repeats of 51–68 residues folded in a structure that resembles a seven-blade pro-

Figure 1. Nek 6 coimmunoprecipitates with a 120-kD protein. Structure of Nercc1 polypeptide. (a) HEK293 cells were transfected with either empty vector (−) or pCM5 Flag–Nek6 (+). Cell lysates were prepared 48 h later, and Nek6 was immunoprecipitated using an anti-Flag antibody. The washed immunoprecipitates were incubated with $\text{Mg}^{2+}$ [$\gamma^{-32}\text{P}]\text{ATP}$. Coomassie stain (left panel) and $32\text{P}$ autoradiography (right panel) of the gel are shown. (b) Cartoon of Nercc1 polypeptide domain structure. [NLS] Nuclear localization signal, [RCC1] RCC1 homology domain, [Gly] polyglycine stretch, [PXXP] proline-rich motifs, [S/TP sites] Ser/Thr-Pro motifs. (c) Alignment of the RCC1 repeats of the human regulator of chromosome condensation (RCC1) with Nercc1 RCC1 domain repeats. Amino acid numbers are shown. Of the 10 residues known to be important for the exchange activity of RCC1 toward Ran [i.e., affecting the $K_{\text{m}}$ or $K_{\text{cat}}$; Jiet al. 1999], Nercc1 lacks a conserved residue corresponding to RCC1 D44, R206, D182, and H270, although a residue of similar charge exists at the position +1 for the latter two; D128 E157 and H304 are conserved, whereas H78, R101 and H410 are substituted by similarly charged residues. Notably, mutation of RCC1 D182 [an asparagine in Nercc1] results in a protein with no measurable GEF activity.
peller (Renault et al. 1998). The Nercc1 RCC1 domain has a 27% identity and 43% similarity to RCC1, and like RCC1 contains seven tandem repeats [Fig. 1c]. We have been unable to detect a protein kinase with this domain organization in lower eukaryotes; interestingly, a hypothetical protein kinase in the genome of Drosophila melanogaster (accession code AAF56344) shows an architecture homologous to Nercc1 kinase, that is, an N-terminal NIMA-related protein kinase domain followed by a series of RCC1 domain repeats.

During the revision of this paper, the sequences of two different mammalian NIMA-family kinases containing an RCC1 domain appeared in GenBank. One, cloned from mouse (accession code AF407579) and zebrafish (AF407580) by Beier and colleagues (S. Liu, W. Lu, T. Obara-Ishibara, I. Drummond, and D.R. Beier. A defect in a novel Nek-family kinase causes cystic disease in the mouse and in zebrafish. unpubl.), has been named Nek8. Mouse Nek 8 is a protein of 698 residues, 32% identical and 49% similar to the first 752 residues of Nercc1. It lacks the C-terminal tail that in Nercc1 spans from the polyglycine region to the end of the protein; this is an important regulatory region in Nercc1 [see below]. A second protein kinase identical in sequence to Nercc1 has been deposited in GenBank, this can be accessed both as Nek8 (AY045850) and Nek9 (NM033116) [Holland et al. 2002].

The Nercc1 protein kinase is expressed in all human cell lines tested, including HEK293, HeLa, and U2OS cells; Nercc1 protein expression is detected in these and other mammalian cell lines, as well as in all mouse tissues tested [see Supplemental Fig. 2].

**Nercc1 interaction with Nek6**

We sought to confirm the association of Nek6 with recombinant Nercc1 by coexpression of GST–Nek6 and Flag–Nercc1 in 293 cells. Full-length Flag–Nercc1 is seen to bind specifically to GST–Nek6, whereas the C-terminally truncated Flag–Nercc1[1–739], despite comparable expression, is unable to bind GST–Nek6 [Fig. 2a]. Reciprocally, the fusion of the Nercc1 kinase C-terminal segment, Nercc1[732–979], to GST is sufficient to enable specific binding of Flag–Nek6 [Fig. 2b]. Further analysis [see below] indicates that the site of Nek6/Nercc1 interaction lies between the Nercc1 amino acids 732 and 891.

**Nercc1 homodimerizes through a coiled-coil domain distinct from the Nek6-binding site**

Nercc1 contains a predicted coiled-coil motif near its C terminus [residues 891–940], a likely candidate for an oligomerization domain [Fig. 3a]. The ability of Nercc1 to form homooligomers is shown by the coprecipitation of Flag–Nercc1 with HA–Nercc1 [Fig. 3b]. The Nercc1(891–940) segment was fused to GST and coex-

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**Figure 2.** Nercc1 binding to Nek6. (a) HEK293 cells were cotransfected with full-length (FL) Flag–Nercc1 or Flag–Nercc1[1–739], and either GST–Nek6 or GST alone. GST fusion proteins were isolated from cell lysates using GSH-agarose beads. Western blots of the GSH-agarose isolate and the cell extract are shown. (b) HEK293 cells were cotransfected with Flag–Nek6 and either GST–Nercc1[732–979] or GST alone. Western blots of the GSH-agarose isolates and the cell extract are shown; a cartoon of the Nercc1 Flag-fusion proteins used is shown below.
pressed with either full-length Flag–Nercc1 or Flag–Nercc1 deleted of its C-terminal 89 residues (Nercc1 residues 1–891). Figure 3c shows that although full-length Nercc1 associates specifically with GST–Nercc1(891–940), deletion of the C-terminal 89 residues of Nercc1 abolishes this association (Fig. 3c, left panel). In addition, Nercc1(1–891) cannot oligomerize with full-length Nercc1 (Fig. 3c, right panel). Thus, Nercc1 oligomerizes through its C-terminal coiled-coil domain. As shown below, this oligomerization is important for the regulation of Nercc1 protein kinase activity.

Gel filtration analysis of both recombinant and endogenous Nercc1 in 293 cells shows that the protein kinase exists in a high molecular mass complex of ~600 kD (Fig. 3d). The larger-than-expected size probably reflects the association of Nercc1 with other proteins such as Nek6,
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and perhaps higher-order homooligomers and/or a complex of asymmetric shape.

The 891–940 deletion does not affect Nek6 interaction with Nercc1; full-length Nercc1 and Nercc1[1–891] both interact with Nek6 with similar affinities. Moreover, GST–Nercc1[891–940] does not bind Nek6 (data not shown). Thus, Nek6 binds to Nercc1 between amino acids 732 and 891, that is, in the region between the end of the Nercc1 RCC1 domain and the beginning of the coiled-coil domain.

Protein kinase activity of Nercc1

The catalytic properties of Nercc1 were studied using immunoprecipitates of Flag-tagged forms of the Nercc1 polypeptide transiently expressed in HEK293 cells. The specificity of the measured protein kinase activity for the Nercc1 polypeptide was verified by the inability of an ATP-binding site mutant of Nercc1 (K81M) to catalyze significant 32P transfer to itself or exogenous substrates (Fig. 4b, lanes 3,4). Thus, the kinase activity described below is caused by Nercc1 and not a contaminating protein kinase (e.g., Nek6). Nercc1 can autophosphorylate and phosphorylate different histones and MBP, whereas β-casein is phosphorylated much less rapidly (data not shown). Phosphoamino acid analysis of Nercc1 autophosphorylation and phosphorylation of histone H3 showed that Nercc1 phosphorylates serine and threonine residues exclusively (data not shown). Recombinant wild-type Nercc1 has low basal activity when extracted from exponentially growing cells; however, preincubation with Mg2+ plus ATP (100 µM) induces Nercc1 autophosphorylation (accompanied by a slowing in electrophoretic mobility) and activation (this activation can be reversed by phosphatase treatment; see below). The rate of in vitro activation is greatly enhanced if Mn2+ replaces Mg2+ (data not shown). Autophosphorylation/autoactivation is time and ATP concentration dependent; importantly, 10 µM ATP fails to enable significant Nercc1 activation even after incubation times of 90 min at 25°C, whereas 100 µM ATP gives maximal activation [10- to 20-fold] by 60 min, with half-maximal activation at 20 min (Fig. 4a). Once activation is complete, Nercc1 catalyzes a robust phosphorylation of H3 at 5 µM to 10 µM ATP. This apparent increase in affinity for ATP after activation enables the design of a Nercc1 kinase assay that reflects the extent of activation achieved. Notably, endogenous Nercc1, immunoprecipitated out of exponentially growing cells by specific antibodies, shows a similar pattern of Mg2+ ATP-dependent autoactivation in vitro. Nercc1 is able to use GTP as a phosphate donor; GTP supports autoactivation, and after maximal autoactivation in vitro, enables the phosphorylation of histone H3 at ∼30% the rate observed with ATP (data not shown).

We next carried out a structure–function analysis of the ability of recombinant Nercc1 kinase to autoactivate in vitro. A series of Nercc1 variants, transiently expressed in HEK293 cells, was assayed for H3 kinase activity after preincubation at 25°C for 30 min with Mg2+ or Mg2+ plus 100 µM ATP, the latter a condition sufficient to enable near-maximal autoactivation of wild-type Nercc1 [Fig. 4a]. After washing away the non-radioactive ATP, the H3 kinase assay was commenced using a concentration of [γ-32P]ATP (5 µM to 10 µM) below that capable of supporting autoactivation of wild-type Nercc1 kinase [Fig. 4b, lanes 1,2]. An ATP-binding loop mutant of Nercc1 [K81M] shows no significant autophosphorylation/kinase activity irrespective of preincubation with Mg2+ ATP [Fig. 4b, lanes 3,4]; this is also true of Nercc1[338–979], which lacks the entire Nercc1 protein kinase domain [data not shown]. Nercc1[1–391], which lacks the RCC1 domain and the C-terminal tail, shows a low basal protein kinase activity but can be modestly activated by preincubation with Mg2+ and 100 µM ATP [Fig. 4b, lanes 5,6, and 4c]; Nercc1[1–391, K81M] is entirely devoid of activity [Fig. 4b, lanes 7,8], as is Nercc1[1–308], which terminates the end of the canonical kinase domain [data not shown]. Nercc1[1–739] retains both the protein kinase domain and the RCC1 domain, but is nevertheless inactive and incapable of autoactivation [Fig. 4b, lanes 9,10]. Deletion of the coiled-coil domain Nercc1[1–889] greatly diminishes the rate of autoactivation [Fig. 4b, lanes 11,12], whereas deleting the proline-rich C-terminal segment but retaining the coiled coil [Δ763–889] permits substantial autoactivation [Fig. 4b, lanes 13,14]. Selective deletion of the RCC1 domain, Nercc1[Δ347–732], results in a very high basal H3 kinase activity as compared with wild-type Nercc1, which is not further increased by preincubation in vitro with Mg2+ plus 100 µM ATP [Fig. 4b, lanes 15,16, and 4c]. A more detailed examination of the time course of activation by ATP [100 µM] shows that although both Nercc1[1–391] and Nercc1[1–889] are capable of autoactivation, the rate is greatly diminished as compared with wild type. Thus deletion of the RCC1 domain produces a mutant with a basal activity similar to the maximal attainable by autoactivation [although the Δ347–732 polypeptide is less stable at 25°C]; however, if the C-terminal dimerization domain is also deleted, as in Nercc1[1–391], the basal activity returns to low levels and autoactivation is severely retarded.

These results suggest a mechanism for Nercc1 kinase regulation (at least in vitro) in which the Nercc1 homodimer is maintained in an inhibited state by the ability of the RCC1 domain to abrogate intramolecular autophosphorylation. The inability of autoactivation to occur at ATP concentrations that enable robust phosphate transfer once activation has occurred suggests that the inhibitory action of the RCC1 domain operates, at least in part, by obstruction of the ATP-binding site. One prediction of this model is that the RCC1 domain and the kinase domain of Nercc1 are likely to interact. The occurrence of such an interaction is shown in Figure 4d, Flag–Nercc1[338–739] expressed in HEK293 cells associates directly with coexpressed HA–Nercc1[1–391], supporting the view that the RCC1 domain may inhibit the kinase domain through a direct interaction. Nercc1 autophosphorylation is likely to occur in trans within the homodimer, as deletion of the RCC domain results in...
Figure 4. Nercc1 autoactivation in vitro. (a) Flag–Nercc1 was immunoprecipitated from HEK293 cells, washed, and incubated at 25°C in phosphorylation buffer for the indicated times with 10 µM or 100 µM ATP. Incubations were terminated by washing, followed by the addition of 10 µM [*32P]*ATP and histone H3 (1 µg/50 µL). After 10 min at 30°C, [*32P] incorporation was stopped by addition of SDS sample buffer, followed by SDS-PAGE and blot transfer. The anti-Flag immunoblot (upper panel), [*32P] autoradiography (middle panel), and the relative quantity of [*32P] incorporated into histone H3 (bottom panel) are shown. (b) Immobilized Flag-tagged Nercc1 variants, isolated after transient expression in HEK293 cells, were washed and incubated in phosphorylation buffer at 25°C for 30 min with Mg2+ and with or without 100 µM ATP. After an additional wash, samples were incubated at 30°C with Mg2+ plus 10 µM [*32P]*ATP and histone H3 (1 µg/50 µL). After 10 min the reaction was stopped by addition of SDS sample buffer followed by SDS-PAGE and blot transfer. The [*32P] autoradiogram (upper panel) and anti-Flag immunoblot (middle panel) are shown. (c) Time course of activation of the H3 kinase activity of wild-type and mutant Nercc1. (■) Flag–Nercc1 (wild-type); ( ●) Flag–Nercc1[Δ446–732]; ( ●) Flag–Nercc1[1–391]; and ( ▲) Flag–Nercc1[1–891], were expressed in HEK293 cells, immobilized on anti-Flag-agarose, washed and incubated at 25°C with Mg2+ plus 100 µM ATP. At the times indicated, samples were washed, followed by addition of Mg2+ plus 10 µM [*32P]*ATP and histone H3 (1 µg/50 µL). After 10 min at 30°C, SDS sample buffer was added, and [*32P] incorporation into H3 was measured (using a PhosphorImager) after SDS-PAGE and blot transfer. [*32P] incorporation is expressed as a percentage of Nercc1 wild-type value at t = 0, that is, no preincubation with 100 µM ATP. (d) The Nercc1 protein kinase domain and RCC1 domain interact in vivo. HEK293 cells were transfected with the HA–Nercc1 protein kinase domain, HA–Nercc1[1–391], and either Flag–Nercc1 RCC1 domain, Flag–Nercc1[338–778], or empty plasmid. Anti-Flag immunoprecipitates were immunoblotted with anti-HA (upper panel) or anti-Flag (middle panel). The expression of HA-Nercc1[1–391] is shown in the lower panel. A cartoon of the Nercc1 variant used in Figure 6 is below.
spontaneous activation in vivo only if the C-terminal tail, that is, the ability to homodimerize, remains intact.

Although this model fits the data for Nercc1 activation in vitro it is obvious that, inasmuch as the intracellular ATP concentration is 2 mM to 5 mM, the low basal activity of recombinant and endogenous Nercc1 kinase must reflect the operation in vivo of one or more negative regulatory inputs, in addition to the inhibition provided by the RCC1-like domain. Such inputs could include protein phosphatases, as well as inhibitory ligands or polypeptides.

Nercc1 binds to Ran

The presence in Nercc1 of a domain homologous to RCC1, a nucleotide exchange factor protein for the Ran GTPase, raises the question of whether Nercc1 binds Ran, and if so, to what functional effect. Prokaryotic recombinant GST and GST fusions with the Nercc1 kinase domain—GST–Nercc1[1–391], the RCC1 domain—GST–Nercc1[338–739], and the C-terminal tail—GST–Nercc1[732–979] were immobilized on GSH-agarose beads and incubated with prokaryotic recombinant Ran that had been preloaded with GTP/βS or GTPγS. Whereas neither GST nor GST–Nercc1[732–979] bind Ran, the GST–Nercc1 kinase domain and RCC1-like domain fusion proteins are able to bind Ran with very high efficiency (Fig. 5a). Both Nercc1 domains bind Ran-GDP to a somewhat greater extent than Ran-GTP, however, the degree of this preference is somewhat variable between experiments.

To examine the interaction between Ran and Nercc1 in vivo, we coexpressed HA–wild-type Ran (wild type) with different Flag–Nercc1 mutants, and probed the Flag immunoprecipitates for the presence of HA–wild-type Ran. Cell lysis and subsequent washes were carried out in the presence of excess Mg2+ to conserve Ran in its nucleotide-bound form. Figure 5b shows that wild-type Ran associates with full-length Nercc1 (both wild-type and K81M) as well as with the isolated Nercc1 catalytic domain fragments, 1–308 and 1–391, as expected from Figure 5a. The specificity of this interaction was assessed by examining the relative ability of Ran to bind the Nercc1 catalytic domain, Nercc1[1–308], or Nek 6, another protein kinase in the NIMA subfamily (Fig. 5c); no binding of HA–Ran to Nek6 is detectable, whereas both the Nercc1 catalytic domain and RCC1 [the Ran GEF, a positive control] bind avidly to Ran.

The deletion of the RCC1 domain, Nercc1[Δ347–732], or the C-terminal tail, Nercc1[1–739], does not detectably impair the binding of HA–Ran, and in contrast to the in vitro results, the isolated Nercc1 RCC1-like domain, Nercc1[338–739], shows very little association with Ran. The inability of the Nercc1 RCC1 domain to bind Ran in vivo is as yet unexplained. The RCC1-like domain may simply bind Ran with lower affinity than the catalytic domain; conversely, access of Ran to the isolated Nercc1 RCC1-like domain in vivo may be obstructed.

We next addressed the relative binding in vivo of Nercc1 to Ran-GDP versus Ran-GTP. A potential confounding element in assessing the interactions of Nercc1 with the different forms of Ran in vivo is the subcellular localization of the various partners; in interphase cells Ran-GDP is located exclusively in the cytoplasm, whereas Ran-GTP is restricted to the nucleus (Kalab et al. 2002), a situation maintained by the nuclear location of the Ran GEF, RCC1, and the cytosolic localization of Ran GAP. We therefore coexpressed either full-length, wild-type Nercc1, which is exclusively cytoplasmic (see below) or a nuclear-targeted form of full-length Nercc1 (NLS-Nercc1) with wild-type Ran or Ran mutants that bind GDP [T24N] or GTP [G19V] exclusively; either Ran or Nercc1 was immunoprecipitated and probed for the association with the other polypeptide. Figure 5d shows that full-length, wild-type Nercc1 binds coexpressed wild-type Ran and Ran T24N to a greater extent than RanG19V. Although this suggests a preference for Ran-GDP, it should be recalled that whereas RanT24N and Nercc1 are both cytoplasmic, RanG19V is exclusively nuclear in localization. However NLS-Nercc1, which is exclusively nuclear (see Fig. 7c below), also does not bind the GTP-locked mutant RanG19V, strongly supporting the conclusion that full-length Nercc1 indeed has a higher affinity for Ran-GDP over Ran-GTP.

Nercc1 is phosphorylated and activated during mitosis and can be phosphorylated in vitro by p34Cdc2

We next examined Nercc1 protein levels and activity during cell cycle progression. The level of Nercc1 protein in HeLa cell extracts remains constant during different phases of the cell cycle [G1/S, G2, M, G1]; however, the Nercc1 polypeptide displays a marked slowing in electrophoretic mobility during mitosis [Fig. 6a], which can be mimicked by treatment in vivo with the protein phosphatase inhibitor calyculin [data not shown]. A similar electrophoretic slowing of Nercc1 occurs in CHO-K1, COS7, U2OS, or HEK293 cells arrested in mitosis [data not shown]. In addition to nocodazole-induced mitotic arrest, we also examined mitotic cells prepared by shake-off from a culture that had been pseudosynchronized in G1/S by thymidine block and then released into the cell cycle [Fig. 6b]. Nercc1 is unmistakably up-shifted in electrophoretic mobility during mitosis (Fig. 6). The deletion of the RCC1 domain, Nercc1[Δ347–732], or the C-terminal tail, Nercc1[1–739], does not detectably impair the binding of HA–Ran, and in contrast to the in vitro results, the isolated Nercc1 RCC1-like domain, Nercc1[338–739], shows very little association with Ran. The inability of the Nercc1 RCC1 domain to bind Ran in vivo is as yet unexplained. The RCC1-like domain may simply bind Ran with lower affinity than the catalytic domain; conversely, access of Ran to the isolated Nercc1 RCC1-like domain in vivo may be obstructed.

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Figure 5. Nercc1 binding to Ran. (a) The binding of recombinant Ran to GST–Nercc1 variants in vitro. GST and GST–Nercc1 variants were purified and immobilized on GSH-agarose. Ran was produced in bacteria as a GST fusion, purified, cleaved from the immobilized GST fusion, and loaded with GDP$^\text{S}$ (GDP) or GTP$^\text{S}$ (GTP). Immobilized GST or GST–Nercc1 fusion proteins were incubated with Ran in the Ran binding buffer containing the indicated nucleotides (100 µM). After extensive washing, the proteins retained on the GSH-agarose were eluted into SDS sample buffer, and analyzed by immunoblot for Flag (upper panel) and GST (lower panel). The Ran input is shown. (b,c) HEK293 cells were cotransfected with HA–Ran and Flag–Nercc1 variants or Flag vector. Cells were extracted into Ran lysis buffer. Anti-Flag immunoprecipitates and aliquots of the extracts were subjected to immunoblot with anti-HA [upper panel] and anti-Flag [middle panel] antibodies. Expression of HA–Ran is shown in the lower panel. (d) HEK293 cells were transfected with Flag–Nercc1 (left) or Flag–NLS-Nercc1 (right) together with either HA–Ran wild type, HA–Ran G19V (constitutively GTP-bound), or HA–Ran T24N (constitutively GDP-bound or nucleotide-free). Cells were extracted into Ran lysis buffer. Anti-Flag immunoprecipitates were immunoblotted with anti-HA [upper panel] and anti-Flag [middle panel]. The expression of the HA–Ran variants is shown in the lower panel.
mitotic shake-off (mitotic cells) versus those that remain attached (non-mitotic cells). The Nercc1 kinase activity in mitotic cells is fourfold to fivefold higher than in exponentially growing cells, despite comparable polypeptide levels; nocodazole-arrested, non-mitotic cells show a small increase in Nercc1 kinase activity, probably caused by contaminating mitotic cells (Fig. 6c). Similar results were obtained with U2OS cells (data not shown). Thus, Nercc1 is activated during mitosis.

To determine whether the observed mitotic Nercc1 activation and change in electrophoretical mobility were caused by phosphorylation, we incubated Nercc1 immunopurified from mitotic cells with phosphatase. As a control, recombinant Flag–Nercc1 preactivated by incubation with 100 µM ATP [γ-32P]ATP (10 µM) and histone H3 (2 µg/50 µL). The reaction was stopped by addition of SDS sample buffer. Shown are an anti-Nercc1 (N1) immunoblot of the immunoprecipitates (upper panel) and the 32P incorporation into Nercc1 (middle panel) and histone H3 (lower panel). 32P incorporation into NlgG immunoprecipitates (background) was quantified by PhosphorImager and subtracted from H3 32P incorporation in anti-Nercc1 immunoprecipitates. The resulting Nercc1 activity was expressed as the percentage of activity in exponentially growing cells. (d) Flag–Nercc1 preactivated by incubation with 100 µM ATP [black bars] and endogenous Nercc1 immunopurified from cells arrested in mitosis by nocodazole (gray bars) were incubated in alkaline phosphatase buffer with no addition (columns 1 and 2) or with 40 U of calf intestine alkaline phosphatase (columns 5, 6) with (columns 3, 4) or without (columns 3, 4) 4 mM EGTA. After washing, Nercc1 activity was assayed and expressed as a percentage of non-phosphatase-treated enzyme (columns 1 and 2). (e) Nercc1 is an in vitro substrate of p34Cdc2. Flag–Nercc1 K81M was produced in HEK293 cells, immunopurified with anti-Flag antibody, and eluted from the immunoprecipitates with Flag peptide. Soluble K81M was incubated at 30°C for the indicated times in phosphorylation buffer containing 100 µM [γ-32P]ATP with and without purified active p34Cdc2/cyclin B from Xenopus MPF (maturation promoting factor). Coomassie staining and 32P autoradiography of Nercc1 K81M are shown. Quantitation of incorporated 32P into Nercc1 K81M was carried out by PhosphorImager.
was incubated with phosphatase, a similar decrease in protein kinase activity was observed. Thus, Nercc1 activation in vitro and in vivo during mitosis is caused by phosphorylation.

Transiently expressed, catalytically inactive forms of recombinant Nercc1 [K81M or 338–979] show an up-shift on SDS PAGE in response to nocodazole, much the same as wild-type Nercc1 [data not shown]. Such inactive Nercc1 mutants cannot autophosphorylate, indicating that the mitotic modification of Nercc1 at least in part is owing to another protein kinase; an obvious candidate is p34\(^{\text{Cdc2}}\). Recombinant full-length Flag–Nercc1 [K81M] eluted from Flag-agarose beads is phosphorylated in vitro by purified Xenopus active p34\(^{\text{Cdc2}}\)/cyclin B (maturation promoting factor, MPF; Fig. 6e). MPF-catalyzed Nercc1 phosphorylation induces an up-shift in Nercc1 similar to that observed in vivo in mitotic cells, and overall \(^{32}\)P incorporation into Nercc1 [K81M] rapidly approaches 1 mole PO\(_4\)/mole protein [Fig. 6e]. Similar results are obtained with cyclin B immunoprecipitates from nocodazole-arrested mitotic HeLa cells, whereas cyclin B immunoprecipitates from non-mitotic cells do not catalyze Nercc1 phosphorylation [data not shown]. Inhibition of the Nercc1-phosphorylating activity in the cyclin B immunoprecipitate from mitotic cells by the CDK inhibitor roscovitine confirms the identity of the kinase activity as Cdc2. Thus, Nercc1 is an in vitro substrate for p34\(^{\text{Cdc2}}\), and this phosphorylation produces a change in Nercc1 electrophoretic mobility similar to that observed in mitotic cells, indicating that p34\(^{\text{Cdc2}}\) contributes to Nercc1 phosphorylation during mitosis.

It is important to point out that phosphorylation of Nercc1 by Cdc2/MPF in vitro does not significantly alter maximal Nercc1 kinase activity toward histone H3. Similarly, the up-shift in Nercc1 mobility that occurs on treatment of cells with calyculin is not accompanied by an increase in Nercc1 kinase activity [data not shown]. Thus, although the autophosphorylation/autoactivation of Nercc1 in vitro is also accompanied by a slowing in electrophoretic mobility, the occurrence of an up-shift is not synonymous with Nercc1 activation. The functional consequences of Cdc2-catalyzed phosphorylation of Nercc1 during mitosis are not yet known.

**Nercc1 cellular localization**

Immunofluorescence studies using the N1 or C1 anti-Nercc1 peptide antibodies showed Nercc1 to have a finely granular cytoplasmic fluorescence in all cell lines tested [HeLa, PtK2, HEK293, NIH3T3]. Nercc1 appears diffusely within the cytoplasm without association with organelles, plasma membrane, or cytoskeletal elements [Fig. 7a; see also Supplemental Fig. 3]. We also performed nonequilibrium sucrose density gradient fractionation of HEK293 cells; Nercc1 sedimentation corresponded to that of other cytoplasmic proteins [e.g., lactate dehydrogenase], and was distinct from a variety of membrane markers, for example, βCOP, a TGN marker. Overlay of Nercc1-enriched fractions with lighter sucrose followed by centrifugation to equilibrium did not result in Nercc1 translocation up into the low-density fraction, as occurs with membrane-bound proteins [data not shown]. Thus Nercc1 is localized in the cytoplasm unattached to cellular membranes.

Notably, Nercc1 immunoreactivity is absent from the nucleus [Fig. 7a]. The lack of nuclear Nercc1 was also evident on overexpression of recombinant Flag-Nercc1 [Fig. 7c; Supplemental Fig. 3]. This was surprising, as Nercc1 contains a classical nuclear localization signal [NLS] that is fully functional when appended to the Nercc1 N terminus or to another polypeptide [Fig. 7c]. Careful examination of immunofluorescence specimens of cells subjected to a variety of treatments including leptomycin B, an inhibitor of CRM-dependent nuclear export [Supplemental Fig. 3], failed to uncover instances of endogenous Nercc1 in nuclei.

We next examined Nercc1 localization in mitotic cells [Fig. 7b]. HeLa cells were blocked in mitosis with nocodazole, or enriched in mitotic cells by double thymidine block followed by release and mitotic shake-off; in both instances, Nercc1 immunofluorescence in mitotic cells was diffusely distributed throughout the cell and conspicuously absent from the chromosomes. To test whether Nercc1 is associated with the mitotic spindle, mitotic HeLa cells were fixed after treatment for 1 min with the nonionic detergent saponin. Eg5, a microtubule-binding motor, is readily visualized on spindle microtubules after this light saponin treatment, whereas endogenous Nercc1 immunofluorescence is completely removed from the cells under these conditions. Thus Nercc1 is not [tightly] associated with the spindle microtubules, and is diffusely distributed through the cell during mitosis.

A series of Flag–Nercc1 mutants was examined for their cellular distribution during transient expression in HeLa cells [Fig. 7c]. Four types of subcellular distribution were observed: cytoplasmic, nucleocytoplasmic, predominantly nuclear, and exclusively nuclear. Overexpressed wild-type Nercc1 is distributed in the cytoplasm like endogenous Nercc1; however, ∼5% of cells show slight nuclear immunofluorescence. The inactive Nercc1 ATP-site mutant [K81M] shows a substantial nuclear component, suggesting that the NLS might be inactivated by autophosphorylation. The Nercc1[1–391] variant, although potentially capable of being activated, is nevertheless exclusively nuclear; this establishes the functionality of the Nercc1 NLS, as does the exclusive nuclear localization of a wild-type Nercc1 to which a copy of its endogenous NLS is fused at its N terminus.

The conversion of Nercc1 to forms that achieve nuclear localization results in the frequent occurrence of micronuclei, multiple nuclei, and lobed nuclear morphologies. Such phenotypes have been associated in other circumstances with lagging chromosomes and chromosome nondisjunction in anaphase [Cimini et al. 2001]. The frequency of these morphologies was highest in cells expressing Nercc1[1–391], that is, an active kinase domain with the nuclear localization; however, expression of the cytoplasmic kinase-inactive variant Nercc1[1–739], as well as the nuclear/cytoplasmic vari-
Nercc1 kinase regulates mitotic progression

Figure 7. Nercc1 cellular localization. (a) Nercc1 immunolocalization in interphase. Specificity of the anti-Nercc1 C1 antibody. Cells were immunostained as described, using anti-Nercc1 C1 antibody (left panels) or the C1 antibody preincubated with the immunizing peptide (right panels). Bar, 10 µm. (b) Immunocytochemical identification of Eg5, Nercc1, and DNA in mitotic HeLa cells, before and after a light (1-min) saponin treatment. Bar, 10 µm. (c) Flag–Nercc1 variants were transfected into HeLa cells, and their localization was visualized using anti-Flag antibody. The subcellular distribution of Flag–Nercc1 was assessed and assigned to four subgroups: cytoplasmic, nucleocytoplasmic, predominantly nuclear, and nuclear. Variants that induced abnormal nuclear morphologies are shown in panel 5; an example of the lobed nuclear morphology observed with several variants is shown.

Nercc1 [K81M] and Nercc1[1–308] have a similar, but less marked effect (Fig. 7c, column 5).

Nercc1 regulates mitotic progression

The structural similarities between Nercc1 and NIMA, the activation of Nercc1 during mitosis, the ability of Nercc1 to bind Ran in vitro and in vivo and to induce abnormal nuclear morphology when targeted to the nucleus, together point to the likelihood that Nercc1 is a mitotic regulator. A preliminary indication was provided by the observation that transfection of a plasmid encoding an eGFP–Nercc1 [K81M] fusion protein appeared to interdict cell division. Thus, time-lapse recordings of
transfected HeLa cells showed that, whereas 78% of 70 cells transfected with a plasmid encoding eGFP alone underwent division within the subsequent 36 h, only 4% of 52 HeLa cells transfected with eGFP–Nercc1[K81M] underwent division, and 85% proceeded to cell death as compared with 18% of the eGFP-transfected cells. Interestingly, even wild-type Nercc1 was somewhat toxic—only 29% of 31 HeLa cells transfected with an eGFP–Nercc1 plasmid proceeded through mitosis. The impact of the nuclear-localized Nercc1[1–391] was even more marked; only 1 of 45 HeLa cells transfected with eGFP–Nercc1[1–391] underwent division, and 72% of these cells were dead after 36 h.

To test directly whether Nercc1 participates in the control of mitotic progression, we used live observation of PtK2 cells after microinjection of affinity-purified anti-Nercc1[peptide] antibodies or purified preimmune IgG. When PtK2 cells were injected with Nercc1 antibodies during interphase, mitosis was never subsequently observed, preventing any conclusion. Therefore, antibody microinjection of cells in prophase was undertaken; such cells were identified by the presence of chromosome condensation and nucleolar disassembly. All such cells microinjected with normal rabbit IgG (5 at 2.5 mg/mL and 15 at 10 mg/mL) completed mitosis normally and produced daughter cells, save one, which had a lagging chromosome; this occurrence is consistent with the frequency of lagging chromosomes reported earlier for PtK cells [Izzo et al. 1998]. In contrast, 14 cells of 30 cells (i.e., roughly 45%) microinjected with affinity-purified anti-Nercc1[peptide] (C1) IgG [2.5 mg/mL] showed mitotic abnormalities of two basic types. The first type involved abnormalities in chromatid segregation associated with abnormal spindle dynamics. Thus, four cells never entered anaphase B (i.e., movement of poles with attached chromosomes in opposite directions), although the spindle was apparently normal, nevertheless, cytokinesis proceeded. In two experiments this resulted in the subsequent trapping of one or more chromosomes in the cytokinetic furrow, creating DNA bridges between the nuclei of daughter cells (Fig. 8a, left column, A). In two other cases, separation of the chromosomes to opposite poles stopped prematurely, chromosome decondensation was observed, and a cytokinetic furrow formed to one side of the spindle, resulting in formation of one daughter cell with 4N DNA and one daughter cell without DNA (Fig. 8a, middle column, B). The second type of abnormality concerned the formation of the mitotic spindle. In eight cells, the spindle was not visible on phase contrast images throughout the whole duration of the recording, and ~40 min after the nuclear envelope breakdown, the chromosomes concentrated in the center of the rounded-up PtK2 cell, with chromosome arms extending far into the cell periphery [Fig. 8a, right column, C]. When fixed immediately after the recording, these cells showed a highly disrupted spindle [Fig. 8b, anti-Nercc1; cf. with a cell microinjected with control IgG, control], although two separated centrosomes were visible, the cells showed an interphase array of microtubules.

Two anti-Nercc1 C1-microinjected cells formed a bipolar mitotic spindle, but anaphase started prematurely, specifically at only 8 min after the last monooriented chromosome acquired biorientation (this is 1 min less than the minimum time recorded for PtK cells, Rieder et al. 1994). Thus, in these cells, anaphase started without congression of chromosomes to the metaphase plate, although cell division proceeded normally from then on [data not shown].

Similar kinds of mitotic abnormalities, including a failure to enter anaphase B, were observed in 4 of 8 cells microinjected with an anti-Nercc1 IgG raised against a peptide sequence from the Nercc1 catalytic domain [E2 antibody, amino acids 80–94]. Microinjection of 10 PtK2 cells with affinity-purified anti-Nercc1 N1 [amino acids 3–18] IgG did not alter the normal progression of mitosis [data not shown].

We also microinjected the Nercc1 C terminus antibody into CF-PAC1 cells in prophase. CF-PAC1 cells are a human cell line that contains levels of Nercc1 comparable to HeLa and HEK293 cells [data not shown] and that has been used previously for microinjection studies of spindle dynamics [Mountain et al. 1999]. Three out of five microinjected cells arrested in prometaphase for 3–10 h with several monooriented chromosomes close to the spindle poles (Fig. 8c, arrows); all five control cells microinjected with normal rabbit IgG showed normal mitosis. Among five CF-PAC1 cells microinjected early in prometaphase with anti-Nercc1 antibody, only one showed a similar mitotic defect, whereas the other four proceeded through mitosis normally.

The occurrence of similar mitotic abnormalities in PtK2 cells microinjected with two independently prepared anti-Nercc1 antibodies, raised against different Nercc1 peptides, and the absence of these phenotypes in response to nonimmune IgG, together with the occurrence of similar abnormalities in CF-PAC1 cells indicate strongly that interference with Nercc1 is the basis for these phenotypes. In summary, microinjection of anti-Nercc1[C-terminal peptide] C1 IgG and anti-Nercc1[catalytic domain peptide] E2 IgG results in the frequent occurrence of abnormal spindle dynamics and abnormal chromosomal congression and segregation, indicating that Nercc1 ordinarily participates in the regulation of these processes.

Discussion

Nercc1 kinase is a novel mitotic regulator

The Nercc1 kinase is a new member of the mammalian branch of the NIMA-like kinases. Perhaps the most consequential aspect of this work is the clear-cut demonstration that Nercc1, like NIMA, is a pleiotropic regulator of mitotic progression, participating in the control of spindle dynamics and chromosome separation. Among the previously characterized mammalian kinases with NIMA-like catalytic domains, only Nek2 has been shown to be clearly involved in any aspect of mitotic regulation, that is, centrosome separation. Although
mammalian cells were shown to respond to NIMA overexpression with chromatin condensation and are blocked in G2 by different nonfunctional versions of NIMA, it has not previously been clear whether functional homologs of NIMA exist in metazoans (O’Connell et al. 1994; Lu and Hunter 1995). None of the described mammalian enzymes with protein kinase domains homologous to NIMA (Nek1–7; Nigg 2001) appears to have functions similar to that of Aspergillus NIMA, that is, control of the G2/M transition (Osmani et al. 1991a; Lu et al. 1991b). envelope organization during and after mitosis (Osmani and Means 1994), chromosome condensation (Osmani et al. 1994; O’Connell et al. 1994), and spindle and nuclear envelope organization during and after mitosis (Osmani et al. 1991b).

We show that microinjection of interfering anti-Nercc1 IgGs into PtK2 and CF-PAC1 cells in prophase induces several types of abnormalities. Many PtK2 cells microinjected with anti-Nercc1 antibody arrest in prometaphase without an observable spindle. Other cells show a bipolar spindle, but defective chromosome orientation and segregation result in aneuploidy after cytokinesis. In a few cells, spindle separation is markedly defective, but cytokinesis proceeds, resulting in the sequestration of the entire spindle and 4N DNA content into only one of the daughter cells. CF-PAC1 cells microinjected in prophase with anti-Nercc1 antibody also show abnormalities in chromosome attachment, but in contrast to PtK2 cells, CF-PAC1 cells microinjected with anti-Nercc1 antibody in prophase arrest in prometaphase with monooriented chromosomes. Inasmuch as these IgG injections were made after chromosome condensation, we have no direct evidence concerning the participation of Nercc1 in this or other earlier steps in G2/M progression. Nevertheless, overexpression of nuclear forms of Nercc1, especially Nercc1(1–391) and Nercc1(1–308), produce markedly abnormal morphologies of the interphase nucleus [Fig. 7c], and these Nercc1 variants collect at peripheral sites near the tips of lobed nuclei. The basis for this appearance is unknown, and its relationship to NIMA-induced chromosomal condensation in mammalian cells will probably require a side-by-side comparison.

The molecular basis for the abnormalities observed when anti-Nercc1 IgG is introduced into mitotic cells is not known. The phenotypes suggest possible roles for Nercc1 in spindle assembly and dynamics as well as in chromosome attachment, alignment at the spindle midzone, and segregation. The occurrence of arrest in prometaphase suggests activation of the spindle checkpoint, whereas the instances in which mitotic progression of antibody-microinjected PtK2 cells continues through anaphase and cytokinesis despite chromosome misalignment and maldistribution indicate that the metaphase-anaphase and mitotic exit checkpoints have been bypassed. A possible explanation is that Nercc1 regulates one or more components of the cell cycle machinery involved in both spindle formation and checkpoint action. Motor proteins have this characteristic and are thus good candidates to be targets for Nercc1 (Brunet and Vennos 2001).

In addition to possible functional homology, Nercc1 has a striking number of structural similarities with NIMA aside from their homologous catalytic domains. Nercc1 has a C-terminal tail containing an NLS, a substantial set of S/TP sites, several PEST regions, and a coiled coil that serves as a homooligomerization motif. In addition, Nercc1 is an in vitro substrate of p34Cdc2 and shows increased activity during mitosis. Nevertheless, Nercc1 also displays some remarkable differences from NIMA, the most conspicuous being the presence of a domain homologous to RCC1 and the ability to bind Ran. Also, whereas NIMA is localized to the nucleus and its levels change greatly during cell cycle progression, Nercc1 is cytoplasmic in interphase and its protein levels do not appear to change during the cell cycle. Thus, it is highly unlikely that Nercc1 is an exact mammalian equivalent of NIMA. More likely, the numerous functions of NIMA characterized in Aspergillus have been parsed out among a number of kinases in mammalian cells, which differ in regulation and perhaps specificity; Nercc1 certainly appears to be one of these.

In this regard, the association of Nercc1 with Nek6 and Nek7 (data not shown) requires comment. The tight association of Nercc1 with these two kinases suggests that they represent elements in a signal transduction cascade or a linked effector unit or possibly both. Apart from the role of Nek6/7 as a candidate p70 S6 kinase (Belham et al. 2001), the cellular responses to Nek6/7 are as yet uncharacterized. When extracted after transient expression in normally cycling cells, Nek 6/7 shows considerable basal activity, in contrast to Nercc1. We showed previously that the spontaneous activity of recombinant Nek6 is reversed by protein phosphatase treatment in vitro. Preliminary experiments indicate that Nercc1 is capable of phosphorylating and activating Nek7 in vitro. It is therefore conceivable that some or all of the phenotypes observed on microinjection of interfering anti Nercc1 IgG are caused by interference with Nek6/7 function.

Regulation of Nercc1 kinase function

Our analysis thus far of the regulation of Nercc1 activity indicates that in vitro, the recombinant Nercc1 polypeptide is a homooligomer that is maintained in an inactive state by occlusion of its catalytic domain by the succeeding RCC1-like domain; this inhibition may occur either in cis or in trans within the dimer. The monomeric variant Nercc1(1–739) also has low activity and is almost entirely resistant to autoactivation in vitro by Mg2+ plus ATP. The further deletion of the RCC1-like domain produces Nercc1(1–391); this variant also shows low basal activity but is capable of significantly greater autoactivation in the presence of Mg2+-ATP than is Nercc1(1–739), although at rates far below wild type. In contrast, a deletion of the RCC1-like domain that leaves the Nercc1 dimer intact, that is, Nercc1(Δ347–732), shows very high basal activity, either through spontaneous activation in vivo, or essentially instantaneous autoactivation in vitro upon addition of Mg2+-ATP. The very strong dependence
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Figure a shows a series of images labeled A, B, and C, each with different time points indicated. The images appear to be micrographs of cells, with varying stages of development or treatment.

Figure b displays two panels labeled 'Control' and 'anti-Nercc'. The images show fluorescent labeling, suggesting a comparison between control and treated conditions.

Figure c contains a set of images with time points labeled from 0 to 60, possibly indicating a time course of a cellular process or response to a treatment.
Nercc1 kinase regulates mitotic progression

The rate of Nercc1 autoactivation in vitro, although the latter conclusion is as yet tentative given the technical difficulties of those experiments. In addition, we have not yet determined whether MPF-catalyzed Nercc1 phosphorylation alters Nercc1 binding to its known interactors, Ran and Nek6/7.

As to the interactions of Ran with Nercc1, our in vitro assays do not as yet show any major impact of Ran on the rate of Nercc1 autoactivation. Nevertheless, we are intrigued by the specific and relatively tight binding of Ran to the Nercc1 catalytic domain in vitro and in vivo, as well as the important role of the Nercc1 RCC1-like domain as a negative regulatory element. The Nercc1 RCC1-like domain also binds Ran specifically, although less tightly than the Nercc1 catalytic domain.

Figure 8. The effect of microinjection of anti-Nercc1 IgG into PtK2 and CF-PAC1 cells. (a) PtK2 cells were microinjected with affinity-purified anti-Nercc1 [C1] IgG (2.5 mg/mL; typically, the volume of microinjected material comprised ∼10% of cell volume) in prophase. Representative phase contrast images from time-lapse recordings are shown. Recorded cells were fixed 3 min after the last image in the sequence shown and stained with Hoechst 33342 DNA stain [lowest image in each panel]. Time in minutes is shown in the lower right-hand corner of the images, with acquisition of the last frame before the onset of anaphase serving as time 0 (a, b). The first image in panel C was taken 2 min after the nuclear envelope breakdown. Bar, 10 μm. (A) Anaphase A starts and proceeds normally, but the poles do not separate. Chromosomes remain trapped in the cytokinetic furrow and a bridge of DNA remains between the daughter cells. (B) An example of an extreme case of the absence of anaphase B phenotype. After moving the chromosomes apart in anaphase A, the substantial further separation typical of anaphase B does not occur, and a cytokinetic furrow separates the daughter cells into one containing all chromosomes and a cytoplast. Hoechst staining confirms the absence of DNA in the right cell. (C) After the nuclear envelope breakdown, the cell fails to form a mitotic spindle, or the spindle collapses soon after formation. Mitotic progression stops in prometaphase. See b. (b) PtK2 cells were microinjected with normal IgG [Control] or anti-Nercc1 [C1] IgG [anti-Nercc1] in prophase. Cells were fixed and stained with Hoechst 33342 DNA (blue), and anti-tubulin antibody (red). Control cells were fixed at metaphase, anti-Nercc1 injected cells failed to enter a normal metaphase, and were fixed at t = 120 min after microinjection. (c) CF-PAC1 cells were microinjected with affinity-purified anti-Nercc1 [C1] IgG in prophase. Representative phase contrast images from a time-lapse recording are shown. The recorded cell was fixed 3 min after the last image in the sequence shown and stained with Hoechst 33342 DNA stain and anti-tubulin [lowest image]. Time after antibody microinjection in minutes is shown in the lower right-hand corner of the images [t = 0 taken 2 min after nuclear envelope breakdown]. Arrows show monooriented chromosomes. Bar, 10 μm.
through the focal generation of Ran-GTP in the vicinity of the chromosome-bound GEF, this generation of Ran-GTP is critical in directing the construction of the mitotic spindle as well as in the reassembly of the nuclear envelope after telophase [Azuma et al. 1999; Hetzer et al. 2000]. The gradient of Ran-GTP formed around the chromosomes by the action of RCC1 [Kalab et al. 2002] controls microtubule polymerization and motor activity and thus promotes aster formation and spindle assembly around chromosomes [for reviews, see Kahana and Cleveland 1999; Heald and Weis 2000]. The mechanism through which Ran controls microtubule polymerization is similar to that of nucleocytoplasmic transport: binding of Ran-GTP to importins/freeves and activates different aster-promoting activities that are inactivated by importin binding [for reviews, see Dasso 2001; Walczak 2001]. How Ran controls other steps involved in aster formation and spindle assembly [e.g., the activity of motor proteins like Eg5] and nuclear envelope formation, or whether Ran controls other aspects of mitosis is presently unknown [for a review on Ran and cell cycle control, see Moore 2001]. Based on the phenotypes elicited by microinjection of anti-Nercc1 IgG into mitotic cells, and the ability of Nercc1 to bind Ran, we propose that the activation of Nercc1 during mitosis [which is ultimately caused by the intramolecular transphosphorylation of the Nercc1 activation loop] and/or the execution of its cellular functions will prove to involve the displacement of Ran-GDP in the vicinity of the chromosomes; this implies that although diffusely distributed, Nercc1 is activated and/or acts primarily in the vicinity of the chromosomes. Certainly the control of protein kinase localization and activation by small GTPases, as first shown for Ras and Raf, is now recognized as a general regulatory mechanism in the regulation of protein kinase function. Reconstituting in vitro the apparatus underlying the cell cycle-dependent regulation of Nercc1 activity will be a substantial challenge.

In conclusion, Nercc1 is a mammalian NIMA-like kinase that acts as a regulator of spindle function and chromosome segregation. Nercc1 is tightly bound to another NIMA-like protein kinase, Nek6/7; the contribution of each kinase to the function of the other is as yet unknown. Nercc1 is activated in mitosis by disinhibition and intramolecular autophosphorylation. It is a target for cyclinB/Cdc2 and binds specifically to the Ran GTPase through its catalytic and novel RCC1-like domain. The significance of Cdc2 phosphorylation and Ran binding in Nercc1 activation, targeting, and function, as well as the immediate cellular substrates of Nercc1 await discovery.

Materials and methods

Protein sequencing, cDNA cloning and manipulation

The cloning of the Nercc1 cDNA and the construction of the different DNA plasmids used in this work is described in the Supplemental Material [available online at http://www.genesdev.org].

Cell culture and transfection

See Supplemental Material [available online at http://www.genesdev.org].

Cell lysis, immunoprecipitation, in vitro binding, and immunoblotting

Cells were rinsed with PBS, flash-frozen in liquid nitrogen, and stored at −70°C. Cell lysis used a buffer containing: 50 mM Tris [pH 7.4], 100 mM NaCl, 1 mM DTT, 1 mM EDTA, 1 mM EGTA, 10 mM β-glycerophosphate, 2 mM Na3VO4, 25 mM calyculin A, 1% TX100, plus protease inhibitors [EDTA-free tablets; Roche]. Protein concentration was determined by the Bradford reagent [BioRad]. Immunoprecipitations were carried out with the indicated antibodies prebound to protein A/G-agarose [Santa Cruz], and washed in the lysis buffer containing 0.5 M LiCl.

Gel filtration

The 293 cells were lysed in 1% TX-100 lysis buffer, ultracentrifuged at 100,000g for 40 min, and loaded to a precalibrated HiPrep 16/60 Sephacryl S-300 High Resolution column [Pharmacia]. Gel filtration was carried out in 50 mM Tris [pH 7.5], 150 mM NaCl, 1 mM DTT buffer at 0.5 mL/min.

Protein kinase assays

Protein kinase assays were carried out after immunoprecipitation; recombinant Nercc1 was isolated using anti-Flag antibodies prebound to protein A/G-agarose beads. Endogenous Nercc1 was immunopurified using N1 antibodies prebound to protein A/G-agarose beads. Complexes were washed sequentially with lysis buffer and phosphorylation buffer [50 mM MOPS at pH 7.4, 1 mM DTT, 1 mM EGTA, 5 mM MgCl2, 10 mM β-glycerophosphate, 25 mM calyculin A]. Nercc1 autoactivation was carried out by incubation of immobilized Nercc1 in phosphorylation buffer plus 100 µM ATP at 25°C for the indicated times. Activation was terminated by washing the immobilized Nercc1 in phosphorylation buffer, and the protein activity achieved was assayed by incubation at 30°C in phosphorylation buffer supplemented with either 10 µM or 100 µM [γ-32P]ATP, with an exogenous substrate, usually histone H3, as indicated. Assays were stopped by addition of electrophoresis sample buffer and boiling, and the proteins were resolved by SDS-PAGE. 32P incorporation was measured with a PhosphorImager system or by liquid scintillation counting, as indicated.

Immunocytochemistry

Cells grown on coverslips were rinsed with PBS, fixed in methanol at −20°C for 15 min, rinsed twice with PBS, and incubated
for 30 min at room temperature with the appropriate dilution of primary antibody in PBS. To visualize endogenous Nercc1, affinity-purified anti-Nercc1 peptide antibodies (N1 and C1) were used at 10 µg/mL. Microtubules were visualized with an αβ-tubulin-specific antibody. Coverslips were washed with PBS, and incubated with labeled secondary antibodies from corresponding species in appropriate combination: fluorescein or rhodamine X-conjugated donkey anti-rabbit, Cy2 or rhodamine X-conjugated donkey anti-mouse (each at 1:450). Incubation was terminated with a rinse in PBS, and the coverslips were mounted on a microscope slide. For immunodetection blocking, Nercc1 C1 or N1 antibody was incubated at 37°C for 30 min with a 15-fold molar excess of immunizing peptide. After centrifugation at 12,000g for 10 min, the mixture was used for immunoblotting or immunochemistry, as indicated.

Microinjections and time-lapse recordings

For real-time observation of the effects of recombinant Nercc1 expression, both wild-type and variant, on cellular morphology and behavior during one cell cycle, HeLa cells grown on 25-mm glass coverslips were transfected with pEGFP-C2 vector or this vector encoding GFP fusions with Nercc1 wild-type, Nercc1[K81M], or Nercc1[1–391] using Fugene [Roche Molecular Biochemicals]. Fugene-containing medium was removed after 12 h, and DMEM supplemented with 10% calf serum and penicillin–streptomycin was added after rinsing. The percentage of transfected cells undergoing division within 24 h was monitored. Using this transfection procedure, cells transfected with empty pEGFP-C2 underwent division at a frequency similar to nontransfected cells. The coverslips (in a Sykes-Moore chamber) were mounted on a microscope stage prewarmed to 37°C; a region with the highest density of transfected cells (GFP-positive) was selected for observation, and phase contrast images were acquired using a 40× 1.0 NA objective every 10 min for 25 h. Light was kept to a minimum during image acquisitions and shuttered between acquisitions.

To observe the effect of anti-Nercc1 antibodies on mitosis, PtK2 cells were grown to subconfluency on 25-mm round glass coverslips placed inside 35-mm cell culture dishes. A cell in prophase was found using phase contrast optics and microinjected in the period between the nuclear disassembly and nuclear envelope breakdown with either 2.5 or 10 µg/mL normal rabbit IgG [Jackson Immunoresearch] for control experiments, or with 2.5 mg/mL rabbit anti-Nercc1 C- or N-terminus or kinase domain antibody with 0.5 µg/mL rhodamine-labeled dextran 3000 [Molecular Probes]. Typically, the volume of microinjected material comprised ~10% of cell volume. Immediately after microinjection, the cover slip with microinjected cells was placed in a Sykes-Moore chamber [Belco Glass] filled with bicarbonate-free DMEM supplemented with 10% fetal Calf Serum. The chamber was transferred onto the stage of a Zeiss Axiocamt 100M microscope maintained at 37°C with the aid of an Air-Therm heater controller [World Precision Instruments] and a custom-made microscope incubator. Microinjected cells were found by rhodamine fluorescence using a maximum possible density neutral density filter (typically, ND 1.0, Chroma Technology). Phase contrast images were acquired every 20 or 30 sec with a Hamamatsu Orca-100 CCD camera driven by Metamorph 4.0 [Universal Imaging Corporation], we used a 100× 1.4 NA objective and light was kept to a minimum during image acquisitions and shuttered between acquisitions.

Acknowledgments

We thank Y. Yin for Northern blots, Y. Lin for the mouse tissue protein membranes, E. Casacuberta for help with ORF prediction, and J. Prendable for help with the manuscript. We are grateful to A. Khodjakov and C. Rieder for help interpreting the anti-Nercc1 microinjection experiments. We are also grateful to I. Macara for the different Ran variant cDNAs and RCC1 cDNA, and to J. Maller for purified MPF. This work was supported in part by DK17776.

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Genes Dev. 2002, 16: Access the most recent version at doi:10.1101/gad.972202

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