

NOMAD Code and Model

*Nuclear Engineering
Engineering and Construction
Department
VEP-NFE-1-A
May, 1985*



VIRGINIA POWER NORTH CAROLINA POWER
WEST VIRGINIA POWER

8506040140 850528
PDR ADDCK 05000280
P PDR

THE VEPCO NOMAD CODE AND MODEL

BY

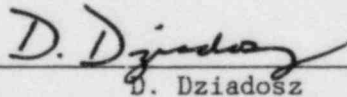
S. M. BOWMAN

NUCLEAR ENGINEERING GROUP
ENGINEERING & CONSTRUCTION DEPARTMENT

VIRGINIA ELECTRIC AND POWER COMPANY
RICHMOND, VIRGINIA

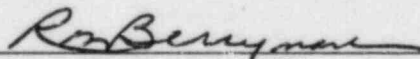
MAY, 1985

Recommended for Approval:



D. Dziadosz
Supervisor, Nuclear Engineering

Approved:



R. M. Berryman
Director, Nuclear Engineering



UNITED STATES
NUCLEAR REGULATORY COMMISSION
WASHINGTON, D. C. 20555

Serial # 85-171

March 4, 1985

Rec'd. MAR 7 1985

Mr. W. L. Stewart, Vice President
Nuclear Operation
Virginia Electric and Power Company
Richmond, Virginia 23261

Nuclear Operations
Licensing Supervisor

Dear Mr. Stewart:

Subject: Acceptance for Referencing of Licensing Topical Report VEP-NFE-1,
"The Vepco NOMAD Code and Model"

We have completed our review of the subject topical report submitted by Virginia Electric and Power Company (VEPCO) letter Serial No. 545. We find the report to be acceptable for referencing in license applications to the extent specified and under the limitations delineated in the report and the associated NRC evaluation, which is enclosed. The evaluation defines the basis for acceptance of the report.

We do not intend to repeat our review of the matters described in the report and found acceptable when the report appears as a reference in license applications, except to assure that the material presented is applicable to the specific plant involved. Our acceptance applies only to the matters described in the report.

In accordance with procedures established in NUREG-0390, it is requested that VEPCO publish accepted versions of this report, proprietary and non-proprietary, within three months of receipt of this letter. The accepted versions shall incorporate this letter and the enclosed evaluation between the title page and the abstract. The accepted versions shall include an -A (designating accepted) following the report identification symbol.

Should our criteria or regulations change such that our conclusions as to the acceptability of the report are invalidated, VEPCO will be expected to revise and resubmit the documentation, or submit justification for the continued effective applicability of the topical report without revision of the documentation.

Sincerely,

Cecil O. Thomas

Cecil O. Thomas, Chief
Standardization and Special
Projects Branch
Division of Licensing

Enclosure: As stated

SAFETY EVALUATION REPORT

Report Title: The Vepco NOMAD Code and Model
Report Number: VEP-NFE-1
Report Date: September, 1983
Originating Organization: Virginia Electric and Power Company
Reviewed By: Core Performance Group, BNL and Core Performance Branch, NRC

INTRODUCTION

The Virginia Electric and Power Company (Vepco) NOMAD code and model are described in references 1 and 2. The NOMAD code is a two-group diffusion theory program with thermal-hydraulic feedback, and was developed to perform one-dimensional axial core analyses in support of reactor startup and cycle operation of the Vepco Surry and North Anna reactors. The accuracy of the code and its ability (in conjunction with a number of other codes) to perform core analyses are demonstrated via comparisons with results from other codes, and with measurements taken at the Surry and North Anna Nuclear Power Stations. The evaluation of the topical report follows.

SUMMARY OF TOPICAL REPORT

NOMAD solves the finite difference form of the 2-group, one dimensional (axial) diffusion equation. The active core is partitioned into 32 intervals, while the top and bottom axial reflectors are each represented by three regions. The solution algorithm for the two-group fluxes employs Gauss elimination for each group equation, with an accelerated power iteration for the coupled equations. The flux at the center of each mesh interval is then obtained by an interpolation scheme based on the finite difference form of the diffusion equation. Since the boundary fluxes are known, the flux at the center of the interval is obtained directly. Simpson's rule of integration is then applied over the interval to obtain the average fast and thermal fluxes in the region. These fluxes are used in the calculation of the region power which is the coupling variable for the thermal-hydraulic feedback.

The moderator temperature is calculated from the moderator enthalpy in the region (which depends on the power, flow rate and inlet enthalpy) and the system pressure via the HOH (reference 3) subroutine. Single phase homogeneous flow is assumed, with no bulk boiling or void formation.

The macroscopic cross sections for each region are then recalculated based on the moderator and fuel temperatures and a new flux calculation is performed. This process continues until the thermal-hydraulic convergence criterion is satisfied.

NOMAD accounts for the time dependent behavior of xenon and iodine by solving the analytical form of the governing equations. A solution for the equilibrium concentrations of xenon and iodine is also available. The time-dependent solution of the equations is performed iteratively for each time interval based on the fluxes from the present and the previous time steps.

Iterations are performed until the flux, thermal-hydraulic and xenon distributions converge.

Basic cross section data for NOMAD are obtained from a 1/4-core PDQ07 One Zone (reference 4) model depletion from BOC to EOC for the particular cycle and unit of interest. At various points in the depletion the relevant flux and concentration files are used to perform re-edit calculations (i.e., the fluxes and eigenvalue are not recomputed) to obtain the flux weighted macroscopic cross sections that result from variations from nominal (reference) conditions in the fuel and moderator temperature, and boron and xenon concentrations.

Reflector cross sections are obtained from the NULIF (reference 5) code, and control rods are represented by core averaged values of the absorption and removal cross sections obtained as the difference between PDQ07 rods-in and rods-out calculations.

The application of the NOMAD model to a desired reactor statepoint involves normalizations to results from more detailed calculations. The Vepco PDQ07

(Discrete (reference 6) and One Zone) and FLAME (reference 7) models for the particular reactor and cycle provide reference data for the NOMAD normalization.

Basically, three normalizations are performed: control rod worth, xenon concentration, and axial power distribution. Control rod worths for each bank inserted alone or for the banks inserted in sequence are normalized to the corresponding worths from PDQ07 Discrete model calculations via input control rod worth normalization factors. The fission yields for iodine and xenon are modified to force agreement of the NOMAD calculated equilibrium xenon concentration to that obtained from the PDQ07 One Zone model at 150 MWD/MTU.

Normalization of the axial power distribution is achieved via the buckling coefficient search option in NOMAD. Axially dependent, radial buckling coefficients which force agreement between the NOMAD axial offset and core midplane power, and corresponding values from the three-dimensional FLAME code, are obtained for a number of core conditions (BOC HZP and HFP, and at each depletion step from 150 MWD/MTU to EOC.) The resultant buckling coefficients are saved for use in subsequent calculations. The HFP ARO buckling coefficients resulting from the above searches are automatically adjusted by NOMAD to account for the insertion of control rods, and for changes in power level.

The NOMAD code and model are capable of performing a variety of reactor physics calculations:

1. The generation of core coverage axial power and burnup distributions.
2. Axial Offset control and load follow maneuver simulations.
3. Peaking factor analyses.
4. Final Acceptance Criteria (FAC) analyses.

5. Criticality searches on boron concentration, control rod bank position, core power level or hot full power inlet enthalpy.
6. Differential and integral control rod bank worths.
7. Xenon worth and reactivity coefficients.

SUMMARY OF REGULATORY EVALUATION

The Vepco one-dimensional NOMAD code and model described in references 1 and 2 are intended for use (along with the previously approved Vepco PDQ07 Discrete and One-Zone models, and the Vepco FLAME model) in performing reload core design and reload key parameter generation. Vepco is currently developing an axial power shape control methodology and the applications of NOMAD in this area will be the subject of a separate Topical Report which Vepco plans to submit for staff review.

The present review considered the information presented in the topical report and in reference 2 (Attachment-1, Response to Questions and Comments on Vepco NOMAD Code and Model, and Attachment-2, Proposed Text Changes to the NOMAD Topical Report). The review also considered the additional NOMAD information provided by Vepco at a meeting with the staff on April 16, 1984.

In general the information presented indicates that the NOMAD code and model are similar in methodology to other approaches currently in use for one-dimensional reactor analyses. The NOMAD diffusion theory finite difference representation together with the prescription for determining the region-centered fluxes has been reviewed and found to be correct and consistent with standard industry practice. The xenon/iodine calculations have also been reviewed and found to provide an accurate technique for tracking the xenon and iodine isotopics.

The NOMAD thermal-hydraulic feedback model employs an energy balance to determine the moderator axial enthalpy distribution. Axial moderator

enthalpies and temperatures generated by the NOMAD thermal-hydraulic feedback model are found to be in excellent agreement with the results from COBRA (reference 8) for a North Anna 120% overpower case. The fuel temperature fit is based on the Vepco PDQ07 thermal-hydraulic feedback model which has been previously reviewed and approved by the staff. It is concluded that the NOMAD thermal-hydraulic model is acceptable.

NOMAD has been normalized to results from more accurate PDQ07 and FLAME calculations. The Vepco PDQ07 and FLAME calculational models have been previously reviewed by the staff and found to be acceptable. From the results presented in references 1 and 2 it is concluded that the adjustments made to NOMAD in the course of the normalization ensures acceptable agreement with the results from these more accurate calculations.

The ability of NOMAD to perform the required reactor physics calculations is demonstrated by comparisons with measured and PDQ07 and FLAME calculated results. Data are presented for differential boron worths, isothermal temperature coefficients, xenon worths (after startup, orderly shutdown and trip), and differential and integral rod worths. Agreement with measurement and/or higher order calculations is acceptable.

A number of axial power distribution comparisons with measured and FLAME calculated data are presented for both North Anna³ units and for Surry Unit-1. Comparisons of flux difference and critical boron data generated by NOMAD in simulations of load reduction and power escalation tests at North Anna Unit-1 are also presented. While the measured data on critical boron for one of the tests is not extensive, the level of agreement between NOMAD and measurement and/or FLAME is reasonable.

Final acceptance criteria analyses were performed for North Anna Units 1 and 2 and the results were compared to results from an accepted and verified vendor model used in the design and licensing of the Surry and North Anna reactors. There were minor differences between the Vepco results and the vendor results for both the 3 case and 18 case analyses.

At maximum values of the peaking factor (F_0) NOMAD results are within 1% of the vendor results and within +1% and -2.5% of the vendor results in the remainder of the core.

REGULATORY POSITION

Based on the review of the NOMAD code and methodology, including comparisons to other calculations and to measurements described above, we find NOMAD represents an acceptable methodology for performing the reactor physics analyses enumerated under the section entitled, "Summary of Topical Report", above. These analyses are related to reload cores similar to those of the Surry and North Anna reactors.

This report may be referenced in licensing submittals by Vepco for the Surry and North Anna reactors.

REFERENCES

1. S. M. Bowman, "The VEPCO NOMAD Code and Model," VEP-NFE-1, Virginia Electric and Power Co. (Sept. 1983).
2. Letter from W. L. Stewart (Vepco) to H. R. Denton (USNRC), "Vepco NOMAD Code and Model Supplemental Information," (July 6, 1984).
3. L. L. Lynn, "A Digital Computer Program for Nuclear Reactor Analysis Design Water Properties," WAPD-TM-680, Westinghouse Electric Corporation (July 1967).
4. J. R. Rodes, "The PDQ07 One Zone Model," VEP-FRD-20A, Virginia Electric and Power Co. (July 1981).
5. P. D. Breneman, "The NULIFP01 Code," NFE Calculational Note PM-13, Virginia Electric and Power Co. (March 1979).
6. M. L. Smith, "The PDQ07 Discrete Model," VEP-FRD-19A, Virginia Electric and Power Co. (July 1981).
7. W. C. Beck, "The Vepco FLAME Model," VEP-FRD-24A, Virginia Electric and Power Co. (July 1981).
8. F. W. Silz, "VEPCO Reactor Core Thermal-Hydraulics Analysis Using the COBRA IIIC/MIT Computer Code," VEP-FRD-33, Virginia Electric and Power Co. (August 1979).

CLASSIFICATION/DISCLAIMER

The data and analytical techniques described in this report have been prepared specifically for application by the Virginia Electric and Power Company. The Virginia Electric and Power Company makes no claim as to the accuracy of the data or techniques contained in this report if used by other organizations. Any use of this report or any part thereof must have the prior written approval of the Virginia Electric and Power Company.

ABSTRACT

The Virginia Electric and Power Company (Vepco) has developed NOMAD, a one-dimensional (axial), two energy group, diffusion theory computer code with thermal-hydraulic feedback, and a calculational model designated as the Vepco NOMAD model.

The model utilizes the Vepco computer codes NOMAD, XSEDT, XSFIT, XSEXP, NULIF, FXYZ, FDELH, and PCEDT. The model also uses data from the Vepco PDQ07 Discrete, PDQ07 One Zone, and FLAME models. The model is used to perform one-dimensional reactor physics analysis in support of reactor startup and cycle operation of the Vepco Surry and North Anna nuclear reactors. The accuracy of the NOMAD model is demonstrated through comparisons with other codes and with measurements taken at the Surry and North Anna Nuclear Power Stations.

ACKNOWLEDGMENTS

The author would like to thank Mr. T. W. Schleicher for his assistance in performing some of the computer calculations and data preparation required for this report and Mrs. Anna Pegram for typing of the manuscript. The author also wishes to express his appreciation to the people who reviewed and provided comments on this report.

TABLE OF CONTENTS

	Page
CLASSIFICATION/DISCLAIMER	i
ABSTRACT	ii
ACKNOWLEDGMENTS	iii
TABLE OF CONTENTS	iv
LIST OF TABLES	v
LIST OF FIGURES	vi
SECTION 1 - INTRODUCTION	1-1
SECTION 2 - CODE DESCRIPTION	2-1
2.1 Introduction	2-1
2.2 Neutron Flux Calculation	2-3
2.3 Thermal-Hydraulic Feedback	2-7
2.4 Xenon Calculation	2-8
2.5 Radial Buckling Coefficient Model	2-10
2.6 Control Rod Model	2-12
2.7 Criticality Search	2-12
2.8 Delta-I Control	2-13
2.9 Boration and Dilution Calculations	2-14
2.10 Final Acceptance Criteria (FAC) Analysis	2-15
2.11 Differential and Integral Rod Worth Calculations	2-17
2.12 Xenon Worth Calculation	2-18
SECTION 3 - MODEL DESCRIPTION	3-1
3.1 Introduction	3-1
3.2 Cross Section Generation	3-2
3.3 Model Normalization	3-4
3.4 1-D/2-D Synthesis	3-6
3.5 FAC Analysis Model	3-7
SECTION 4 - USER INFORMATION	4-1
4.1 Input Description	4-1
4.2 Error & Warning Messages	4-9
4.3 Execution Time	4-10
4.4 Output	4-10
4.5 I/O Units	4-13
SECTION 5 - RESULTS	5-1
5.1 Introduction	5-1
5.2 Reactivity Parameters	5-1
5.3 Thermal-Hydraulic Feedback	5-2
5.4 Axial Power Distribution	5-2
5.5 Differential and Integral Rod Worths	5-2
5.6 Load Follow Maneuver Simulation	5-3
5.7 FAC Analysis	5-4
SECTION 6 - SUMMARY AND CONCLUSIONS	6-1
SECTION 7 - REFERENCES	7-1

LIST OF TABLES

Table	Title	Page
3-1	Macroscopic Cross Section Variable Dependence	3-8
5-1	Reactivity Coefficients Comparison	5-5
5-2	Comparison of NOMAD and COBRA Moderator Enthalpy and Temperature Distributions	5-6
5-3	Rod Swap Comparison, Part 1	5-7
5-4	Rod Swap Comparison, Part 2	5-8
5-5	N1C2 70% Load Reduction Test Power and D-Bank History	5-9
5-6	N1C3 Shutdown/Return to Power Case 1 Power and D-Bank History	5-10
5-7	N1C3 Shutdown/Return to Power Case 2 Power and D-Bank History	5-12

LIST OF FIGURES

Figure	Title	Page
2-1	NOMAD Code Flow Diagram	2-19
2-2	Axial Mesh Points and Regions	2-21
2-3	Axial Region Center and Boundary Mesh Points	2-22
3-1	Vepco NOMAD Model Flow Diagram	3-9
5-1	Xenon Worth After Startup, North Anna Unit 1 Cycle 3 . . .	5-14
5-2	Xenon Worth After Shutdown, North Anna Unit 1 Cycle 3 . .	5-15
5-3	Xenon Worth After Trip, North Anna Unit 1 Cycle 3 . . .	5-16
5-4	Xenon Worth After Startup, Surry Unit 1 Cycle 6	5-17
5-5	Xenon Worth After Shutdown, Surry Unit 1 Cycle 6	5-18
5-6	Xenon Worth After Trip, Surry Unit 1 Cycle 6	5-19
5-7	Axial Power Comparison, N1C2 HZP BOC	5-20
5-8	Axial Power Comparison, N1C2 HFP ARO Eq. Xe. BOC	5-21
5-9	Axial Power Comparison, N1C3 HZP BOC	5-22
5-10	Axial Power Comparison, N1C3 HFP ARO Eq. Xe. BOC	5-23
5-11	Axial Power Comparison, N1C4 HZP BOC	5-24
5-12	Axial Power Comparison, N1C4 HFP ARO Eq. Xe. BOC	5-25
5-13	Axial Power Comparison, N2C2 HZP BOC	5-26
5-14	Axial Power Comparison, N2C2 HFP ARO Eq. Xe. BOC	5-27
5-15	Axial Power Comparison, S1C6 HZP BOC	5-28
5-16	Axial Power Comparison, S1C6 HFP ARO Eq. Xe. BOC	5-29
5-17	Differential Rod Worth Comparison, North Anna Unit 1 Cycle 3	5-30
5-18	Integral Rod Worth Comparison, North Anna Unit 1 Cycle 3	5-31
5-19	Differential Rod Worth Comparison, North Anna Unit 1 Cycle 4	5-32
5-20	Integral Rod Worth Comparison, North Anna Unit 1 Cycle 4	5-33
5-21	Differential Rod Worth Comparison, North Anna Unit 2 Cycle 2	5-34

5-22	Integral Rod Worth Comparison, North Anna Unit 2 Cycle 2	5-35
5-23	Differential Rod Worth Comparison, Surry Unit 1 Cycle 6	5-36
5-24	Integral Rod Worth Comparison, Surry Unit 1 Cycle 6	5-37
5-25	Differential Worth of Control Banks A thru D in Overlap Mode, North Anna Unit 1 Cycle 3	5-38
5-26	Integral Worth of Control Banks A thru D in Overlap Mode, North Anna Unit 1 Cycle 3	5-39
5-27	Differential Worth of Control Banks A thru D In Overlap Mode, North Anna Unit 1 Cycle 4	5-40
5-28	Integral Worth of Control Banks A thru D in Overlap Mode, North Anna Unit 1 Cycle 4	5-41
5-29	Differential Worth of Control Banks A thru D In Overlap Mode, North Anna Unit 2 Cycle 2	5-42
5-30	Integral Worth of Control Banks A thru D in Overlap Mode, North Anna Unit 2 Cycle 2	5-43
5-31	Differential Worth of Control Banks A thru D in Overlap Mode, Surry Unit 1 Cycle 7	5-44
5-32	Integral Worth of Control Banks A thru D in Overlap Mode, Surry Unit 1 Cycle 7	5-45
5-33	N1C2 70% Load Reduction Test, Axial Flux Difference	5-46
5-34	N1C2 70% Load Reduction Test, Critical Boron Concentration	5-47
5-35	N1C3 Shutdown/Return to Power Case 1, Axial Flux Difference	5-48
5-36	N1C3 Shutdown/Return to Power Case 1, Critical Boron Concentration	5-49
5-37	N1C3 Shutdown/Return to Power Case 2, Axial Offset	5-50
5-38	N1C3 Shutdown/Return to Power Case 2, Critical Boron Concentration	5-51
5-39	$F_Z(Z)$ Results, North Anna Unit 1 Cycle 4	5-52
5-40	$F_Q(Z)$ Results, North Anna Unit 1 Cycle 4	5-53
5-41	$F_Z(Z)$ Results, North Anna Unit 2 Cycle 2	5-54
5-42	$F_Q(Z)$ Results, North Anna Unit 2 Cycle 2	5-55

SECTION 1 - INTRODUCTION

The purposes of this report are to describe a reactor analysis computer code and model which were developed at Virginia Electric & Power Company (Vepco) and to demonstrate the accuracy of this model by comparing analytical results generated by the model to results from other codes and to actual measurements from Surry Units No. 1 and 2 and North Anna Units No. 1 and 2.

The code to be described is a one-dimensional (axial), two energy group, diffusion theory (with thermal-hydraulic feedback) computer code and is named the NOMAD code. The model to be described is designated as the Vepco NOMAD model. In addition to NOMAD, the model uses the Vepco computer codes XSED¹, XSFIT¹, XSEXP¹, FXYZ², FDELH³, PCEDT³, and NULIF⁴. The model also utilizes data from the Vepco PDQ07 Discrete⁵, PDQ07 One Zone⁶ and FLAME⁷ models. A detailed description of the input requirements, functioning, physical models, and output capabilities of these codes can be obtained from the referenced code manuals or reports.

The types of reactor physics calculations which can be performed within the general capabilities of the Vepco NOMAD model include:

1. Core average axial power and burnup distributions
2. Axial offset
3. Peaking factors ($F_Q(Z)$, $F_{XY}(Z)$, $F_Z(Z)$)
4. Final Acceptance Criteria (FAC) Analysis
5. Load follow maneuver simulation
6. Criticality searches on boron concentration, control rod bank position, core power level, or hot full power (HFP) inlet enthalpy
7. Differential control rod bank worths
8. Integral control rod bank worths as a function of rod bank position.

The remainder of this report describes the Vepco NOMAD code, the purposes and interrelationships of the other computer codes which comprise the Vepco NOMAD model, the specific modeling of a reactor core with these codes, and the comparisons of calculated results with appropriate results obtained with the Vepco PDQ07 Discrete and One Zone models, the Vepco FLAME model, and with core measurements obtained from the Surry and North Anna Nuclear Power Stations.

SECTION 2 - CODE DESCRIPTION

2.1 Introduction

The Vepco NOMAD (Nuclear Operations Model for Analysis in One Dimension) code provides a relatively simple and inexpensive method for calculating axial power distributions and core reactivity. The calculation contains three levels of iteration: a source or flux calculation is performed during the inner iterations, thermal-hydraulic feedback in the second iteration level, and the xenon concentration in the outer iterations. The two outer levels of iteration are optional.

The neutron flux is calculated by solving the finite difference form of the two-group diffusion equations using Gauss elimination. A Chebyshev polynomial scheme is used to accelerate convergence.

The thermal-hydraulic feedback model accounts for the effect of the nonuniform fuel and moderator temperature distributions on the flux distribution. Single phase flow with no bulk boiling is assumed. Successive relaxation is used to accelerate convergence.

The NOMAD code provides two methods for calculating the axial xenon distribution. The equilibrium xenon calculation is based on the present flux distribution. The xenon depletion calculation is based on the flux distributions from the present and the previous timesteps using an iterative technique. The first method is applicable to fuel cycle design calculations, whereas the second is used for load follow maneuvers or xenon transients. Using the xenon distribution obtained, flux and thermal-hydraulic calculations are performed again. This process continues until the flux, thermal-hydraulic, and xenon convergence criteria are satisfied.

The NOMAD code also contains the following capabilities:

1. Radial buckling calculation and normalization

2. Criticality search on a selected variable
3. Delta-I control
4. Boration and dilution calculations
5. Final Acceptance Criteria (FAC) Analysis
6. Differential and integral rod worth calculations
7. Xenon worth calculation.

Figure 2-1 is a simplified flow diagram of the calculations performed by NOMAD.

The remainder of this section describes in greater detail the models used in the flux, thermal-hydraulic, and xenon calculations, and the methods employed in the seven calculations listed above.

2.2 Neutron Flux Calculation

NOMAD calculates the neutron flux by solving the following finite difference diffusion equations:

$$-\frac{d}{dz} \left[\frac{D_i(z) d\phi_i(z)}{dz} \right] + \Sigma_i(z)\phi_i(z) = \frac{\chi_i G(z)}{\lambda} + \Sigma_{r,i-1}(z)\phi_{i-1}(z), \quad (2.2-1)$$

i=1,2

where

i = the neutron energy group,

$D_i(z)$ = diffusion coefficient for group i at position z,

ϕ = neutron flux,

$\Sigma_i(z) = D_i(z)B^2(z) + \Sigma_{a,i}(z) + \Sigma_{p,i}(z)$ = the total cross section,

B^2 = radial buckling,

Σ_a = absorption cross section,

Σ_r = removal cross section,

Σ_p = poison cross section,

χ_i = fraction of fission neutrons born in group i ($\chi_1=1$, $\chi_2=0$),

$G(z) = \sum_{j=1}^2 \nu \Sigma_{f,j}(z)\phi_j(z)$ = the fission source,

Σ_f = fission cross section,

ν = number of neutrons per fission, and

λ = eigenvalue = K_{eff} .

Equation 2.2-1 is, of course, simply a set of neutron balance equations, where neutron losses on the left-hand side must be balanced by neutron gains on the right-hand side. The first term on the left-hand side describes the leakage from a unit volume by neutron migration in the z-direction, and the second term represents all other losses. These include absorption (Σ_a and Σ_p), leakage in the radial directions (DB^2),

and scattering out of the energy group ($\Sigma_{r,i}$). The sources on the right-hand side are fission ($\chi_i G(z)/\lambda$), and scattering into the group from the next-higher energy group ($\Sigma_{r,i-1}$).

The z -axis in the region of solution, $0 \leq z \leq Z$, is subdivided into axial regions by mesh points at which the solution is to be determined. At some interior point z_n , integrate from $z_{n-} = z_n - h_{n-}/2$ to $z_{n+} = z_n + h_{n+}/2$, where h_{n-} and h_{n+} are the heights of the axial regions below and above point z_n , respectively. (See Figure 2-2.)

The approximations

$$\left. \frac{d\phi_i(z)}{dz} \right|_{z_{n-}} = \frac{\phi_i(z_n) - \phi_i(z_{n-1})}{h_{n-}}, \quad \left. \frac{d\phi_i(z)}{dz} \right|_{z_{n+}} = \frac{\phi_i(z_{n+1}) - \phi_i(z_n)}{h_{n+}}, \quad (2.2-2)$$

$$\begin{aligned} \int_{z_{n-}}^{z_{n+}} \Sigma_i(z) \phi_i(z) dz &= \Sigma_i(z_{n-}) \int_{z_{n-}}^{z_n} \phi_i(z) dz + \Sigma_i(z_{n+}) \int_{z_n}^{z_{n+}} \phi_i(z) dz \\ &= [\Sigma_i(z_{n-}) h_{n-}/2 + \Sigma_i(z_{n+}) h_{n+}/2] \phi_i(z_n) = \bar{\Sigma}_i^{n,n} \phi_i, \end{aligned} \quad (2.2-3)$$

and

$$\frac{D_i(z_{n\pm})}{h_{n\pm}} = \bar{D}_i^{n\pm} \quad (2.2-4)$$

are used to obtain the difference equation

$$\begin{aligned} -\bar{D}_i^{n+} \phi_i^{n+1} + (\bar{D}_i^{n+} + \bar{D}_i^{n-} + \bar{\Sigma}_i^n) \phi_i^n - \bar{D}_i^{n-} \phi_i^{n-1} \\ = \chi_i/\lambda \sum_{j=1}^2 \bar{\Sigma}_{f,j}^n \phi_j^n + \bar{\Sigma}_{r,i-1}^n \phi_{i-1}^n. \end{aligned} \quad (2.2-5)$$

From the form of the equations it can be seen that the macroscopic

cross-sections are required to be constant in each half-interval. In fact, the code assumes these parameters are constant in each axial region.

The code also assumes that the three top and bottom axial regions are reflector regions and the flux at the outer boundaries of these regions is zero. The system of equations given by Equation 2.2-5 and the zero flux boundary conditions, may be expressed in matrix form as

$$\begin{matrix} M & \phi & = & 1/\lambda & F & \phi & + & R & \phi \\ \hline & = & - & & = & - & & = & - \end{matrix} \quad (2.2-6)$$

where M is a tridiagonal matrix.

NOMAD solves this system of equations by the method of power iterations using Gaussian elimination. The rate of convergence is accelerated by replacing the p th iterate of ϕ (i.e., ϕ^p) with a linear combination of ϕ^p and the previous iterate, ϕ^{p-1} ,

$$\phi^{\sim p} = \phi^p (1 + \theta^p) - \phi^{p-1} \theta^p, \quad (2.2-7)$$

where θ^p is an acceleration parameter computed on the basis of Chebyshev polynomials.

The eigenvalue for iteration n is calculated as

$$\lambda = \lambda \frac{\sum_{k=1}^N \sum_{j=1}^2 v \Sigma_{f,j}(z_k) \phi_j^n(z_k)}{\sum_{k=1}^N \sum_{j=1}^2 v \Sigma_{f,j}(z_k) \phi_j^{n-1}(z_k)}, \quad (2.2-8)$$

where N is the total number of mesh points.

The converged solution to Equation 2.2-6 gives the fast and thermal fluxes at the mesh points between each axial region. Next, NOMAD calculates the fluxes at the center of each region. For a region of height $2m$, label the top and bottom mesh points as t and b , respectively, and the region center as c . (See Figure 2-3.)

Using the approximations

$$\left. \frac{d\phi_i(z)}{dz} \right|_b = \frac{\phi_i(c) - \phi_i(b)}{m}, \quad \left. \frac{d\phi_i(z)}{dz} \right|_t = \frac{\phi_i(t) - \phi_i(c)}{m}, \quad (2.2-9)$$

and

$$\int_b^t \Sigma_i(z) \phi_i(z) dz = \Sigma_i(c) \phi_i(c) \cdot 2m, \quad (2.2-10)$$

and integrating Equation 2.2-1 yields

$$\begin{aligned} D_i(c) \left[\frac{2\phi_i(c) - \phi_i(t) - \phi_i(b)}{2m^2} \right] + \Sigma_i(c) \phi_i(c) \\ = \chi_i/\lambda \sum_{j=1}^2 \nu \Sigma_{f,j}(c) \phi_j(c) + \Sigma_{r,i-1}(c) \phi_{i-1}(c). \end{aligned} \quad (2.2-11)$$

Since the fluxes at each region boundary, $\phi_i(b)$ and $\phi_i(t)$, are known, NOMAD solves for the region center flux $\phi_i(c)$ directly. The code then integrates $\phi_i(b)$, $\phi_i(c)$, and $\phi_i(t)$ using Simpson's method to obtain the average fast and thermal fluxes in the region.

Finally, the relative power is calculated for each region Z

$$P(Z) = \frac{\kappa \Sigma_{f1}(Z) \bar{\phi}_1(Z) + \kappa \Sigma_{f2}(Z) \bar{\phi}_2(Z)}{\sum_{Z=1}^R \kappa \Sigma_{f1}(Z) \bar{\phi}_1(Z) + \kappa \Sigma_{f2}(Z) \bar{\phi}_2(Z)} \cdot R \quad (2.2-12)$$

where

$P(Z)$ = relative power in axial region Z

κ = energy per fission (watt-sec)

$\bar{\phi}_i(Z)$ = average group i flux in region Z

R = total number of axial regions with fuel.

2.3 Thermal-Hydraulic Feedback

The thermal-hydraulic feedback model uses an energy balance to calculate the moderator enthalpy as a function of axial position

$$\text{Enthalpy out} = \text{Enthalpy in} + \text{Power} / \text{Flow Rate},$$

where

Enthalpy out = moderator enthalpy exiting the region (BTU/lbm)

Enthalpy in = moderator enthalpy entering the region (BTU/lbm)

Power = power produced in the region (BTU/hr)

Flow Rate = core moderator flow rate (lbm/hr).

Single phase, homogeneous flow is assumed with no bulk boiling or void formation. The moderator enthalpy and system pressure are input to the HOH subroutine⁸, which calculates the corresponding moderator temperature.

The thermal-hydraulic feedback model calculates the fuel temperature rise above the moderator temperature as a function of relative power and burnup

$$\text{Fuel temp}(Z) = \text{Mod. temp}(Z) + (\text{DGEFPD} \cdot \text{Burnup}(Z) + \text{FTFO}) \cdot \text{RPD}(Z) \cdot \text{PR},$$

where

Fuel temp(Z) = fuel temperature in axial region Z

Mod. temp(Z) = moderator temperature in region Z

DGEFPD = fuel temperature vs. burnup coefficient
(°R/EFPD • Relative Power)

Burnup(Z) = burnup of region Z (EFPD)

FTFO = fuel temperature vs. relative power coefficient
(°R/Relative power)

RPD(Z) = relative power density in region Z
 PR = core relative power (fraction of full power).

This fuel temperature fit is based on the one used in the Vepco PDQ07 thermal-hydraulic feedback model⁹.

NOMAD then recalculates the macroscopic cross sections based on the new fuel and moderator temperatures, performs another flux calculation, and performs another thermal-hydraulic calculation. This process continues until the thermal-hydraulic convergence criteria is satisfied.

2.4 Xenon Calculation

NOMAD calculates the iodine and xenon concentrations for each axial region using an analytic solution to the iodine and xenon rate equations. This solution is simply an integration of the iodine and xenon rate equations which assumes that the flux and the cross-sections remain constant over the time interval for which the calculation is performed. Prior to calculating the iodine and xenon concentrations, NOMAD normalizes the fast and thermal fluxes (in neutrons /cm²-sec) to the core power level

$$\phi_2^N(Z) = \frac{\text{POWDEN} \cdot \text{PR} \cdot \text{RPD}(Z)}{\kappa \Sigma_{f1}(Z) \cdot \phi_1^N(Z) / \phi_2^N(Z) + \kappa \Sigma_{f2}(Z)} \quad (2.4-1)$$

$$\phi_1^N(Z) = \phi_2^N(Z) \cdot \phi_1^N(Z) / \phi_2^N(Z) \quad (2.4-2)$$

where

$\phi_i^N(Z)$ = normalized flux for group i in region Z (neutrons/cm²-sec)

POWDEN = power density (watts/cc)

PR = core relative power (fraction of full power)

RPD(Z) = relative power density in region Z

$\phi_i^N(Z)$ = region average relative flux for group i in region Z.

NOMAD then uses these normalized fluxes to calculate the iodine and xenon concentrations

$$I(Z)_{i+1} = [I(Z)_i - \gamma_I \Sigma_f \phi / \lambda_I] e^{-\lambda_I(t_{i+1} - t_i)} + \gamma_i \Sigma_f \phi / \lambda_I \quad (2.4-3)$$

$$Xe(Z)_{i+1} = \left[Xe(Z)_i - \frac{(\gamma_I + \gamma_{Xe}) \Sigma_f \phi}{LX} + \frac{\lambda_I I(Z)_i - \gamma_I \Sigma_f \phi}{\lambda_I - LX} \right] e^{-LX(t_{i+1} - t_i)} - \left[\frac{\lambda_I I(Z)_i - \gamma_I \Sigma_f \phi}{\lambda_I - LX} \right] e^{-\lambda_I(t_{i+1} - t_i)} + \frac{(\gamma_I + \gamma_{Xe}) \Sigma_f \phi}{LX} \quad (2.4-4)$$

where

$I(Z)_{i+1}$ = iodine concentration in region Z at step i+1

γ_I = iodine fission yield

λ_I = iodine decay constant

t_{i+1} = time (seconds) at step i+1

$\Sigma_f \phi = \kappa \Sigma_{f1}^N(Z) \phi_1^N(Z) + \kappa \Sigma_{f2}^N(Z) \phi_2^N(Z) / K_{AVG}$

K_{AVG} = average energy per fission

$Xe(Z)_{i+1}$ = xenon concentration in region Z at step i+1

γ_{Xe} = xenon fission yield

$LX = \lambda_{Xe} + \sigma_{a1}^{Xe} \phi_1(Z) + \sigma_{a2}^{Xe} \phi_2(Z)$

λ_{Xe} = xenon decay constant

σ_{aj}^{Xe} = xenon absorption cross section, group j.

To calculate the equilibrium iodine and xenon concentrations, the exponential terms in Equations 2.4-3 and 2.4-4 are set to zero.

Each xenon depletion is actually performed in two substeps. During the first substep, the xenon is depleted for 55% of the depletion time using the flux from the previous timestep. The second substep, which depletes the remaining 45% of the depletion time, is performed iteratively with the flux calculation at the present timestep. Thus, the xenon is depleted with the flux from the previous timestep for 55% of the depletion and with the flux from the present timestep for 45% of the depletion.

After NOMAD calculates the iodine and xenon concentrations, the macroscopic cross-sections are adjusted and the flux and thermal-hydraulic calculations are performed again. The iodine and xenon concentrations are calculated with the new fluxes. This process continues until the xenon distribution converges.

2.5 Radial Buckling Coefficient Model

The radial buckling model accounts for radial leakage and compensates for the radial dimensions which are neglected in a one-dimensional axial model. The buckling coefficients, which are used to calculate the radial buckling as a function of core height, are adjusted to obtain agreement with a three-dimensional code for axial offset and relative power at the core midplane. The equations for the radial buckling function expressed in terms of buckling coefficients are:

$$\text{BUK1}(Z) = B0 \cdot (1 + \text{BTILT} \cdot (Z - Z0) / \text{HT}) \cdot \text{CZ}(Z) \quad (2.5-1)$$

$$\text{BUK2}(Z) = \text{BTH} \cdot \text{BUK1}(Z) \quad (2.5-2)$$

$$\begin{aligned} \text{CZ}(Z) &= \cos(\text{BMID} \cdot \text{PI} \cdot (Z - Z0) / \text{HT}) && \text{if } \text{BMID} > 0.05 \\ &= 1./(\cos(\text{BMID} \cdot \text{PI} \cdot (Z - Z0) / \text{HT})) && \text{if } \text{BMID} < -0.05 \\ &= 1.0 && \text{otherwise,} \end{aligned} \quad (2.5-3)$$

where

$\text{BUK1}(Z)$ = Fast group buckling

BUK2(Z) = Thermal group buckling
 B0 = Buckling amplitude coefficient
 BMID = Buckling curvature coefficient
 BTILT = Buckling tilt coefficient
 BTH = Thermal-to-fast group buckling ratio
 HT = Active core height
 Z = Axial position
 Z0 = HT / 2
 PI = 3.1415927.

For a positive BMID, the function is a convex curve. For a negative BMID, the function is a concave curve. When BMID is near zero, the function is a straight line. BTILT adjusts the slope of the curve. If it is positive, the buckling is greater in the top half of the core. If it is negative, the buckling is greater in the bottom half of the core. B0 and BTH must always be positive so that the buckling function is positive.

NOMAD has an automated buckling coefficient search option. The search iterates on BTILT, BMID and B0, respectively, until the axial offset, midplane power, and eigenvalue converge on the target values (eigenvalue target is 1.0) or until a maximum number of iterations have been performed and a warning is printed. When a buckling coefficient search is completed, the coefficients are written to a dataset with the core average burnup (in EFPH). This buckling coefficient dataset may be read and used in subsequent calculations by NOMAD. If the core average burnup lies between two burnups in the buckling coefficient table, linear interpolation is performed to determine the coefficients for that step.

NOMAD automatically adjusts the buckling coefficients at power levels below 100% power to compensate for changes in the radial buckling as a function of core power. Since the dependence of the cross sections

on moderator temperature are derived from a two-dimensional model, this adjustment is necessary to obtain consistent agreement between NOMAD and a three-dimensional model, where the moderator temperature varies in the axial direction.

2.6 Control Rod Model

NOMAD accounts for the control rod cross sections in the top reflector and any rodged fuel regions. NOMAD adds the control rod macroscopic cross sections to the region macroscopic cross sections. If a particular control rod bank is located at step 225 (top of fuel) or less, then the top reflector is completely rodged with respect to that bank. If the bank is positioned at a higher step, the top reflector is partially rodged, and the rod cross sections are volume-weighted by the fraction of the reflector region which is rodged.

The code inserts each control rod bank by step (228 steps = completely withdrawn, 0 steps = completely inserted). The distance per rod step is 0.625 inches in NOMAD. The fuel region which is partially rodged (where the tip of the control rod bank is located) is handled in the same manner as the partially rodged reflector region, by volume-weighting the rod cross sections.

The NOMAD control rod model includes the modeling of four control banks, two shutdown banks, and one part length bank, and the capability to move all four control banks in overlap.

2.7 Criticality Search

The criticality search option in NOMAD searches for the value of a selected variable (e.g., boron concentration, control rod bank position, core power level, HFP inlet enthalpy) which will give the desired target eigenvalue. The search takes the errors from the two previous guesses and

uses linear interpolation or extrapolation to guess what value of the selected variable will give an error of zero. NOMAD performs another eigenvalue calculation with the new value of the selected variable. The search continues until the eigenvalue converges on the target value.

Since a control rod bank can only be inserted in discrete steps, the criticality convergence criterion may not be satisfied when a control rod search is performed. In this case, the code optionally performs a critical boron search after the control rod bank position is adjusted as near to critical as possible.

2.8 Delta-I Control

In order to simulate a load follow or other maneuver where the reactor is required to operate within a certain delta-I band, a delta-I control option is available that automatically adjusts the control rods to keep the delta-I within its operating band. Delta-I is defined as:

$$\text{Delta-I(\%)} = \frac{\text{Power(top)} - \text{Power(bottom)}}{\text{Power(top)} + \text{Power(bottom)}} \cdot \text{PR} \cdot 100, \quad (2.8-1)$$

where

Power(top) = relative power in top half of core

Power(bottom) = relative power in bottom half of core

PR = core relative power level.

When delta-I is outside the operating band, NOMAD moves D-bank from the bite position (216 steps) to the rod insertion limit by increments of 20 steps. Each of these points is used to determine the delta-I as a function of D-bank position by a cubic least squares fit. The cubic equation is then solved to find the D-bank position which will adjust delta-I to the designated value. If the desired delta-I cannot be

achieved, then delta-I is adjusted to the nearest possible value. The user may request that the code adjust delta-I to (1) the target delta-I (the center of the operating band) or (2) the nearest edge of the operating band. Adjusting delta-I to the edge of the band requires less boration or dilution to accomplish. The eigenvalue calculation is then performed at the adjusted D-Bank position. NOMAD automatically performs a critical boron search after the rods are moved to re-establish criticality.

2.9 Boration and Dilution Calculations

Boration and dilution calculations are important when studying a possible load follow maneuver to insure that the water processing system can handle the rapid changes in boron concentration. NOMAD performs boration and dilution calculations for every step following the first criticality search if the boration/dilution parameters are input. NOMAD solves the following equations¹⁰:

$$\text{WATER}(J) = -\text{SYSMAS} / \text{H2ODEN} \cdot \ln \left[\frac{1 + (\text{BOR}(J) - \text{BOR}(J-1))}{(\text{BOR}(J-1) - \text{Cin})} \right] \quad (2.9-1)$$

$$\text{TIMEMN} = \text{WATER}(J) / \text{LDRATE}, \quad (2.9-2)$$

where

WATER(J) = water processed (gallons) at step J

SYSMAS = total primary coolant system mass (lbm)

H2ODEN = water density in letdown line (8.2 lbm/gal)

BOR(J) = boron concentration at step J

BOR(J-1) = boron concentration at step J-1

Cin = boron concentration in letdown line

TIMEMN = minimum time required to perform boration/dilution

LDRATE = letdown rate.

WATER(J) is multiplied by -1 in a dilution case in order to distinguish between boration and dilution cases.

If TIME MN is greater than the time between steps J-1 and J, NOMAD calculates the maximum achievable change in the boron concentration and prints a warning message to indicate that the minimum time required for the boration or dilution is greater than the time allowed. If instructed, the code then performs another criticality search on control rod bank position, core power level, or inlet enthalpy at the maximum (or minimum) boron concentration achievable in the boration (or dilution).

2.10 Final Acceptance Criteria (FAC) Analysis

NOMAD is capable of performing Final Acceptance Criteria (FAC) analysis. One part of this capability is the average power distribution calculation for a load follow depletion. The code integrates the axial flux distributions over the timesteps specified to obtain the average flux distribution:

$$\bar{\phi}_i(Z) = \frac{\sum_{j=1}^T \phi_{ij}(Z) t_j}{\sum_{j=1}^T t_j}, \quad i=1,2 \quad (2.10-1)$$

where

$\phi_{ij}(Z)$ = group i flux in region Z for timestep j (neutrons/cm²-sec)

t_j = length of timestep j

T = total number of timesteps specified.

This flux distribution is substituted into Equation 2.2-12 to obtain the average power distribution and Equations 2.4-3 and 2.4-4 to obtain the average iodine and xenon distributions. The load follow depletion is then performed using these power and xenon distributions.

To perform a FAC analysis, NOMAD combines the axial power distributions that it calculates in the load follow calculations with the $F_{XY}(Z)$ data input to the code to determine $F_Z(Z)$, $F_{XY}(Z)_{calc.}$, $F_{XY}(Z)_{allowable}$, and $F_Q(Z)_{calc.}$ NOMAD performs the following sequence at each step in the FAC analysis case. First, it determines the rod configuration (i.e., ARO, D in, D+C in) for each axial region. Then the code selects the $F_{XY}(Z)$ that corresponds to the axial level and the rod configuration of each region. NOMAD checks each $F_{XY}(Z)$ to insure that it is not less than the minimum $F_{XY}(Z)$ allowed for that rod configuration as specified in the user input. If the reactor is not at full power, NOMAD adjusts the $F_{XY}(Z)$ as follows:

$$F_{XYREL} = F_{XY}(Z) \cdot (1 + ADJUST \cdot (1 - PR)), \quad (2.10-2)$$

where

$F_{XYREL} = F_{XY}(Z)$ adjusted for the core relative power level

$F_{XY}(Z) = F_{XY}$ at axial region Z for 100% power

$ADJUST = F_{XY}$ power adjustment factor

(e.g., 0.3 for North Anna and Surry)

PR = core relative power level.

Next, NOMAD calculates the $F_Q(Z)$ for this case:

$$F_{QTEST} = F_{XYREL} \cdot RPD(Z) \cdot PR \cdot F_{QGRID}, \quad (2.10-3)$$

where

$F_{QTEST} = F_Q(Z) \cdot PR$ for this step

$RPD(Z)$ = relative axial power in region Z

F_{QGRID} = correction factor for grids \cdot uncertainty factor

$$= 1.025 \cdot UF$$

$UF = 1.03$ (3 case FAC analysis)

$UF = 1.00$ (18 case FAC analysis).

If FQTEST is greater than the previous $F_Q(Z)$, then the following values are saved:

$$F_Q(Z)_{\text{calc.}} = \text{FQTEST} \quad (2.10-4)$$

$$F_{XY}(Z)_{\text{calc.}} = \text{FX YREL} \quad (2.10-5)$$

$$F_{XY}(Z)_{\text{allowable}} = F_Q(Z)_{\text{limit}} / (F_Z(Z) \cdot \text{PR} \cdot \text{FQGRID}) \quad (2.10-6)$$

$$F_Z(Z) = \text{RPD}(Z) \quad (2.10-7)$$

Once the entire load follow simulation has been completed and the final values for $F_Z(Z)$, $F_{XY}(Z)_{\text{calc.}}$, $F_{XY}(Z)_{\text{allowable}}$, and $F_Q(Z)_{\text{calc.}}$ have been obtained, NOMAD checks for any limit violations for $F_{XY}(Z)_{\text{calc.}}$, $F_{XY}(Z)_{\text{allowable}}$, and $F_Q(Z)_{\text{calc.}}$ and flags them in the FAC ANALYSIS RESULTS output.

2.11 Differential and Integral Rod Worth Calculations

Differential and integral rod worth calculations are available in NOMAD. The control rod banks may be inserted or withdrawn in any order chosen by the user (e.g., single bank, multiple banks in overlap, multiple banks together, etc.). The differential worth is calculated as follows:

$$\text{DIFF}(I) = (1/\text{RKEF3} - 1/\text{RKEF1}) \cdot 1\text{E}+5 / \text{ISTEPS}, \quad (2.11-1)$$

where

$\text{DIFF}(I)$ = differential rod worth for case I (pcm/step)

RKEF3 = Keff for case I+1

RKEF1 = Keff for case I-1

ISTEPS = number of steps rods moved from case I-1 to I+1.

The differential worth is not calculated for the first or last case of the rod worth sequence.

NOMAD calculates the integral worth as follows:

$$RINT(I) = (1/RKEF2 - 1/RKEF) \cdot 1E+5, \quad (2.11-2)$$

where

RINT(I) = integral rod worth for case I (pcm)

RKEF2 = Keff for case I

RKEF = Keff for 1st case of rod worth sequence.

The integral worth is calculated for every step of the rod worth sequence.

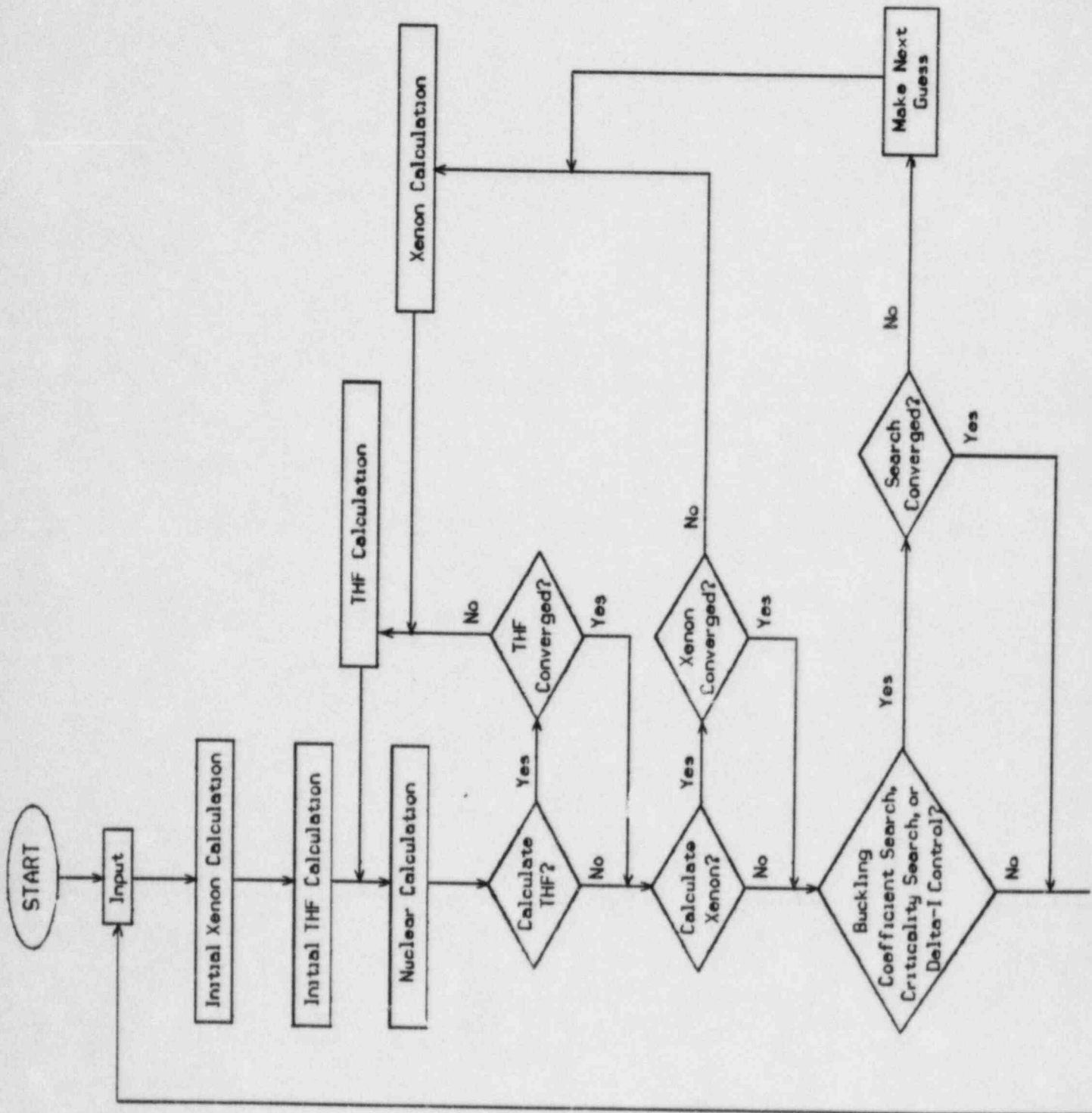
2.12 Xenon Worth Calculation

NOMAD can automatically calculate the xenon worth at any selected timestep(s). When the xenon worth option is on, NOMAD saves the calculated xenon distribution in a separate array, resets the xenon distribution to zero and performs another eigenvalue calculation. It calculates the xenon worth from the two eigenvalues:

$$\text{Xenon worth} = \frac{\text{Keff (no xenon)} - \text{Keff (w/xenon)}}{\text{Keff (no xenon)} \cdot \text{Keff (w/xenon)}} \cdot 1E+5. \quad (2.12-1)$$

The program then restores the saved xenon distribution. Thus, the xenon worth at any timestep of a problem can be determined without interrupting the flow of the other calculations being performed.

FIGURE 2-1
NOMAD Code Flow Diagram



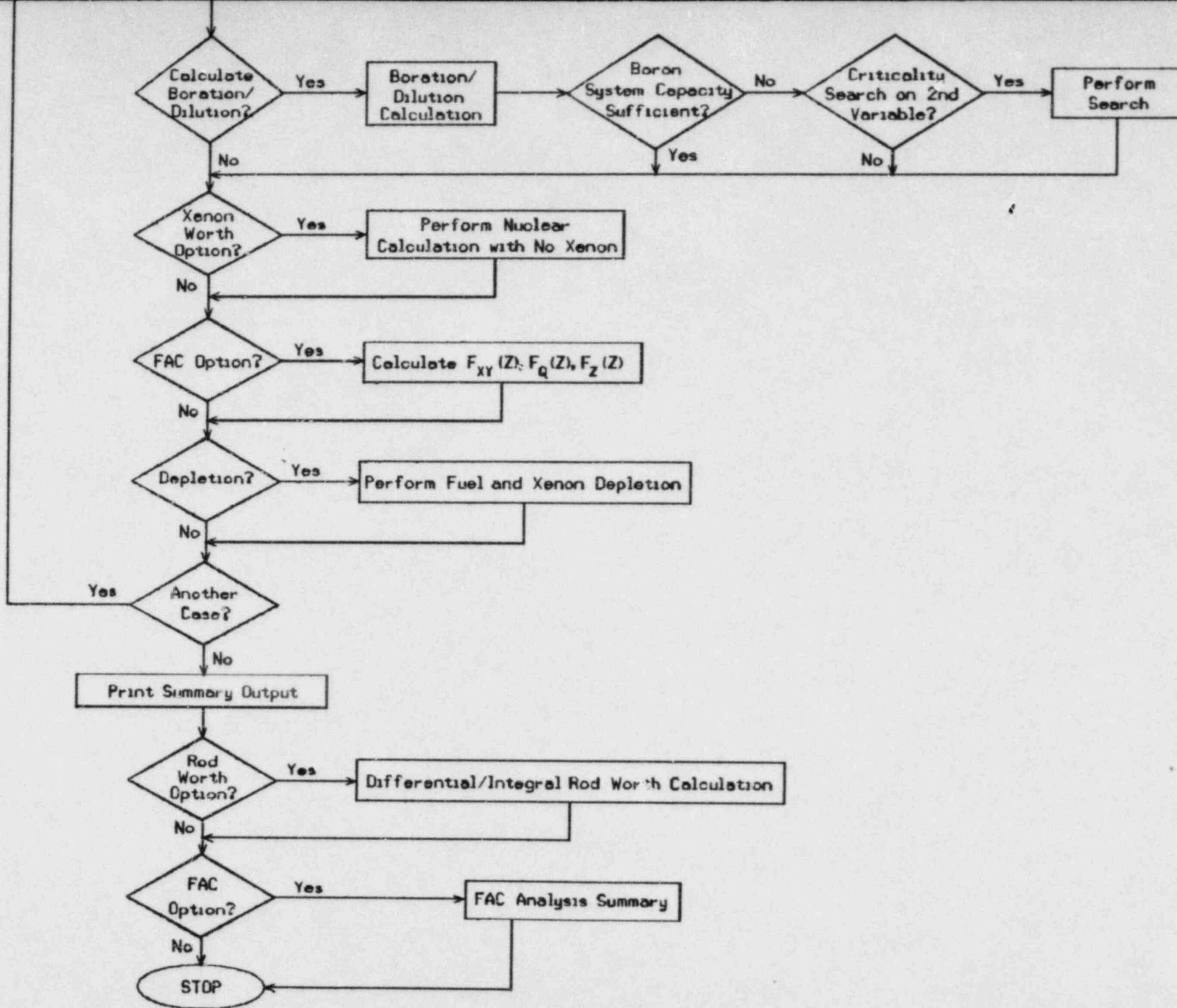


FIGURE 2-1 (Continued)

FIGURE 2-2
Axial Mesh Points and Regions

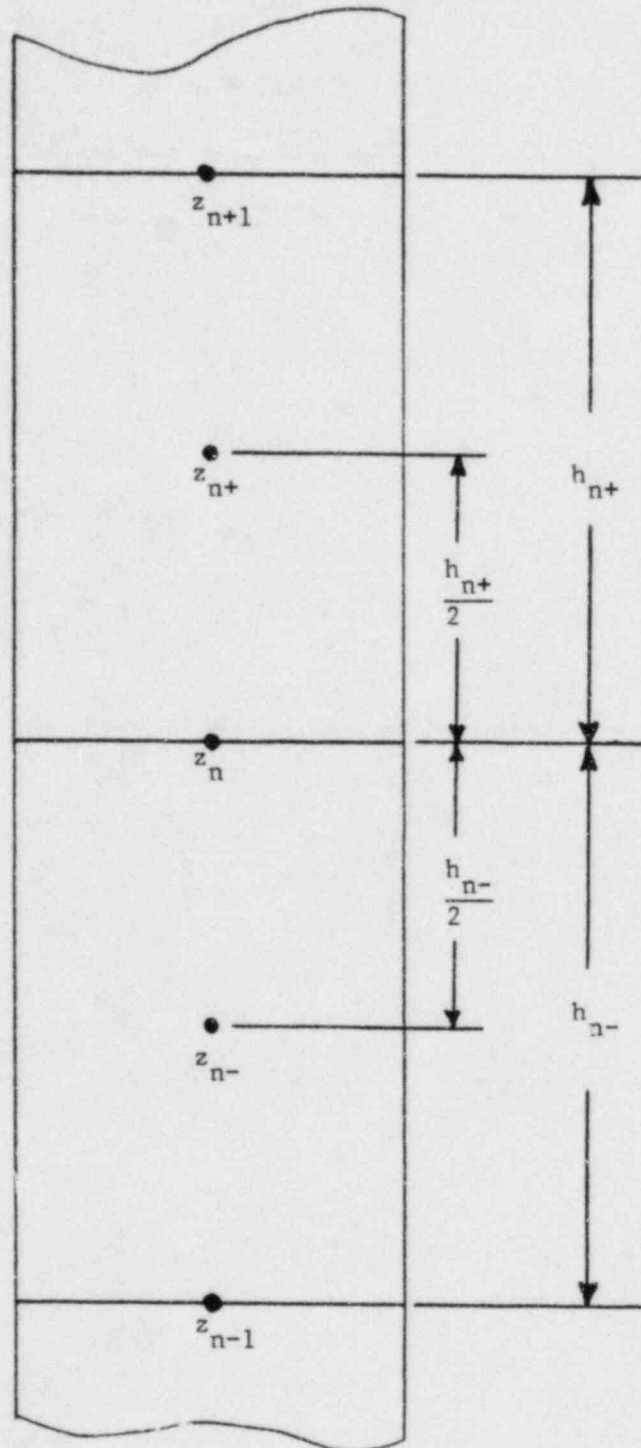
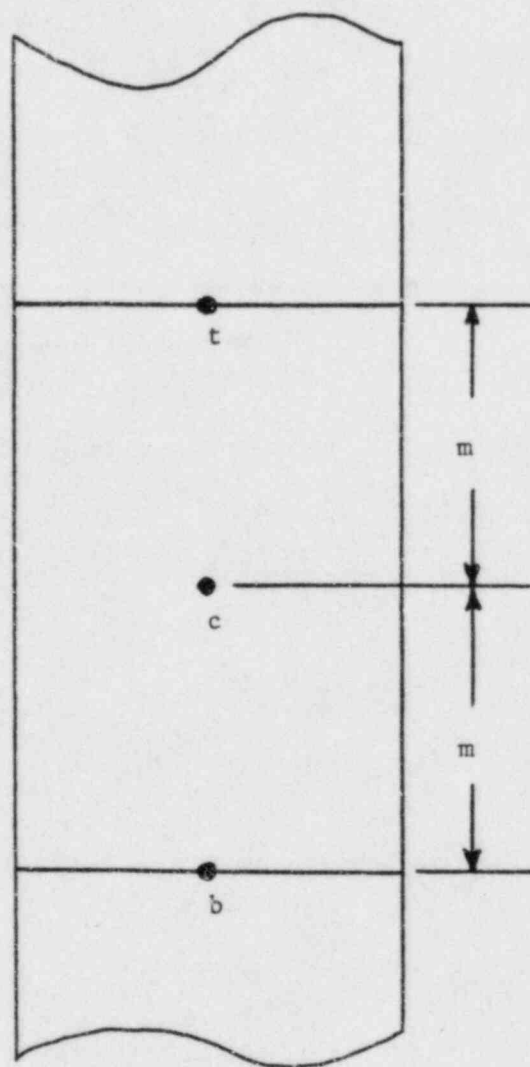


FIGURE 2-3
Axial Region Center and Boundary Mesh Points



SECTION 3 - MODEL DESCRIPTION

3.1 Introduction

The Vepco NOMAD model is used to calculate axial power distributions and core reactivity for one-dimensional geometries in which the core is represented by 32 axial fuel regions and three top and three bottom reflector regions. The method used by the Vepco NOMAD model to perform these calculations is a finite difference solution of the two energy group diffusion theory equations. Moderator and fuel temperature effects are accounted for by thermal-hydraulic feedback.

The Vepco NOMAD model incorporates several calculational steps. First, a quarter core PDQ07 One Zone⁶ depletion is performed and the flux and concentration files at each burnup step are saved for the particular unit and cycle being studied. Then PDQ performs flux-weighted macroscopic cross section calculations for a series of change cases at each burnup step from 150 MWD/MTU to EOC. The core average macroscopic cross sections from these calculations are then processed for input into the NOMAD computer code. These cross-sections, as well as the cycle normalization data (i.e., BOC core average axial burnup distribution, equilibrium iodine and xenon concentrations, integral control rod bank worths, and axial offset and core midplane power at each depletion step), are then used by NOMAD to perform an iterative, two-group finite difference diffusion theory calculation for the neutron flux as a function of core height. The method of solution comprises three levels of iteration: neutron flux, thermal-hydraulic feedback, and xenon concentration. The neutron flux calculation is performed first based on initial guesses of fuel and moderator temperatures and xenon concentrations. Then a new set of fuel and moderator temperatures are calculated. Using these new temperatures, another flux calculation is performed. Once the flux and temperatures have both converged, a new set

of xenon concentrations is calculated. Using the new xenon concentrations, the flux and thermal-hydraulic feedback calculations are performed again. This process continues until the convergence criteria for all three levels are satisfied.

Several interrelated computer codes are used to perform the calculations outlined above. The computer codes comprising the Vepco NOMAD model and their interrelationships are presented in the flow diagram in Figure 3-1. The NOMAD computer code is the principal analytical tool in the Vepco NOMAD model. The other codes provide either input data or data manipulation. The PDQ07 One Zone model and the XSEDT¹ code are used to generate core average macroscopic cross sections at different core conditions. The XSFIT¹ and XSEXP¹ codes process these data for use by NOMAD. NULIF⁴ is used to calculate the top and bottom reflector macroscopic cross sections. These generally remain the same from cycle to cycle. The Vepco PDQ07 One Zone, PDQ07 Discrete⁵, and FLAME⁷ models supply cycle normalization data. The FXYZ² code provides $F_{XY}(Z)$ input to NOMAD for FAC analysis. FDELH³ and PCEDT³ perform 1-D/2-D synthesis of NOMAD and PDQ07 Discrete or One Zone results.

The remainder of this section describes in greater detail the input to and functioning of the computer codes used in the Vepco NOMAD model.

3.2 Cross Section Generation

The Vepco NOMAD code requires the following two group macroscopic cross sections for the solution of the axial flux and power distributions:

$$D_1, \Sigma_{a1}, \Sigma_{r1}, v\Sigma_{f1}, \kappa\Sigma_{f1}, D_2, \Sigma_{a2}, v\Sigma_{f2}, \text{ and } \kappa\Sigma_{f2}.$$

These cross sections actually consist of base macroscopic cross sections and polynomial coefficients that adjust the base cross sections for changes in the fuel and moderator temperatures and the boron and xenon concentrations. The cross sections are generated from the PDQ07 One Zone

model for that unit and cycle.

The PDQ07 One Zone model is depleted to EOC and the flux and concentration files are saved at each burnup step. A series of restart calculations are performed at each burnup step from 150 MWD/MTU to EOC. (The BOC step is not included because there is no xenon present.) Using the input flux and concentration files, PDQ only performs the flux-weighted macroscopic cross section calculations (i.e., no flux or eigenvalue calculation). The cases contain variations in the fuel and moderator temperatures and the boron and xenon concentrations which should include all core conditions encountered during reactor operation. The range covered for each variable is:

Fuel Temperature	331 to 2052 degrees Fahrenheit (N. Anna)
	487 to 2326 degrees Fahrenheit (Surry)
Moderator Temperature	543 to 613 degrees Fahrenheit (N. Anna)
	526 to 596 degrees Fahrenheit (Surry)
Boron Concentration	0 to 1800 ppm
Xenon Concentration	0 to 2.00 E-08 atoms/bn-cm.

The XS EDT code copies the core average macroscopic cross sections obtained from each of these calculations to a dataset which is read by the XS FIT code.

The variables upon which each macroscopic cross section have been found to be dependent are listed in Table 3-1. The XS FIT code analyzes the PDQ07 macroscopic cross sections and generates base macroscopic cross sections and polynomial coefficients which express these cross sections in terms of these variables:

$$\text{SIGt} = \text{SIGtbase} + (\text{IV1} \cdot \text{COEFt1} + \dots + \text{IVx} \cdot \text{COEFtx})$$

where

SIGt = macroscopic cross section type t for a certain set
of independent variables

SIGtbase = base macroscopic cross section type t

IVx = value of independent variable x

COEFtx = coefficient for SIGt versus independent variable x
= $\Delta \text{SIGt} / \Delta \text{IVx}$.

It then compares the PDQ07 cross sections to those calculated with the polynomial coefficients to verify the accuracy of the coefficients.

These base cross sections and polynomial coefficients are passed to the XSEXP code which smooths several of the fast group polynomial coefficients and calculates base cross sections and polynomial coefficients beyond the lower and upper burnup extremes using a linear least squares extrapolation. These extrapolated cross sections are needed to account for axial regions with burnups less than the core average at BOC and greater than the core average at EOC. The XSEXP codes writes this final set of base cross sections and polynomial coefficients to a dataset which is read by NOMAD.

3.3 Model Normalization

The Vepco NOMAD model for a particular unit and cycle must be normalized to the Vepco PDQ07 Discrete, PDQ07 One Zone, and FLAME models for the same unit and cycle.

The BOC axial burnup distribution from the Vepco FLAME model is input to the NOMAD model in the cycle/geometry deck. There is a one-to-one correspondence between the fueled axial regions in the NOMAD model and the axial nodes in the FLAME model. The NOMAD BOC axial power distributions at HZP and HFP are normalized to the FLAME BOC axial power distributions using the buckling coefficient search option in the NOMAD code. This search finds a combination of buckling coefficient values which give a

buckling distribution that forces the axial offset and the power at the core midplane to match those from FLAME for the same conditions.

The NOMAD model xenon parameters are normalized to the PDQ07 One Zone model. The NOMAD model is depleted from BOC to 150 MWD/MTU using the xenon parameters from the previous cycle. The fast and thermal xenon microscopic absorption cross sections are assumed to remain constant. The Iodine 135 and Xenon 135 fission yields are modified to force the NOMAD equilibrium iodine and xenon concentrations to agree with those from the PDQ07 One Zone model at 150 MWD/MTU:

$$\gamma_I(\text{new}) = \gamma_I(\text{old}) \cdot I^{135}(\text{PDQ}) / I^{135}(\text{NOMAD}) \quad (3.3-1)$$

$$\gamma_{Xe}(\text{new}) = [\gamma_{Xe}(\text{old}) + \gamma_I(\text{old})] \cdot \frac{Xe^{135}(\text{PDQ})}{Xe^{135}(\text{NOMAD})} - \gamma_I(\text{new}) \quad (3.3-2)$$

The NOMAD model is depleted again from BOC to 150 MWD/MTU with the new fission yields to verify that the concentrations now agree with the PDQ07 One Zone model.

The axial power distributions obtained from the NOMAD model are normalized to the FLAME model results for the remainder of the cycle by performing buckling coefficient searches at each depletion step from 150 MWD/MTU to EOC. These buckling coefficients are saved in a table which is input to the NOMAD code for all subsequent calculations.

The NOMAD model control rod cross sections are normalized by forcing agreement between the NOMAD and PDQ07 Discrete rod bank integral worths. In the case of rod swap worth calculations, the NOMAD bank worths are normalized to the PDQ07 Discrete bank worths for each bank inserted alone. Otherwise, they are normalized to the PDQ07 worths for the banks inserted in sequence (e.g., D in, D+C in, D+C+B in., etc.).

The core average cross sections Σ_{a1} , Σ_{a2} , and Σ_{r1} from the PDQ07 "Rod bank out" case are subtracted from the same cross sections for the "Rod

bank in" case. These values are input to the NOMAD code for that bank, the control rod normalization is set to 1.0, and the integral bank worth is calculated with NOMAD. The PDQ07 bank worth is divided by the NOMAD bank worth, and the control rod normalization is multiplied by that ratio. The bank worth is calculated again with NOMAD to verify that the worth now agrees with the PDQ07 Discrete model. This process is repeated for each rod bank.

3.4 1-D/2-D Synthesis

Prior to the execution of the 1-D/2-D synthesis option in NOMAD, a 2-D PDQ07 case is run for each of the rodded configurations present in the synthesis case (e.g., ARO, D in, D+C in, etc.). The IFM average power files from these PDQ07 cases are saved for input to the FDELH code. An input dataset is created for FDELH, omitting the axial power sharing values. The job and case ID's for the IFM files are listed in the order that the rodded configurations occur from bottom to top of core. NOMAD reads this input dataset and re-writes it with the axial power sharings which it calculates. FDELH subsequently reads this data and performs the 1-D/2-D synthesis:

$$F_p(x,y) = P_1 \cdot F_{p1}(x,y) + P_2 \cdot F_{p2}(x,y) + P_3 \cdot F_{p3}(x,y), \quad (3.4-1)$$

where

$F_p(x,y)$ = relative power for fuel in location (x, y)

P_n = axial power sharing from NOMAD for rod configuration n

$F_{pn}(x, y) = F_p(x, y)$ for rod configuration n.

The PCEDT code then performs a power census edit which provides the percentage of pins in the core whose relative power is greater than the specified value for percentage values of 1%, 2%, ..., 9%, 10%, 20%, ..., 80%, and 90%.

3.5 FAC Analysis Model

The Vepco FXYZ code provides NOMAD with $F_{XY}(Z)$ values at each burnup step for each different rodded configuration which appears in the load follow calculations. It calculates these based on the three-dimensional power distributions obtained from FLAME and the $F_{\Delta H}$ data from the PDQ07 Discrete model

$$F_{XY}(Z) = \left[\frac{F_Q(X,Y,Z)}{P(Z)} \cdot \frac{F_{\Delta H}(X,Y)}{RPD(X,Y)} \cdot F_{CON} \cdot F_{XEN} \right]_{\text{Maximum}} \quad (3.5-1)$$

where

- $F_Q(X,Y,Z)$ = relative power in node (X,Y,Z) from FLAME
- $P(Z)$ = core average axial power in plane Z from FLAME
- $F_{\Delta H}(X,Y)$ = peak pin power in assembly (X,Y) from PDQ07 Discrete
- $RPD(X,Y)$ = relative power in assembly (X,Y) from FLAME
- $F_{CON} = F_E \cdot F_U$
- F_E = engineering heat flux hot channel factor = 1.03
- F_U = measurement uncertainty factor = 1.05
- F_{XEN} = radial xenon re-distribution correction factor = 1.03.

If $F_{\Delta H}(X,Y)$ is less than $RPD(X,Y)$, a value of 1.0 is substituted for that ratio. NOMAD reads the $F_{XY}(Z)$ data from a file where FXYZ stores them. Usually only data for ARO and D in rod configurations are necessary for FAC analysis. NOMAD then performs the FAC analysis calculation as previously described in Section 2.10.

TABLE 3-1
MACROSCOPIC CROSS SECTION VARIABLE DEPENDENCE

<u>Cross Section</u>	<u>Fuel Temp.</u>	<u>Mod. Temp.</u>	<u>Boron Conc.</u>	<u>(Boron • Mod. Temp.)</u>	<u>Xenon</u>	<u>Xenon²</u>	<u>(Boron • Xenon)</u>
D_1	X	X	X				
Σ_{r1}	X	X	X	X			
Σ_{a1}	X	X	X	X			
$\nu\Sigma_{f1}$	X	X	X				
$\kappa\Sigma_{f1}$	X	X	X				
D_2	X	X	X	X	X		
Σ_{a2}	X	X	X	X	X	X	X
$\nu\Sigma_{f2}$	X	X	X	X	X		
$\kappa\Sigma_{f2}$	X	X	X	X	X		

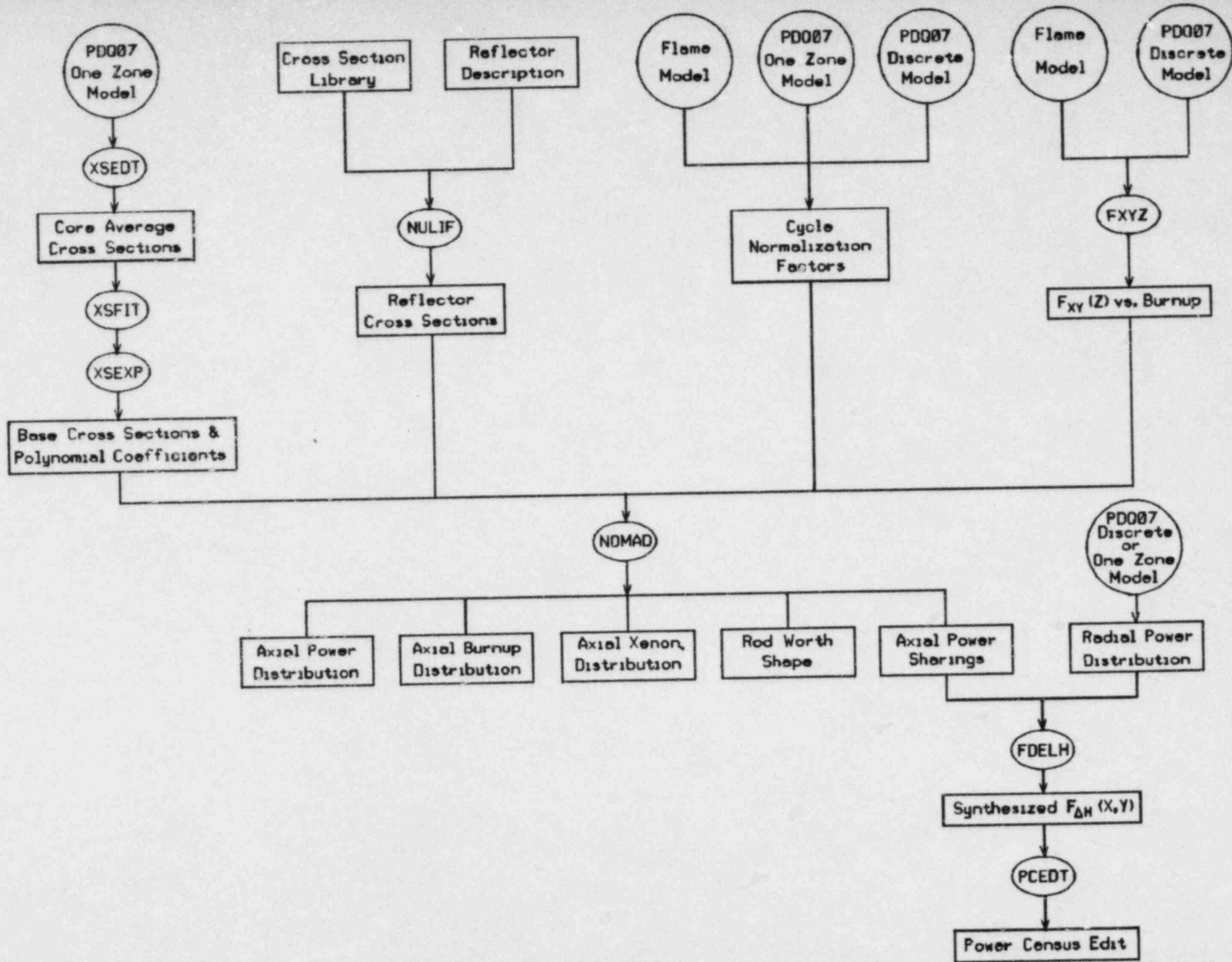


FIGURE 3-1
Vepco NOMAD Model 1 Flow Diagram

SECTION 4 - USER INFORMATION

4.1 Input Description

NOMAD input is read as a series of numbered and categorized cards. The 010 through 060 cards are required at the beginning of every job. However, the 040 card can be omitted if a buckling coefficient table is input. The 070 through 090 cards are optional and should only be input when needed. Each card is free format. The data for a single card number may be continued onto subsequent cards, but the card number only appears on the first card. A 500 card is always the last card for each case (or step), and the user may run up to 749 dependent cases following the independent (i.e., first) case. Any of the numbered input cards may appear in a dependent case, but each case must end with a 500 card.

During the delta-I control and the criticality search options, the power, control rod positions, or boron concentration may be changed by the code. To use the values calculated by the code in the previous step, the user inputs a negative value for the appropriate variable(s) on the 500 card.

A detailed description of each input card begins on the following page.

010 CARD -- Title (1 line, 80 characters maximum)

020 CARD -- General Parameters

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	NRGNS	INTEGER	Number of axial regions
2	EPS	REAL	Eigenvalue convergence limit
3	IWRITE(1)	INTEGER	Temp., power, and burnup edit: 0 => Do not write edit 1 => Print & fiche edit 2 => Fiche edit
4	IWRITE(2)	INTEGER	Macroscopic cross-section edit
5	IWRITE(3)	INTEGER	Iodine, xenon and flux edit
6	IWRITE(4)	INTEGER	Axial xenon and RPD plot file
7	IWRITE(5)	INTEGER	Axial power sharing edit and flux squared edit (must =1 for 1-D/2-D synthesis)

030 CARD -- Thermal-Hydraulic Data

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	ITMAX	INTEGER	Maximum number of thermal iterations
2	TAU	REAL	Thermal relaxation parameter
3	THCON	REAL	Thermal convergence criterion
4	TMO	REAL	Moderator reference temperature (°Fahrenheit)
5	TFO	REAL	Fuel reference temperature (°Rankine)
6	DGEFPD	REAL	Fuel temperature vs. burnup coefficient (°R / (°FPD • relative power))
7	FTFO	REAL	Fuel temp. vs. relative power coefficient (°R / relative power)
8	ENTHIN(1)	REAL	HZP inlet enthalpy (BTU/lbm)
9	ENTHIN(2)	REAL	HFP inlet enthalpy (BTU/lbm)
10	HEIGHT	REAL	Core height (centimeters)
11	POWER	REAL	HFP core power level (watts)
12	FLORAT	REAL	Core flow rate (lbm/hour)
13	SYSPR	REAL	System pressure (psia)
14	POWDEN	REAL	Power density (watts/cc)

040 CARD -- Buckling Coefficient Input

(Not required if buckling coefficient table is used)

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	BO	REAL	Buckling amplitude
2	BMID	REAL	Buckling curvature coefficient
3	BTILT	REAL	Buckling tilt coefficient
4	BTH	REAL	Thermal buckling fraction
5	RPDMID	REAL	Target power at midplane
6	AXOFFT	REAL	Target axial offset
7	ILOCK	INTEGER	Buckling coefficient table flag. Use this card instead of table: 0 => For this step only 1 => For subsequent steps

050 CARD -- Control Rod Cross Sections

(Seven cards: control D-A, shutdown B&A, PL rods)
(Number 050 appears on the first card only)

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	RODWTH(1,I)	REAL	Fast macroscopic absorption cross section
2	RODWTH(2,I)	REAL	Thermal macroscopic absorption cross section
3	RODWTH(3,I)	REAL	Fast macroscopic removal cross section
4	RODWTH(4,I)	REAL	Control rod normalization
5	IOVRLP(I)	INTEGER	Bank overlap (1st 3 cards only)
6	RDBANK(I)	REAL	Bank name (8 characters in ' ')

060 CARD -- Delta-I Control and Criticality Search Parameters

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	IDCNTL	INTEGER	Delta-I control option. If delta-I out of allowed band: -1 => adjust to edge of band 0 => do not adjust +1 => adjust to target value
2	TARGET	REAL	Target delta-I (%)
3	DELTIL	REAL	Lower delta-I band (%)
4	DELTIH	REAL	Upper delta-I band (%)
5	RILHFP	REAL	D-bank HFP insertion limit (steps)
6	RILHAF	REAL	D-bank insertion limit at 50% power (steps)
7	EIGEN	REAL	Criticality search target eigenvalue
8	EPS2	REAL	Criticality search convergence limit
9	ICRIT	INTEGER	Criticality search variable: 1 => Boron concentration (ppm) 2 => D-bank position (steps) -2 => D-bank followed by boron conc. 3 => Power (%) 4 => HFP inlet enthalpy (BTU/lbm)
10	CRTMAX	REAL	Maximum value of ICRIT
11	CRTMIN	REAL	Minimum value of ICRIT

070 CARD -- Core Average Fixed Parameters (Optional Card)

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	IFX1	INTEGER	Fixed fuel and moderator temperature flag: 0 => Do not fix 1 => Fix
2	IFX2	INTEGER	Fixed burnup flag
3	IFX3	INTEGER	Fixed xenon flag
4	IFX4	INTEGER	Time flag: -1 => continuous clock +1 => 24 hour clock
5	TMFX	REAL	Moderator temp. (°F)
6	TFFX	REAL	Fuel temperature (°R)
7	EFPDFX	REAL	Burnup (EFPD)
8	XENFX	REAL	Xenon concentration (Atoms/bn-cm)
9	TIMEFX	REAL	Initial clock time (hours)

080 CARD -- Boration / Dilution Input (Optional Card)

<u>VARIABLE</u>	<u>NAME</u>	<u>TYPE</u>	<u>DESCRIPTION</u>
1	SYSMAS	REAL	Primary system mass (lbm)
2	CBBOR	REAL	Boration line boron conc. (ppm)
3	CBDIL	REAL	Dilution line boron conc. (ppm)
4	FLOBOR	REAL	Boration line flow rate (gpm)
5	FLDIL	REAL	Dilution line flow rate (gpm)
6	NXCRT	REAL	If boron system incapable of maintaining criticality: 0 => Print warning only 1 => Adjust D-bank position 2 => Adjust power level 3 => Adjust inlet enthalpy

090 CARD -- FAC Analysis Input (Optional Card)

VARIABLE	NAME	TYPE	DESCRIPTION
1	FQGRID	REAL	Grid correction factor: 1.056 => 3 case 1.025 => 18 case
2	ADJUST	REAL	F_{XY} power adjustment factor
3	NMAPS	INTEGER	Number of $F_{XY}(Z)$ maps input
4	FXYMIN(I) (I=1,NMAPS)	REAL	Minimum $F_{XY}(Z)$ values (• F_U • F_E) for each $F_{XY}(Z)$ map
5	MPBURN(I) (I=1,NMAPS)	INTEGER	Burnup (MWD/MTU) at which each $F_{XY}(Z)$ map is printed
6	FQLIMX(J)	REAL	X-coordinate for F_Q limit curve (core height in feet)
7	FQLIMY(J)	REAL	Y-coordinate for F_Q limit curve ($F_Q(Z)$ limit) [Repeat variables 6 & 7 in pairs for J=1,4]
8	FXYLMX(J)	REAL	X-coordinate for F_{XY} limit line (core height in feet)
9	FXYLMY(J)	REAL	Y-coordinate for F_{XY} limit line ($F_{XY}(Z)$ limit) [Repeat variables 8 & 9 in pairs for J=1,4]

100 CARD -- Recovery File Options (Optional Card)

VARIABLE	NAME	TYPE	DESCRIPTION
1	DFILE	REAL	Recovery file flag SAVE => Save new recovery file RESTORE => Restore old file
2	FNAME	REAL	8 character file name
[Note: 1 space must be placed between variables 1 and 2]			

500 CARD -- Case Card

VARIABLE	NAME	TYPE	DESCRIPTION
1	TIME	REAL	Depletion time interval (hours)
2	PCTPOW	REAL	Core average power (%) ¹
3	IRDPOS(1)	INTEGER	D-Bank position (steps) ¹
4	IRDPOS(2)	INTEGER	C-Bank position (steps) ¹
5	IRDPOS(3)	INTEGER	B-Bank position (steps) ¹
6	IRDPOS(4)	INTEGER	A-Bank position (steps) ¹
7	IRDPOS(5)	INTEGER	SB-Bank position (steps) ¹
8	IRDPOS(6)	INTEGER	SA-Bank position (steps) ¹
9	IRDPOS(7)	INTEGER	PL-Rods position (steps) ¹
10	BORON	REAL	Boron Concentration (ppm) ¹
11	IXEN	INTEGER	Xenon Option: -1 => Xenon from previous step 0 => Xenon depletion 1 => No xenon 2 => Equilibrium xenon
12	IOPT	INTEGER	Case Option: 0 => Static case 1 => Depletion 2 => 1st step of rod worth sequence 3 => Criticality search 4 => 1st step of FAC analysis 5 => Xenon worth calculation 6 => Buckling coefficient search 7 => Frozen THF 8 => 1st step of average power distribution calculation -8 => Final step of average power distribution calculation; perform load follow depletion

¹ If a negative value is input for any of these variables, the value from the end of the previous case is used.

NOMAD also reads a cycle/geometry deck. The data in this deck remains constant for a particular unit and cycle. The cycle deck contains reflector macroscopic cross sections, xenon parameters, axial region dimensions, and the BOC axial burnup distribution.

The cycle/geometry deck is read in free format in the following order:

D_1	Σ_{a1}	Σ_{r1}	(Bottom reflector fast cross sections)
D_2	Σ_{a2}		(Bottom reflector thermal cross sections)
D_1	Σ_{a1}	Σ_{r1}	(Top reflector fast cross sections)
D_2	Σ_{a2}		(Top reflector thermal cross sections)
Xe σ_{a1}	Xe σ_{a2}	γ_I	γ_{Xe} (Iodine and xenon parameters)
Height(cm)	BOC burnup(=0.0)		(Region 1 -- bottom reflector)
Height(cm)	BOC burnup(=0.0)		(Region 2 -- bottom reflector)
Height(cm)	BOC burnup(=0.0)		(Region 3 -- bottom reflector)
Height(cm)	BOC burnup(EFPD)		(Region 4 -- bottom fuel region)
.	.	.	.
.	.	.	.
.	.	.	.
Height(cm)	BOC burnup(EFPD)		(Region NRGNS-3 -- top fuel region)
Height(cm)	BOC Burnup(=0.0)		(Region NRGNS-2 -- top reflector)
Height(cm)	BOC Burnup(=0.0)		(Region NRGNS-1 -- top reflector)
Height(cm)	BOC Burnup(=0.0)		(Region NRGNS -- top reflector)

4.2 Error & Warning Messages

NOMAD prints error messages whenever it detects an error in input or execution. If it is an input error, it will terminate after checking the remainder of the input for that case for errors. If the error occurs during execution, the job is terminated.

The code also prints warning messages for less severe problems, such as a criticality search not converging. Execution continues after a warning message is printed.

NOMAD returns a condition code of zero for a successful job

completion in most cases. Other programmed return codes are:

- 11 - Power sharing calculated for 1-D/2-D synthesis
- 333 - Job terminated, recovery file could not be found
- 444 - Job terminated, thermal-hydraulic feedback did not converge
- 555 - Job terminated, maximum number of cases exceeded
- 666 - Job terminated, buckling search did not converge
- 777 - Job terminated, cross section file error
- 888 - Job terminated, FAC analysis input error
- 999 - Job terminated, input error.

4.3 Execution Time

Approximate execution (CPU) times for NOMAD for several types of cases are given below:

HZP, no xenon	0.1 seconds
HFP, eq. xenon	1.1 seconds
HFP, xenon depletion, criticality search	0.7 seconds
HFP, eq. xenon, criticality search	3.4 seconds
Buckling coefficient search (10 depletion steps)	4 - 6 minutes
FAC analysis (72 hour load follow)	5 - 8 minutes.

4.4 Output

NOMAD offers the user flexibility in its output control. The output options are IWRITE(1) - IWRITE(5) on the 020 card. All five of these options are turned off and on by values of 0 and 1, respectively. All edits are written to the printer except IWRITE(4), which writes to a plot file.

The temperature, power, and burnup edit is a one page edit which includes the k-effective, delta-I, axial offset, number of iterations to reach convergence, and cycle average burnup. It also prints for each axial region and the core average the values of each of the following parameters: k-infinity, moderator enthalpy (BTU/lbm), moderator temperature ($^{\circ}$ F), fuel temperature ($^{\circ}$ R), relative power density (RPD), and fuel burnup (EFPH). This edit is generally the most useful one in design calculations. The macroscopic cross section edit is a two page edit which provides the fast and thermal cross sections for each axial region and the core average. This edit is recommended only for debugging purposes or special applications.

The iodine, xenon, and flux edit lists the xenon and iodine concentrations and the fast and thermal fluxes by axial region and core average. This edit is required for normalization of the xenon model. The axial power sharing edit gives the fraction of the axial power which exists in each axial segment with a different rod configuration. This edit is necessary to perform 1-D/2-D synthesis. The flux squared sharing edit gives the fraction of the fast group flux squared which exists in each axial segment with a different rod configuration. This edit is an importance weighting function which may be used in performing rod swap worth calculations.

These edits are normally printed once per case, but they may appear more than once when a criticality search or a delta-I control adjustment is performed. The first occurrence of each edit contains the results of the nuclear calculation prior to any variable adjustments. The final occurrence contains the results of the final nuclear calculation after all searches have been completed.

In addition to the output options selected, NOMAD always prints the following output:

- (1) Input card image listing
- (2) Cycle/geometry deck card image listing
- (3) Buckling coefficients & distribution and reflector cross sections
- (4) Input summary
- (5) 1-D analysis case summary.

NOMAD offers two options for writing output to a dataset for creating plots. The first is IWRITE(4), which writes the axial xenon and relative power distributions to a file. Each distribution is labeled with the cycle burnup for that step, so that several distributions may be saved in the same file. The user is advised to not use this option in any job where IOPT = 2, 4, or 5 (rod worth, FAC analysis, or xenon worth). These options also write data to the same plot file. The plot output for these three options is similar to the printed output for them.

The second plot option writes most of the data which appears in the 1-D Analysis Case Summary to a file. The data includes the case number, time, power, boron, D-bank position, delta-I, peak power, k-effective, and xenon. If the boration/dilution calculations are performed, the water processed per step is written instead of the xenon concentration. This option is exercised by assigning a dataset to the appropriate unit (see Section 4.5).

4.5 I/O Units

The I/O units used by NOMAD are:

<u>Unit No.</u>	<u>Description</u>
2	Buckling coefficient table (input)
3	Cycle/geometry input deck
4	Cross section input
5	Card input
6	Printer and microfiche
8	$F_{XY}(Z)$ input
9	Case summary plot file
10	Plot file for: RPD, xenon dist., rod worth, FAC analysis, xenon worth
11	Buckling coefficient table (output) ¹
12	Microfiche
13	Recovery file
14	Scratch disk space
21	FDELH code input for 1-D/2-D synthesis.

¹Data is written to unit 11 only when a buckling coefficient search is performed.

SECTION 5 - RESULTS

5.1 Introduction

The purpose of this section is to demonstrate the predictive capability of the NOMAD model for calculations of axial power distributions, delta-I and axial offsets, critical boron concentrations, differential and integral control rod worths, load follow maneuvers, and core peaking limits for FAC analysis. This section presents: 1) reactivity parameter comparisons to measured data from the Surry and North Anna Nuclear Power Stations and to other Vepco codes; 2) a thermal-hydraulic feedback calculation comparison to COBRA¹¹; 3) axial power distribution comparisons to measured data from Surry and North Anna; 4) differential and integral control rod worth comparisons to measured data and the Vepco FLAME model; 5) comparisons of load follow maneuver simulations of delta-I, axial offset, and critical boron concentration to measured data from North Anna; and 6) FAC analysis results obtained with NOMAD.

5.2 Reactivity Parameters

Differential boron worths were calculated with NOMAD at BOC, HZP and EOC, HFP. The BOC, HZP values are compared to measured data and the EOC, HFP values are compared to the Vepco PDQ07 Discrete model in Table 5-1. Isothermal temperature coefficients obtained with NOMAD at BOC, HZP also are compared to measured data in Table 5-1.

Figures 5-1 through 5-6 compare NOMAD xenon worths after startup, orderly shutdown, and trip to results obtained with XETRN¹², Vepco's zero-dimensional xenon transient code. Note that XETRN assumes the xenon worth is a linear function of the xenon concentration, whereas NOMAD calculates the xenon worth directly from the eigenvalues.

5.3 Thermal-Hydraulic Feedback

The accuracy of the thermal-hydraulic feedback model has been verified by direct comparison with the COBRA code. Both NOMAD and COBRA were run for a North Anna 120% overpower case. The COBRA model used four 17 x 17 fuel assemblies to represent the core. COBRA also used as input the axial power distribution calculated by NOMAD. The system pressure, core flow rate, and inlet enthalpy were identical for both COBRA and NOMAD. Table 5-2 shows a comparison of the moderator enthalpy and moderator temperature distributions calculated by NOMAD and COBRA.

5.4 Axial Power Distribution

Axial power distribution comparisons between the Vepco NOMAD model and measurements are presented in Figures 5-7 through 5-16 for BOC, HZP, no xenon and BOC, HFP, equilibrium xenon conditions. Representative axial power distributions comparisons are shown for North Anna 1 Cycle 2, North Anna 1 Cycle 3, North Anna 1 Cycle 4, North Anna 2 Cycle 2, and Surry 1 Cycle 6.

The NOMAD predictions attempt to simulate the actual core conditions. However, NOMAD does not represent the spacer grids in order to increase calculation efficiency. The accuracy which is compromised is insignificant.

5.5 Differential and Integral Rod Worths

The Vepco NOMAD model predictions and startup physics measurements for differential and integral control rod bank worths for B-bank are compared in Figures 5-17 through 5-24. B-bank was the rod swap reference bank for each of these cycles and its worth was measured by boron dilution.

In addition, NOMAD results for Banks A through D moving in overlap

are compared to the Vepco FLAME model in Figures 5-25 through 5-32. No measurements were performed at these conditions.

Integral control rod bank worths were calculated for banks measured by rod swap using an importance weighting technique. NOMAD performed a critical boron search with the reference bank fully inserted. Another bank was then fully inserted, and NOMAD performed a criticality search on the reference bank position and calculated the flux-squared sharings. These flux-squared sharings were then used to determine a weighted average of the PDQ07 Discrete bank worths for the bank inserted alone and the bank inserted with the reference bank. These synthesized bank worths and the measured worths are compared in Table 5-3. The critical position of the reference bank predicted by NOMAD for each bank fully inserted is compared to measurement in Table 5-4.

5.6 Load Follow Manuever Simulation

NOMAD's load follow simulation capability has been verified by comparison to three sets of measured data for load follow type cases. The first set of data consists of hourly delta-I readings and two critical boron measurements from a 70% load reduction test performed near the end of North Anna Unit 1 Cycle 2. The power and D-bank history for this case is listed in Table 5-5. Figures 5-33 and 5-34 compare NOMAD results to the measured delta-I and critical boron concentrations.

Two additional sets of measured data were recorded near the end of North Anna Unit 1 Cycle 3 during power escalations following reactor trips. The first incident occurred on April 16-20, 1982, and the second on April 30 - May 2, 1982. The power and D-bank histories for these two cases are given in Tables 5-6 and 5-7. The negative times listed in Table 5-6 are simply the number of hours before the comparisons in Figures 5-35 and 5-36 begin. The data prior to 0.0 hours was not plotted because it was at HFP, equilibrium conditions (delta-I is virtually

constant) or low power levels (no delta-I data available). Hourly readings of delta-I and eight critical boron measurements were taken during the first case. Results from the NOMAD simulation are plotted versus these data in Figures 5-35 and 5-36, respectively. During the second case, both ex-core delta-I readings and INCORE axial offset measurements were performed, since delta-I cannot be measured accurately at low power levels. (The INCOREs were performed on only a limited number of assemblies each time.) The delta-I readings have been converted to axial offsets in order to compare NOMAD results to both types of data in Figure 5-37. Figure 5-38 plots the NOMAD critical boron concentrations versus thirteen measured values for this case.

5.7 FAC Analysis

Standard three and eighteen case FAC (CAOC) analyses were performed with the Vepco NOMAD model for North Anna 1 Cycle 4 and North Anna 2 Cycle 2. The Vepco NOMAD model results were found to be consistent with the reload analysis results from an accepted and verified vendor model which has been used in the design and licensing of the Surry and North Anna reactors. Both models indicated minor technical specification violations near the core bottom in the three case analyses and no violations in the eighteen case analyses.

Figures 5-39 through 5-42 show the NOMAD results for the eighteen case analyses. The $F_Q(Z)$ plots in Figures 5-40 and 5-42 contain the radial xenon re-distribution factor and an uncertainty factor of 10.9%, which includes the engineering hot channel factor, the measurement uncertainty factor, and the grid correction factor.

TABLE 5-1
REACTIVITY COEFFICIENTS COMPARISON

Differential Boron Worth, BOC HZP (pcm/ppm)

Unit/Cycle	NOMAD	PDQ Discrete	Measured	%Difference ¹
N1C2	-9.15	-9.10	-8.88	3.04
N1C3	-8.15	-8.08	-8.54	-4.57
N1C4	-7.70	-8.04	-8.25	-6.67
N2C2	-8.97	-8.91	-8.46	6.03
S1C6	-8.27	-8.31	-8.78	-5.81
S1C7	-8.43	-8.44	-8.44	-0.12

Differential Boron Worth, EOC HFP (pcm/ppm)

Unit/Cycle	NOMAD	PDQ Discrete	%Difference ²
N1C2	-9.06	-9.43	-3.92
N1C3	-8.17	-8.47	-3.54
N1C4	-7.90	-8.62	-8.35
N2C2	-8.91	-9.40	-5.21
S1C6	-8.81	-9.31	-5.37
S1C7	-8.93	-9.16	-2.51

Isothermal Temperature Coefficient, BOC HZP (pcm/°F)

Unit/Cycle	NOMAD	PDQ One Zone	Measured	Difference ³
N1C2	-5.07	-3.87	-2.36	-2.71
N1C3	-3.99	-3.40	-4.36	0.37
N1C4	-4.90	-3.52	-4.92	0.02
N2C2	-5.29	-3.27	-2.27	-3.02
S1C6	-4.28	-3.79	-2.32	-1.96
S1C7	-6.49	-5.68	-5.85	-0.64

¹ % Difference = (NOMAD - Measured) / Measured x 100

² % Difference = (NOMAD - PDQ07)/PDQ07 x 100

³ Difference = NOMAD - Measured

TABLE 5-2

COMPARISON OF NOMAD AND COBRA
MODERATOR ENTHALPY AND TEMPERATURE DISTRIBUTIONS

Position (inches)	Enthalpy (BTU/lbm)		Moderator Temperature (°F)	
	NOMAD	COBRA	NOMAD	COBRA
4	544.9	545.1	548.4	548.0
8	546.8	546.9	549.9	549.5
12	549.1	549.3	551.8	551.4
16	551.8	552.0	554.0	553.5
20	554.7	554.8	556.3	555.8
24	557.7	557.8	558.7	558.2
28	560.8	560.9	561.1	560.6
32	563.9	564.0	563.6	563.0
36	567.0	567.1	566.0	565.5
40	570.2	570.2	568.4	567.8
44	573.3	573.4	570.9	570.3
48	576.5	576.6	573.3	572.6
52	579.6	579.7	575.7	575.0
56	582.8	582.9	578.1	577.3
60	586.0	586.1	580.4	579.7
64	589.2	589.3	582.8	582.1
68	592.4	592.5	585.2	584.5
72	595.7	595.8	587.5	586.9
76	598.9	599.0	589.9	589.3
80	602.2	602.3	592.2	591.7
84	605.5	605.6	594.6	594.1
88	608.8	608.9	596.9	596.5
92	612.2	612.3	599.2	598.9
96	615.5	615.6	601.5	601.3
100	618.9	619.0	603.8	603.7
104	622.3	622.3	606.1	606.4
108	625.6	625.7	608.4	608.3
112	629.1	629.1	610.7	610.6
116	632.5	632.5	612.9	612.9
120	635.5	635.9	615.2	615.1
124	639.1	639.2	617.3	617.4
128	642.4	642.4	619.4	619.5
132	645.3	645.4	621.3	621.4
136	648.0	648.0	623.0	623.0
140	650.1	650.1	624.4	624.3
144	651.6	651.5	625.3	625.2

TABLE 5-3

ROD SWAP COMPARISON, PART 1

Integral Bank Worths (pcm)

North Anna Unit 1 Cycle 3

<u>Bank</u>	<u>NOMAD/PDQ</u>	<u>Measured</u>	<u>%Difference</u>
D	1048	1089	-3.76
C	839	777	7.98
A	620	722	-14.13
SB	1006	919	9.47
SA	1096	1238	-11.47

North Anna Unit 1 Cycle 4

D ¹	N/A	N/A	N/A
C	808	843	-4.15
A	479	562	-14.77
SB	980	1023	-4.20
SA	997	1094	-8.87

North Anna Unit 2 Cycle 2

D	1010	1015	-0.49
C	780	757	3.04
A	757	812	-6.77
SB	713	664	7.38
SA	912	948	-3.80

Surry Unit 1 Cycle 6

D	1228	1234	-0.49
C	819	815	0.49
A	538	551	-2.36
SB	1018	1013	0.49
SA	1093	1137	-3.87

¹D-bank worth was measured by a combination of rod swap & dilution.

TABLE 5-4

ROD SWAP COMPARISON, PART 2

Reference Bank Critical Position (Steps)

North Anna Unit 1 Cycle 3

<u>Bank</u>	<u>NOMAD</u>	<u>Measured</u>	<u>%Difference</u>
D	143	143	0
C	117	103	14
A	97	97	0
SB	143	119	24
SA	158	170	-12

North Anna Unit 1 Cycle 4

D ¹	N/A	N/A	N/A
C	154	163	-9
A	114	122	-8
SB	182	190	-8
SA	186	201	-15

North Anna Unit 2 Cycle 2

D	186	195	-9
C	171	168	3
A	167	175	-8
SB	166	155	11
SA	181	189	-8

Surry Unit 1 Cycle 6

D	179	175	4
C	123	110	13
A	93	83	10
SB	154	138	16
SA	166	157	9

¹D-bank worth was measured by a combination of rod swap & dilution.

TABLE 5-5

N1C2 70% LOAD REDUCTION TEST
POWER AND D-BANK HISTORY

Time (Hours)	Power (%)	D-bank (steps)
00.00	98.8	228
01.00	82.9	190
02.00	56.2	186
03.00	32.8	150
04.00	29.5	152
05.00	29.2	166
06.00	29.2	162
07.00	29.6	157
08.00	30.0	156
09.00	30.6	145
10.00	30.5	143
11.00	30.3	145
12.00	30.2	141
13.00	30.2	140
14.00	29.9	138
15.00	29.5	137
16.00	29.5	137
17.00	28.9	136
18.00	29.3	136
19.00	29.0	136
20.00	27.9	136
21.00	28.4	137
22.00	28.4	137
23.00	27.5	145
24.00	27.3	151
25.00	27.2	155
26.00	27.2	157
27.00	26.7	160
28.00	26.7	160
29.00	26.7	160
30.00	26.7	160

TABLE 5-6

N1C3 SHUTDOWN/RETURN TO POWER CASE 1
POWER AND D-BANK HISTORY

Time (Hours)	Power (%)	D-bank (steps)
-23.00	100.9	215
-22.00	100.4	215
-21.00	100.6	215
-20.00	100.5	215
-19.00	100.6	215
-18.00	100.8	215
-17.00	100.7	215
-16.00	100.6	215
-15.00	100.5	215
-14.00	100.0	215
-13.00	99.9	214
-12.00	100.6	215
-11.00	100.5	215
-10.00	100.6	215
-9.00	0.0	0
-8.00	0.0	0
-7.00	0.0	0
-6.00	0.2	47
-5.00	1.2	156
-4.00	1.9	179
-3.00	0.0	0
-2.00	0.2	185
-1.00	4.9	193
0.0	17.7	197
1.00	23.5	186
2.00	27.1	160
3.00	29.7	160
4.00	30.1	160
5.00	30.2	160
6.00	29.8	160
7.00	30.7	160
8.00	29.3	158
9.00	29.2	151
10.00	29.4	143
11.00	29.4	143
12.00	29.4	142
13.00	30.9	141
14.00	45.7	141
15.00	48.9	144
16.00	49.0	159
17.00	48.7	170
18.00	48.4	178
19.00	48.0	181
20.00	47.5	182
21.00	47.3	183
22.00	47.6	184
23.00	47.7	184
24.00	47.3	182
25.00	47.7	180

TABLE 5-6 (Continued)

26.00	47.7	178
27.00	48.8	177
28.00	49.0	175
29.00	49.0	172
30.00	48.4	170
31.00	48.9	169
32.00	48.4	168
33.00	48.4	168
34.00	47.7	168
35.00	48.3	168
36.00	48.4	167
37.00	48.8	165
38.00	55.6	172
39.00	69.9	181
40.00	73.4	183
41.00	73.2	183
42.00	76.9	190
43.00	87.7	205
44.00	92.0	209
45.00	99.6	211
46.00	99.7	211
47.00	100.2	211
48.00	99.6	211
49.00	99.8	211
50.00	100.0	211
51.00	100.1	211
52.00	99.8	211
53.00	100.0	211
54.00	100.3	211
55.00	100.2	211
56.00	100.2	211
57.00	100.2	211
58.00	0.0	0
59.00	0.0	0
60.00	0.0	0
61.00	1.4	88
62.00	5.1	128
63.00	26.9	200
64.00	28.3	182
65.00	28.1	172
66.00	30.1	160
67.00	30.0	160
68.00	30.0	160
69.00	29.8	160
70.00	44.3	170
71.00	50.4	161
72.00	49.5	161
73.00	47.2	161
74.00	47.6	161
75.00	47.2	161
76.00	47.1	158
77.00	47.3	158

TABLE 5-7

N1C3 SHUTDOWN/RETURN TO POWER CASE 2
POWER AND D-BANK HISTORY

Date	Time (Hours)	Power (%)	D-bank (steps)
4 30 82	16.20	100.3	218
4 30 82	18.60	99.7	218
4 30 82	18.85	99.8	218
4 30 82	20.72	100.2	218
4 30 82	21.50	95.2	209
4 30 82	21.78	92.1	206
4 30 82	22.06	88.3	200
4 30 82	22.35	83.6	190
4 30 82	22.63	75.8	180
4 30 82	23.15	66.0	171
4 30 82	23.70	58.2	158
5 1 82	0.50	44.0	127
5 1 82	1.30	30.1	110
5 1 82	2.08	18.2	110
5 1 82	2.85	3.2	90
5 1 82	3.87	2.3	145
5 1 82	4.70	2.0	162
5 1 82	5.77	2.1	173
5 1 82	6.78	2.3	177
5 1 82	7.81	2.5	174
5 1 82	8.84	2.2	169
5 1 82	9.88	0.0	0
5 1 82	10.88	0.0	0
5 1 82	11.90	0.0	0
5 1 82	12.95	0.0	0
5 1 82	13.20	0.0	91
5 1 82	13.53	0.0	91
5 1 82	13.95	1.3	91
5 1 82	14.27	1.3	91
5 1 82	14.95	1.7	81
5 1 82	15.73	1.7	81
5 1 82	15.95	1.3	69
5 1 82	16.85	1.8	61
5 1 82	17.58	1.8	61
5 1 82	17.85	1.8	59
5 1 82	18.95	1.6	59
5 1 82	19.95	2.1	59
5 1 82	21.00	1.8	45
5 1 82	22.07	1.8	45
5 1 82	23.07	1.7	45
5 1 82	23.97	2.1	45
5 2 82	1.00	2.2	45
5 2 82	2.02	2.3	45
5 2 82	3.07	2.2	45
5 2 82	3.95	2.4	45
5 2 82	4.37	2.4	45
5 2 82	4.98	2.2	45
5 2 82	6.01	2.3	45
5 2 82	6.35	2.3	45

TABLE 5-7 (Continued)

5	2	82	6.95	2.3	45
5	2	82	8.00	2.8	46
5	2	82	8.17	7.3	50
5	2	82	8.53	7.3	50
5	2	82	9.05	7.3	64
5	2	82	9.17	17.6	63
5	2	82	9.44	22.6	63
5	2	82	10.14	29.8	68
5	2	82	10.55	29.8	68
5	2	82	11.00	31.8	69
5	2	82	12.25	31.8	71
5	2	82	13.05	31.8	71
5	2	82	13.25	29.1	71
5	2	82	14.22	39.9	108
5	2	82	15.19	51.6	155
5	2	82	16.21	64.8	174
5	2	82	17.31	67.1	181
5	2	82	18.21	68.7	185
5	2	82	19.21	69.2	185
5	2	82	20.21	73.9	193
5	2	82	21.24	83.0	201
5	2	82	22.24	91.1	208
5	2	82	23.24	96.1	214
5	2	82	23.87	98.7	214

XENON WORTH AFTER STARTUP

NORTH ANNA UNIT 1 CYCLE 3

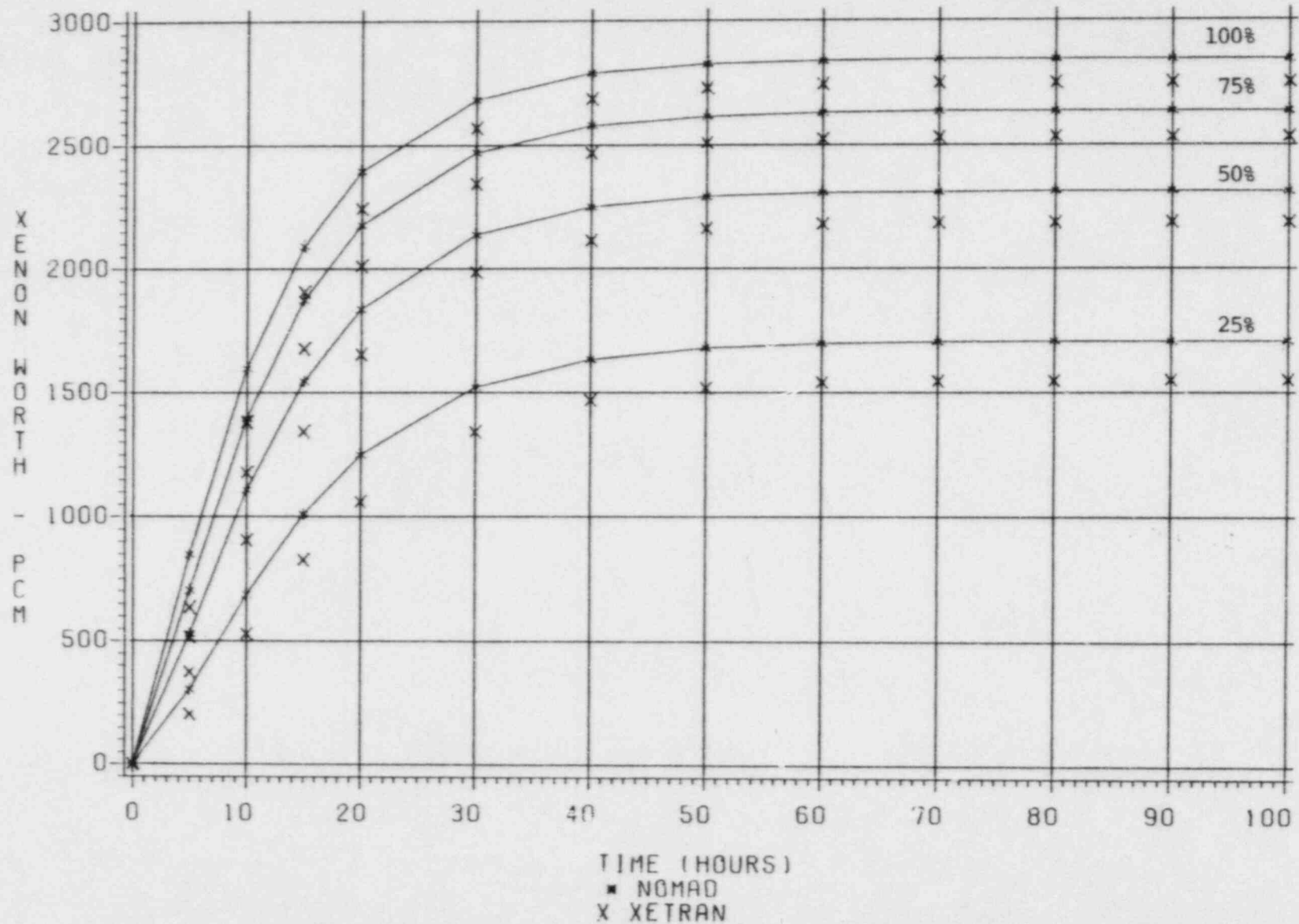


FIGURE 5-1

XENON WORTH AFTER SHUTDOWN

NORTH ANNA UNIT 1 CYCLE 3

FIGURE 5-2

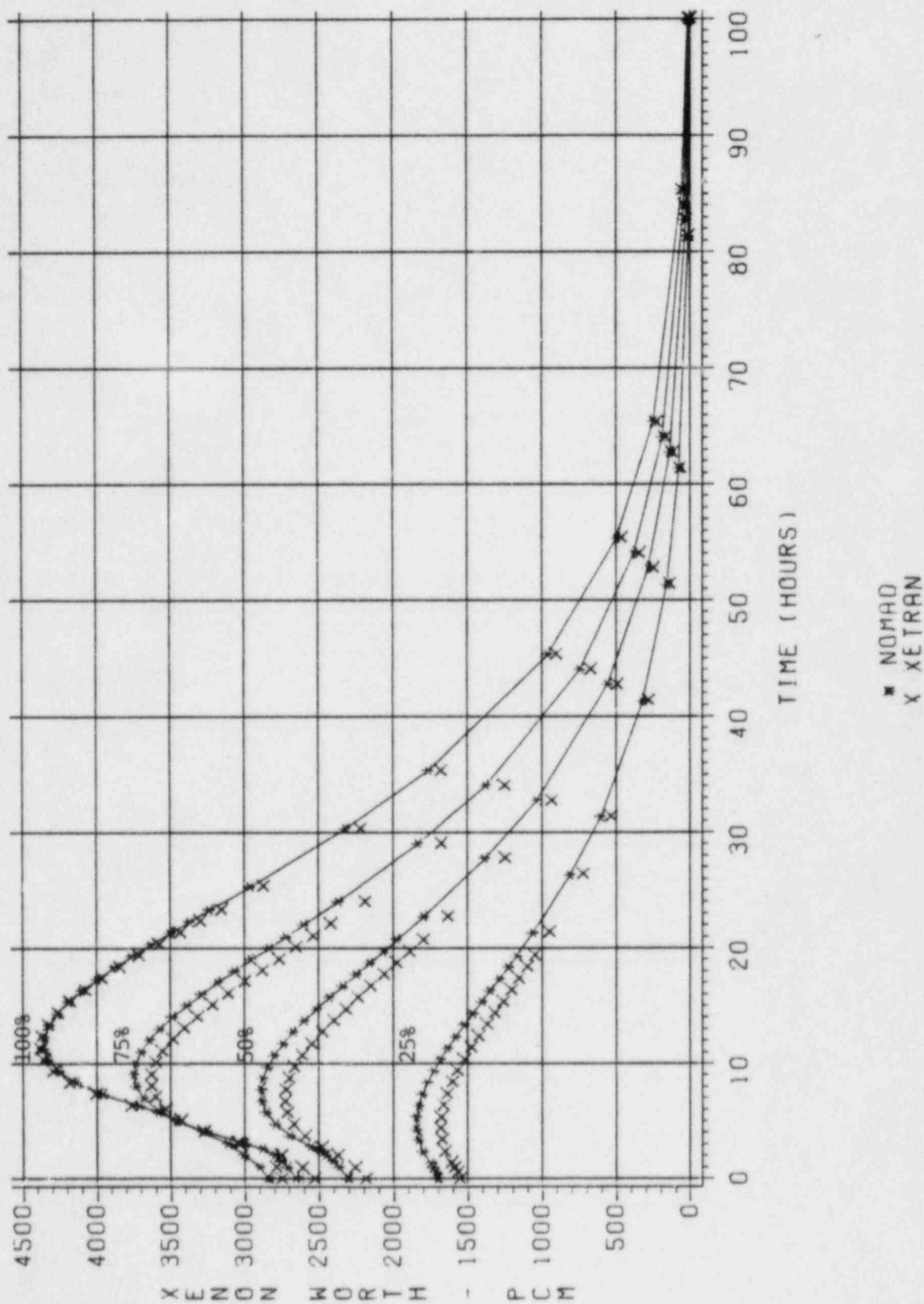
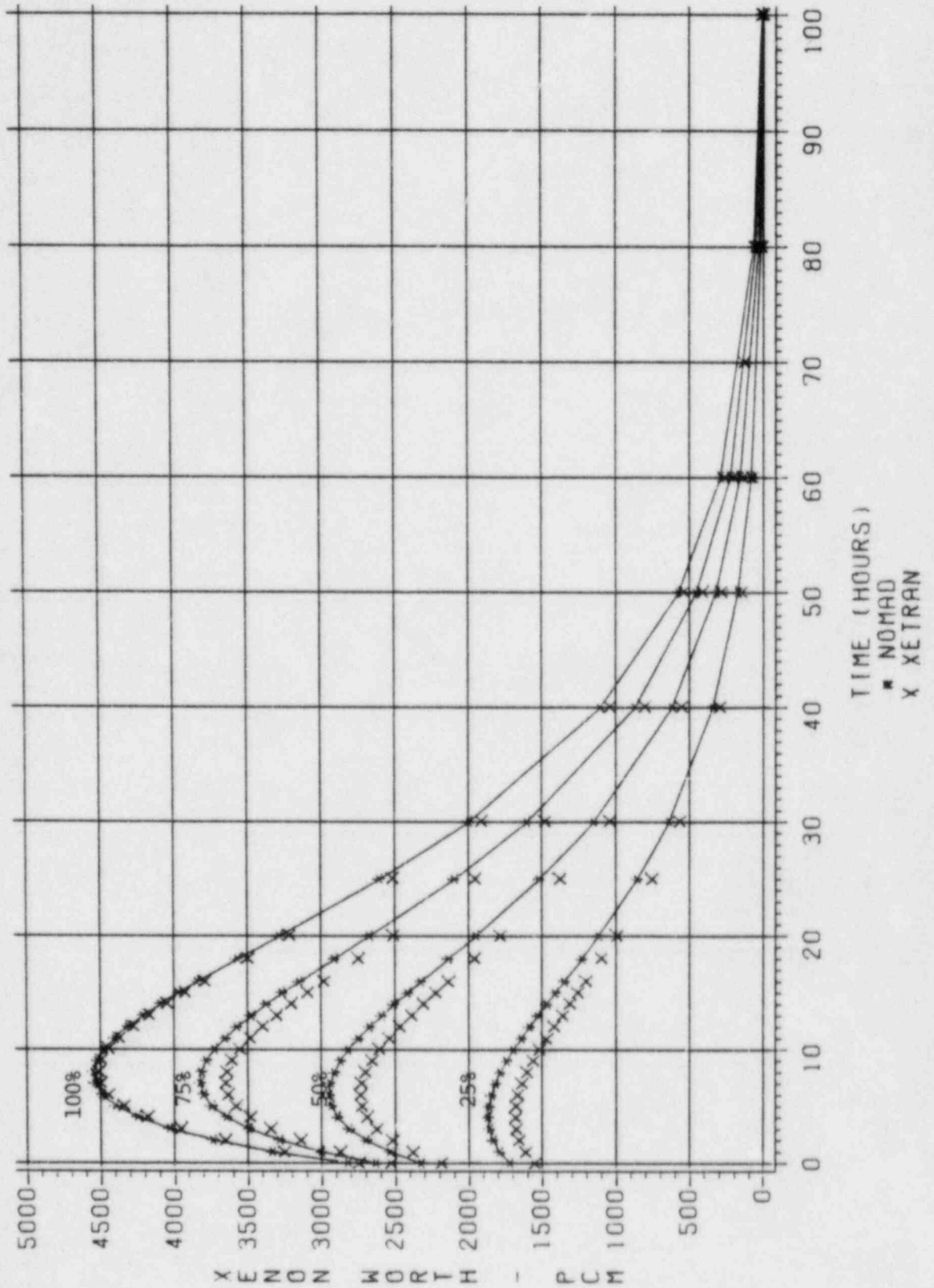


FIGURE 5-3

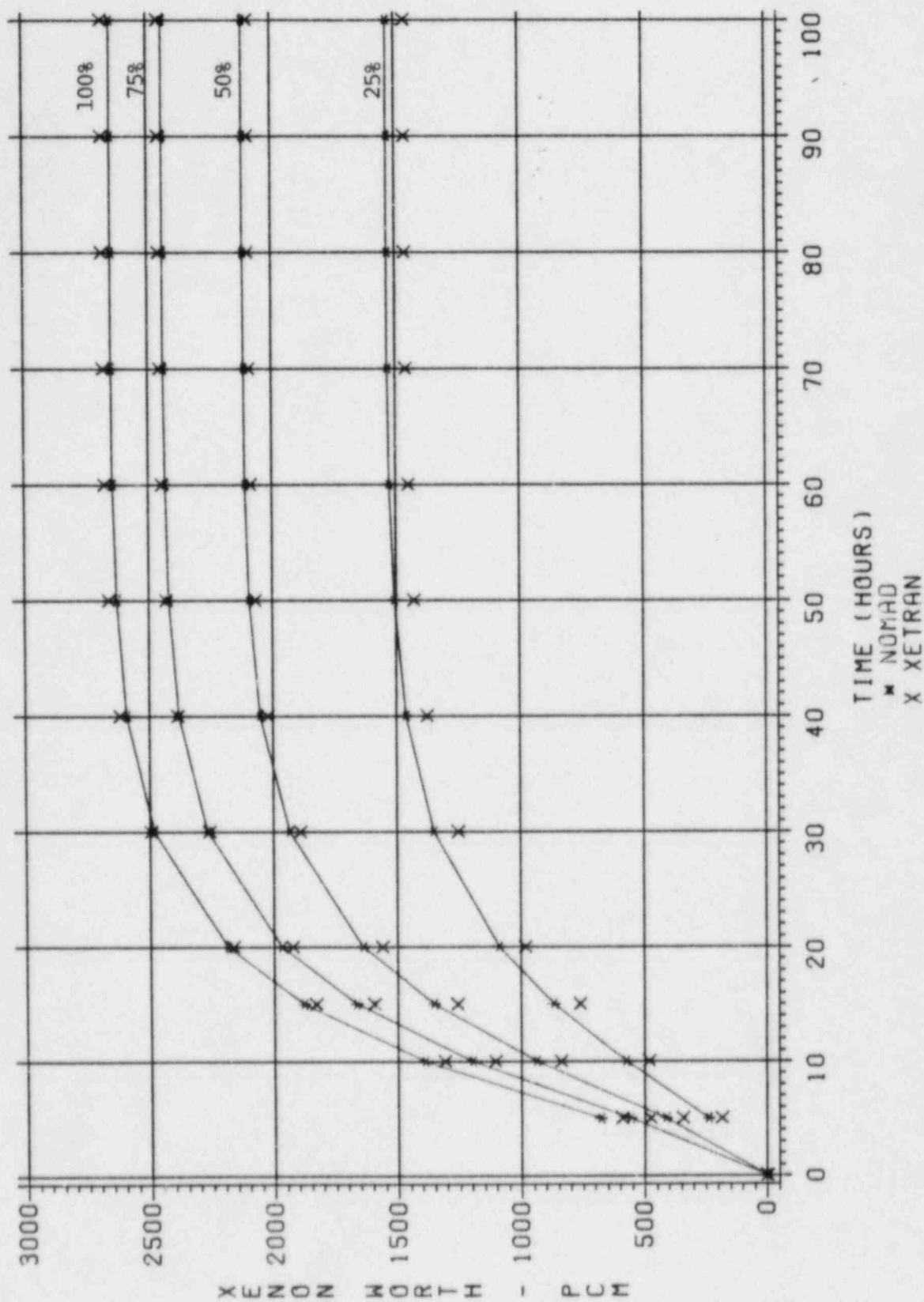
XENON WORTH AFTER TRIP NORTH ANNA UNIT 1 CYCLE 3



XENON WORTH AFTER STARTUP

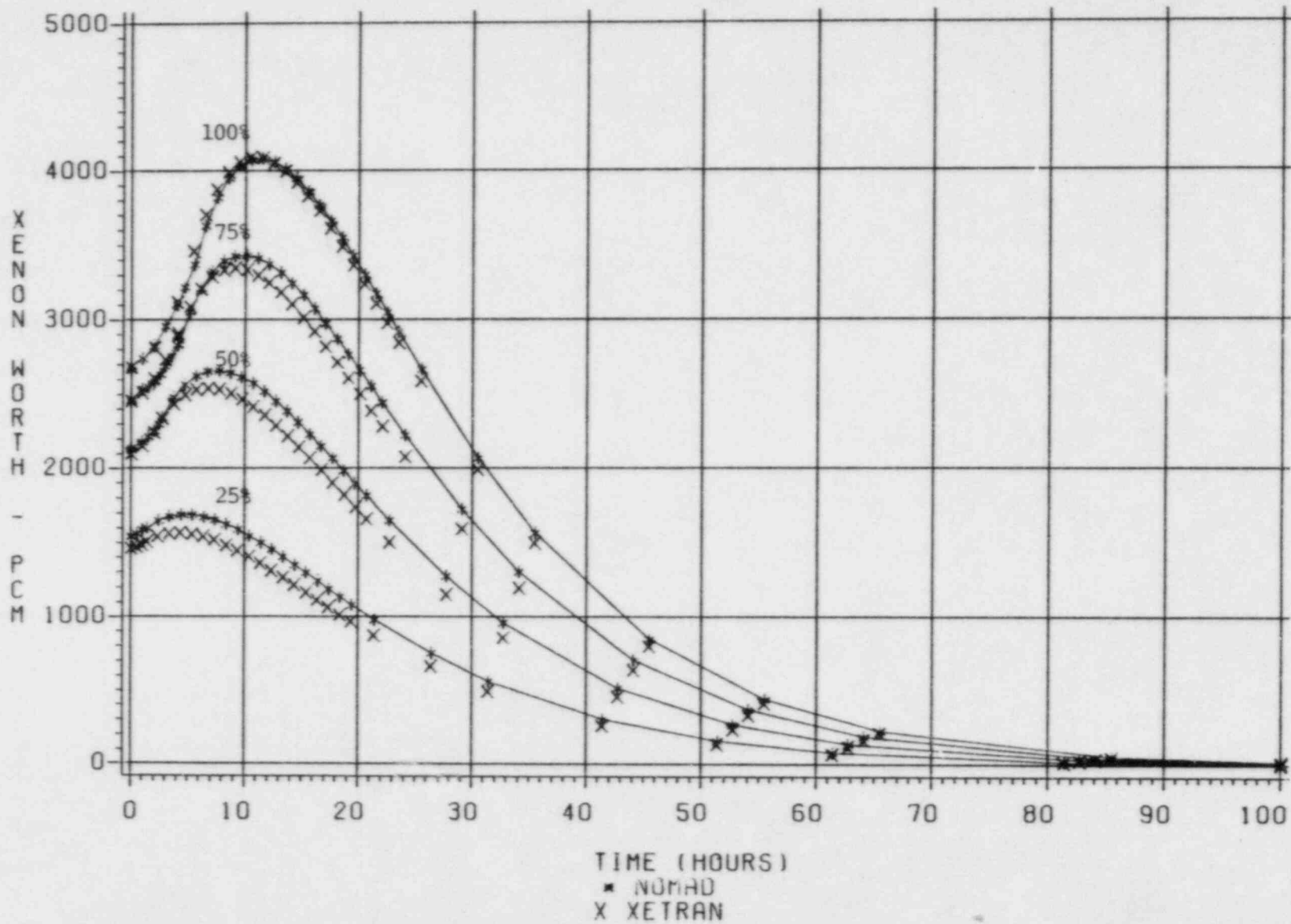
SURRY UNIT 1 CYCLE 6

FIGURE 5-4



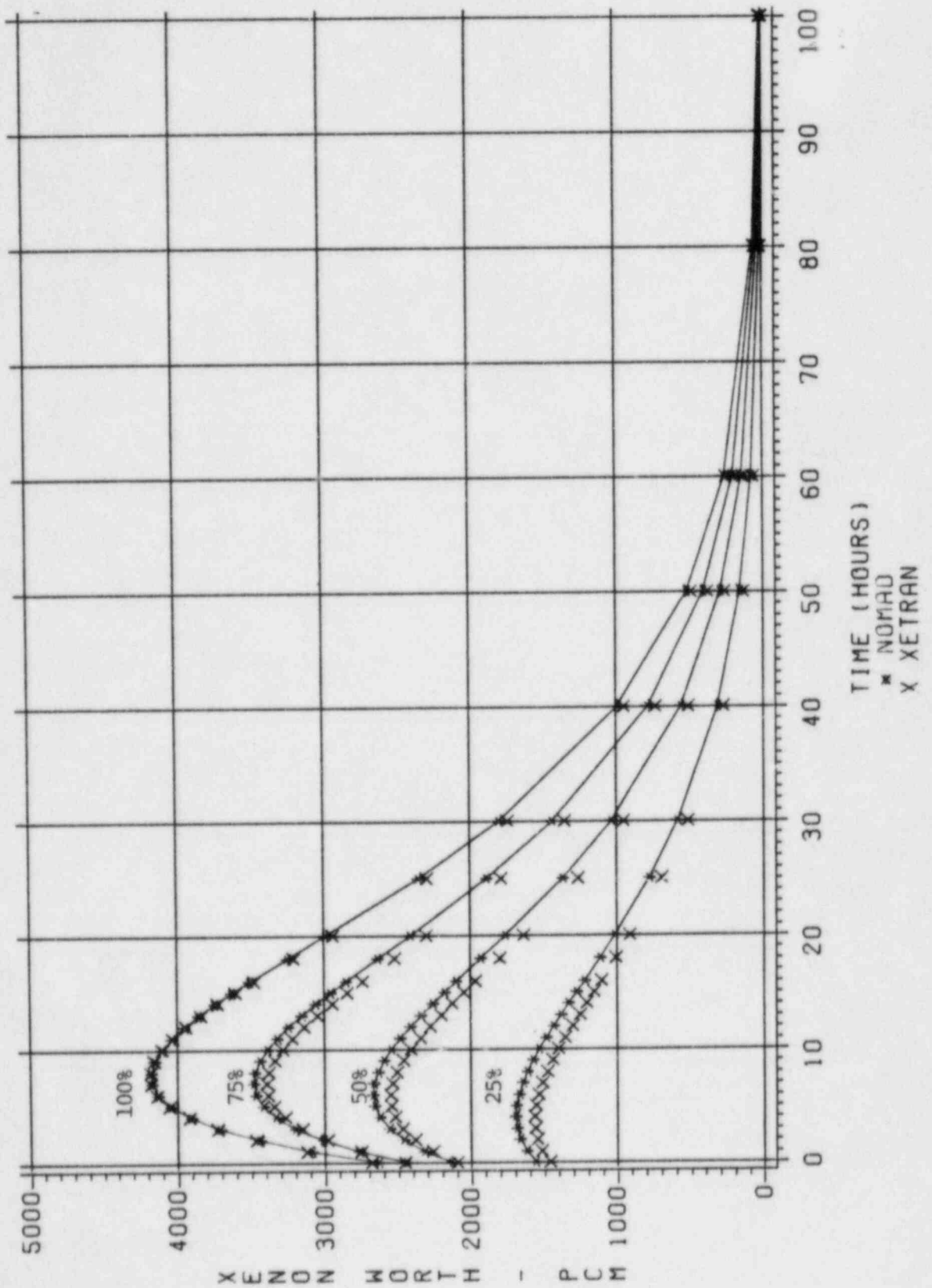
XENON WORTH AFTER SHUTDOWN

SURRY UNIT 1 CYCLE 6



XENON WORTH AFTER TRIP SURRY UNIT 1 CYCLE 6

FIGURE 5-6



AXIAL POWER COMPARISON

N1C2 HZP BOC

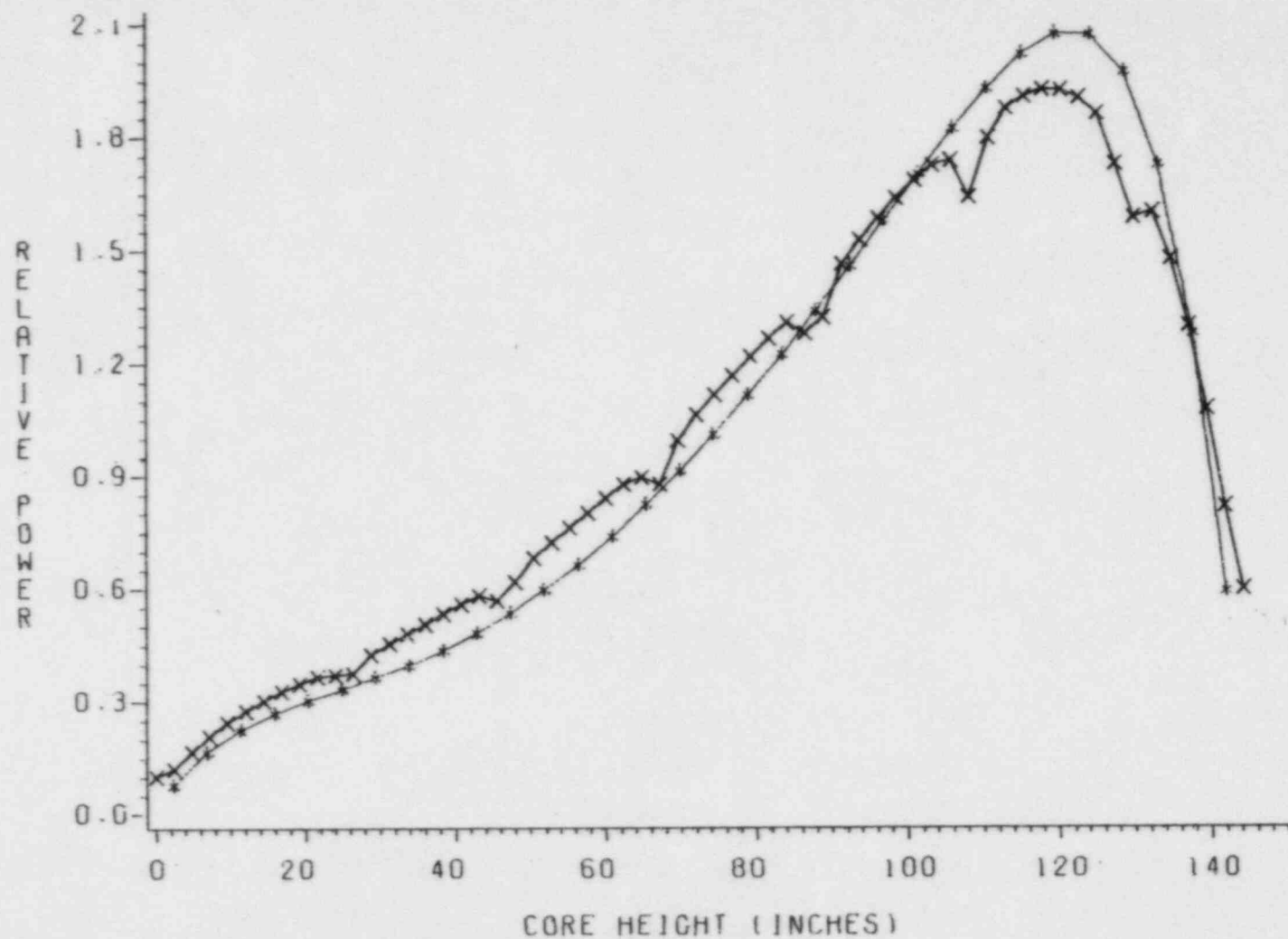


FIGURE 5-7

AXIAL POWER COMPARISON

N1C2 HFP ARO EQ. XE. BOC

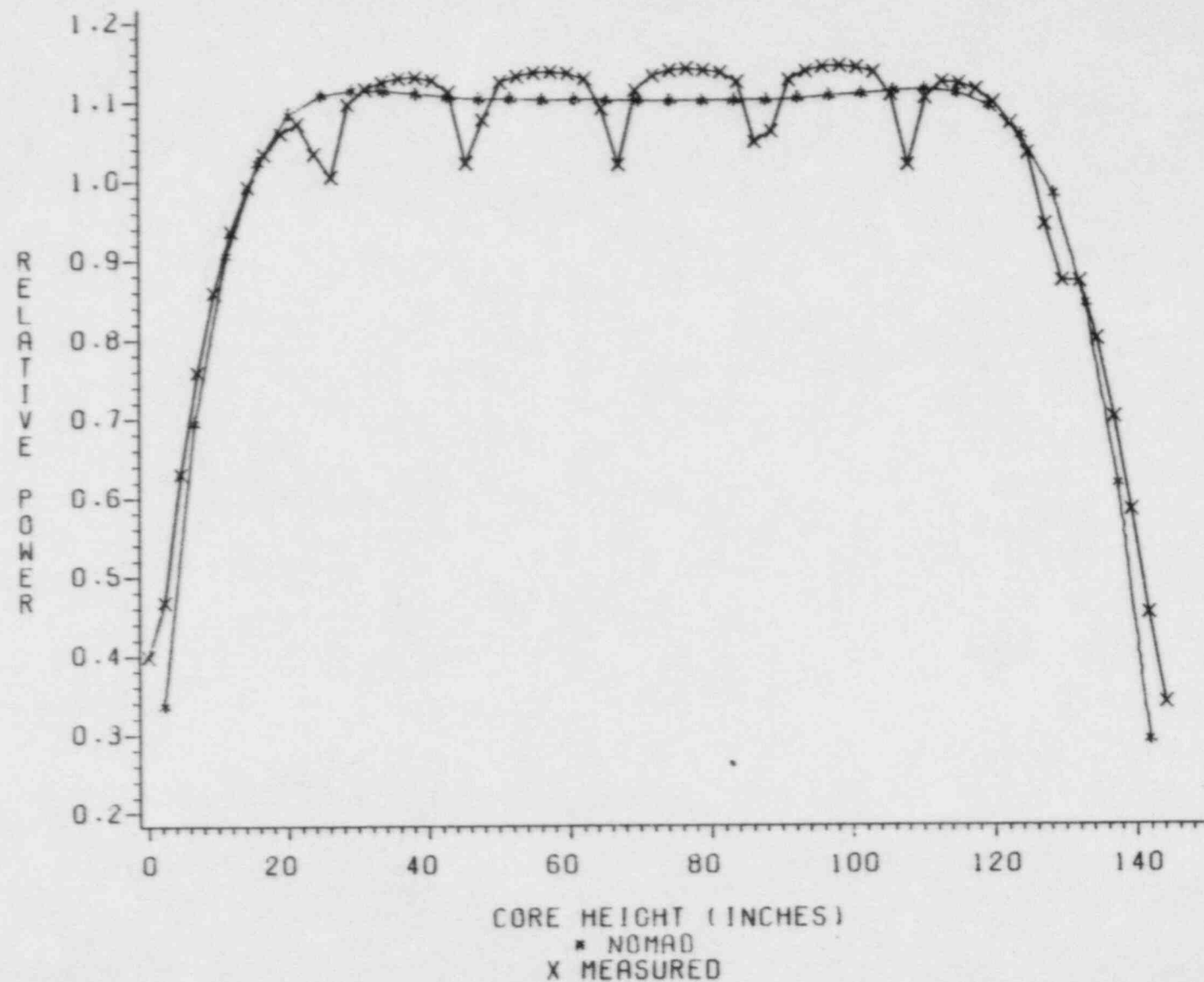


FIGURE 5-8

AXIAL POWER COMPARISON

NIC3 HZP B0C

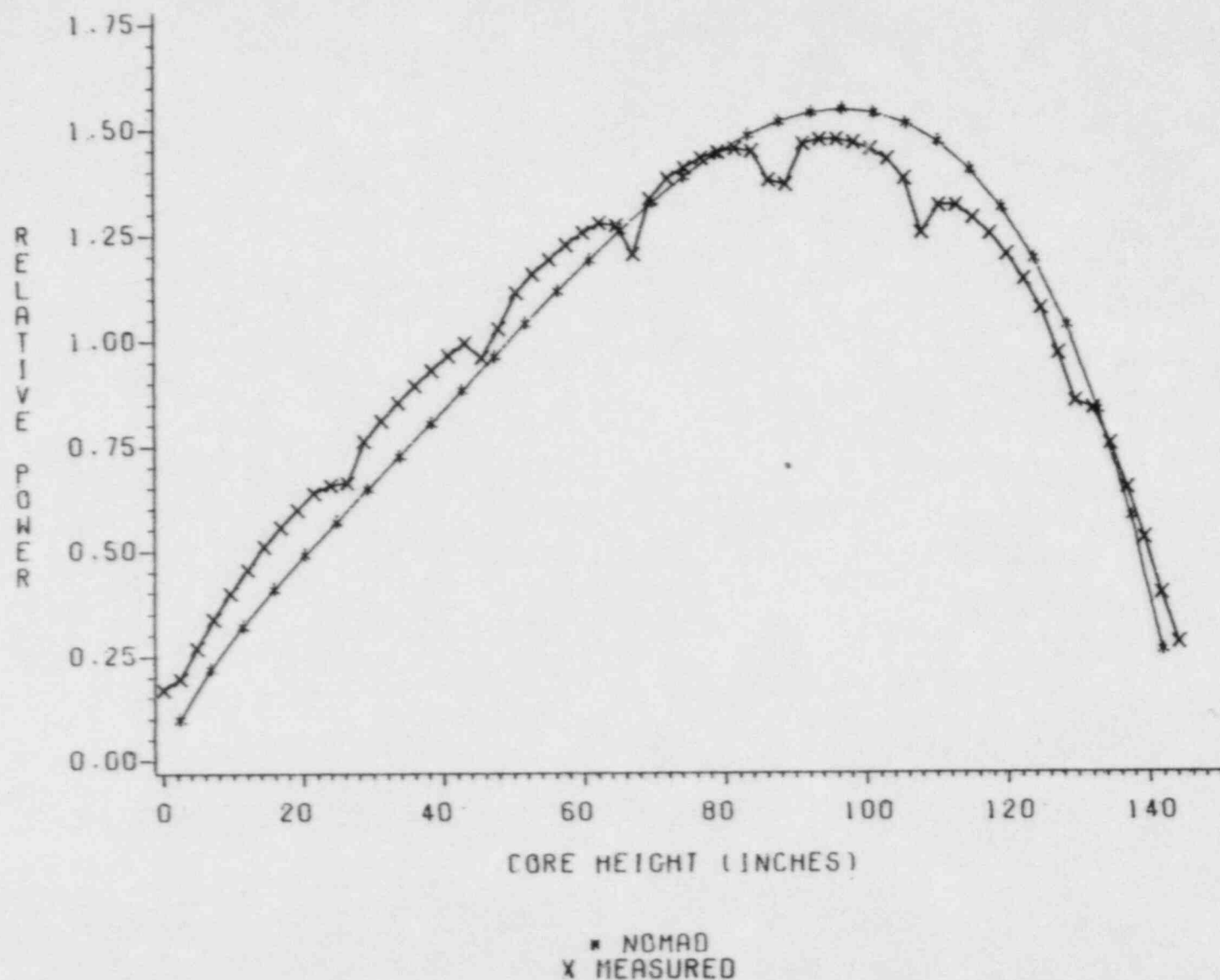


FIGURE 5-9

AXIAL POWER COMPARISON

N1C3 HFP ARO EQ. XE. BOC

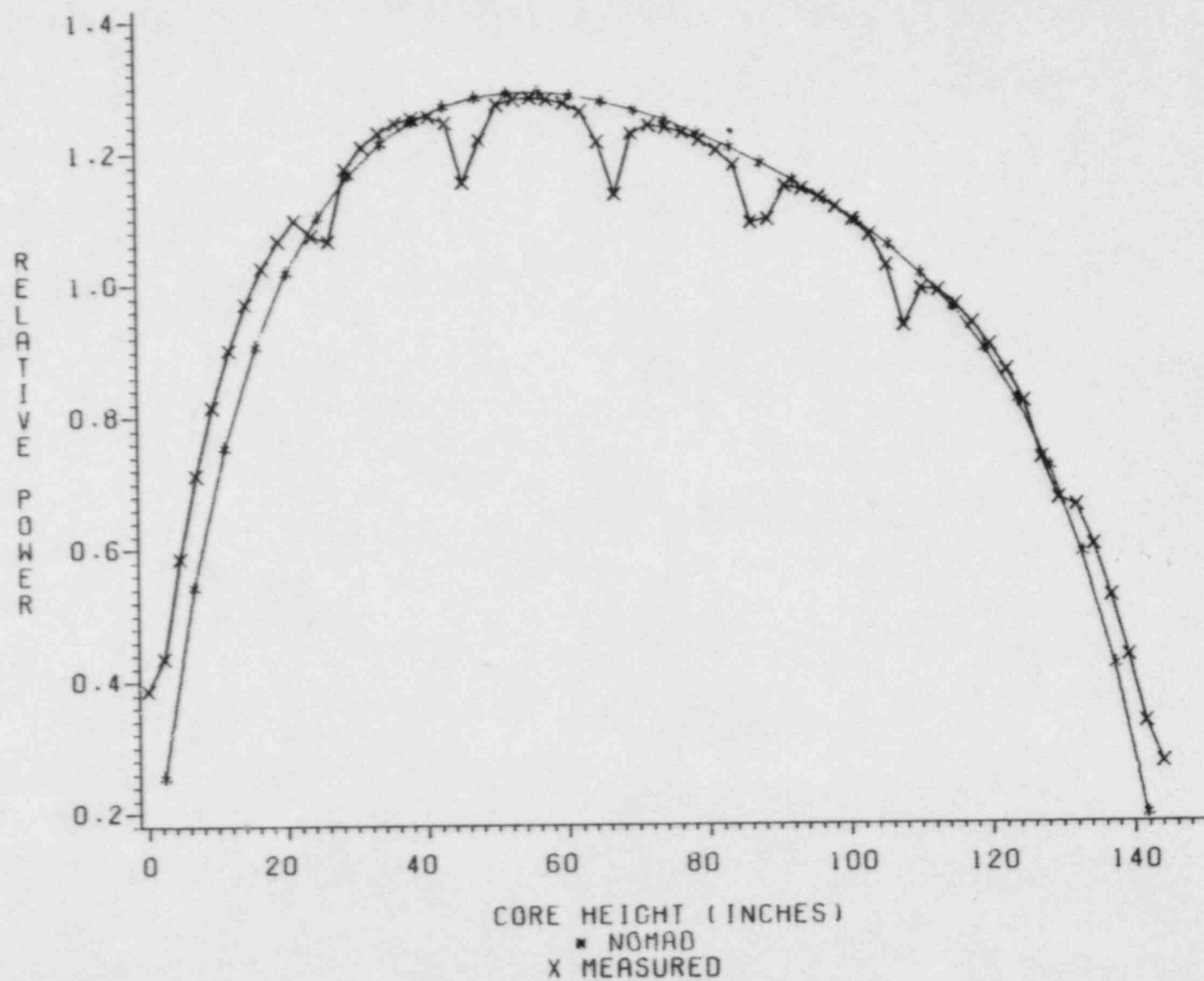


FIGURE 5-10

AXIAL POWER COMPARISON

N1C4 HZP BOC

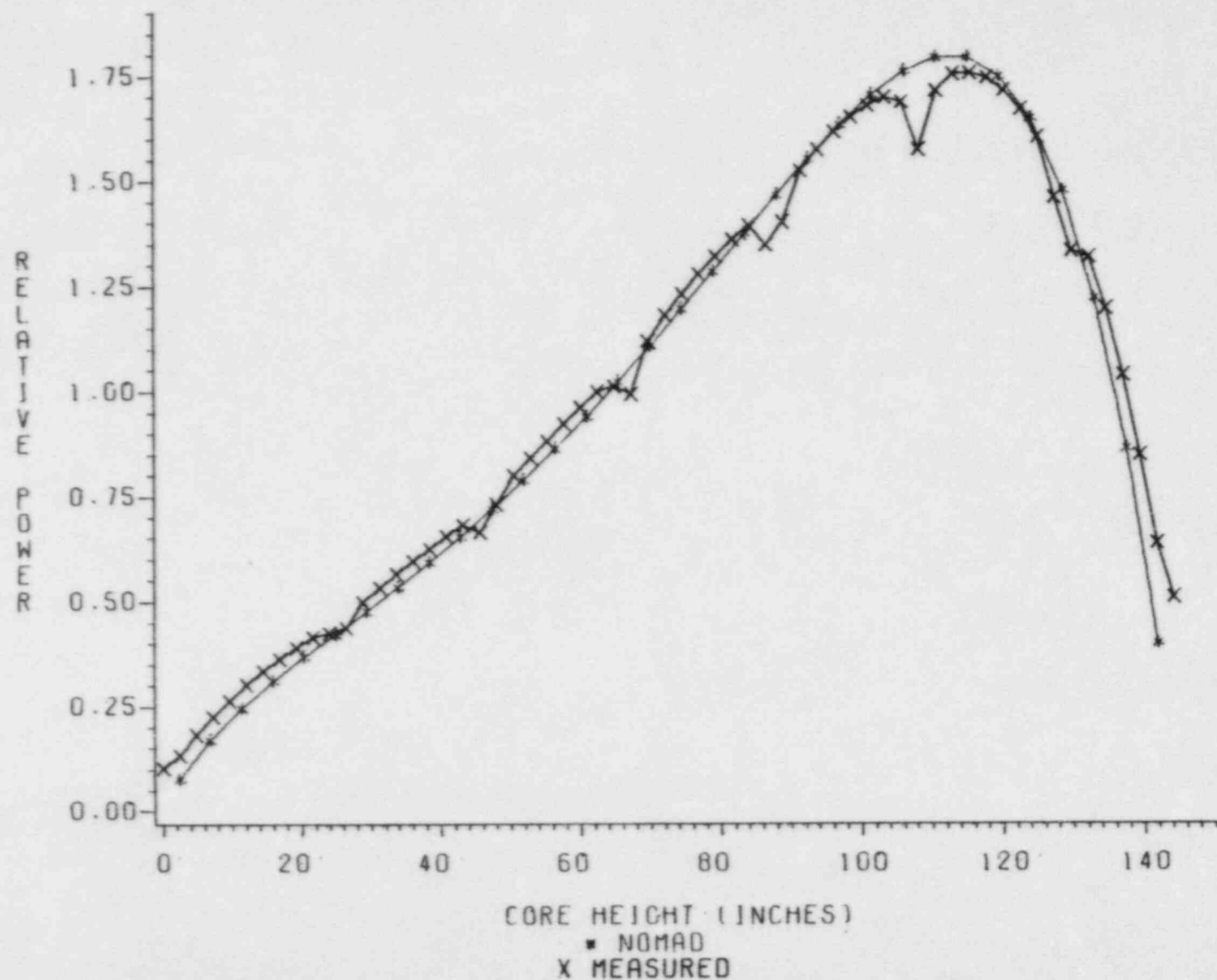


FIGURE 5-11

AXIAL POWER COMPARISON

NIC4 HFP ARO EQ. XE. BOC

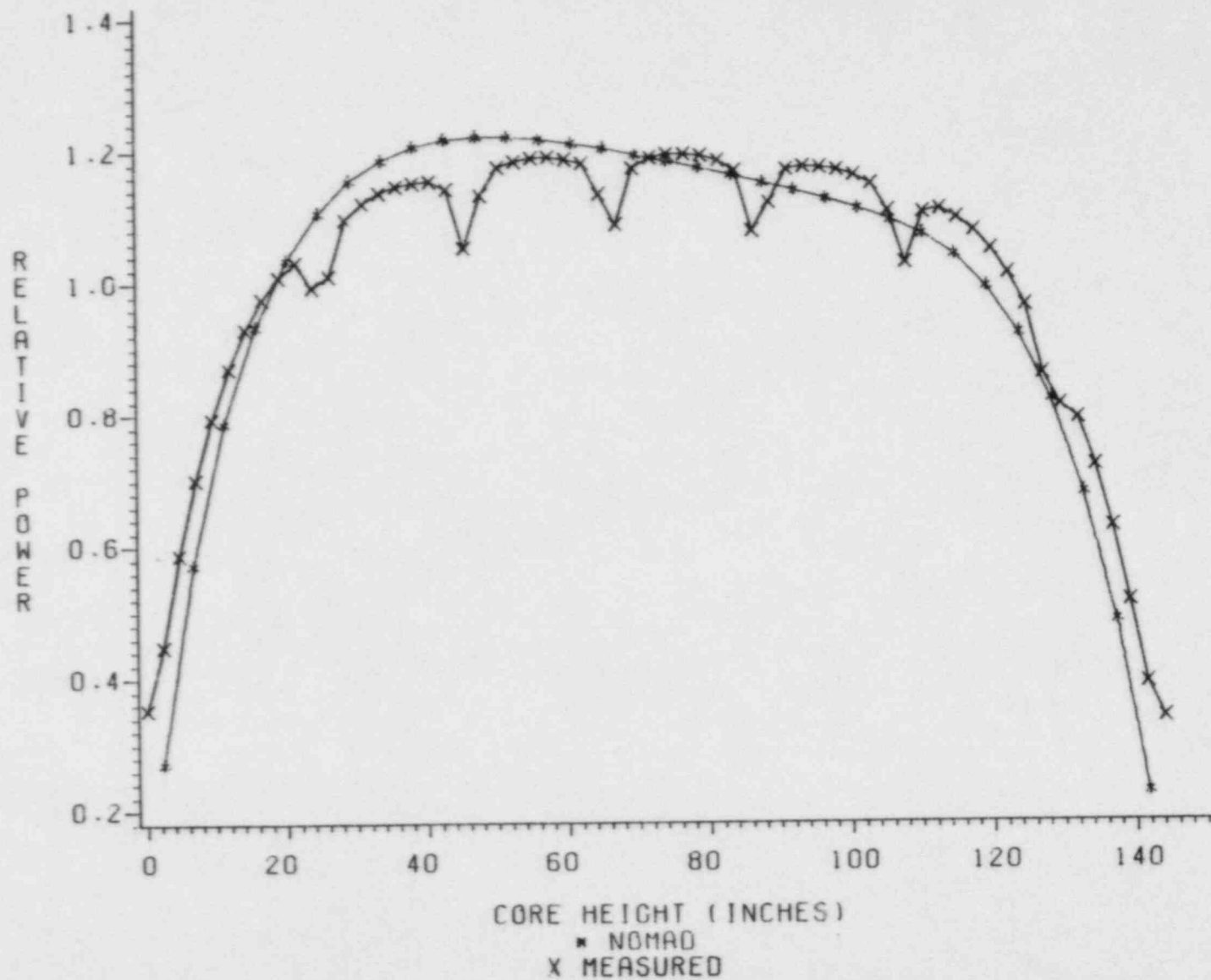


FIGURE 5-12

AXIAL POWER COMPARISON

N2C2 HZP BOC

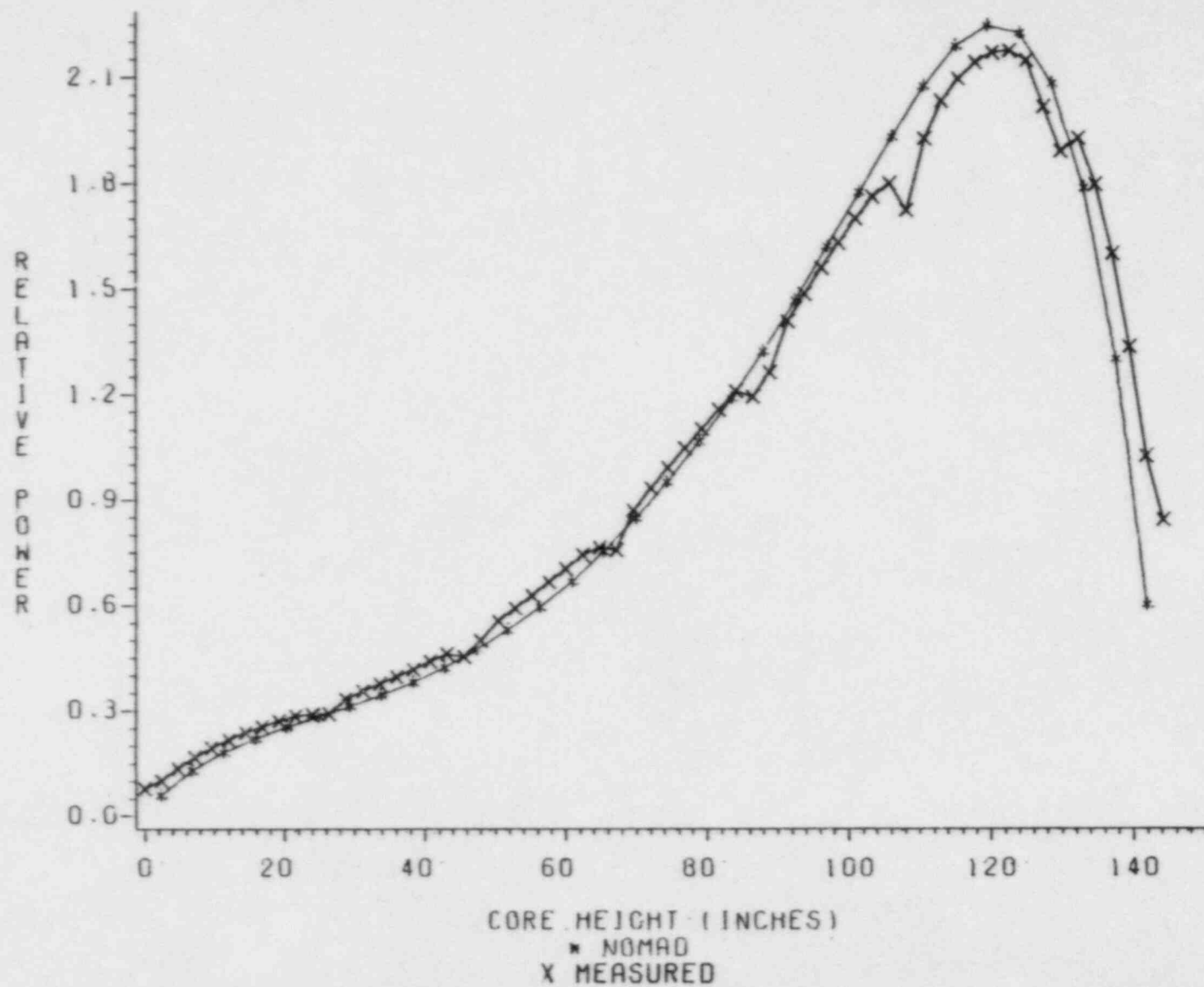


FIGURE 5-13

AXIAL POWER COMPARISON

N2C2 HFP ARO EQ. XE. BOC

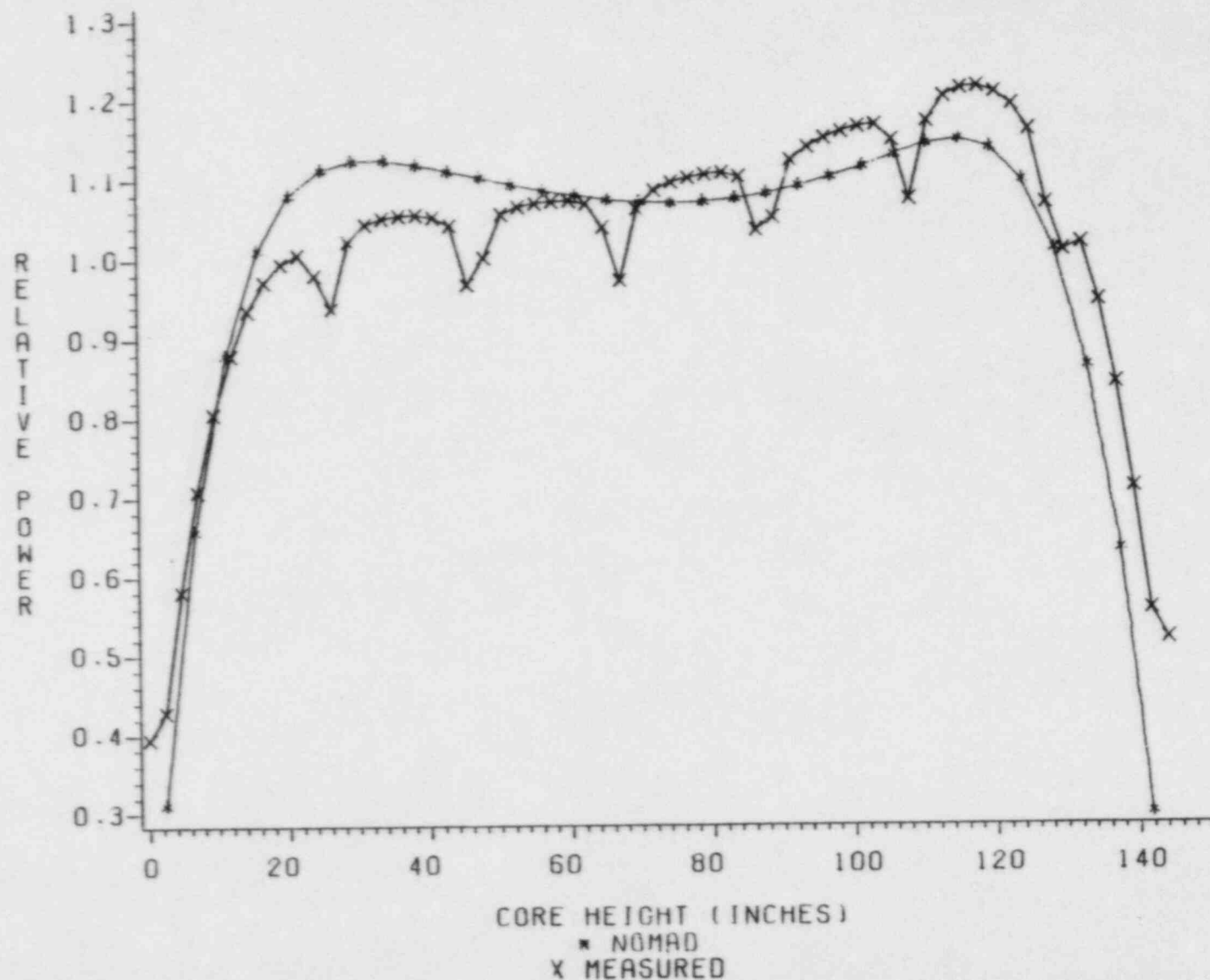


FIGURE 5-14

AXIAL POWER COMPARISON

S1C6 HZP B0C

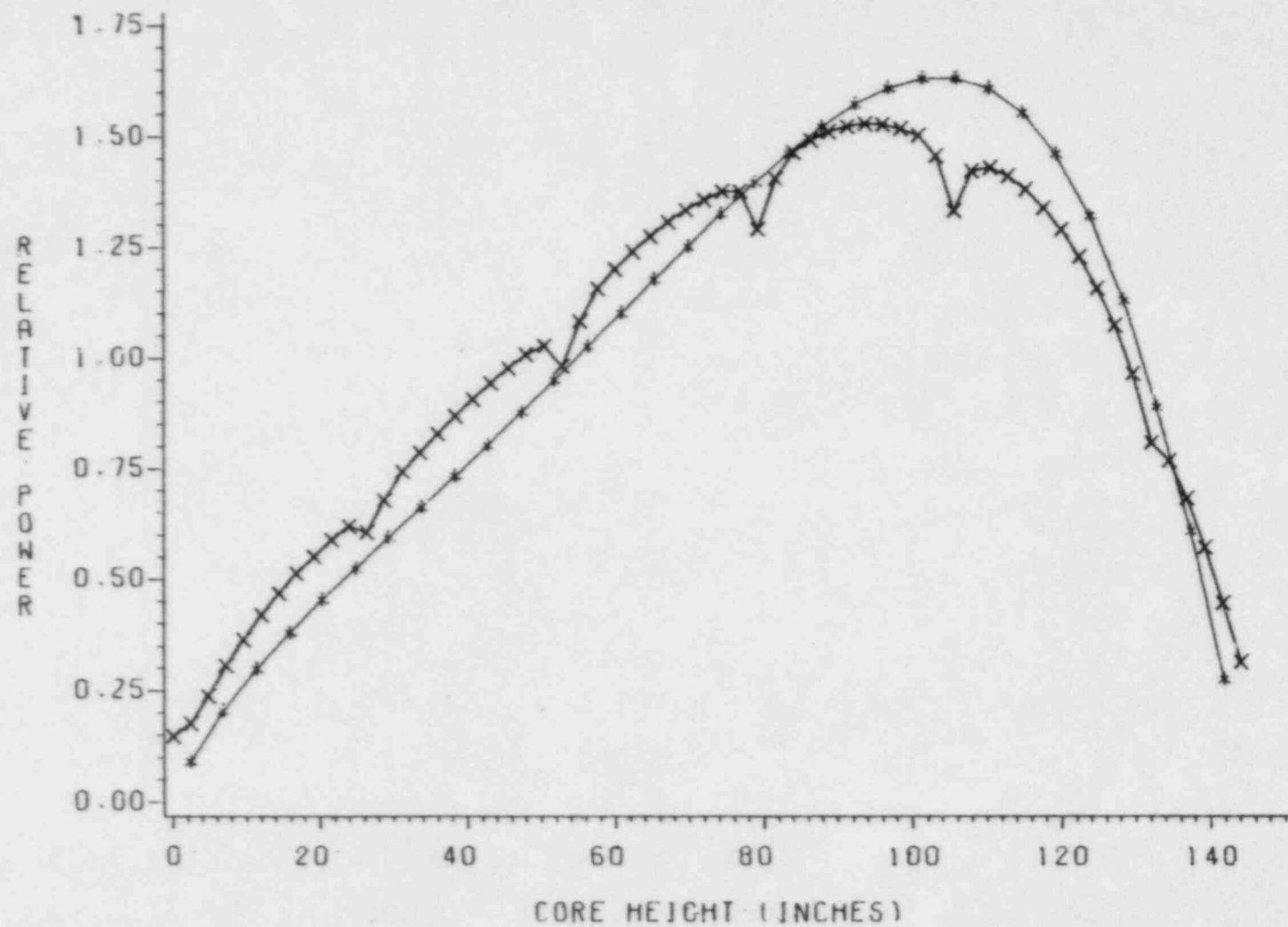
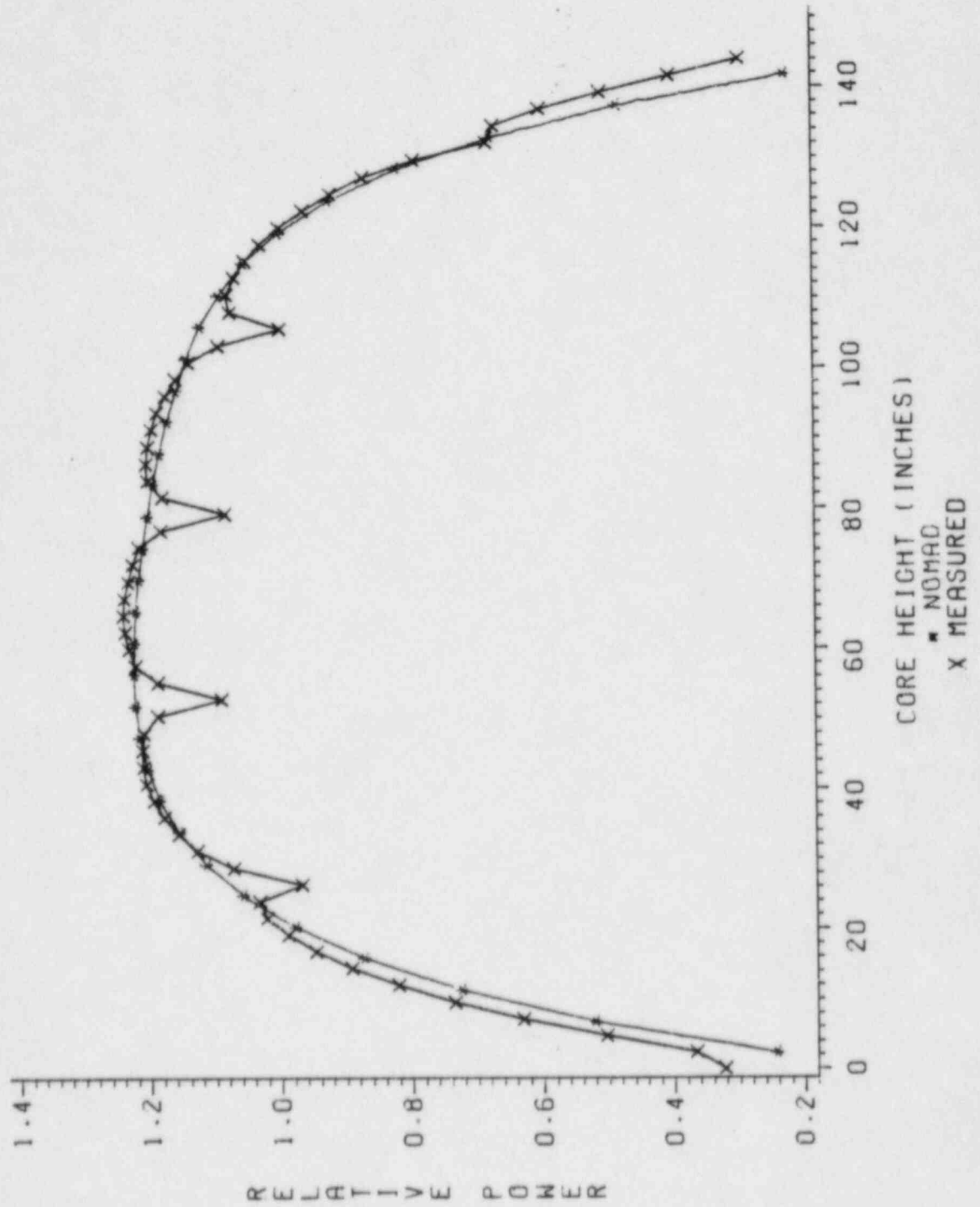


FIGURE 5-15

FIGURE 5-16

AXIAL POWER COMPARISON

S1C6 HFP ARO EQ. XE. BOC



DIFFERENTIAL ROD WORTH COMPARISON

NORTH ANNA UNIT 1 CYCLE 3

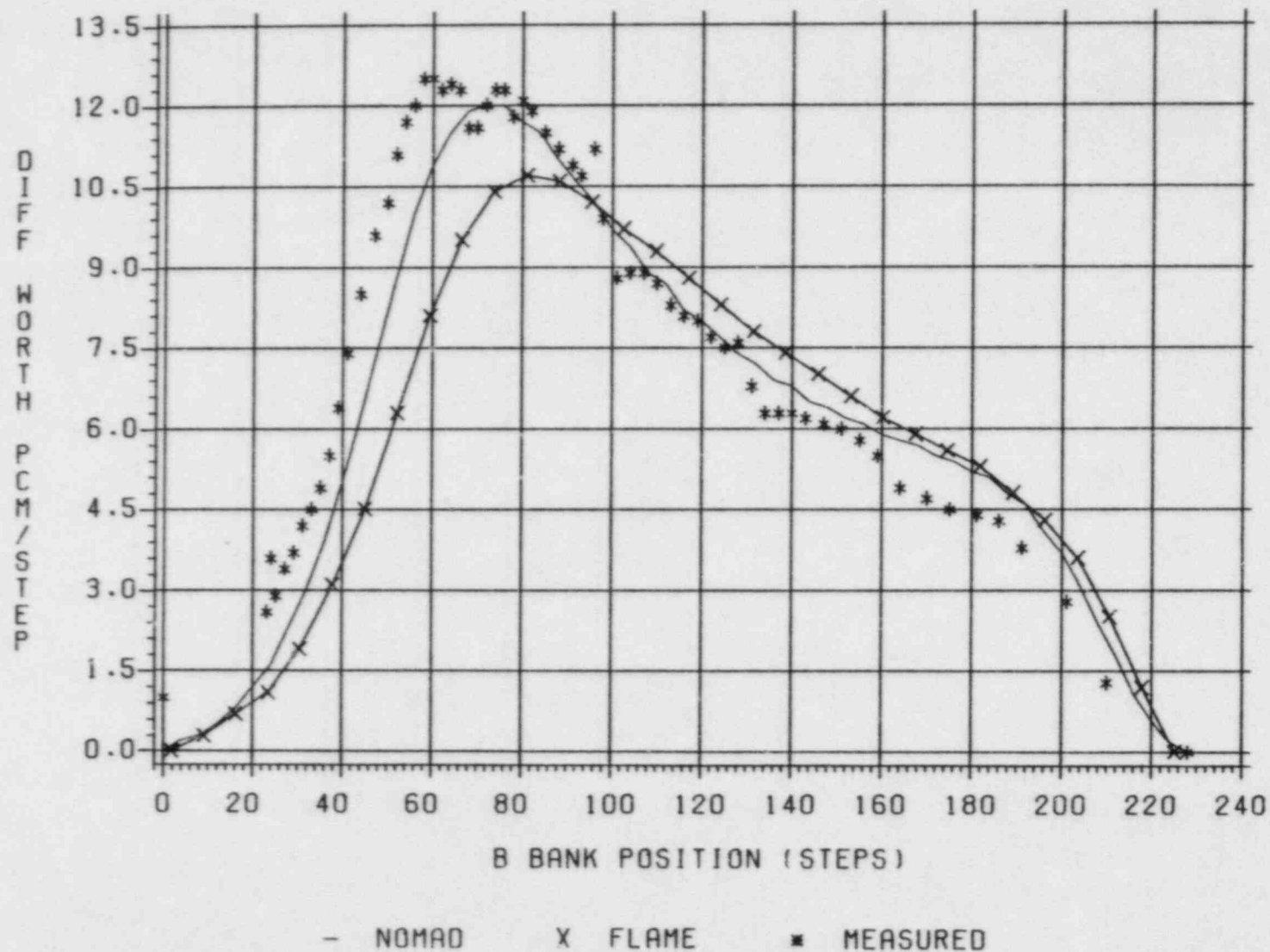
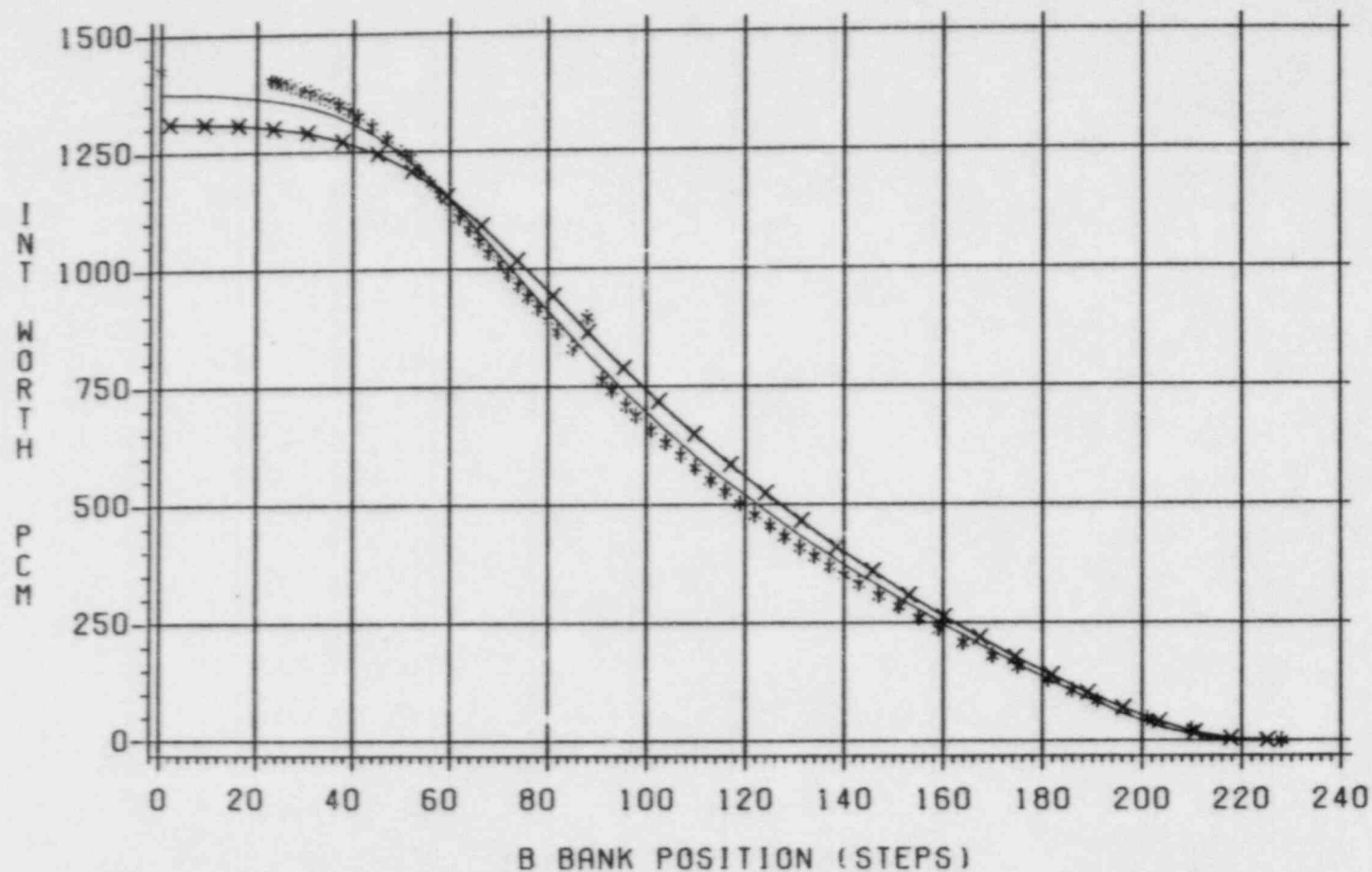


FIGURE 5-17

INTEGRAL ROD WORTH COMPARISON

NORTH ANNA UNIT 1 CYCLE 3

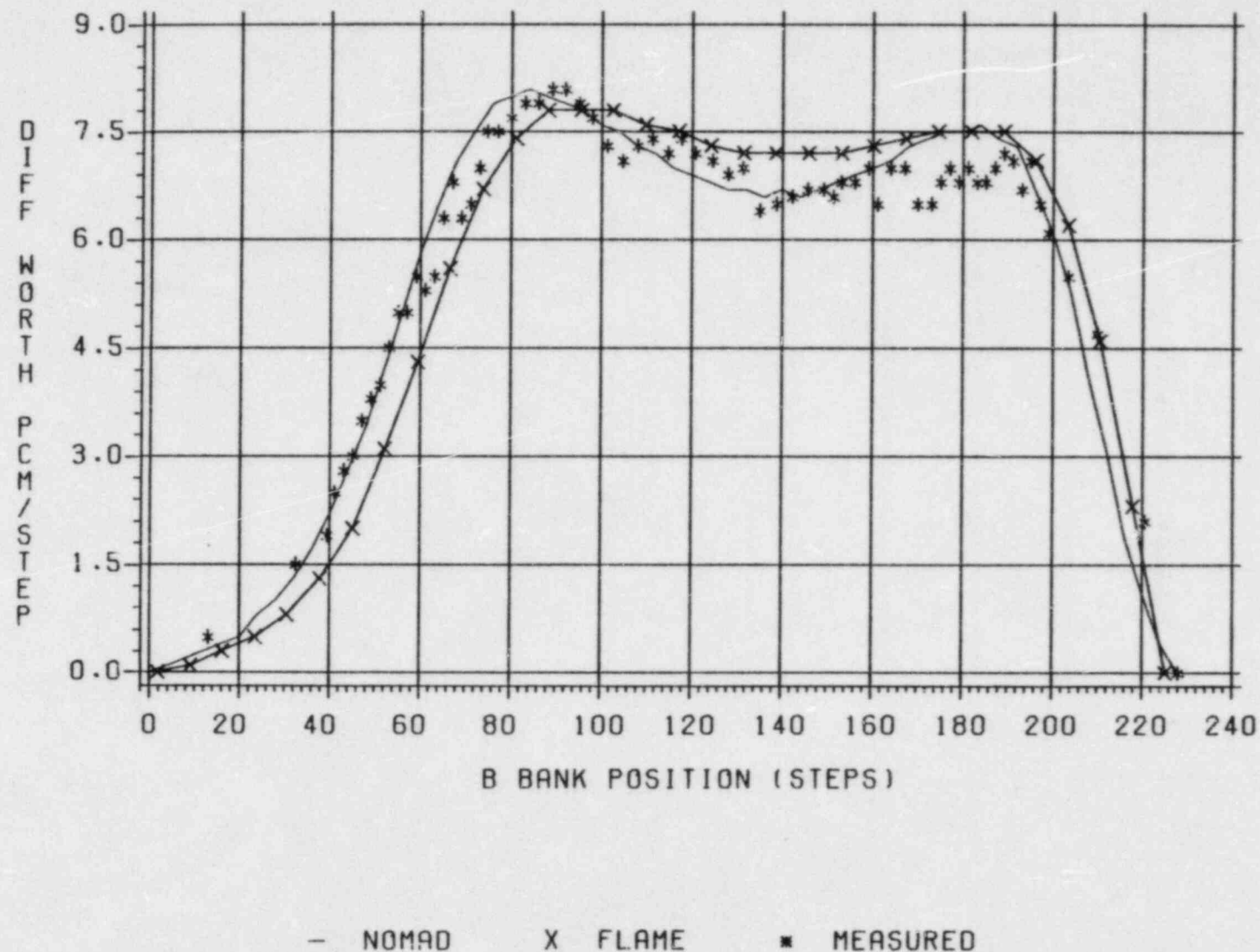


— NOMAD X FLAME * MEASURED

FIGURE 5-18

DIFFERENTIAL ROD WORTH COMPARISON

NORTH ANNA UNIT 1 CYCLE 4



INTEGRAL ROD WORTH COMPARISON

NORTH ANNA UNIT 1 CYCLE 4

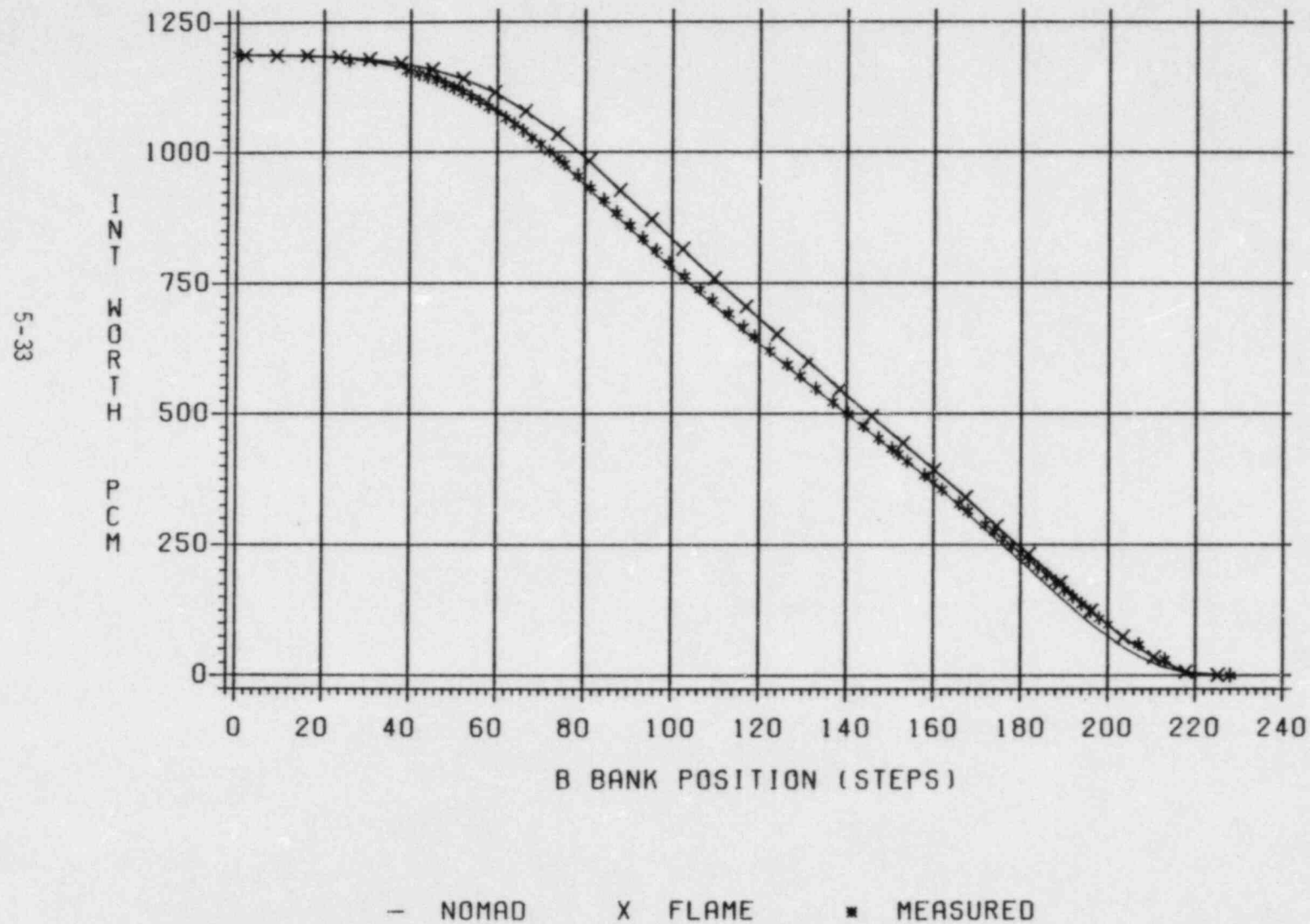


FIGURE 5-20

DIFFERENTIAL ROD WORTH COMPARISON

NORTH ANNA UNIT 2 CYCLE 2

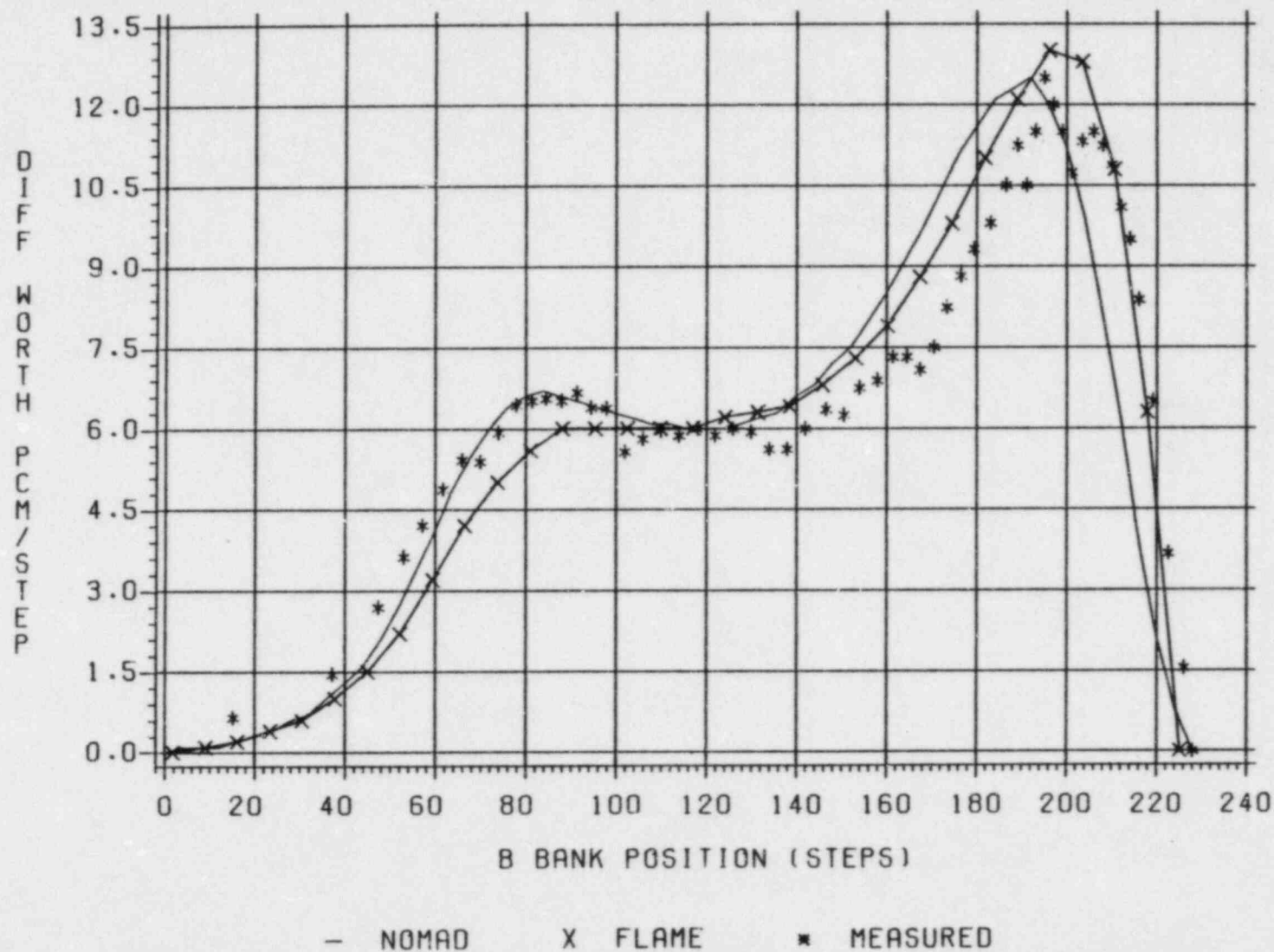


FIGURE 5-21

INTEGRAL ROD WORTH COMPARISON

NORTH ANNA UNIT 2 CYCLE 2

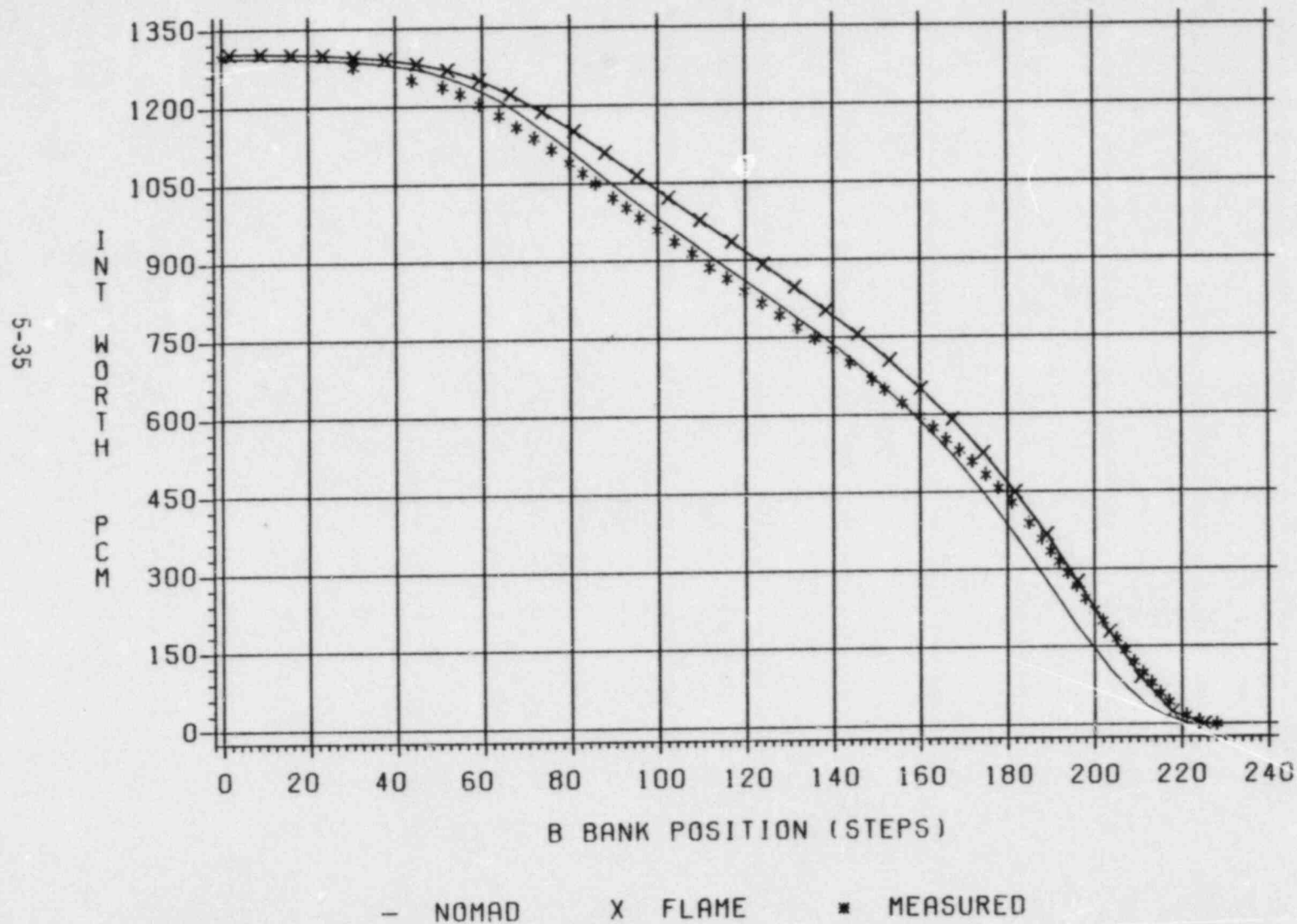


FIGURE 5-22

DIFFERENTIAL ROD WORTH COMPARISON

SURRY UNIT 1 CYCLE 6

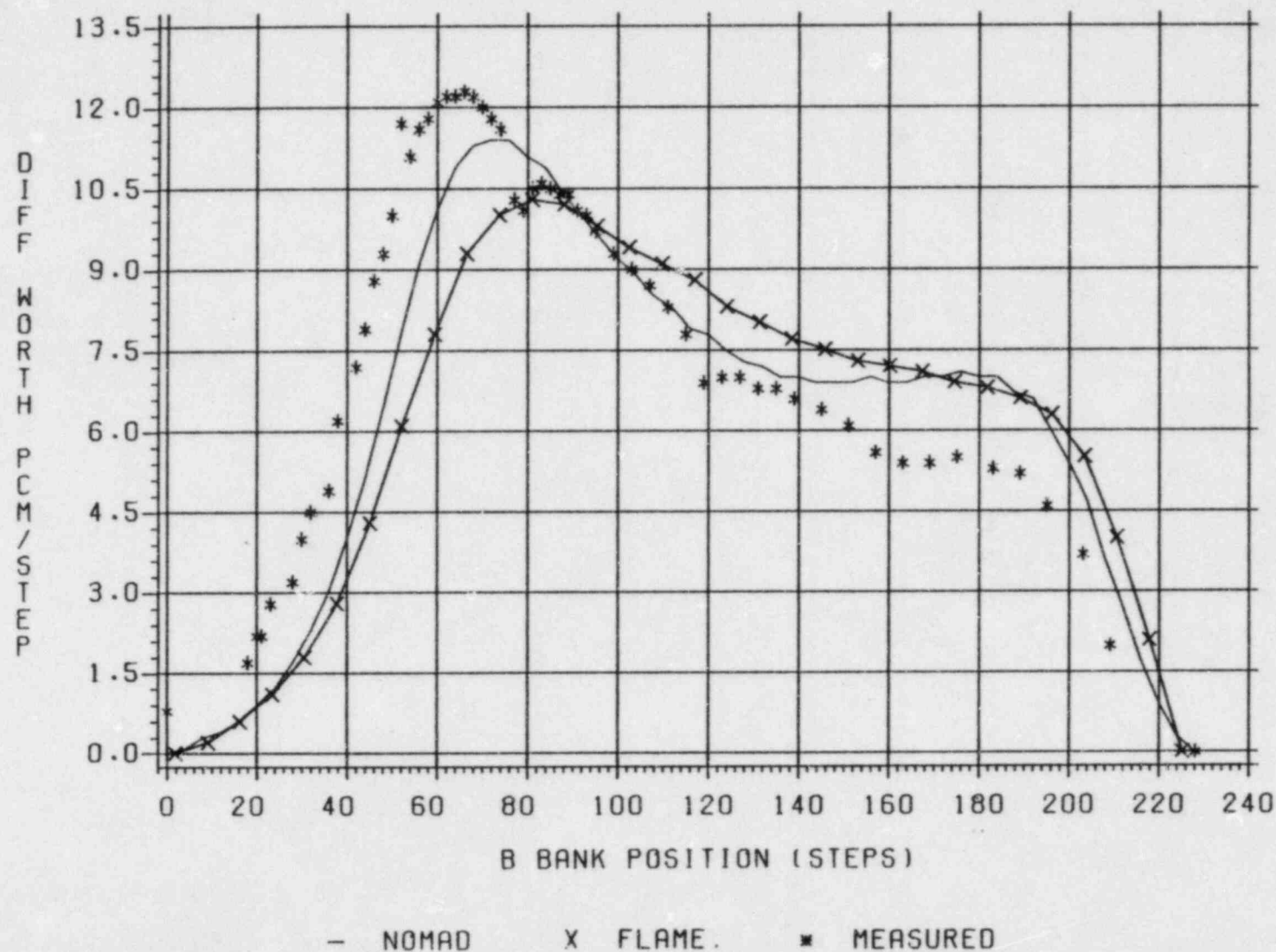
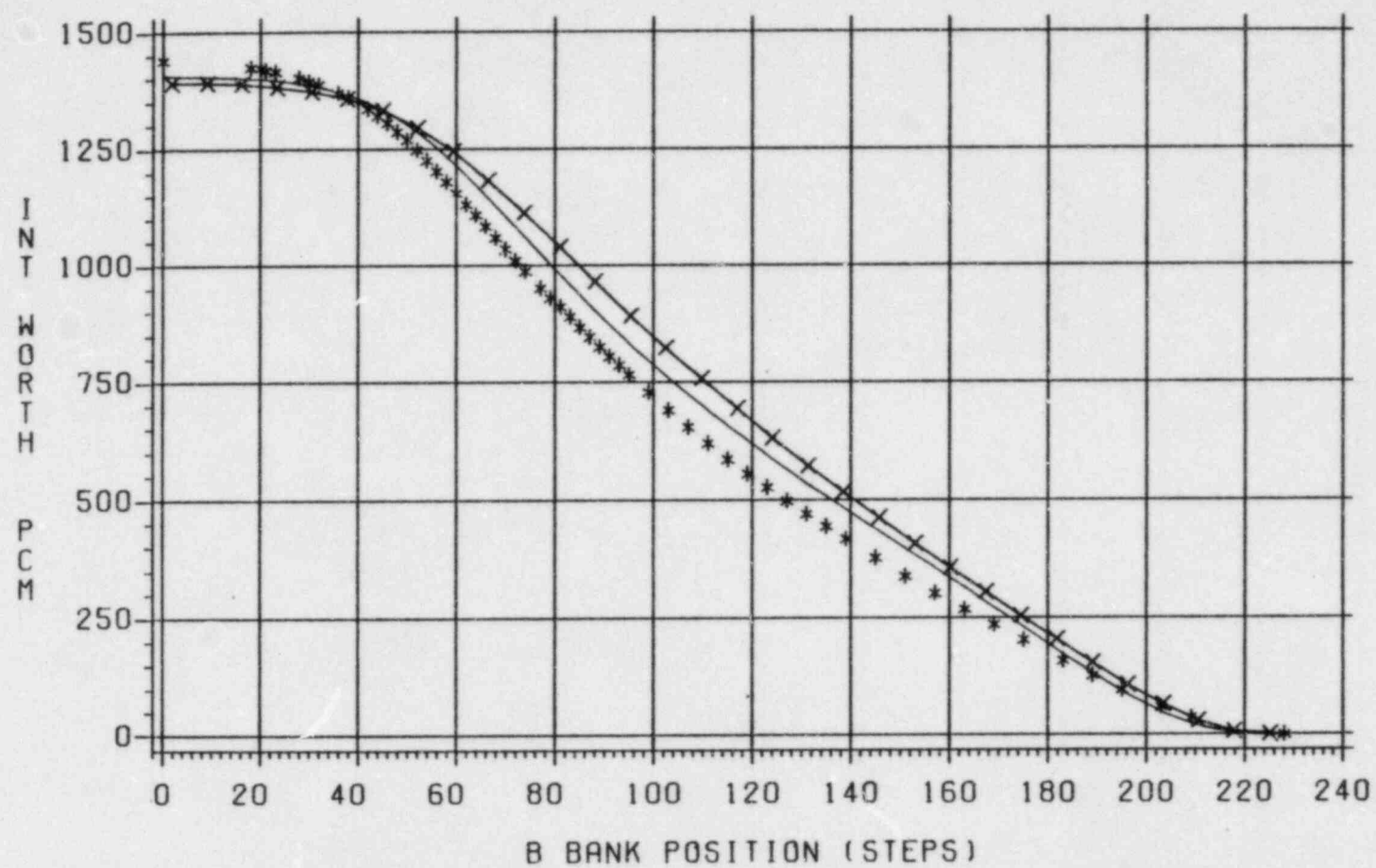


FIGURE 5-23

INTEGRAL ROD WORTH COMPARISON

SURRY UNIT 1 CYCLE 6



— NOMAD x FLAME * MEASURED

FIGURE 5-24

5-37

FIGURE 5-25
 DIFFERENTIAL WORTH OF CONTROL BANKS
 A THROUGH D IN OVERLAP MODE
 NORTH ANNA UNIT 1, CYCLE 3

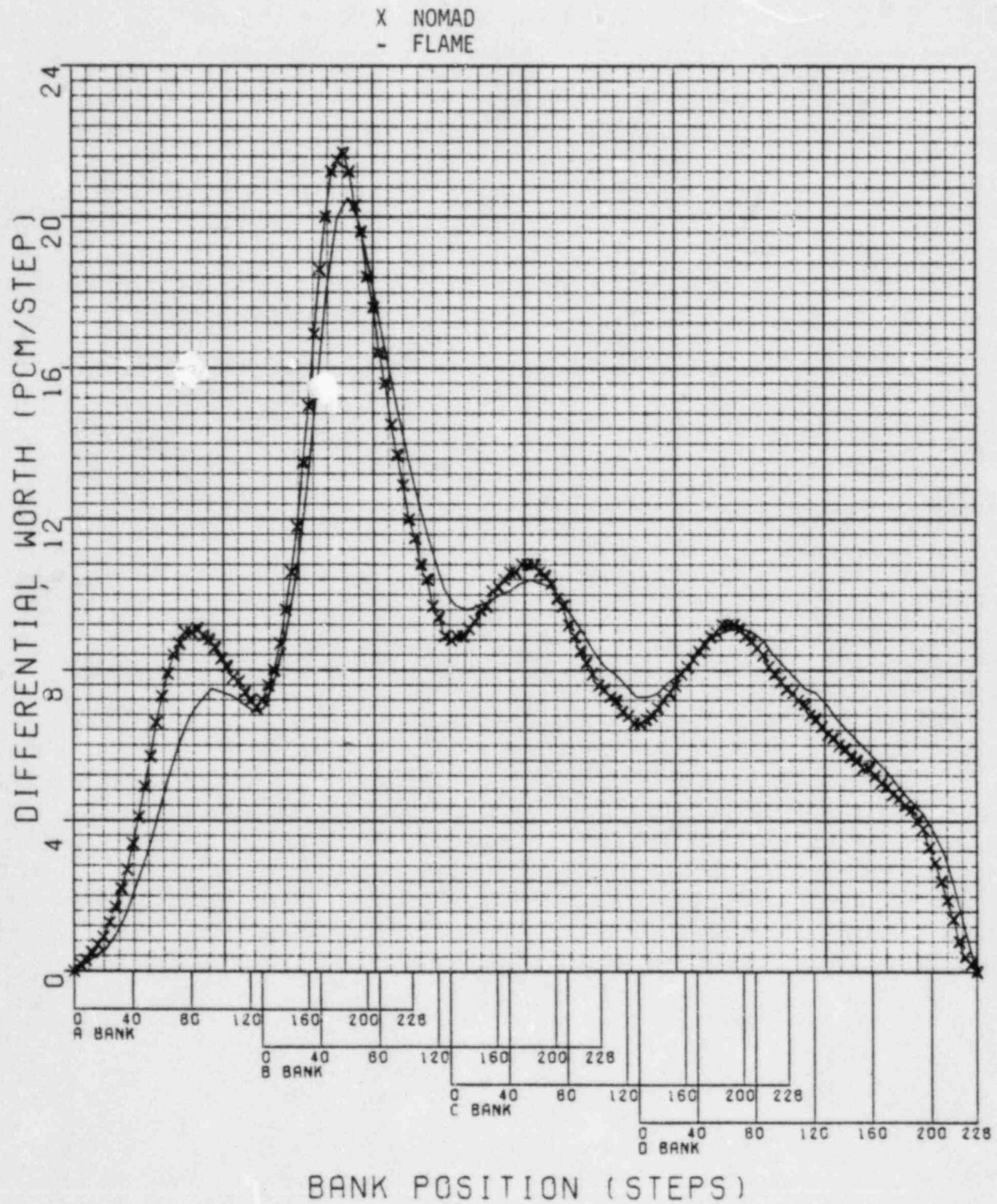


FIGURE 5-26
 INTEGRAL WORTH OF CONTROL BANKS
 A THROUGH D IN OVERLAP MODE
 NORTH ANNA UNIT 1, CYCLE 3

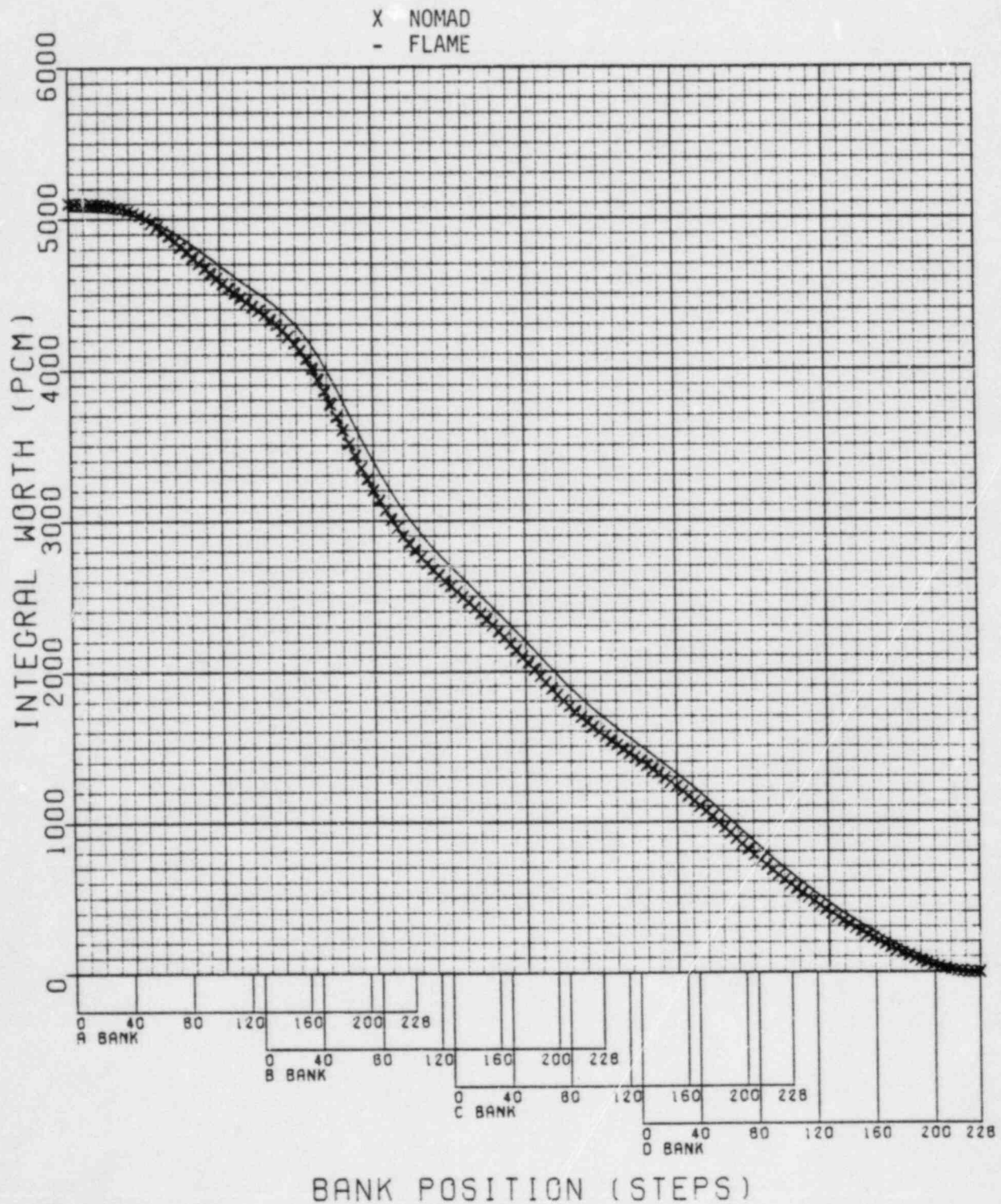


FIGURE 5-27
DIFFERENTIAL WORTH OF CONTROL BANKS
A THROUGH D IN OVERLAP MODE
NORTH ANNA UNIT 1, CYCLE 4

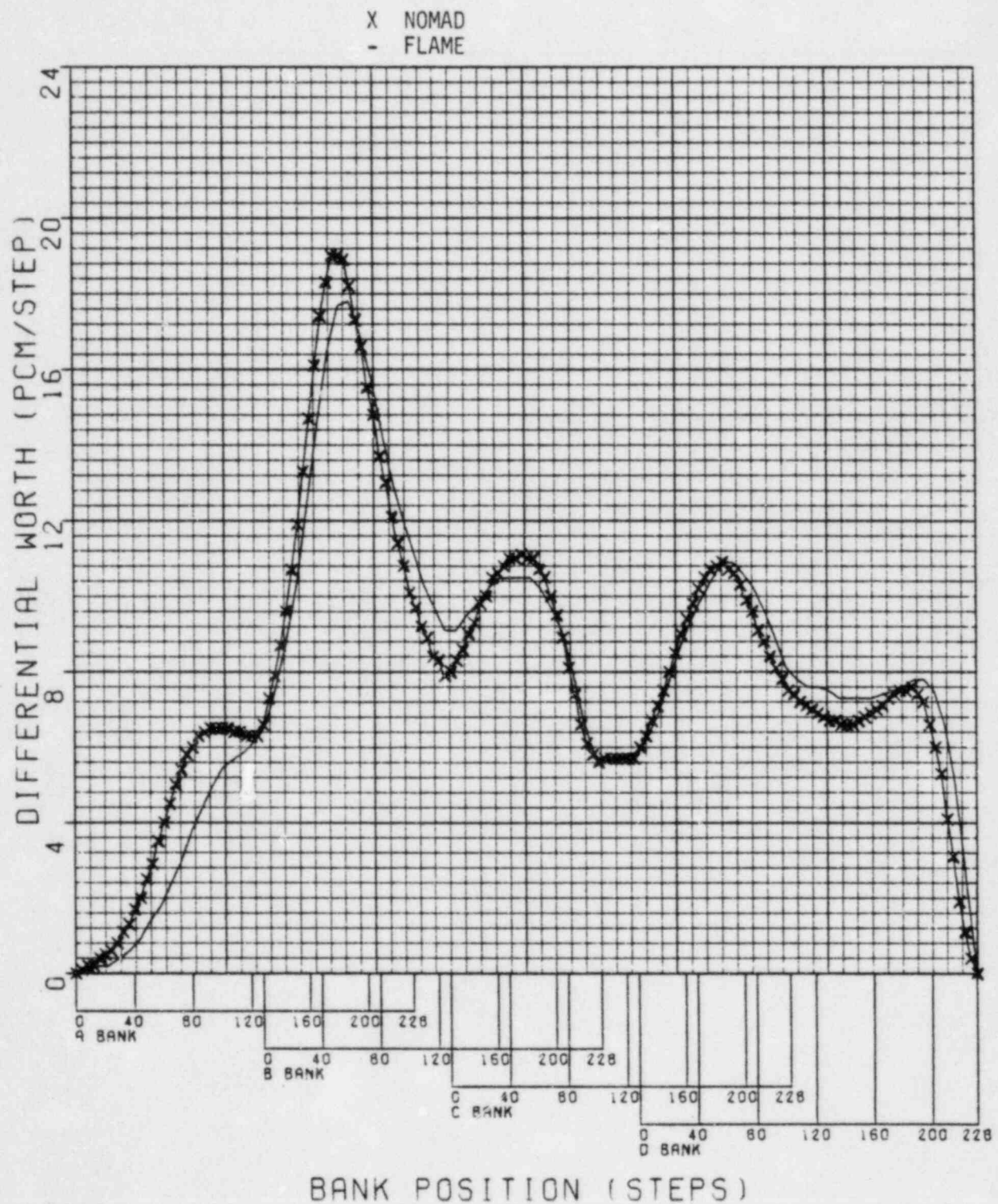


FIGURE 5-28
 INTEGRAL WORTH OF CONTROL BANKS
 A THROUGH D IN OVERLAP MODE
 NORTH ANNA UNIT 1, CYCLE 4

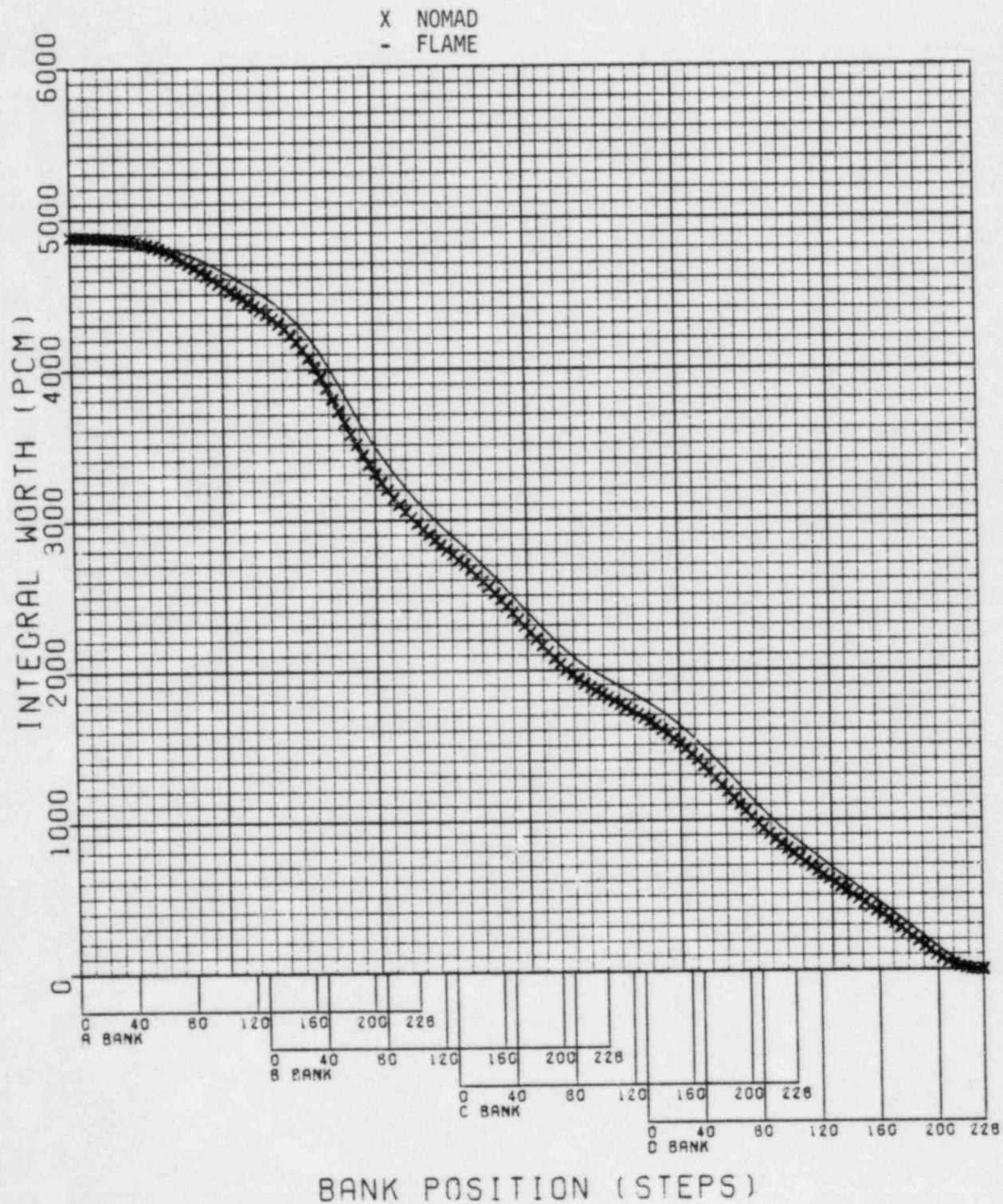


FIGURE 5-29
DIFFERENTIAL WORTH OF CONTROL BANKS
A THROUGH D IN OVERLAP MODE
NORTH ANNA UNIT 2, CYCLE 2

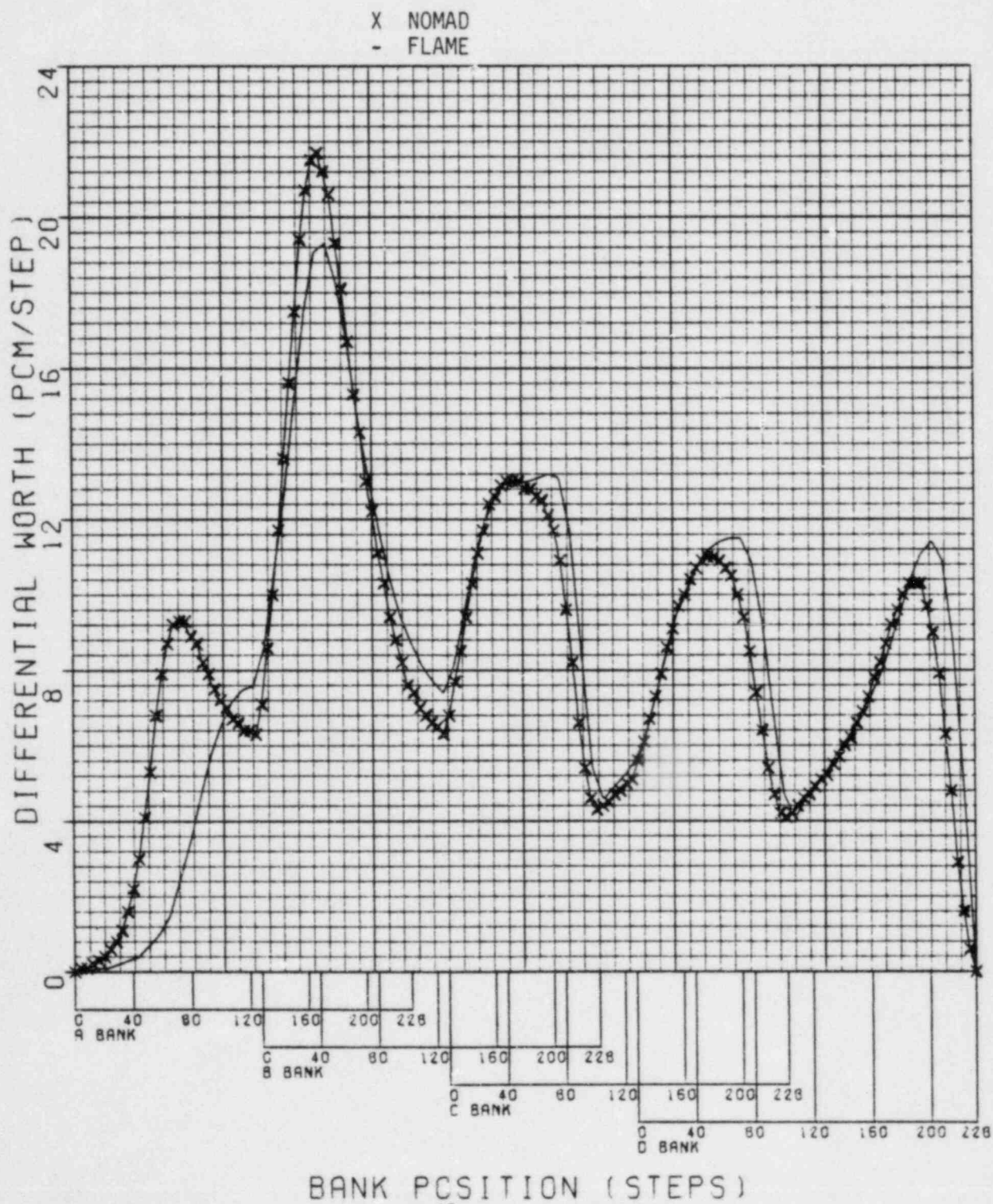


FIGURE 5-30

INTEGRAL WORTH OF CONTROL BANKS
A THROUGH D IN OVERLAP MODE
NORTH ANNA UNIT 2, CYCLE 2

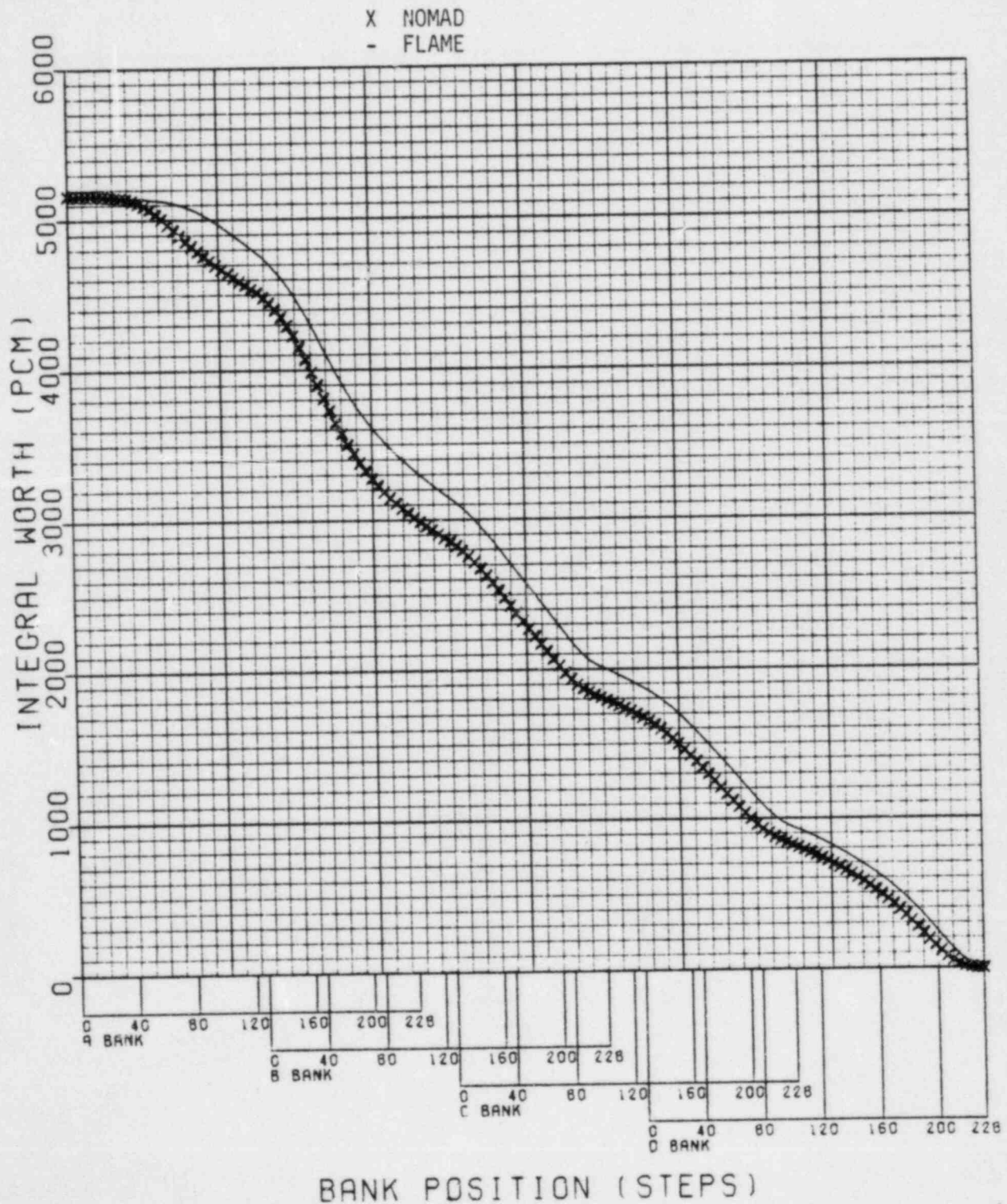


FIGURE 5-31

DIFFERENTIAL WORTH OF CONTROL BANKS
A THROUGH D IN OVERLAP MODE
SURRY UNIT 1, CYCLE 7

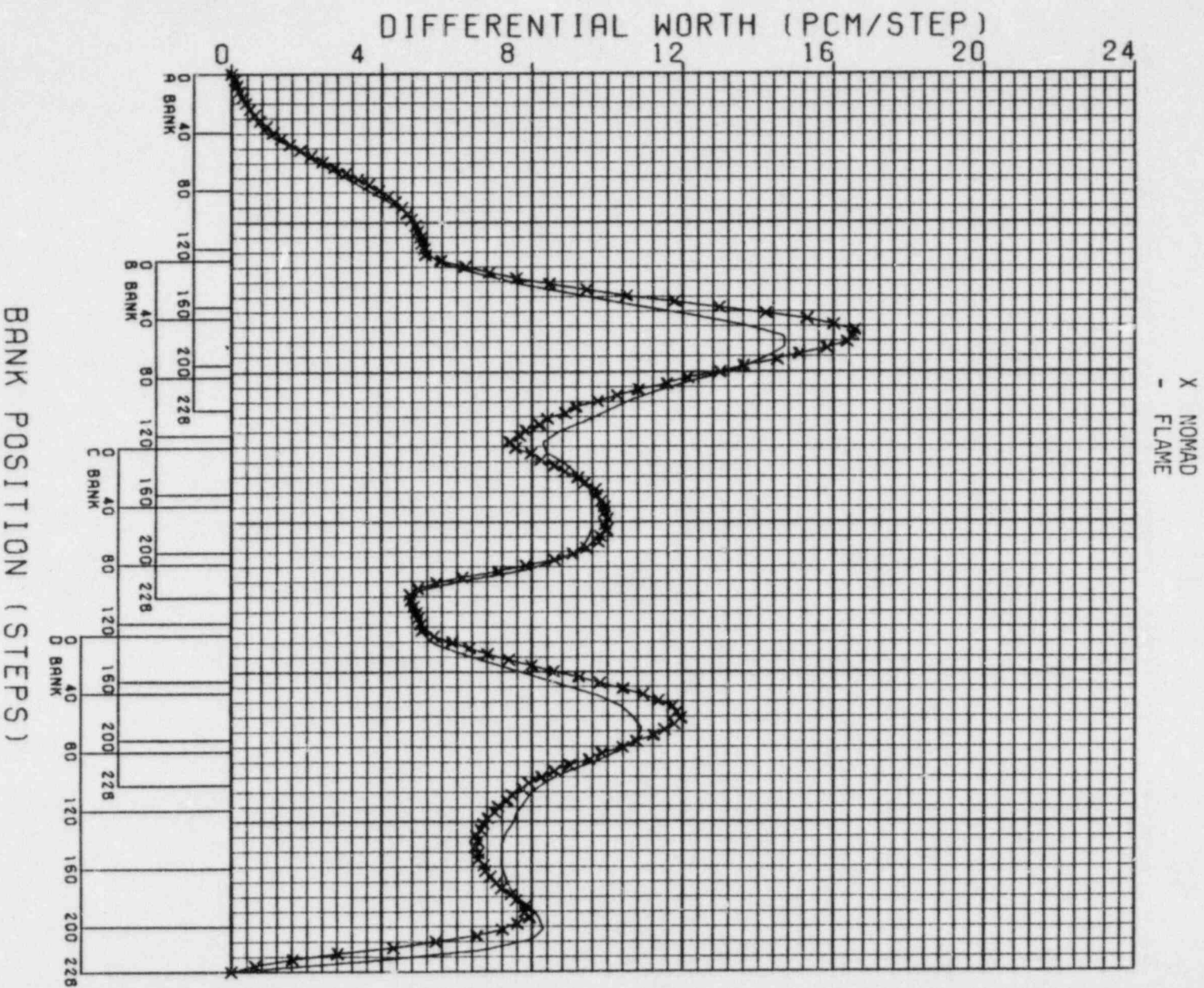
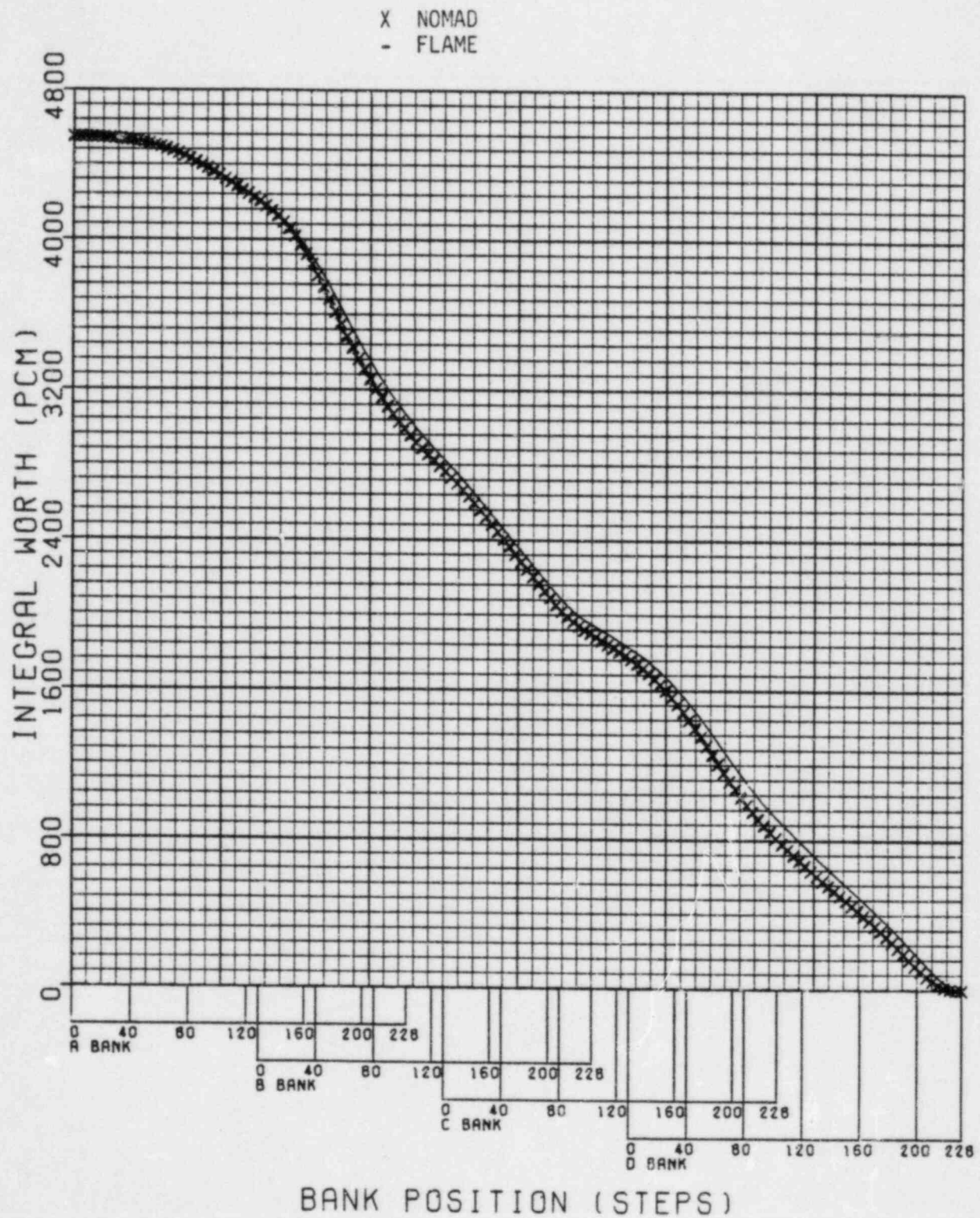


FIGURE 5-32

INTEGRAL WORTH OF CONTROL BANKS
A THROUGH D IN OVERLAP MODE
SURRY UNIT 1, CYCLE 7



N1C2 70% LOAD REDUCTION TEST
AXIAL FLUX DIFFERENCE

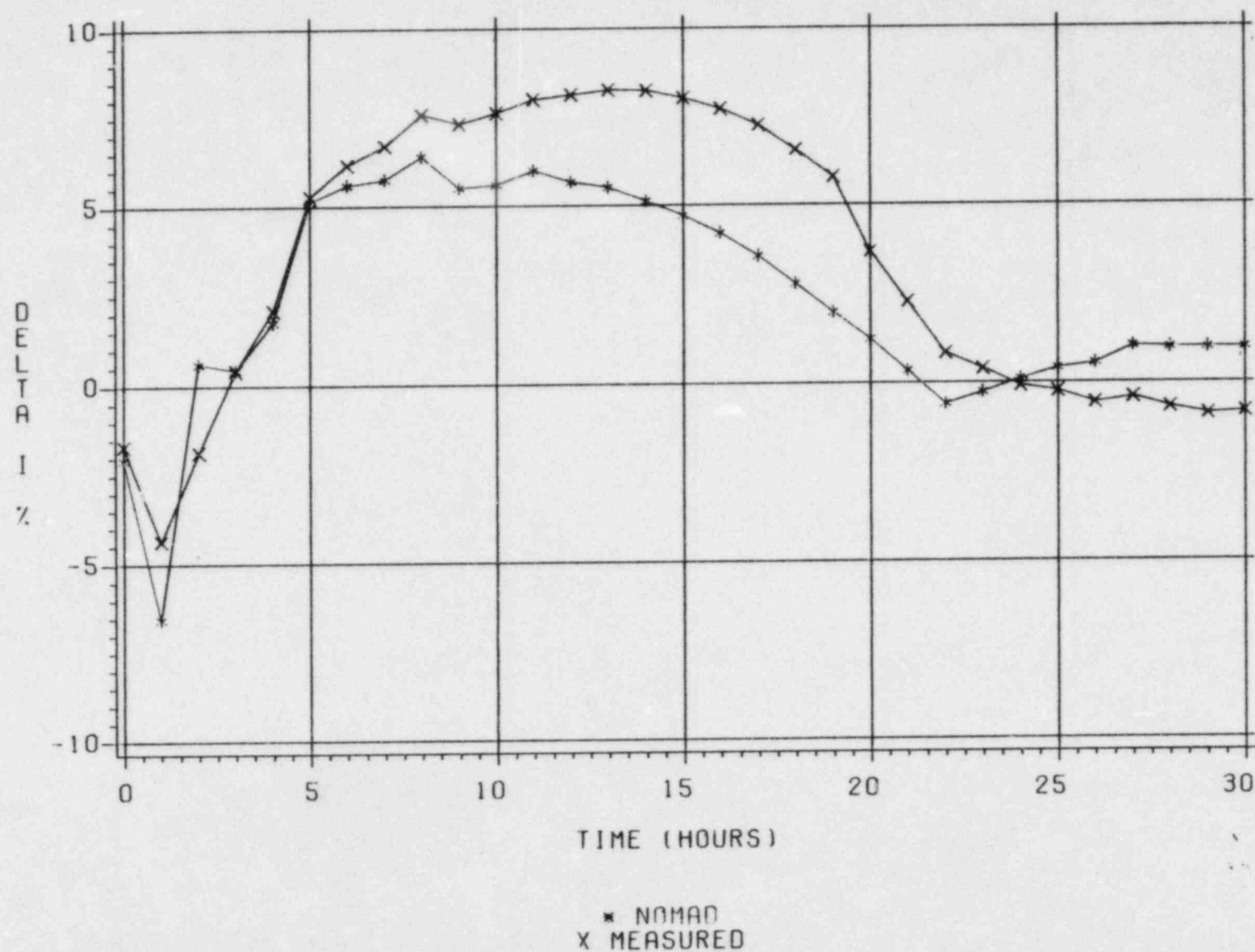


FIGURE 5-33

N1C2 70% LOAD REDUCTION TEST
CRITICAL BORON CONCENTRATION

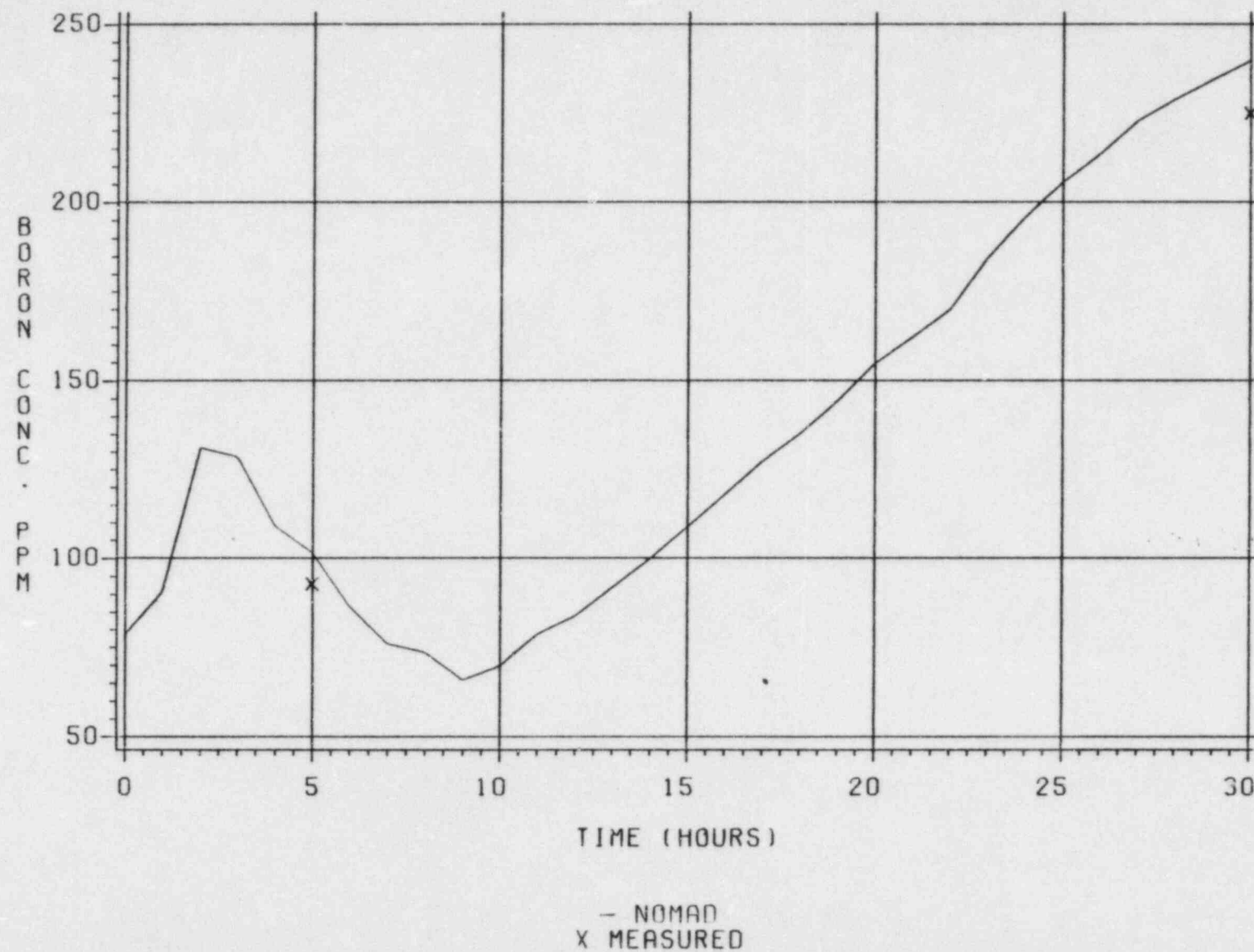


FIGURE 5-34

N1C3 SHUTDOWN / RETURN TO POWER (CASE 1)
AXIAL FLUX DIFFERENCE

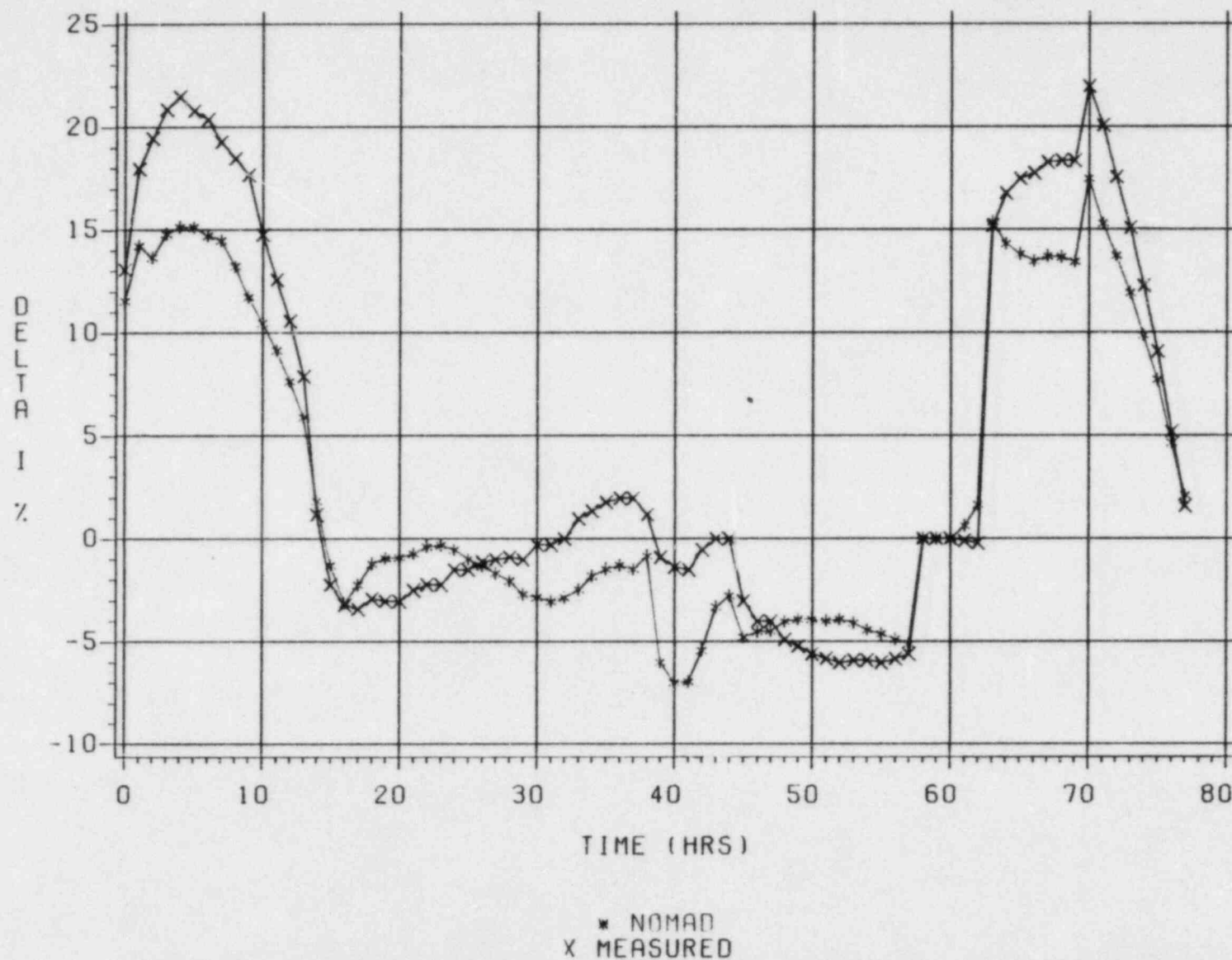
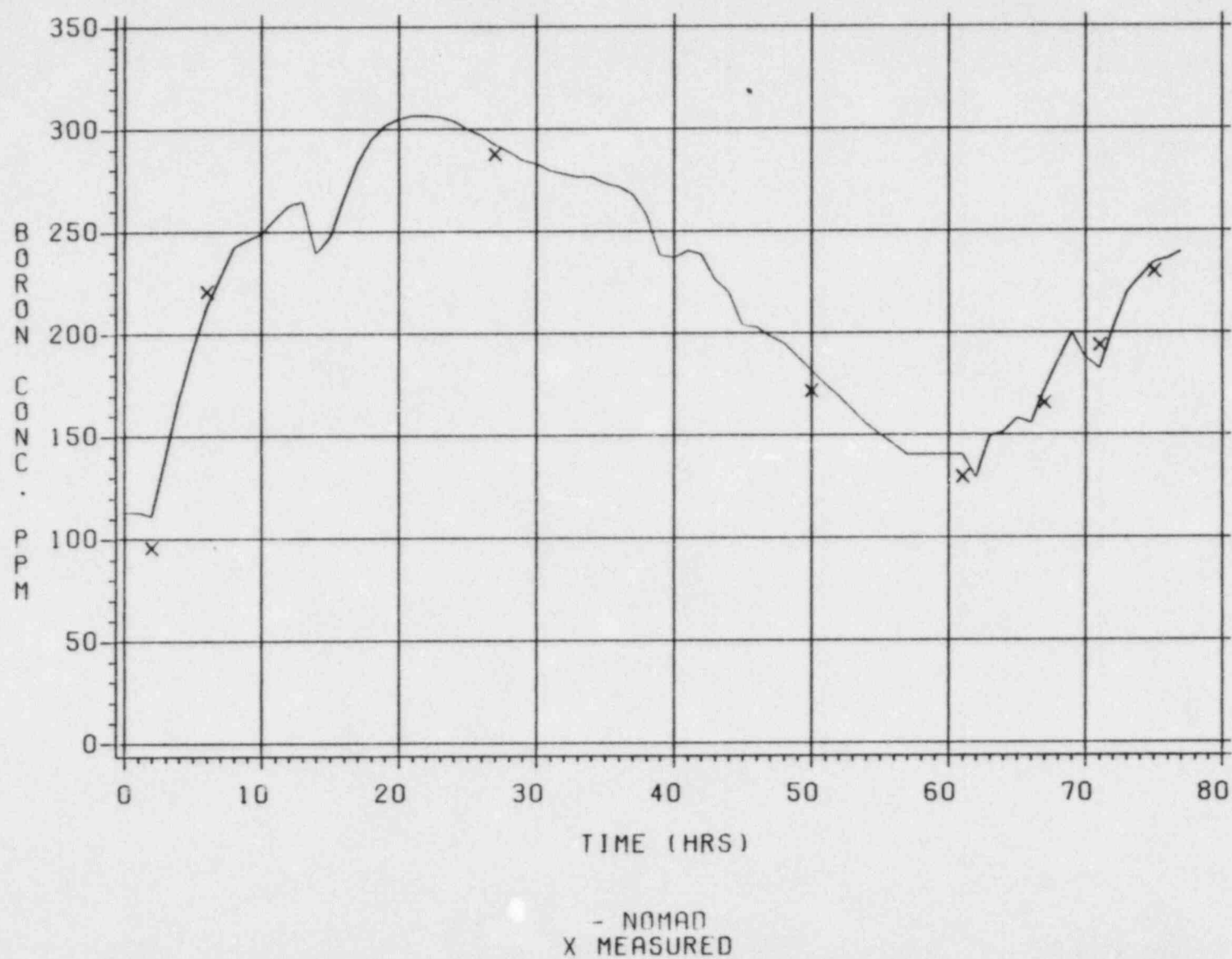


FIGURE 5-35

N1C3 SHUTDOWN/RETURN TO POWER (CASE 1)
CRITICAL BORON CONCENTRATION



N1C3 SHUTDOWN/RETURN TO POWER (CASE 2)

AXIAL OFFSET

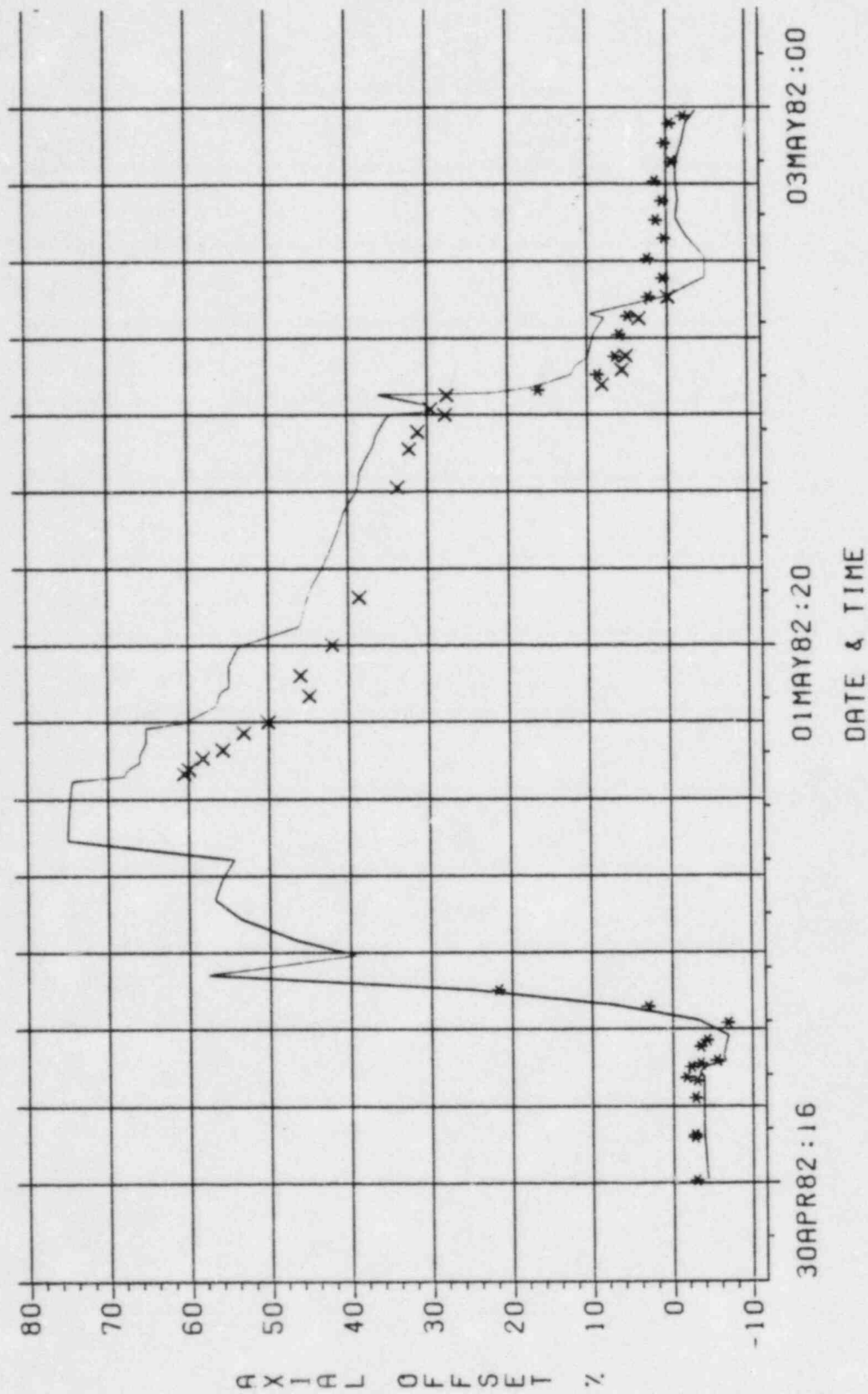


FIGURE 5-37

5-51

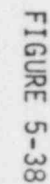


FIGURE 5-39

FZ(Z) RESULTS
 NORTH ANNA UNIT 1, CYCLE 4
 (18 Case FAC Analysis)

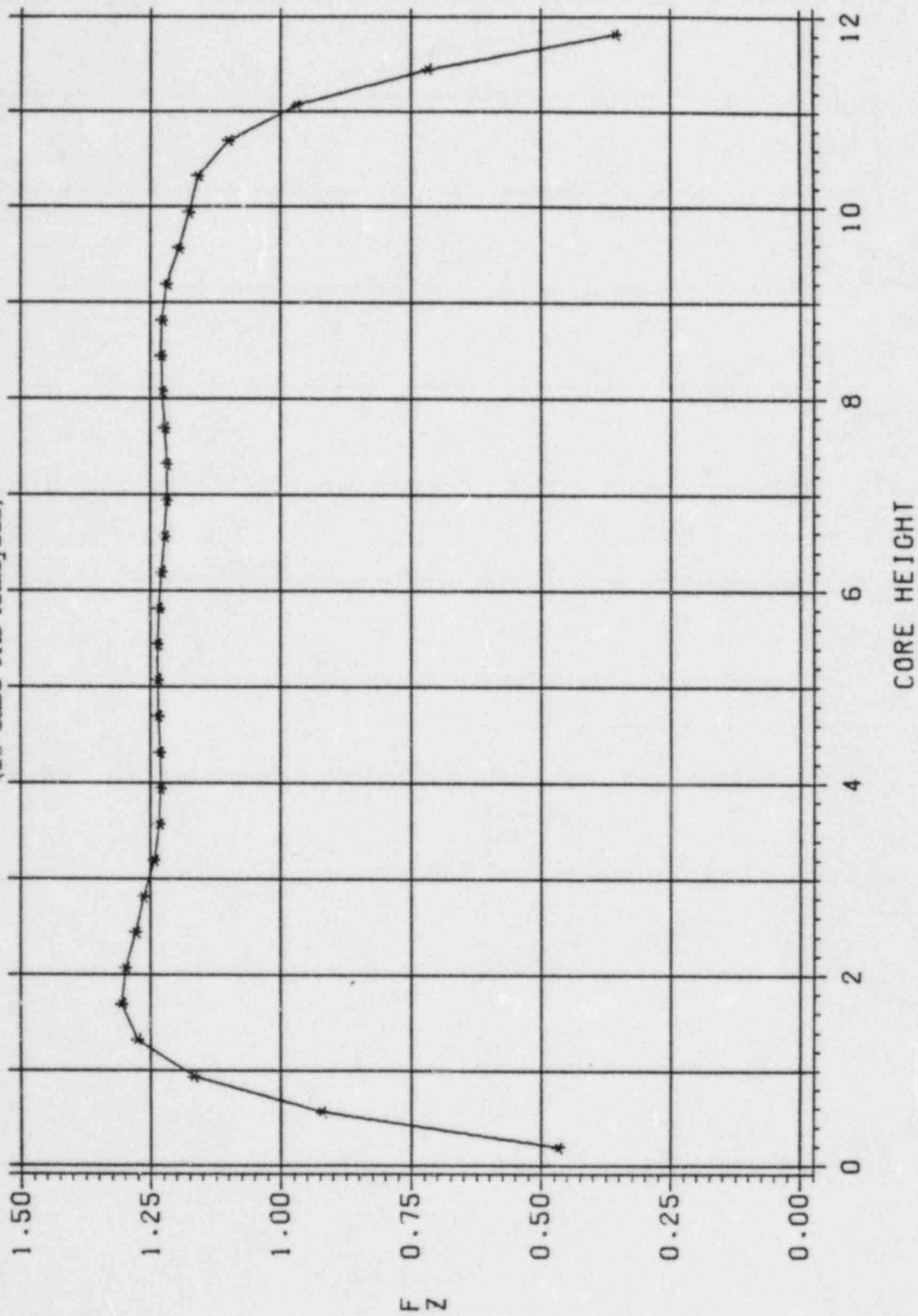


FIGURE 5-40

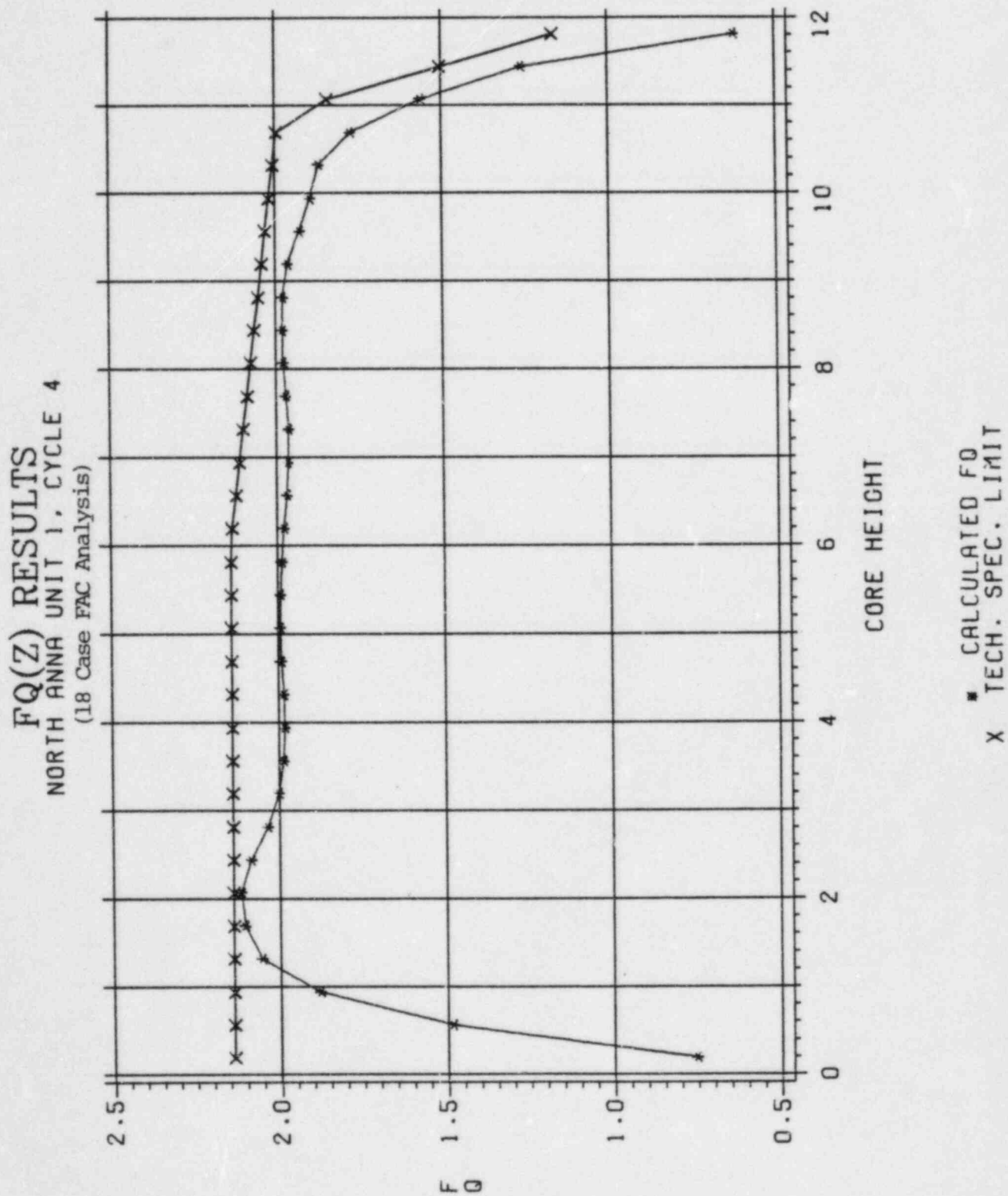


FIGURE 5-41

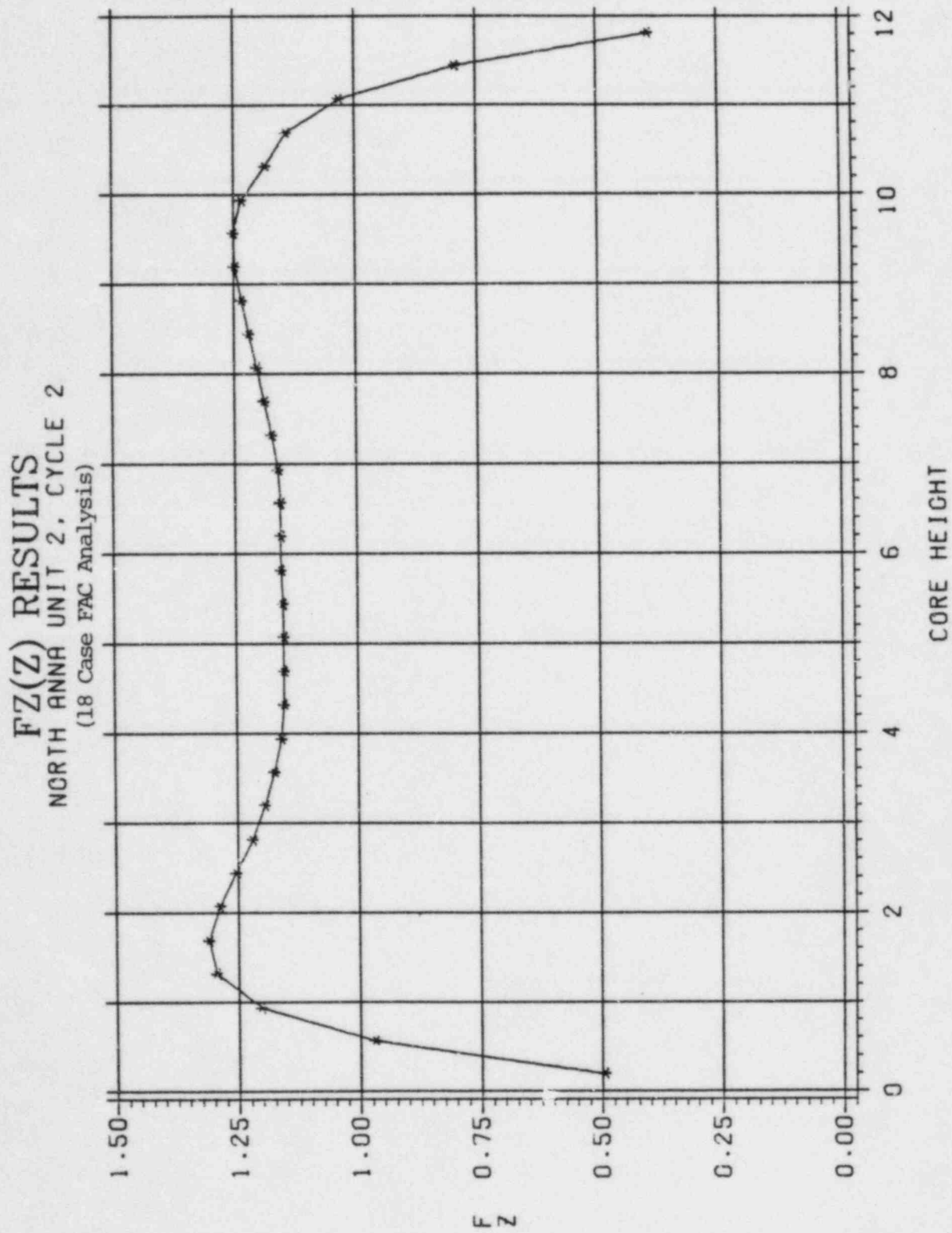
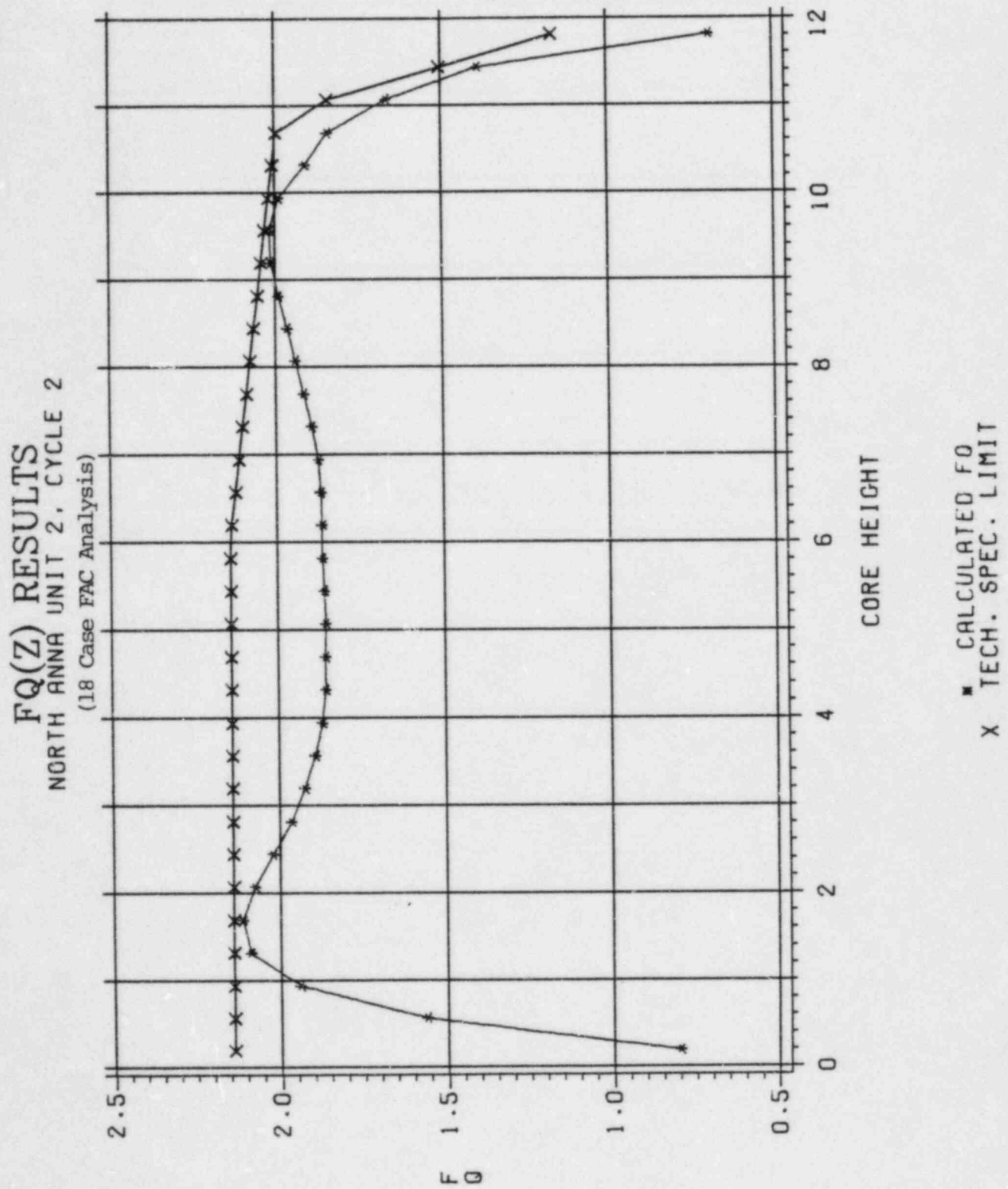


FIGURE 5-42



SECTION 6 - SUMMARY AND CONCLUSIONS

The Vepco NOMAD code and model are operational at Vepco for the purpose of performing one-dimensional reactor physics analyses and supporting the evaluation of core performance. The model consists of NOMAD with the NULIF, XSED, XSFIT, XSEXP, FXYZ, FDELH, and PCEDT codes providing either input or data manipulation. The accuracy of the Vepco NOMAD model has been established through extensive comparison of calculations to measurements from the units at the Surry and North Anna Nuclear Power Stations and to calculations from other Vepco codes. The results of these comparisons indicate that the Vepco NOMAD model (which includes normalization to the Vepco PDQ07 Discrete, PDQ07 One Zone, and FLAME models) provides the capability to predict core peaking factors, axial power distributions, differential and integral rod worths, and load follow maneuvers as well as perform Final Acceptance Criteria (FAC) Analysis.

Verification of and improvements to the Vepco NOMAD code and model will continue to be made as more experience is obtained through the application of the model to the units at the Surry and North Anna Nuclear Power Stations.

SECTION 7 - REFERENCES

1. S. M. Bowman, "VEPCO 1-D Code Development: Final Report", NFE Technical Report No. 250, March, 1983 (Virginia Electric and Power Co.).
2. D. A. Daniels, "Fxy(Z) Synthesis Methodology and Computer Code", NFE Technical Report No. 180, January, 1981 (Virginia Electric and Power Co.).
3. J. G. Miller, "The FDELHP01 and PCEDTP01 Codes", NFE Technical Report No. 201, July, 1981 (Virginia Electric and Power Co.).
4. P. D. Breneman, "The NULIFP01 Code", NFE Calculational Note PM-13, March, 1979 (Virginia Electric and Power Co.).
5. M. L. Smith, "The PDQ07 Discrete Model", VEP-FRD-19A, July, 1981 (Virginia Electric and Power Co.).
6. J. R. Rodes, "The PDQ07 One Zone Model", VEP-FRD-20A, July, 1981 (Virginia Electric and Power Co.).
7. W. C. Beck, "The Vepco FLAME Model", VEP-FRD-24A, July, 1981 (Virginia Electric and Power Co.).
8. L. L. Lynn, "A Digital Computer Program for Nuclear Reactor Analysis Design Water Properties", WAPD-TM-680, July, 1967 (Westinghouse Electric Corporation).
9. J. G. Miller, "Thermal-Hydraulic Feedback Input to the Two-Dimensional PDQV2 Code", NFE Calculational Note PM-19, June, 1979 (Virginia Electric and Power Co.).
10. Course Notes "Basic PWR Physics Course", September, 1980 (Westinghouse Electric Corporation).
11. F. W. Sliz and K. L. Basehore, "VEPCO Reactor Core Thermal-Hydraulics Analysis Using the COBRA IIIc/MIT Computer Code", VEP-FRD-33-A, October, 1983.
12. D. A. Daniels, "XETRN and SMTRN", NFE Technical Report No. 112, February, 1980 (Virginia Electric and Power Co.).