

A nucleoporin is required for induction of Ca^{2+} spiking in legume nodule development and essential for rhizobial and fungal symbiosis

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Nuclear-cytoplasmic partitioning and traffic between cytoplasmic and nuclear compartments are fundamental processes in eukaryotic cells. Nuclear pore complexes mediate transport of proteins, RNAs and ribonucleoprotein particles in and out of the nucleus. Here we present positional cloning of a plant nucleoporin gene, *Nup133*, essential for a symbiotic signal transduction pathway shared by *Rhizobium* bacteria and mycorrhizal fungi. Mutation of *Nup133* results in a temperature sensitive nodulation deficient phenotype and absence of mycorrhizal colonization. Root nodules developing with reduced frequency at permissive temperatures are ineffective and electron microscopy show that *Rhizobium* bacteria are not released from infection threads. Measurement of ion fluxes using a calcium-sensitive dye show that *Nup133* is required for the Ca^{2+} spiking normally detectable within minutes after application of purified rhizobial Nod-factor signal molecules to root hairs. Localization of NUP133 in the nuclear envelope of root cells and root hair cells shown with enhanced yellow fluorescent protein fusion proteins suggests a novel role for NUP133 nucleoporins in a rapid nuclear-cytoplasmic communication after host-plant recognition of symbiotic microbes. Our results identify a component of an intriguing signal process requiring interaction at the cell plasma membrane and at intracellular nuclear and plastid organelle-membranes to induce a second messenger.

legume symbiosis | nucleoporin | nuclear pore | plant-microbe interaction

Two of the nutrients plants require in high amounts for optimal growth are nitrogen and phosphate. Plant roots mine the soil for nutrients, and 80–90% of all land plants have symbiotic interaction with arbuscular mycorrhizal fungi that increase the root surface and supply plants with phosphate (1, 2). Extending the repertoire for nutrient acquisition, legumes also enter into symbiosis with nitrogen fixing bacteria, collectively called rhizobia (3). Nitrogen fixed in the root nodule organs that house the nitrogen-fixing bacteria is estimated to be equal to the amount used around the world in chemical fertilizers. Genetic studies in the model legume *Lotus japonicus* have, in addition to *Nup133*, defined six loci (*SymRK*, *Castor*, *Pollux*, *Sym15*, *Sym24*, and *Sym30*) required for both mycorrhizal colonization and rhizobial induced root nodule development (4, 5). The fungal symbiotic signal molecule triggering this signal transduction pathway is unknown. In contrast, both the rhizobial Nod-factors and corresponding plant receptor kinases required for perception have been identified. In *Lotus*, NFR1 and NFR5 are predicted receptor kinases that operate upstream of the common pathway. Presumably they funnel Nod-factor signaling into the signal transduction pathway shared with mycorrhizal fungi (6, 7). Nod-factor perception is a key feature in rhizobial symbiosis determining host-bacteria recognition, initiation of infection thread formation, and root nodule inception. Purified Nod-factors trigger a variety of responses in roots, such as cell divisions forming

nodule primordia, activation of gene expression, root hair deformation, and calcium oscillations in root hair cells (8, 9). Functional NFR1/NFR5 receptor kinases are required for these responses, whereas common pathway mutants usually display an attenuated response to Nod-factor. Several lines of evidence suggested that calcium acts as secondary messenger, and the analysis of mutants of *Lotus* and/or *Medicago* indicates that calcium spiking is a component of the common pathway. Analysis of mutants has shown that a LRR protein kinase (SYMRRK/NORK) and a cation channel(s) (CASTOR/POLLUX/DMI1) are required for the induction of calcium spiking. A predicted calcium calmodulin protein kinase (DMI3) probably integrates the calcium fluctuations to activate induction of downstream genes (10–15). Interestingly, CASTOR and POLLUX proteins were localized to root cell plastids, implicating ion signaling through plastids in the early signal transduction after signal perception (14). Characterization of nodulation deficient *nup133* mutants reported here add a unique component to this pathway and focus attention on the role of nuclear associated calcium spiking in Nod-factor mediated signaling.

Materials and Methods

Plant Material. Isolation of *nup133-1*, *nup133-2* mutants previously called *sym3-1* and *sym3-2* were described in ref. 16, and *nup133-3* and *nup133-4* (previously called *sym3-3* and *sym45*) were also isolated in the ecotype Gifu B-129 background. Plants were grown with or without *M. loti* strain NZP2235, TONO, and R7A. The root hair curling procedure has been described (6).

Electrophysiology. Seedlings of *L. japonicus* were germinated, mounted, and microinjected with Oregon Green-488 BAPTA-1 (Molecular Probes) essentially as described (9). Fluorescence was imaged by using a Nikon TE2000 inverted microscope coupled to a Hamamatsu Photonics digital charge-coupled device (CCD) camera. The excitation wavelength of 488 nm with an 11-nm bandpass was selected by using an Optoscan Monochromator (Cairn, Faversham, Kent, U.K.), and an emission filter of 545 (± 15) nm was used. Images covering the protruding part of the root hair, including the entire nuclear region, were collected every 5 s with a 200-ms exposure using METAFLUOR software, and derivative traces were generated by using Microsoft EXCEL. The data presented were transformed to first derivative traces (in arbitrary units) as de-

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Abbreviation: eYFP, enhanced yellow fluorescent protein.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession nos. AJ890251, AJ890252, and AP008949–AP008952).

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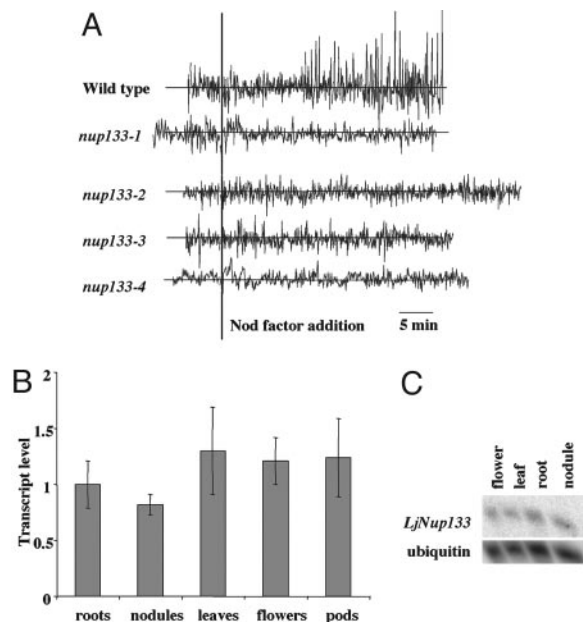


Fig. 2. Calcium oscillations in root hairs in response to purified Nod-factor. (A) Each trace is from a single root hair using seedlings of wild-type *L. japonicus* or mutants carrying the *nup133-1*, *nup133-2*, *nup133-3* or *nup133-4* alleles. Seven, four, six, and three root hairs were analyzed for each allele, respectively. An additional 10 analyses were done with *nup133-3* grown at 10°C and assayed at 15°C, but the traces were similar to that shown. Root hairs were injected with the calcium-sensitive dye Oregon green-488 BAPTA-1 and, after ≈ 20 min, Nod factor was added at 10^{-8} M. The data are graphed showing typical traces of the first derivatives of fluorescence intensity (in arbitrary units) between 5 s sequential time points. There was no observed calcium spiking in any of the mutants under the conditions tested. (B and C) Transcript level of *Nup133* in wild-type plants. (B) Quantitative RT-PCR results are shown as fold increase compared to roots. Error bars represent a 95% confidence interval. (C) Northern hybridization showing similar expression levels of *Nup133* in different *Lotus* tissues

cells normally assayed by us and others (8, 9, 15, 17) for this phenotype.

Taken together, the phenotypic characterization of a temperature-dependent nonnodulation phenotype, the impaired mycorrhizal colonization (4, 5), the lack of normal Nod-factor induced alkalization in the root hair space, and the lack of Nod-factor induced calcium spiking in developing root hairs, lead us to conclude that *NUP133* is part of the common pathway required to mount a response to signals from rhizobial and mycorrhizal symbionts.

Positional Cloning of *Nup133*. To characterize the gene molecularly, *Nup133* was isolated by using a positional cloning approach. On the genetic map of *Lotus*, the *Nup133* locus is located on the short arm of chromosome I (20). Subsequent fine mapping in an F_2 population and genotyping of 822 mutant plants identified markers delimiting *Nup133* to a 0.06 cM region of 22 kb (Fig. 5, which is published as supporting information on the PNAS web site). Located within this region are four genes predicted to encode a pentatricopeptide repeat-containing protein, a chlorophyll a/b binding protein, a Ca^{2+} transporting ATPase, and a homologue of an *Arabidopsis* expressed protein, respectively (Fig. 5). Considering the suggested role for calcium as second messenger, the search for mutant alleles was initially focused on sequencing the Ca^{2+} transporting ATPase gene. Contrary to our expectations, this gene was not mutated. Subsequent sequencing of gene regions corresponding to the homologue of the *Arabidopsis* expressed protein in *nup133-1*, *nup133-2*, *nup133-3*, and *nup133-4* identified the *Nup133* gene.

Short deletions shifting the reading frame, leading to premature stop codons, were found in *nup133-1*, *nup133-2*, and *nup133-4* (Table 2). A retrotransposon is inserted in *nup133-3*. All four mutant alleles were complemented by the cloned *Nup133* using *Agrobacterium rhizogenes* to generate transgenic roots (19). The wild-type gene, including a 1.8-kb promoter region and a 3.6-kb 3' region was introduced into *nup133-1*, *nup133-2*, *nup133-3*, and *nup133-4* mutant plants via *A. rhizogenes* and the nodulation phenotype scored after inoculation with strain NZP2235 of *M. loti*. All mutants were complemented with high efficiency for nodulation (Fig. 1E and Table 3, which is published as supporting information on the PNAS web site). Complementation for mycorrhizal colonization was tested and obtained in the *nup133-1* mutant. Complementation was scored as normal colonization and abundant arbuscule formation in transgenic roots.

Sequencing of full-length cDNAs isolated from a *Lotus* leaf library determined the transcription start site at least 62 bp upstream of the start codon and a 3' untranslated region of 210 nucleotides (Fig. 5). Alignment of genomic and cDNA sequences defined eight exons in *Nup133* (Fig. 5). The *Lotus Nup133* cDNA encodes a conceptual protein of 1,309 aa, corresponding to 146.5 kDa. Southern hybridization indicates that *NUP133* is encoded by a single gene in both the small genome of *Lotus japonicus* and the large genome of pea (data not shown). In the fully sequenced *Arabidopsis* and rice genomes, *Nup133* homologs are single functionally uncharacterized genes in each species. The *Arabidopsis* predicted protein At2g05120 is 54% identical, and the predicted rice protein (AAN52748) is 47% identical to *NUP133*. Less similar, with 20% global identity, are yeast and human *Nup133* nucleoporins (Fig. 6, which is published as supporting information on the PNAS web site). Three-way alignment of human, yeast, and *Lotus* proteins identifies a set of 59 conserved amino acids positioned along the length of the proteins suggesting common origin of corresponding genes. This finding is supported by conservation of the two first intron positions among 25 introns of the human gene and eight introns of plant *Nup133* genes. The *Lotus NUP133* protein was identified as a *Nup133* nucleoporin by PSI-BLAST. After two iterations, the *Lotus NUP133* amino acid sequence aligned to the entire length of *Nup133* proteins from mouse and human with *E* values of 0 and overall identities of $\approx 20\%$. The N-terminal domain of human *Nup133* consists of an α/β domain with a seven bladed β -propeller fold, and the C-terminal domain was predicted to be all α -helical (21). Secondary structure prediction and fold recognition on *Lotus NUP133* suggest a similar overall structure and recognize the β -propeller structure of human *Nup133* within the first 600 N-terminal residues of *Lotus NUP133*. From secondary structure predictions and an alignment with human *Nup133*, the N-terminal α/β domain and C-terminal α helical domain of *Lotus NUP133* were estimated to localize between residues 50–535 and 555–1309, respectively. The positions of β -propeller blades suggested by secondary structure prediction are shown on the alignment of N-terminal domains of *Lotus NUP133* and human *Nup133* proteins in Fig. 3.

Expression of *Nup133* in Different Plant Organs. The symbiotic mutant phenotype suggests a function for *Nup133* in root tissues. To test this prediction, expression of *Nup133* in different plant organs was determined by Northern and quantitative RT-PCR analysis. Results of both analyses demonstrate that *Nup133* transcripts are present in all organs tested (Fig. 2B and C). No significant induction was detected in roots several days after inoculation with *M. loti* (data not shown). The observed expression in all organs tested and the similarity to a nucleoporin would predict a general function for *Nup133* and a mutant phenotype affecting overall plant development. For example, mutation of the *PAUSED* gene (encoding an homolog of nuclear export receptor for tRNA) in *Arabidopsis* affected shoot and apical meristem growth, leaf development, and lateral root formation (22). No such general effect was observed

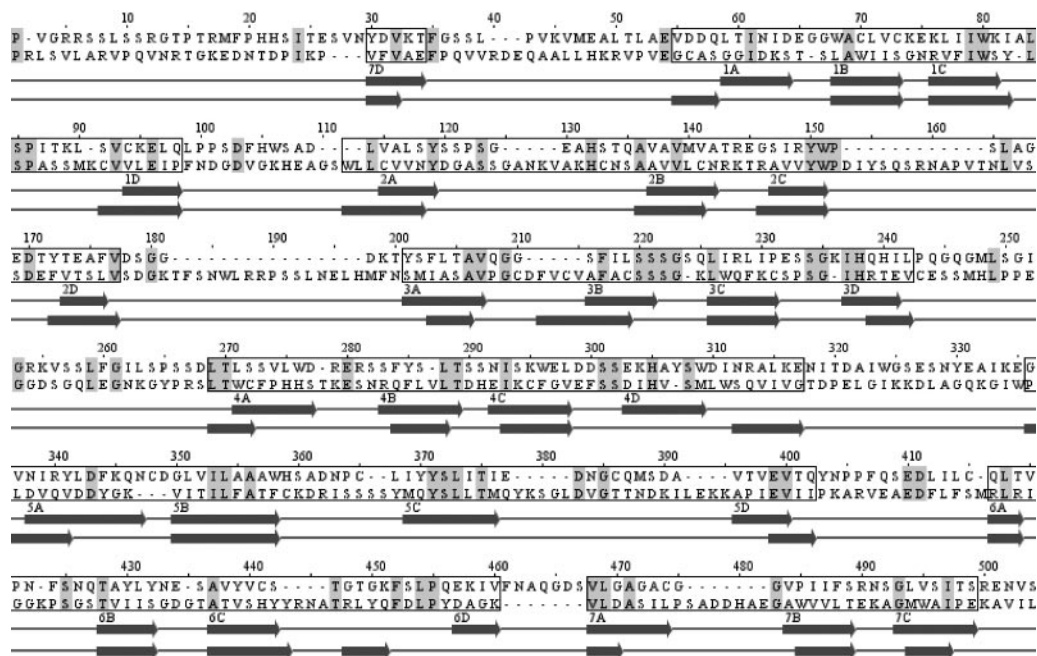


Fig. 3. Alignment of the N-terminal 50–535 residues of *Lotus* NUP133 to human Nup133 (top sequence). Amino acid residues marked in gray are identical. Propeller blades are boxed and β strands named from 1 (A–D) to 7 (A–D) after position in the human Nup133 propeller are shown below. Human Nup133 strand annotations are depicted above the *Lotus* NUP133 strand predictions. Although *Lotus* NUP133 has quite a few insertions compared to human Nup133, the N-terminal domain of *Lotus* NUP133 is predicted to adopt a seven-bladed β -propeller fold, similar to the human Nup133 propeller.

with the *nup133* mutants, but generally the four mutant lines have a lower number of seeds in mature pods than wild type.

Localization of NUP133 Protein in the Nuclear Rim. Yeast and mammalian nuclear pore complexes contain ≈ 30 proteins identified mainly by proteomics. Although nuclear pores serve the same purpose in nuclear–cytoplasmic trafficking, global conservation of nucleoporin sequences is within a range of 10–20% identities, which is surprisingly low. Corroborating this observation, a search of the *Arabidopsis* genome using human nucleoporins and nuclear pore associated protein sequences, has so far identified only four *Arabidopsis* encoded proteins with $\approx 20\%$ identities and 30–40% similarity to nucleoporins (23). Neglecting this lack of conservation between organisms, we attempted functional characterization of NUP133 protein in yeast. A GFP–NUP133 N-terminal fusion expressed from a galactose-inducible promoter was transformed into a temperature-sensitive yeast strain deleted for the *NUP133* gene. Complementation was not achieved, and the GFP fusion protein was not localized to the nuclear rim as expected for a nucleoporin (Fig. 7, which is published as supporting information on the PNAS web site). These results suggest that *Lotus* NUP133 and yeast Nup133p are too diverse for functional complementation. In contrast, an eYFP–NUP133 N-terminal fusion protein expressed in transgenic *Lotus* roots complemented *nup133-1* mutants, and the fusion protein was clearly localized to the nuclear rim in root cells (Fig. 4 D and G–L) as well as nuclear rim in root hair cells (Fig. 4 E and F). In Fig. 4 E–K, the nuclear rim localization is illustrated by corresponding sets of light microscopy and confocal images. Analysis of confocal stack images of consecutive sections of nuclei shows nuclear rim localization of the fusion protein in the complete nuclear sphere. In accordance with the NUP133 similarity to nucleoporins, the punctuate appearance of eYFP–NUP133 fluorescence (Fig. 4 H, I, and K) is comparable to the punctuate localization characteristic for proteins residing in nuclear pore complexes of yeast and mammalian cells. In control experiments, expression of eYFP alone resulted in an even distribution of fluorescent protein within nuclei (Fig. 4 A–C).

Discussion

The collective evidence obtained by genetic and physical mapping, sequencing of mutant alleles, and successful complementation

unequivocally identified the nucleoporin *Nup133* gene. The presence of only one gene copy in *Lotus*, pea, and the fully sequenced *Arabidopsis* and rice genomes, together with conservation of two intron positions, suggest that plant *Nup133* genes are closest ho-

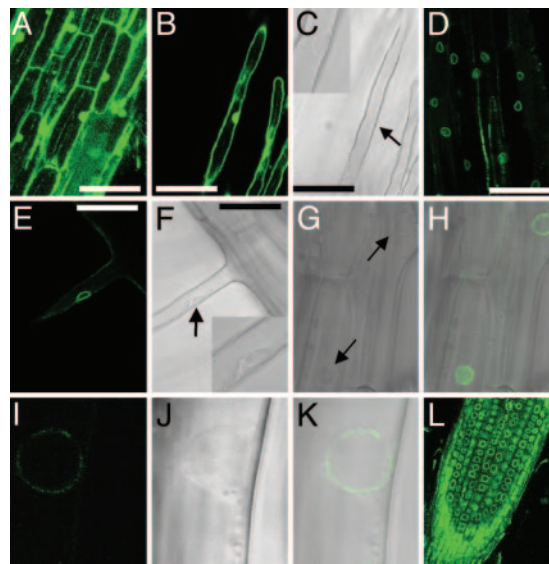


Fig. 4. Subcellular localization of eYFP–NUP133 fusion protein in the nuclear rim. (A and B) Fluorescent image of a control eYFP localization in *nup133-1*. (C) Light microscopy of B with nucleus marked by arrow and highlighted in insert. (D, E, H, I, K, and L) Fluorescent images of eYFP–NUP133 fusion protein showing nuclear rim localization in *nup133-1* mutant background. (E–K) Single cell localisation of eYFP–NUP133 in the nuclear rim. (E) Root hair cell showing nuclear rim localization of eYFP–NUP133. (F) Corresponding light microscopy with nucleus marked by arrow and highlighted in *Inset*. (H) Fluorescent overlay of G light microscopy with nuclei marked by arrows. Note the surface view of one nucleus showing punctate localization of eYFP–NUP133 fusion protein. (K) Overlay of corresponding fluorescent image of eYFP–NUP133 (I) and light microscopy image (J). Note the punctate eYFP–NUP133 localization. (L) Localization of eYFP–NUP133 in nuclear rims of cells in a transgenic root tip; note the cell files. (A and D) Epidermal cells. (B, C, E, and F) Root hair cells. (Scale bar: 50 μm .)

mologs of yeast and mammalian Nup133 genes. Further evidence from protein sequence conservation in *Lotus* NUP133, comparable to the general level of conservation in the NUP133 protein family and the overall α/β - α domain structure together with the predicted N-terminal seven-bladed β -propeller domain, support the identification of *Lotus* NUP133 as a member of the Nup133 nucleoporin family. Localization of eYFP fusion proteins at the nuclear rim makes it most likely that NUP133 and its homologs are components of the nuclear pore complex in plants. The punctuate localization of eYFP-NUP133 fluorescence, characteristic for nuclear pore proteins, supports this interpretation (Fig. 4 H, I, and K).

In yeast, Nup133p is a constituent of the Nup84p complex and in mammals the corresponding protein is a component of the Nup107-160 complex residing in the nuclear basket structure of the nuclear pore (24–27). Studies in yeast and HeLa cells suggest a function in export of mRNA, nuclear pore assembly and distribution (18, 28). Such a general function for the *Lotus* NUP133 nucleoporin is difficult to reconcile with the predominantly symbiotic phenotype observed. Interestingly, *Drosophila* has one example of cell-specific developmental effects in a nucleoporin mutant that is also affected in the response to fungal and bacterial pathogens. Mutation of *members only*, the gene encoding a nucleoporin homologous to mammalian Nup88, leaves Dorsal and Dif transcription factors in the cytoplasm upon pathogen infection, and innate immune response is compromised in *Drosophila* larvae (29). In *Arabidopsis*, recently published results show that a putative nucleoporin 96 protein is required for both basal resistance against bacterial pathogens and for R-gene-mediated pathogen resistance (30). These parallels are tantalizing and might have implications for the emerging comparative studies of IL-1/Toll receptor-mediated innate immune response in mammals/flies, inducible disease resistance in plants, and symbiotic interactions (31). A general role in plant-microbe interaction may also explain the absence of widespread developmental changes in *Lotus nup133* mutants. In yeast, Nup133p is involved in controlling the RanGTPase gradient across the nuclear envelope that, in turn, regulates import processes and nuclear pore assembly (32). Deletion of *NUP133* leads to reduced binding of the RanGTPase nuclear carrier Ntf2p protein to nuclear pore complexes. In light of these results, RanGTPase interaction could be the key to understanding NUP133 function during symbiosis and a target to take into consideration in pharmacological studies using agonist and antagonists to study early signal transduction downstream of Nod-factor perception (33). An alternative, more speculative, possibility is a NUP133 involvement in nuclear

calcium oscillations (34). Atomic force microscopy of *Xenopus* nuclear pores showed that the distal ring of the nuclear basket opens and closes in a calcium-dependent fashion and was suggested to serve as a calcium-sensitive iris-like diaphragm (35). Cryoelectronic tomography studies of *Xenopus* nuclear pore architecture also revealed an interesting temperature-dependent correlation between plugging of nuclear pores and nucleocytoplasmic transport (36).

Functional analysis of nuclear pores, nucleoporins, and nuclear trafficking in plants is in its infancy and we expect *Lotus nup133* mutants to provide inroads to these studies. Further impact is expected from characterization of the first *Arabidopsis* nucleoporin (Nup96) mutant isolated in a parallel effort (30). We find it very interesting that both the *Lotus* and the *Arabidopsis* nucleoporin genes identified are involved in plant-microbe interaction. This finding indicates an essential function for nuclear-cytoplasmic trafficking in plant responses to microbes and directs attention toward a role for a plant complex corresponding to the Nup107-160 subcomplex that, in mammalian nuclear pores, contains both Nup133 and Nup96 proteins. In symbiotic studies, one of the future challenges is to explain the NUP133-dependent calcium spiking mounted by wild-type plants 10 min after Nod-factor application and the intriguing paucity of Ca^{2+} spiking in *nup133* mutants carrying weaker alleles that unexpectedly initiate nodule development albeit at reduced frequency. Signaling involving both plastids through CASTOR/POLLUX and the nucleus through NUP133 makes the early symbiotic signal transduction pathway unusual. To our knowledge, a developmental signaling process dependent on participation of these two subcellular compartments in the first stages of signal amplification has not previously been found in plants. We expect that further studies of this pathway will provide insights into eukaryotic processes coordinating nuclear and organellar activities as well as a better understanding of plant-microbe interactions.

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