A re-calculation of criticality property of ²³¹Pa using new nuclear data

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An interesting paper published by an earlier Bhabha Atomic Research Centre (BARC) team in Current Science¹ and elsewhere²⁻⁷ dealt with data of criticality of minor actinide nuclides and their relevance to the long-livedfission-waste problem. of The purpose the present article is to communicate the changes in the results of criticality calculation of ²³¹Pa due to the use of more recent and improved neutron-nuclear interaction cross-section data. The new results of criticality for ²³¹Pa based on improved basic data differ significantly from those published earlier by the BARC team.

AN interesting study of criticality properties of minor actinide nuclides and their relevance to the long-livedfission-waste problem has been published in the period 1989–1991 in several journals and conferences¹⁻⁷ by Bhabha Atomic Research Centre (BARC). In this paper, we reassess the value of critical radius due to the use of improved input nuclear data of ²³¹Pa. It may be noted that in the case of ²³¹Pa, a value of 162.31 kg corresponding critical calculated to a radius of 13.61 cm $(\text{density} = 15.37 \text{ g/cm}^3)$ has been presented in earlier reports¹⁻⁷. Our finding which is drastically different from the earlier BARC result is that with the use of the new Japanese nuclear data8 even the infinite medium of ²³¹Pa is subcritical.

Using improved nuclear data and methods in simulation of nuclear systems is an important aspect of any serious nuclear programme. The generation and use of accurate nuclear data are considered fundamentally important, as accurate nuclear data are essential inputs to simulate nuclear interactions to obtain engineering parameters. New concepts can be studied with greater confidence if the scientific basis is sound. In the case of minor actinides, such as ²³¹Pa studied in this paper, the nuclear data status is very uncertain compared to the major isotopes of the uranium fuel cycle. The isotope ²³¹Pa occurs in nature due to decay of ²³⁵U actinide (4n + 3) series with a natural total world inventory of about 120 g. In thorium-fuelled reactors, production of ²³¹Pa takes place at several hundred grams per year per GWe of installed capacity⁷. The isotope ²³¹Pa has a long half-life of 32760 ± 110 years for alpha activity. It is produced primarily by fast neutrons through the (n, 2n)reaction in ²³²Th followed by beta decay. It constitutes a considerable source of radio-toxicity. In advanced concepts such as the Energy Amplifier (EA) proposed by Carlo Rubbia *et al.*⁹, production of ²³¹Pa takes place in significant quantities and a net stockpile of the order of 5 kg of ²³¹Pa will persist during the whole life time of an EA plant. Criticality experiment has not been conducted with pure individual actinide isotope such as ²³¹Pa thus far as earlier noted by Clayton¹⁰.

Comments on the earlier BARC calculations of criticality

The earlier study¹ employed the multigroup transport theory approach using the DTF-IV code¹¹ to solve the onedimensional Boltzmann neutron transport equation using the discrete ordinates method to give neutron density as a function of position, angle and energy inside a given medium. This code was used to compute the infinite medium multiplication factors and bare critical masses for spherical fast systems of actinides, using a 35 group cross-section set¹². Further, it may be noted from Srinivasan *et al.*⁷ that for ²³¹Pa a special BARC-assembled nuclear data file was used. Srinivasan et al.⁷ stated that the cross-section set for ²³¹Pa was derived by combining BARC-evaluated data in the 1 to 20 MeV energy region with the ²³³Pa data of the JENDL-2 file below 1 MeV. Srinivasan et al.⁷ also stated that the nuclear data of ²³¹Pa above 1 MeV was based¹³ on BARC calculations of theoretical nuclear models and a complete file for ²³¹Pa was assembled artificially by taking the data of ²³³Pa from JENDL-2 below 1 MeV. In our opinion, this approach of constructing a complete data file by merging data of one isotope above 1 MeV with that of another isotope below 1 MeV is fundamentally unacceptable. The neutron-nuclear interaction cross-sections change drastically as we go from one isotope to another of the same element. The discussions by the BARC team indicate that the earlier BARC study⁷ had an awareness that the calculated critical mass for ²³¹Pa could be very uncertain due to uncertain data used at that time. The discussion⁷ also

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CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999

indicates that the American National Standard Institute (ANSI)¹⁴ critical mass value is 750 kg for ²³¹Pa, 'based on three group diffusion theory calculations by Wu and Ruby¹⁵ and not so carefully evaluated data' (as quoted in Srinivasan

*et al.*⁷) and the Trombay criticality formula-based value is 463 kg. We note further that the ANSI critical mass is presented for ²³¹Pa by the American Nuclear Society (ANS)-based value reported by Wu and Ruby¹⁵. In the ANSI report¹⁴, it is mentioned as '750 kg (indicated)' expressing some caution that the ANS value may need a re-assessment as Wu and Ruby¹⁵ used a diffusion calculation and less experimental data were then available. The revision by BARC of the ANS result is not substantiated by our improved and rigorous investigations presented in this paper. Here, we are not questioning the use of DTF code or the way the code was used, but only the error due to the use of wrong nuclear data.

New calculations of criticality of ²³¹Pa

We used the JENDL-3.2 basic evaluated data file⁸ released in 1994 and available over the internet¹⁶ from the Nuclear Data Section of the International Atomic Energy Agency (IAEA), Vienna. This database provides a comprehensive neutron transport cross-section library for all neutroninduced reactions in 10^{-5} eV to 20 MeV, in ENDF/B-VI format¹⁷. The material identification number in the file is MAT = 9131. These computerized data files contain recommended values for use in application calculations and are coded in the ENDF/B format. The numerical values are obtained based on an evaluation of existing experimental data supplemented by theoretical model-based predictions and systematics^{8,18} and thus represent the best data available in the world in electronic form for application calculations.

A graphical inter-comparison of all the neutron reaction data of JENDL-3 and ENDF/B-VI for all main isotopes of thorium fuel (²³⁰Th, ²³²Th, ²³¹Pa, ²³³Pa, ²³²U, ²³³U, and ²³⁴U) are available in the handbook by Ganesan and MacLaughlin¹⁹. Generally, for these isotopes, the Japanese data, JENDL-3.2 is considered superior because the evaluations of nuclear data for these isotopes were funded well in Japan and significant efforts were made to incorporate the current status of theoretical predictions and available experiments in the creation of JENDL-3.2. On the other hand, as mentioned in the comments section of the electronic file of ENDF/B-VI for these isotopes is a carry over from their earlier ENDF/B-V created around 1981.

Our calculations were performed as follows: A multigroup cross-section set in 69 groups in the WIMS format²⁰ was generated in collaboration with the IAEA Nuclear Data Section using the NJOY97 code system²¹. The multi-group set was generated following cross-section our specifications by McLaughlin from the IAEA Nuclear Data Section, using NJOY97. This task was successfully carried out within the scope of the IAEA Co-ordinated Research Programme (CRP) entitled, 'Final Stage of the WIMS Library Update Project'. The interested reader can get more details on this IAEA-CRP at the web-site http://www-



Figure 1. Comparison of neutron-induced absorption crosssections in 69 energy groups for ²³¹Pa.



Figure 2. Comparison of neutron-induced fission cross-sections in 69 energy groups for 231 Pa.

CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999

rcp.ijs.si/~wlup. The generation of complete WIMS library for ²³¹Pa involves several steps, viz. resonance reconstruction, Doppler broadening of neutron crosssection curves for all reaction channels, calculation of selfshielding factors in the resolved and unresolved resonance region for various dilutions, generation of self-shielded multigroup cross-sections, and finally, multigroup transfer matrices for elastic and inelastic cross-sections. These multigroup data are in the WIMS format in the 69 groups covering the energy region 10^{-5} eV to 10 MeV. These stateof-art calculations were all performed strictly in accordance with the internationally established ENDF/B conventions and procedures for ²³¹Pa for the first time.

The detailed intermediate outputs and the complete 69 group cross-section set in WIMS format derived from JENDL-3.2 are available free of cost upon request from Ganesan. These calculations were also performed using the American evaluated nuclear data file ENDF/B-VI available from the IAEA Nuclear Data Section²². Figures 1–3 present a graphical comparison of the cross-sections in the two files in multigroup form for fission, absorption and 'nubar', the total number of neutrons released per fission for the infinite medium taking self-shielding into account. For clarity, the ratio has also been shown in the graphs separately.

The calculations of infinite medium multiplication factor (K_{∞}) were performed using three different computer codes ITRAN1D (ref. 23), DTF-IV (ref. 11) and NEWMURLI (refs 24, 25) using the 69 group cross-sections for ²³¹Pa. The ITRAN1D is a multigroup integral transport theory code based on interface current approach for three (sphere, slab,



Figure 3. Comparison of effective total neutrons released per fission 'nubar' in 69 energy groups for ²³¹Pa.

CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999

cylinder)⁰²⁵ computers. The computer code DTF-IV is a multigroup integro differential transport theory code based on S_N method. The code NEW MURLI is based on first flight collision probability method. The results were identical with all the above mentioned codes. We believe that improvements in processing approximations such as increasing the number of energy groups or Legendre order for the scattering matrices will not change the conclusions of this paper qualitatively.

Some additional discussions are presented in this section to provide further insight into our study. With the new 69 group cross-section data derived from JENDL 3.2, our calculated K_{eff} for a sphere of radius 13.61 cm is 0.6029 and the calcutated infinite modium multiplication factor is 0.9729. These new results differ significantly from earlier results¹⁻⁷ in which the critical radius was 13.61 cm. We repeated, with Figu 694groupmorosses ection medite devilvied of pomu Edd DFe/Branh, calculations of infinite medium multiplication factor using the ITRAN/DTF-IV/ NEWMURLI codes. The effective one group cross-sections were also obtained by collapsing the 69 group effective cross-sections including self-shielding with the calculated fluxes obtained by transport calculations in the infinite medium of ²³¹Pa. The effective nubar is obtained by collapsing with the product of fission crosssection

and the flux. Table 1 compares the effective one group values for infinite medium of ²³¹Pa and break-up into components.

The infinite medium multiplication factor, K_{∞} , obtained in the earlier BARC studies¹⁻⁷, is 2.199, which is significantly different from our present values of 0.9729 (JENDL-3.2) and 0.9410 (ENDF/B-VI) presented in Table 1. Note that in the JENDL-3.2 file, the value of fission spectrum average of eta is 1.88, which is much less than the value of eta (the infinite medium multiplication factor) in earlier studies of 2.199. Figure 4 compares the normalized neutron spectra in an infinite medium of ²³¹Pa with that of virgin fission neutron spectrum. It should be further stressed, as seen in Figure 4, that the infinite medium neutron spectrum, due to moderation by mainly neutron inelastic scattering in ²³¹Pa, is softer than the virgin fission neutron spectrum. The value of 2.199 for infinite medium multiplication factor obtained in earlier studies¹⁻⁷ indicates that the cross-sections used

Table 1. Values for infinite medium of ²³¹Pa

	Basic file		
Quantity	JENDL-3.2	ENDF/B-VI	
Fission cross-section $S_f(\text{barns})$ Capture $S_c(\text{barns})$ $S_{n,2n}(\text{barns})$ Absorption [capture + fission – $(n, 2n)$] S_a (barns)	0.4270 0.6465 0.0018 1.072	0.5846 0.9724 0.0043 1.553	
Number of neutrons per fission n Infinite medium multiplication factor $K_{\infty} = nS_{y}S_{a}$	2.442 0.9729	2.499 0.9410	

there for ²³¹Pa are very different from the values given in JENDL-3.2 and ENDF/B-VI.

For a sphere with radius of 13.61 cm, we performed Monte Carlo calculations of $K_{\rm eff}$ employing the JENDL-3.2-based continuous energy data and using the MCNP $code^{26}$. For consistency, the WIMS data derived by us from the same JENDL-3.2 was used in the multigroup transport theory calculations using the DTF code. The DTF calculations gave а K_{eff} of 0.60288 while the Monte Carlo calculations resulted in a value of 0.60137 ± 0.0034 . K_{∞} was also calculated with the JENDL-3.2 file using the MCNP code and was found to be 0.9727 ± 00114 . The error quoted in the case of the MCNP calculations is the statistical uncertainty in the Monte Carlo simulation. These results which are summarized in Table 2 provide additional confidence in our calculations.

Conclusions and recommendations

A summary of the results is presented in Tables 1 and 2. Unfortunately, the breakup of the earlier BARC result for K_{∞} of 2.199, into components of fission, absorption, etc. or the detailed basic data actually used there are not available. The only basic data that is available from the previous study is the 'nubar' value given in Table 1 of Srinivasan *et al.*¹. In that table, the total number of neutrons released per fission, nubar, is given as 2.598 corresponding to the neutron energy range 0.6 to 1.1 MeV (9th energy group of the 35 group set). The values in the basic evaluated data files JENDL-3.2 and ENDF/B-VI are seen by us to be 2.296 and 2.380 at the energy of 0.85 MeV. It may be noted that the evaluation of 'nubar' is based on Bois-Frehaut's semiempirical formula²⁷ in JENDL-3.2 and on Howerton's semiempirical formula²⁸ in ENDF/B-VI. It is not clear how the high value of 2.598 for nubar was arrived at in the earlier study.

We conclude that ²³¹Pa does not become critical even for infinite mass. This conclusion is based upon rigorous calculations and analyses using improved data files JENDL-3.2 and ENDF/B-VI and improved processing of these files. These data files provide the state-of-the-art nuclear data based on more recent measurements and state-of-the-art theoretical calculations of cross-sections that are based on improved models and systematics.

In the earlier BARC study⁵, it is stated (within quotes): 'Although ²³¹Pa produced in large quantities in Th/²³³U reactors would be classified as a fissible nuclide on account of even number of neutrons, surprisingly, its K_{∞} value is as high as 2.2 (higher than that of ²³⁵U), indicating that ²³¹Pa is a very good nuclear fuel.' We do not agree with this earlier conclusion in view of our results.

Table 2.	Summary	of results	for ²³¹ Pa
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Description	Earlier BARC study	Present BARC study based on JENDL-3.2
$K_{\rm eff}$ of a sphere of radius of 13.61 cm	1.000 (using the DTF and 35 group data)	0.60288 (using the DTF and 69 group data)
$K_{\rm eff}$ of a sphere of radius of 13.61 cm using Monte Carlo method (MCNP)	Not available	0.6014 ± 0.0034
Infinite medium multiplication factor, K_{∞}	2.199 (DTF)	0.9729 (DTF) 0.9727 ± 0.00114 (MCNP)

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Molecular basis of antifungal toxin production by fluorescent *Pseudomonas* sp. strain EM 85 – A biological control agent

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RHIZOSPHERE-competent bacteria and fungi are the most preferred candidates as biological control agents. Soil pseudomonads are important among them as they colonize the underground growing plant organs efficiently and survive in a variety of diverse conditions¹. The biological activities by which these pseudomonads bring about disease control include rhizosphere colonization, antibiosis, iron chelation by siderophore production, production of volatile compounds, induction of systemic resistance and competition for nutrients². Genetic improvement as well as modification of these biocontrol strains for improved efficiency and consistency of performance in different conditions require a thorough understanding of the mechanism of biocontrol at the molecular level. Complementation of specific mutants defective in antifungal property, generated either through transposon or NTG mutagenesis have been employed for identification of gene(s) responsible for antifungal toxin biosynthesis in *Pseudomonas* spp. by many workers^{3–}

A fluorescent <i>Pseudomonas</i> sp. (strain EM 85) was found to inhibit growth of many soil-borne plant pathogenic fungi and effectively control fungal diseases. The genetics governing this character was probed by mutagenesis followed by a functional complementation analysis after construction of a genomic library of the wild-type strain using the cosmid vector pLAFR 1. A cosmid clone, pANF 17 was able to complement the antifungal toxin production	in a defective mutant. The chromosomal origin of the DNA fragment in the cosmid clone was confirmed by a Southern blot. Sub-clones of the three <i>Eco</i> RI fragments from pANF 17, however, failed in the complementation test. The complementing cosmid was found to be stably maintained and expressed in the defective mutant under laboratory plate assay and <i>in vivo</i> conditions as evidenced by the extraction and detection of the toxin and biological control experiment.
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CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999

RESEARCH ARTICLES

Pseudomonas sp. strain EM 85 was isolated from maize rhizosphere, at the Division of Microbiology, IARI, New Delhi, and is reported to have biological control property against charcoal rot, foot rot and collar rot of maize caused by Macrophomina phaseolina, Fusarium graminearum and F. moniliforme respectively9. The strain also controlled damping-off of cotton caused by Pythium ultimum and Rhizoctonia solani. It is also reported that this isolate possesses multiple antifungal properties and produces siderophores, HCN, fluorescent pigments and ammonia⁹. Possession of multiple antifungal properties makes any biological control agent a better candidate for efficient field performance. However, understanding the exact mechanism of fungal growth inhibition is essential to improve the strain through genetic manipulation. Here, an attempt is made to understand the molecular basis of antifungal property of the fluorescent Pseudomonas sp. strain EM 85 through mutagenesis and functional complementation.

Materials and methods

Media, culture conditions, bacterial strains, fungal cultures and plasmids

Bacterial strains were grown in Luria-Bertani (LB)/Agar (LA) at 30 or 37°C. Antibiotics were added into the medium as and when required, at the following

concentrations: ampicillin, 50 mg; tetracycline, 10 mg; streptomycin, 100 mg; kanamycin 50 mg per one litre of the medium. Fungi were grown on potato dextrose agar (PDA). The bacterial strains, fungal cultures, plasmids and their derivatives used in this study are described in Table 1.

Identification of the biological control bacteria

The taxonomic position of the bacterial strain was identified by carrying out biochemical tests for bacterial identification¹⁰. The GC content of the total DNA of the bacterial strain was found by observing the melting temperature of the DNA¹¹.

Screening of Pseudomonas strain EM 85 for antagonism against plant pathogenic fungi

Screening was done on PDA medium according to the method of Carruthers *et al.*⁷ Actively growing *Pseudo-monas* EM 85 culture was spotted using a loop on the edge of a PDA plate. A fungal agar disc of 5 mm diameter was cut out from an actively growing fungal culture grown on PDA and kept at the centre of the PDA plate inoculated with bacterial culture. Plates were incubated for two to five days and observed for zone of fungal growth inhibition around the bacterial streak.

Chemical mutagenesis of Pseudomonas strain

Table 1. Bacteriai suains, rungai curtures, plasinius and plasiniu derivatives used		
Organisms	Relevent character	Reference
Bacterial strains		
Escherichia coli		
HB 101	F ⁻ , Pro, Leu, thi, LacY, recA, LamB, Str ^r	25
DH5a	F ⁻ , lacZAM15, hsdR1, rec A 1, gyr A 96, thi-1, rel A	12
Pseudomonas		
Strain EM 85	Antifungal toxin producing (Aft ⁺), Amp ^r	9
Strain AN 21	NTG derivative of EM 85 (Aft), Amp ^r	This study
Plasmids		
pLAFR 1	Wide host range cosmid vector, Tet ^r , Ori (RK 2), Tra ⁻	26
pRK 2013	Kan ^r , Tra ⁺ , mobilizing plasmid	27
pKT 230	RSF 1010 derivative Km ^r Str ^r	21
pANF 17	pLAFR 1 cosmid clone from EM 85 gene bank complementing	
1	AN 21 mutant phenotype	This study
pANF 17–1	pKT 230 carrying 11.5 kb <i>Eco</i> RI fragment from pANF 17	This study
pANF 17–2	pKT 230 carrying 9 kb <i>Eco</i> RI fragment from pANF 17	This study
pANF 17–3	pKT 230 carrying 3 kb EcoRI fragment from pANF 17	This study
Fungal cultures (All virulent strains		
obtained from ITCC, New Delhi)		
Rhizoctonia solani		
Fusarium oxysporum f. sp. ciceri		
Fusarium moniliforme		
Fusarium solani		
Macrophomina phaseolina		

 Table 1.
 Bacterial strains, fungal cultures, plasmids and plasmid derivatives used

EM 85 and screening for defective mutants

Pseudomonas strain EM 85 was incubated in 10 ml LB to log phase. To this NTG @ 100 μ g/ml was added and incubated at 30°C for 2 h while shaking. The cells were washed thrice with sterile water and resuspended in 10 ml LB. Mutations were fixed by incubating the broth overnight and the mutant bank was stored at – 70°C after addition of glycerol to a final concentration of 25%. *R. solani* and single colonies of *Pseudomonas* from the mutant bank were co-inoculated on PDA plates for screening defective mutants. Wild-type strain served as a control in the screening tests.

Construction of genomic library of the wild-type Pseudomonas strain EM 85

Plasmid extraction, restriction digestion, agarose gel electrophoresis, preparation and transformation of competent cells, etc. were done according to Sambrook et al.¹², unless otherwise mentioned. Total genomic DNA of strain EM 85 was isolated from four 1 ml log phase culture aliquots using Wizard $^{\rm TM}$ genomic DNA isolation kit (Promega Corporation, Madison, USA). One mg each of genomic DNA was restricted with varying concentrations of EcoRI (0.125, 0.25, 0.5 and 1 unit) in a total reaction volume of 20 µl, for 1 h at 37°C to find out the enzyme dose which can produce maximum amount of DNA having 17-23 kb size. Restriction digestion was stopped by addition of 10 mM EDTA, followed by heat inactivation at 65°C for 15 min. After standardizing the dose of the enzyme, 200 µg of genomic DNA was restricted with appropriate quantity of EcoRI. The partially restricted genomic DNA was carefully layered over a 10-40% sucrose density gradient in a 38 ml polypropylene tube. It was subjected to centrifugation in a Beckman SW 28 rotor for 24 h at 24,000 rpm at 20°C in a Beckman ultracentrifuge. The gradient tube was punctured at the bottom and fractions of 500 µl were collected. Fractions containing DNA fragments of 17-23 kb were pooled after analysing the fractions on a 0.4% agarose gel and subjected to dialysis for 24 h against $T_{10}E_1$ buffer at room temperature with seven changes of buffer. The dialysed DNA was then extracted with butanol and ethanol precipitated and dissolved in 200 µl of TE buffer.

Cosmid vector pLAFR1 was restricted to completion with *Eco*RI and dephosphorylated with calf intestine alkaline phosphatase (CIAP). Two μ g each of genomic DNA fragments and linearized, dephosphorylated vector DNA were allowed to ligate at 16°C for 18 h. *Giga Pack II* (Stratagene) was used for packaging cosmid clones in I heads and the instructions given by the manufacturer were followed. *E. coli* HB 101 strain was incubated for 16 h at 37°C in 50 ml of LB containing 10 mM MgSO₄ and 0.2% (w/v) maltose. The cells were harvested in 10 mM MgSO₄

and the OD of the suspension adjusted to 0.5 with the same. A 1 : 1 dilution of the l phage mixture was made with SM buffer. 100 μ l each of the diluted phage and HB 101 cell suspension were mixed and incubated at room temperature for 30 min. 200 μ l of LB was added into this and incubated at 37°C for 1 h. The mixture was pelletted and resuspended in 50 μ l of fresh LB. The suspension was serially diluted and spread plated onto LB plates containing tetracycline. Colonies were pooled after incubation, in small amounts of LB and stored at -70° C after addition of DMSO @ 88 μ l/ml.

Screening of genomic library for cosmid clones complementing Aft⁻ phenotype

Cosmid clones were conjugally mobilized into the Aft mutant by triparental mating. Individual colonies of AN 21 and HB 101 (pRK 2013) were incubated in 5 ml each of LB containing antibiotics. Cells from 1 ml culture of AN 21 and HB 101 (pRK 2013) were harvested, washed twice with fresh LB, and dissolved in 0.5 ml LB. 100 µl of genomic library stock was spotted onto LA Tet plates and incubated overnight. The patch was scrapped and dissolved in 0.5 ml LB and washed once with fresh LB. The mutant, helper and the genomic library were mixed well in a ratio of 1:1:1 and spotted on a fresh plain LA plate and incubated for 16 h at 37°C. Different dilutions of the conjugal mixture were plated on LA plates, containing ampicillin and tetracycline. The transconjugants of AN 21 coming up after 24 h of incubation were individually checked for production of antifungal toxin against R. solani on PDA.

Isolation of cosmid DNA complementing Aft⁻ phenotype and sub-cloning

Total plasmid was extracted from transconjugant of AN 21 exhibiting Aft⁺ activity by alkaline lysis plasmid miniprep method¹². This was used to transform competent cells of DH5a and the transformants were selected on LA Tet plates. A few transformed colonies were picked up and the plasmids in them were individually mobilized into the mutant strain AN 21 by triparental mating, as described earlier. Individual transconjugants were again checked for antifungal toxin production against R. solani. Plasmid isolation was carried out from the tranformants of DH5a containing the cosmid carrying aft⁺ DNA fragment. For subcloning, pANF 17 was restricted to completion with EcoRI and subjected to electrophoresis on a 0.75% low melting point agarose under cold room condition. The three genomic DNA fragments of molecular weight 11.5, 9 and 3 kb were eluted separately using DNA cleaning kit (Bangalore Genei (P) Ltd). Plasmid vector pKT 230 was used for sub-cloning. Isolation of plasmid, linearization with EcoRI, dephosphorylation of the vector and ligation

RESEARCH ARTICLES

reaction were performed according to Sambrook *et al.*¹² and the procedure described earlier.

Southern hybridization

1 µg each of total genomic DNA and the native plasmid DNA of the wild-type strain Pseudomonas EM 85 was electrophoresed on a 0.4% agarose gel. The DNA in the gel was depurinated in 0.2 N HCl for 10 min and then denatured with several volumes of 0.2 N NaOH and 0.6 M NaCl for 20 min with gentle shaking. The gel was then soaked in 25 mM sodium phosphate buffer, pH 6.8 as described by Bittner et al.¹³. The denatured DNA was transferred to nylon membrane in LKB 2005 transphor unit (Pharmacia, UK). The nylon membrane was air dried and baked in vacuum at 80°C for 2 h using a gel drier and prehybridized at 65°C in a hybridization oven (Amersham Life Sciences, UK) in 25 ml of prehybridization buffer with constant rotation for 6 h. The 11.5 kb EcoRI fragment from pANF 17 was eluted and radiolabelled with [a³²P]dCTP using the labelling kit supplied by Promega Corporation, USA, as per the instructions given by the manufacturer. The labelled probe DNA was added to the prehybridization buffer and hybridization was carried out at 65°C with constant rotation for 18 h. The membrane was washed under stringent conditions, i.e. thrice with solutions containing $2 \times SSC$ and 0.2% SDS at 65°C. The membrane was exposed to X-ray film in a cassette for 3 days at -70° C.

Extraction of antifungal compound from Pseudomonas strain EM 85 and derivatives

Antifungal toxin was extracted according to the method given by Levy and Eyal¹⁴ with slight modification. Ten PDA plates each were heavily inoculated with EM 85, AN 21 and AN 21 (pANF 17) by cross streaking on all directions. After incubation, agar was scrapped from plates, made into small pieces, put inside a 11 flask and 2 volumes of acetone was added. After shaking for a period of 24 h, the contents were passed through muslin cloth and centrifuged. The supernatant was collected and acetone evaporated in a rotary evaporator. The aqueous fraction was extracted twice with diethyl ether and the ether fraction evaporated to dryness. The dried material was dissolved in 1 ml of acetonitrile and used for bioassay against R. solani on PDA. Wells of 10 mm diameter were made towards the edge in PDA plates by removing agar disc from the medium using a cork borer. The wells were then partially sealed with molten soft agar. When the agar solidified, 100 µl of crude antifungal toxin was put carefully into the well and allowed to percolate. Bioassay against *R. solani* was performed by co-inoculating agar disc of *R. solani* growth on the centre of the plate. Growth inhibition was observed after 2 days of incubation.

In vivo biological control activity

The ability of the *Pseudomonas* strains EM85 (Aft⁺), AN 21 (Aft⁻) and AN 21 carrying pANF 17 (Aft⁺) to control *R*. *solani* induced damping-off of cotton was assessed. A completely randomized design was employed. The treatments were (i) *R. solani* alone, (ii) *R. solani* + EM 85, (iii) *R. solani* + AN 21, (iv) *R. solani* + AN 21 (pANF 17), and (v) No inoculation.

Pots of 12 inch diameter were filled with soil @ 12 kg/ pot. One week prior to sowing, fungal inoculum @ 100 g/pot was applied to the upper 10 cm of the pots. Fungal inoculum was prepared in sand, maize powder mixture (9 : 1). Bacterial inoculum was prepared in LB. Seeds were soaked in water for 10 h prior to sowing. 10 seeds/pots were sown. Four replications were kept in each treatment. 3 ml of bacterial inoculum was added to each seed after sowing and covered with soil. Observations on damping-off were made one week after sowing. The data were statistically analysed¹⁵.

Results and discussion

The results of the various biochemical, morphological and physiological tests showed that the strain belongs to the genus *Pseudomonas* and falls in the fluorescence group. The organism was gram negative, motile rod, having positive reactions for fluorescent pigment production, oxidase, catalase, urease and negative reactions for ONPG, arginine hydrolase, indole, MR and VP. The strain was able to utilize carbohydrates like glucose, galactose, arabinose, xylose, fructose, sucrose, trehalose, mannose and unable to use rhamnose, ribose, lactose and inulin. The T_m value of the DNA was found to be $82 \pm 1^{\circ}$ C and the GC content of the total genomic DNA was 68.6 mole%, which falls under the GC content range of the genus *Pseudomonas*¹⁶.

Inhibition of fungal growth on a nutritionally rich medium like PDA by the bacterial strain is an indication of strong antagonistic property. The strain EM 85 was found to inhibit the growth of many soil-borne fungi under PDA plate assay (Table 2). For further studies, *R. solani* was selected due to its faster growth on PDA medium. The phenotype of the wild-type strain was designated as Aft^+ (Antifungal toxin positive) with respect to the fungal growth inhibition.

Incubation with nitrosoguanidine @ 100 µg/ml of the Pseudomonas EM 85 culture for 2 h was found to give a percentage killing of 99.42 and hence probability of finding a particular type of mutant in the surviving population was high. Though nitrosoguanidine induces multiple notoriously close-linked mutations¹⁷, it has been used for generation of bacterial mutants in genetic experiments^{3,6}. On screening the mutant bank, several colonies showed varying degrees of fungal growth inhibition as measured by the distance between the bacterial streak and the growing edge of the fungus. A colony was found to be completely defective in antifungal toxin production and was designated as AN 21. This mutant was subjected to detailed analysis for finding out the nature of the mutation, and has been found to be isogenic to the wild-type strain in all characters examined, like colony morphology, cell shape and size, growth rate, production of siderophores, fluorescent pigments, HCN and ammonia, except for fungal growth inhibition¹⁸. It was found that the fungus grew over the mutant bacterial streak after two days of incubation, whereas the wild-type strain showed a clear zone of inhibition around the colony even after one week of incubation (Figure 1). This showed that mutation affected the antifungal toxin biosynthesis gene(s) in AN 21 and the same was selected for further genetic analysis.

Construction of gene banks employing cosmid cloning is advantageous because concatamers of foreign DNA having 37-52 kb size, i.e. 75 to 105% the size of the lambda DNA can be packaged into I phage heads. Thus, the total genomic library can be represented in lesser number of independent cosmid clones¹⁹. Wide range cosmid vector pLAFR 1 was used here for the construction of the genomic library since the wild-type and the mutant Pseudomonas strains were susceptible to tetracycline, a marker possessed by the comid vector, enabling easy selection. Out of the four different enzyme concentrations used for partial restriction of the total genomic DNA, 0.25 units of enzyme/µg of the DNA was found to yield maximum amount of fragments of size 17-23 kb. Packaging of the ligated vector-insert molecule in I heads and subsequent transfection of the phage to E. coli HB 101 resulted in the formation of around 25,000 bacterial colonies. Plasmid isolation and restriction analysis of eight randomly picked clones showed presence of insert fragments in all of them.

 Table 2.
 Distance of inhibition zone on phytopathogenic fungi by strain EM 85

Fungal culture	Distance of inhibition (mm)* (from the growing edge of the fungus)
Rhizoctonia solani Fusarium oxysporum f. sp. ciceri	7.0 6.5
Fusarium moniliforme Fusarium solani Macrophomina phaseolina	6.5 6.0 5.5

*Average of 10 independent measurements

CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999

The average insert size has been calculated as 26.06 kb (Table 3). Assuming that the EM 85 genome is similar to *E. coli* genome (approximately 4000 kb), the number of clones to be screened to get a specific DNA fragment of interest in the constructed genomic library with 99% probability was found to be 704.5 or around 705 colonies²⁰.

Mobilization of individual gene bank clones separately into the mutant requires lot of manpower as well as material. So mobilization рп masse of the genomic library into AN 21 was performed for complementation experiments. A colony of AN 21 carrying a cosmid clone was thus obtained which possessed antifungal toxin production. The complementing clone was named as pANF 17. A new approach was used to isolate the cosmid clone responsible for complementing the defective character. Total plasmid from the transconjugant of AN 21 showing Aft⁺ phenotype, i.e. AN 21 carrying the aft^+ cosmid clone, was isolated, used for transformation of competent cells of



Figure 1. Growth inhibition of *Rhizoctonia solani* by *Pseudomonas*. A, Wild-type strain EM 85; B, Mutant strain AN 21; C, Mutant strain AN 21 carrying complementing cosmid pANF 17.

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Clone	No	o. of DNA	A Total molec	ular

Clone no.	No. of DNA fragments	Total molecular weight (kb)
1	4	32.86
2	2	25.74
3	2	24.53
4	2	25.05
5	2	26.06
6	3	19.78
7	2	28.72
8	2	26.06
Average		26.06

E. coli DH5a and selected on LA Tet plates. It was made sure that the cosmid clone was brought back to the *E. coli* background. It was important to prove that the restoration of Aft^+ activity by the transconjugant of AN 21 was not by any reversion, but by the complementation of the character by the cosmid clone inside. Hence, the complemented cosmid clone pANF 17 in *E. coli* background was again conjugally mobilized into the Aft^- phenotype AN 21 and the transconjugant was tested for production of antifungal toxin against *R. solani*. The suppression of fungal growth by the transconjugant thus obtained confirmed the ability of pANF 17 to functionally complement the antifungal toxin production by the defective mutant AN 21 (Figure 1).

Plasmid DNA was extracted from *E. coli* DH5a harbouring pANF 17 and restricted to completion with *Eco*RI. It was found that the cosmid contained a DNA fragment of size 23.5 kb and had two internal cut sites of *Eco*RI yielding DNA fragments of molecular weights 11.5, 9 and 3 kb (Figure 2). Restriction digestion with *SalI*, *Bgl*II and *Bam*HI showed that there were 3, 2 and 3 internal cut sites respectively, in the inserted DNA fragment in the complementing clone.

The origin of the clone pANF 17 was ascertained by a Southern blot experiment. The strain EM 85 carried a native plasmid of size approximately 25 kb. Since the total DNA extraction may also include this, for confirming the origin of the complementing cosmid clone, it was necessary to take the plasmid DNA also for Southern blot experiment. The labelled 11.5 kb *Eco*RI fragment of pANF 17 cosmid clone was found to hybridize with the total genomic DNA of the wild-type strain EM 85. There was no signal on the lane with the native plasmid (Figure 3). It was thus confirmed that the gene(s) responsible for Aft⁺ phenotype was carried by the chromosome of the *Pseudomonas* strain EM 85 and

the clone pANF 17 was originally derived from the wild-type genome of EM 85.

pANF 17 on restriction digestion with EcoRI released the inserted DNA fragment in pLAFR 1. The gene(s) responsible for antifungal toxin production might be located anywhere in the 23.5 kb fragment. Sub-cloning is an important molecular tool in finding out the smallest possible DNA segment conferring any character. Plasmid pKT 230, a derivative of RSF 1010 which is having wide host range, has been used for sub-cloning since it was able to multiply in all gram negative bacterial backgrounds²¹. Biparental mating was done to mobilize the sub-cloned plasmids pANF 17-1, 17-2 and 17-3 into AN 21 and the transconjugants were tested for antifungal toxin production. No complementation of Aft⁻ phenotype was observed in these transconjugants. This may be due to a break in the continuous coding region of the aft gene, i.e. EcoRI might be having a cut site within the aft gene, thus the individual sub-clones are not in a position to functionally complement the defective character. Sub-cloning with DNA fragments obtained by restriction digestion of pANF 17 with other enzymes or combination of two enzymes may give further details in this regard.

The crude antibiotic from EM 85 and AN 21 (pANF 17) inhibited the mycelial growth of *R. solani* in PDA plate assay, whereas that from AN 21 had no inhibitory effect. This shows that the cosmid clone was stable in the mutant background and expressed. This again affirms that the cosmid clone pANF 17 carried gene(s) or any essential part(s) of gene(s) which determine antifungal toxin production by the *Pseudomonas* strain EM 85.

Pre-emergent damping-off was observed on the emerging seedlings of cotton after one week of sowing. The diseased plants failed to grow. Hence, the number of surviving plants in each treatment was noted for the assessment of disease suppressive ability of the strains (Figure 4). *In vivo* biocontrol efficiency of the mutant and the wild-type strain differed significantly compared to the control treatment. The disease suppression was 57.89% in the case of the wild-type strain and 52.63% in the case of AN 21 carrying the complementing clone pANF 17. There was no significant difference between the diseasecontrolling ability of the wild-type strain EM 85 and AN 21

Figure 2. Agarose gel electrophoresis of complementing clone. Lane 1, pANF 17 restricted with *Eco*RI; lane 2, *Eco*RI linearized pLAFR 1; and lane 3, 1 *Hind*III + *Eco*RI MW marker. **Figure 3.** Southern blot of DNA from *Pseudomonas* strain EM 85 with radiolabelled 11.5 kb *Eco*RI fragment from pANF 17; A, Total genomic DNA; B, Native plasmid DNA.

CURRENT SCIENCE, VOL. 77, NO. 5, 10 SEPTEMBER 1999



Figure 4. Damping-off suppression by *Pseudomonas* on cotton seedlings. A, *R. solani* + EM 85; B, *R. solani* + AN 21; C, *R. solani* + AN 21 carrying pANF 17; D, *R. solani* alone; E, No inoculation with pathogen (control).

*Mean of four replications having 10 seeds each. CD at P (0.05) 1.23.

having pANF 17. In vivo biological control ability of antifungal toxin defective mutants carrying complementing clones have been reported earlier $^{22-24}$. The results show that the complementing clone pANF 17 was stably maintained in the mutant background and was expressed in the pot culture conditions efficiently. The strain EM 85 was reported to produce multiple antifungal compounds and a preliminary study of the chemical nature of the antifungal compound was done by Pal⁹. A detailed analysis of the structure of the compound will further help identify the relative importance of the same in biological control conditions by detecting the chemical in in vivo systems, especially in the rhizosphere.

The possibility of using the cosmid clone pANF 17 to engineer Aft⁻ *Pseudomonas* strains to Aft⁺ phenotype by heterologous expression is open when finer regulatory mechanisms as well as expression studies are made on the *aft* genes.

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