Computer Code Abstracts

TRIFIDO

1. Name of Code: TRIFIDO, a code for calculating kinetic parameters of subcritical neutron multiplying assemblies using pulsed-neutron experimental data.

2. Computer for Which Code is Designed: IBM/360-65 or BULL-625. Programming Language Used: FORTRAN IV.

3. Nature of Physical Problem Solved: The code calculates decay constant and the population of the fundamental prompt-neutron mode extrapolated to initial time, using pulsed neutron experimental data. These data are the resulting time profile of the neutron density of a subcritical multiplicative assembly which is repetitively pulsed with short bursts of neutrons. The time profile is measured by an appropriate detector and recorded with a time analyzer. With the calculated parameters the code determines the values of $k_{sp}$/L and the reactivity by means of the Garelis-Russell method, and the reactivities using the Gozani and Sjöstrand methods.

4. Method of Solution: Least-squares weighted fit for the decay constant and the extrapolated-population calculations; iteration for the Garelis-Russell method.

5. Restrictions on the Complexity of the Problem: The code is presently restricted to 256 experimental data points (for example, those provided by a TMC multi-channel analyzer). This number is adequate for most pulsed neutron measurements. The restriction can be easily overcome, if necessary, by changing a dimension card. Minor parameter adjustments are needed according to the multiplying systems used and to the experimental procedure.

6. Typical Machine Time: 3.60 sec for 1 case in a BULL GE-625 computer, 16.92 sec for 10 cases; 70% of these times on an IBM/360-65 computer.

7. Unusual Features of the Program: It calculates several pairs of decay constants and extrapolated populations of the fundamental prompt-neutron mode varying the starting analysis channel (initial time). Since the code selects as fundamental prompt mode values those which provide the best agreement between fitted and experimental values, it can proceed directly to calculation of $k_{sp}$/L and reactivity. Thus, all these kinetic parameters are obtained in only one computer run. To provide visual checking, the code draws a graph of the experimental data, corrected for noise and dead-time effects, and the fitted values.

8. Status: In use.


10. Operating System or Monitor Under Which Program is Executed: Gecos 3 on BULL GE-625, OS/360 or IBM/360.


12. References:

- T. GOZANI, EIR-Bericht Nr. 79 (April 1965).

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ANCON
A Point-Kinetics Program with Nonlinear Feedback

1. Name of Code: ANCON

2. Computer for Which Code is Designed: CDC-6600. Because ANCON is written entirely in FORTRAN and does not require any special system routines or equipment, it should be operable on most computers.

3. Nature of Physical Problem Solved: ANCON solves the point-reactor kinetic equations including thermal feedback. Lump-type heat balance equations are used to represent the thermodynamics, and the heat capacity of each lump can vary with temperature. Thermal feedback can be either a linear or a nonlinear function of lump temperature, and the impressed reactivity can be either a generalized polynomial function or a sinusoidal function.

4. Method of Solution: The system of coupled first-order differential equations is solved by a method based on continuous analytic continuation. The basic procedure consists of expanding all the dependent variables except reactivity in Taylor series, with a truncation error criterion, over successive intervals on the time axis. Variations of the basic procedure are used to increase the efficiency of the method in...