





This work was supported by the US Department of Energy, Division of Reactor Research and Technology, and the Electric Power Research Institute.

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LA-9303-M, Vol. I (ENDF-324) Manual

Issued: May 1982

The NJOY Nuclear Data Processing System, Volume I: User's Manual

R. E. MacFarlane D. W. Muir R. M. Boicourt



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THE NJOY NUCLEAR DATA PROCESSING SYSTEM, VOLUME I: USER'S MANUAL

by

R. E. MacFarlane, D. W. Muir, and R. M. Boicourt

ABSTRACT

The NJOY nuclear data processing system is a comprehensive computer code package for producing cross sections for neutron and photon transport calculations from ENDF/B-IV and -V evaluated nuclear data. This user's manual provides a concise description of the code, input instructions, sample problems, and installation instructions.

I. INTRODUCTION TO VOL. I

The NJOY¹⁻³ nuclear data processing system is a comprehensive computer code package for producing pointwise and multigroup neutron and photon cross sections from ENDF/B-IV and -V evaluated nuclear data.⁴ This document provides a concise description of the (10/81) version of the code, operating instructions, code installation instructions, and sample problems. Full descriptions of theory, methods, and programming details are given in subsequent volumes of this report.

II. CODE DESCRIPTION

The NJOY code consists of a set of modules, each performing a well-defined processing task. The RECONR module reconstructs pointwise (energy-dependent) cross sections from ENDF/B resonance parameters and interpolation schemes. BROADR Doppler-broadens and thins pointwise cross sections. UNRESR computes effective self-shielded pointwise cross sections in the unresolved-resonance region. HEATR generates pointwise heat production cross sections (kerma factors) and radiation-damage-energy production cross sections. THERMR produces

incoherent inelastic energy-to-energy matrices for free or bound scatterers, coherent elastic cross sections for hexagonal materials, and incoherent elastic GROUPR generates self-shielded multigroup cross sections, cross sections. group-to-group neutron scattering matrices, and photon production matrices from pointwise input. GAMINR calculates multigroup photon interaction cross sections and kerma factors and group-to-group photon scattering matrices. ERRORR produces multigroup covariance matrices from ENDF/B uncertainties. COVR reads the output of ERRORR and performs covariance plotting and output formatting operations. DTFR formats multigroup data for transport codes, such as $DTF-IV^5$ and $ANISN.^6$ CCCCR formats multigroup data for the CCCC standard⁷ interface files ISOTXS. BRKOXS, and DLAYXS. MATXSR formats multigroup data for the MATXS cross-section interface file. The ACER module prepares libraries for the Los Alamos continuous-energy Monte Carlo code MCNP.⁸ POWR prepares libraries for the EPRI-CELL and EPRI-CPM codes.* Finally, MODER changes ENDF/B "tapes" and other ENDF-like NJOY interface files back and forth between formatted (that is, BCD or ASCII) and blocked-binary modes. NJOY incorporates and improves upon the features of its direct ancestor, MINX.⁹ It also includes and extends the photon production capabilities of LAPHANO, 10 the photon interaction capabilities of GAMLEG, 11 the heating capabilities of MACK, 12 the covariance capabilities of PUFF, 13 and the thermal capabilities of FLANGE-II¹⁴ and HEXSCAT.¹⁵

The methods used in these modules are described in detail in subsequent volumes of this report. The following brief account will make the general flow of the code clear. RECONR reads an ENDF/B tape and produces a common energy grid for all reactions (the union grid) such that all cross sections can be obtained to within a specified tolerance by linear interpolation. Resonance cross sections are calculated with the methods of RESEND,¹⁶ but a new method of choosing the energy grid is used that incorporates control of the number of significant figures generated and a resonance-integral criterion to reduce the number of grid points generated for some materials. Summation cross sections (for example, total, inelastic) are reconstructed from their parts. The resulting pointwise cross sections are written onto a "point-ENDF" (PENDF) tape for future use. BROADR reads a PENDF tape and Doppler-broadens the data using the method of SIGMA1,¹⁷ modified for better behavior at high temperatures and low

^{*}EPRI-CELL and EPRI-CPM are proprietary codes. Additional information can be obtained from the Electric Power Research Institute, 3412 Hillview Ave, Palo Alto, California 94394.

energies. The union grid allows all resonance reactions to be broadened simultaneously, resulting in a great savings of processing time. After broadening, the summation cross sections are again reconstructed from their parts. The results are written out on a PENDF tape for future use. UNRESR uses the methods of ETOX¹⁸ to produce effective self-shielded pointwise cross sections, versus temperature and background cross section, in the unresolved-resonance region. The results are added to the PENDF tape in a special format. HEATR computes both heating and radiation-damage-energy production using momentum balance (for capture) or energy balance (for all other reactions). The ENDF/B photon production files are used in both methods when available. The heating results are added to the PENDF tape using ENDF/B reaction numbers in the 300 series, and the radiation damage results use the special identifier 444. THERMR produces pointwise cross sections in the thermal range. Bragg edges in coherent scattering are produced using the method of $HEXSCAT^{15}$ with an improved treatment at high energies. Energy-to-energy incoherent scattering matrices can be computed for free scattering or for bound scattering using a precomputed form factor $S(\alpha,\beta)$ in ENDF format. The secondary angle and energy grids are determined adaptively so as to represent the function to a desired precision by linear interpolation; the angular representation is converted to one based on equally probable angles. Elastic incoherent scattering is represented using equally probable angles computed analytically. The results for all the processes are added to the PENDF tape using special formats and reaction numbers.

GROUPR processes the pointwise cross sections produced by the modules described above into multigroup form using the Bondarenko flux-weighting model.¹⁹ As an option, a pointwise flux solution can be generated for a heavy absorber in a light moderator. Self-shielded cross sections, scattering matrices, and photon production matrices are all averaged in a unified way, the only difference being in the function that describes the "feed" into secondary group g' with Legendre order ℓ from initial energy E. The feed for two-body scattering is computed using a center-of-mass Gaussian integration scheme, which provides high accuracy even for small Legendre components of the scattering matrix. Special features are included for delayed neutrons, the coupled angle and energy dependence of the thermal scattering matrix, and the discrete scattering angles arising for thermal coherent reactions. Prompt fission is treated with a group-to-group matrix. The results are written in a special "groupwise-ENDF" format

(GENDF) for later use by the output formatting modules. GAMINR uses a specialized version of GROUPR. Coherent and incoherent form factors²⁰ are processed in order to extend the useful range of the results to lower energies. Photon heat production cross sections are also generated. The results are saved on a GENDF tape. ERRORR can either produce its own multigroup cross sections using the methods of GROUPR or start from a precomputed set. The cross sections and ENDF covariance data are combined in a way that includes the effects of deriving one cross section from several others. Special features are included to process covariances for data given as resonance parameters or ratios (for example, fission \overline{v}). The COVR module uses the widely available DISSPLA* plotting software to make publication-quality plots²¹ of covariance data; it also provides a site for user-supplied routines to prepare covariance libraries for various sensitivity systems. DTFR is a simple reformatting code that produces cross-section tables acceptable to most discrete-ordinates codes. It also converts the GROUPR fission matrix to χ and $\bar{v}\sigma_f$ and prepares a photon production matrix if desired. The user can define edit cross sections that are any linear combination of the cross sections on the GENDF tape. This makes complex edits such as gas production possible. DTFR also contains system-dependent plotting routines for the cross sections, P_n scattering matrices, and photon production matrices. CCCCR is also a straightforward reformatting code. All of the CCCC-IV 7 options are supported. In the cross-section file (ISOTXS), the user can choose either isotope χ matrices or isotope χ vectors collapsed using any specified flux. The BRKOXS file includes self-shielding factors for elastic removal. Note that some of the cross sections producible with NJOY are not defined in the CCCC-IV files. MATXSR reformats GENDF data into the MATXS file format, which is suitable for input to the TRANSX postprocessor program. The MATXS format uses flexible naming conventions that allow it to store all NJOY data types except delayed neutron and delayed photon spectra.

In the reference version of the code, each module is a separate overlay. The main overlay (NJOY) simply calls in each primary overlay (for example, RECONR, GROUPR) as requested by the user's input commands. The NJOY level also contains utility routines used by all other modules (for example, free-form input, storage allocation, and ENDF/B input/output). The code can easily be decomposed into 15 independent programs and a user library.

^{*}DISSPLA is a proprietary graphics software package. For more information, contact Integrated Software Systems Corp., 4186 Sorrento Valley Blvd., San Diego, California 92121.

III. NJOY INPUT INSTRUCTIONS

All NJOY input is in free form. A subroutine FREE has been included among the NJOY utilities to provide this capability. Fields on the input cards are delimited by any character not used for another purpose (+,-,numeral,E,H,*,R,/). For exponent fields, the E must be present, and spaces are not allowed before the E. Decimal points are not required after numbers. Hollerith entries may use nHSTRING or *STRING*. The form nR causes the following number to be repeated n times. The (/) terminates the input for one call to FREE (it may involve more than one card) leaving any unread variables unchanged. This feature is often used to default variables from the right. The part of the input line to the right of the (/) can be used for comments if desired.

As an example of when the (/) is useful, in several NJOY routines a record of Hollerith information is constructed from user input. This is accomplished by calling FREE with NZA = 17 (the number of Hollerith words required to fill the 66 columns available for an ENDF/B "comment"). The array is preset to 17 blank words, so that the user need not blank-fill the line explicitly. Instead, he can write *MESSAGE*/ where the (/) terminates the process of replacing the default blanks with actual input ("MESSAGE" in this example).

The user should be cautioned that if the (/) is omitted from an input data block that is incomplete, as in the Hollerith example above, FREE will go on reading successive input data cards until the expected NZA words are found, usually resulting in an error condition. For this reason, if the user is uncertain whether he has supplied enough input parameters to "satisfy" a particular call to FREE, it is good practice to use a (/) at the end of the input data for that data block.

Some input examples follow.

LEC	GAL	ILLEGAL
12 12.	1.2E1	1.2+1 1.2 E1
U235	4HU235	4RU235 (does not mean "right-justify")
5R0	2R 1.2E2	

The input to the NJOY sample problems, Section IV, provides additional illustrations of the syntax of free-form input.

The flow through the modules of NJOY is controlled by module names and input/output logical unit numbers. A module typically reads data from an input unit, modifies it, and writes the results on an output unit. Sometimes auxiliary inputs or multiple outputs are required. The output of one module is often the input for another.

RECONR 21 22 . [input for RECONR] *BROADR* 22 25 . [input for BROADR]

Because the files on most units are in ENDF/B format, the modules can be connected in many ways. The files can be saved at any point for later restart. Other combinations will be found in the sample problems.

NJOY provides for a special blocked-binary mode for the ENDF/B files. Such files are indicated with negative unit numbers. The MODER module can be used to convert back and forth between formatted (that is, BCD or ASCII) and blockedbinary modes. The user may assign unit numbers from 20 through 99 for linking modules. Many modules also accept 0 as a unit number, which means that the corresponding file is not used. Unit numbers from 10-19 are reserved for scratch files, and units 5-7 are used for system input and output files.

Detailed input instructions are included as comment cards at the start of each module (overlay); the current set of comment-card instructions is reproduced in this report as Appendix A. The following sections supplement the material in Appendix A by providing expanded descriptions of the significance of certain input parameters. Additional details will be found in subsequent volumes of this report.

A. NJOY Module

IOPT... When operating in a time-sharing environment (IOPT = 1), the code routes input prompts and short output messages to the terminal (TTY). The regular long output is still available for the system printer.

B. RECONR Module

NENDF... This is the unit containing an ENDF/B-IV or -V tape. It is recommended that binary mode be used whenever possible (that is, use MODER).

ERR... A reasonable value is 0.005 ($\frac{1}{2}$ %). For materials with many resonances, such as ⁹³Nb and ²³⁸U, it may be necessary to relax (increase) this somewhat in order to reduce the running time.

ENDF/B tapes are available from the National Nuclear Data Center (NNDC) at Brookhaven National Laboratory, Upton, New York.

TEMPR... Resonances can be constructed directly at an elevated temperature using the $\psi \chi$ resonance shapes (with some loss in accuracy and great savings in time) for single-level Breit-Wigner or Adler-Adler representations. In general, it is recommended that resonance reconstruction be performed at zero Kelvin (TEMPR = 0); BROADR can then be used to Doppler-broaden to the desired temperature with great accuracy.

NDIGIT... All energies are adjusted to have exactly this number of significant digits before the cross section is computed, and no two energies are allowed to be equal. For formatted output, NDIGIT must be less than or equal to 7; larger values are allowed with binary output, if permitted by the word length of the computer.

ERRMAX, ERRINT... These parameters control an optional resonance-integral thinning procedure, which helps reduce the number of energy points generated during resonance reconstruction. The option can be turned off by specifying ERRMAX = ERR.

C. BROADR Module

ISTART... Since Doppler broadening is comparatively expensive, it is often useful to "restart" from previous data at a lower temperature. For example, in attempting to produce cross sections at 300, 900, and 2100 K, the job runs out of time while doing the last temperature. Set ISTART = 1 and TEMP1 = 900, and continue the job using the output of the aborted run as input. (If ISTART = 0, none of the input cross sections are copied to the output.)

ISTRAP... "Bootstrap" refers to using the output of one broadening run as the input for a subsequent higher temperature run. If thinning was used, the second run can be much faster than the first. However, errors will accumulate.

ERRTHN... Cross sections become smoother with Doppler broadening and can be thinned. It is recommended that ERRTHN \leq ERR be used. (See Item B above.)

THNMAX... Broadening and thinning are not normally performed above 1 MeV or the inelastic threshold, whichever is lower. In some cases, it is useful to further reduce this energy, for example to the top of the resolved-resonance energy range or to an energy that avoids sharp steps or triangular representations of resonances found for some evaluations (for example, ENDF/B-V lead). D. HEATR Module

NPK,MTK... Usually only the total kerma is needed (NPK = 0). For structural metals, MTK = 444 can be requested to obtain the total radiation damage production cross section. Other partials are useful in connection with IPRINT = 2.

NQA,MTA,QA... Because of lack of uniformity in evaluation practices, or because of the varying energy values of the separate isotopes in an element, it is sometimes necessary to override the evaluator's Q-values.

IPRINT = 2... This option is provided mainly for evaluators. Computed kerma factors are compared with limits obtained from kinematics, and the capture photon spectrum is compared to the available energy. If this option reveals many violations of energy balance, the results of both heating and gamma dose calculations must be held in doubt.

LOCAL... If photon production data are available, the energy carried away by photons is normally subtracted from the energy available for local heating (LOCAL = 0); the energy appears elsewhere in the system as a result of photon interactions. If photon transport is not treated, this energy should be included in the local heating (LOCAL = 1).

E. THERMR Module

NENDF,MATDE... The thermal ENDF data (only MF = 7 is used) are available on a special set of tapes (320-325).²² Each material has a special MAT number to be used for MATDE and a special list of temperatures. No tape or MATDE value is required for free gas scattering.

NIN,MATDP... The MAT numbers on the input PENDF tape are the MAT numbers from the main evaluation, not the thermal MAT number. The temperatures on the PENDF tape must include the desired values from the thermal tape.

NATOM... For example, 2 for hydrogen in water.

MTREF... Thermal data generated by this module are written onto the PENDF tape using MT = MTREF, which must be in the range 201-250. The MATXSR module recognizes the MT-numbers given in Table I.

F. GROUPR Module

NTEMP, TEMP... The requested temperatures must occur on the input PENDF tape.

NSIGZ,SIGZ... One or more "background" cross sections must be specified in barns for the calculation of self-shielded group constants using the Bondarenko formalism. For infinite dilution, NSIGZ = 1, and by convention, SIGZ = 1.E10. If unresolved data have been added to the PENDF tape using UNRESR, the NSIGZ SIGZ values in GROUPR do not have to agree with the σ_0 values used in UNRESR. In both codes, SIGZ is read in, in descending order, with $\sigma_0 = 1.E10$ first.

WGHT... If the user wishes to supply his own weight function, he must use a "TAB1" record. This is a particular ENDF/B data structure which, if a single

TABLE I

ENDF/B-V	ENDF/B-IV	CONTENTS
221	201	free gas
222	202	H in H ₂ 0
223	203	inelastic H in CH,
224	204	elastic H in CH ₂ 2
225	205	inelastic H in ŹrH
226	206	elastic H in ZrH
227	207	benzene
228	208	D in D ₂ 0
229	209	inelastic graphite
230	210	elastic graphite
233	219	inelastic Be
234	220	elastic Be
231	229	inelastic BeO
232	230	elastic BeO
235		inelastic Zr in ZrH
236	-	elastic Zr in ZrH
	_	

REACTION NUMBERS FOR THERMAL SCATTERING

interpolation scheme INT is employed, has the following form.

card 1 0. 0. 0 0 1 N card 2 N INT cards 3,4,... (E(I),WGHT(I),I=1,N)/

Here N is the number of energy-weight pairs, and INT specifies the functional form to be used to connect the points. For example, INT = 2 specifies that WGHT is a linear function of E between points and INT = 5 specifies that log(WGHT) is a linear function of log(E) between points. The (/) shown after the final weight is <u>required</u>. This is a departure from the normal NJOY convention where (/) is needed only if the reading of a data block is to be terminated prematurely.

MFD,MTD,MTNAME... GROUPR requires that the user specify each reaction type to be processed using its ENDF/B "MT-number." Further, it is illegal to request a reaction type that is absent in the evaluation. The MT-numbers used in a given evaluation can be obtained from the "dictionary" (MF1, MT451), a list of all the "files" (MF-numbers) and "sections" (MT-numbers) for this material (MAT-number). The user should first examine the list of sections of file 3. Each section describes a different nuclear reaction, so the user can select from this list the reactions he wishes to process. A table of ENDF/B MT-numbers is reproduced here as Appendix B. In order to process reaction "vectors" (as opposed to group-to-group transfer matrices), one inputs data cards such as

> 3 103 *(N,P)*/ 3 105/ .

Here MFD=3 specifies a vector, MTD=103 is the section to be processed, and "(N,P)" is the user-supplied name for the reaction. This name is only used to label the listing and is optional as shown by the second line. There are several special MT-numbers recognized by NJOY while processing vectors.

<u>MT</u>	Meaning			
252 253 258 259 452 455 455	 μ (average scattering cosine) ξ (average log decrement) mean lethargy mean reciprocal velocity total fission yield delayed fission yield prompt fission yield 			

The first three are computed from fundamental definitions, not file 3. The last three are computed using MF1 yields and MF3 fission cross sections.

To process one matrix reaction (that is, group-to-group scattering), use use MFD=6 as in this example:

6 16 *(N,2N)*/.

The reaction types with scattering data are most easily found under MF=4 in the dictionary. Caution: MF4, MT103-150 are charged particle angular distributions and should not be requested as matrices.

As a convenience feature, a consecutive sequence of MTDs can be specified as follows.

3 51 *FIRST INELASTIC LEVEL*/ 3 -76 *HIGHER INELASTIC LEVELS*/ 6 51 *FIRST INELASTIC LEVEL*/ 6 -76 *HIGHER INELASTIC LEVELS*/

All values of MTD from 51 through 76 will then be processed into both vectors and matrices.

Fissionable materials introduce some additional complexity. GROUPR produces a prompt fission group-to-group matrix that can be converted into the traditional $v\sigma_f$ and χ vectors by later modules (for example, DTFR). For simple evaluations, it is only necessary to make this request.

3	18	*FISSION	XSEC*/
6	18	*FISSION	MATRIX*/

In several important evaluations, however, the evaluator has divided the fission process into parts: MT19, direct fission (n,f); MT20, second-chance fission (n,n')f; MT21, third-chance fission (n,2n)f; and MT38, fourth-chance fission (n,3n)f. The procedure makes possible a more accurate representation of the high-energy portion of the fission spectrum when fission is induced by neutrons with energies above 5 or 6 MeV. For such evaluations (for example, 235 U, 238 U, 239 Pu), the following input is recommended.

3	18	*TOTAL FISSION*/
3	19	*(N,F)*/
3	20	*(N,N)F*/
3	21	*(N,2N)F*/
3	38	*(N,3N)F*/
6	19	*(N,F)*/
6	20	*(N,N)F*/
6	21	*(N,2N)F*/
6	38	*(N.3N)F*/

Note that 6/18 is omitted. A subsequent module, such as DTFR or MATXSR, can add the partial matrices to obtain the total fission matrix.

A final complication of fission is the existence of delayed neutrons from fission. For those materials that contain delayed neutron data, the user should request these.

3 455 *DELAYED NUBAR*/ 5 455 *DELAYED CHI*/

A later module can add the delayed data to the prompt matrix in order to obtain "steady-state" values for $\bar{v}\sigma_f$ and χ .

The matrix representation of fission is very general, but the results are bulky and expensive, especially for large group structures (for example, the 620-group dosimetry set). For such cases, the short-cut fission spectrum option (MFD=5) can be used. The MTD-value selects the parameters as follows.

MTD <u>Requested</u>	Spectrum from MF/MT	Incident Energy	
18 or 452	5/18	1 MeV	
19 or 456	5/19	1 MeV	
455	5/455	2 MeV	

The incident energy can be changed at line 407 of GROUPR.

If the evaluation includes photon production data, GROUPR will prepare a neutron-to-photon transfer matrix for each reaction requested. These reactions are identified in the material dictionary by the presence of MF=12 (photon

production yields), MF13 (photon production cross sections), or both. For each reaction, input a data card with MFD=MF+4; for example,

16 102 *CAPTURE GAMMA PRODUCTION*/
17 3 *NON-ELASTIC GAMMA PRODUCTION*/ .

The ENDF/B-V format includes multiplicities (MF9) or cross sections (MF10) for the production of radioactive nuclides, including isomers. These files are converted into multigroup form by requesting MFD=90+level or MFD=100+level, respectively; for example,

90 16 *N2N GROUND STATE*/ 91 16 *N2N ISOMER PRODUCTION*/.

Several examples of GROUPR input will be found in the sample problems. G. GAMINR Module

MFD=-1... This option provides the user with a standard list of reaction types suitable for all elements: vectors for 501, 502, 504, 516, 602, and 621; matrices for 502, 504, and 516. Note that 621 is assigned specially for the photon heat production cross section.

H. ERRORR Module

NPEND,NGOUT... The user must supply either a PENDF tape, NPEND, or a GENDF tape, NGOUT, with processed pointwise or groupwise data produced by another NJOY module. Group cross sections need not be in the same group structure as the requested output covariances, although, if the input group structure is much coarser than the output structure, rather crude approximations will be made in deriving an effective set of fine-group cross sections from the coarse input data.

NIN,NOUT... Although ERRORR lacks an explicit multimaterial loop, the same effect can be achieved by executing a series of ERRORR runs, with the output covariance tape NOUT of one run becoming the input covariance tape NIN of the next. Only two unit numbers need be employed, with the data going back and forth between them until the multimaterial library is complete. To save time, binary files should be used for this purpose.

IREAD... For Version V, the list of reactions for which covariances are produced is constructed in one of several ways, depending on the value of IREAD. If IREAD=0, the reaction list is assumed to be identical to the list of sections contained in file 31 or file 33 (see MFCOV below). If IREAD=1, only the "covariance" reactions specifically named by the user will be included in the reaction list. For certain applications, this option can save considerable execution

In either case, IREAD=0 or 1, a complete set of covariance matrices is time. output. That is, covariance matrices are produced for every reaction-pair combination that can be formed from the given reaction list. For those reaction pairs where the evaluator has not specified the covariances, the output matrix contains only zeros. If IREAD=2, the reaction list is initially constructed in the same way as for IREAD=0. This list is then supplemented with a list of extra reactions $MAT1_k/MT1_k$ (MAT1_k #MATD), specified by the user. The output in this case contains matrices for all of the $(MATD/MT_i; MATD/MT_i)$ combinations, as before, plus matrices for all $(MATD/MT_i; MAT1_k/MT1_k)$ combinations. However, covariances among the <u>extra</u> reactions, that is $(MAT1_k/MT1_k; MAT1_m/MT1_m)$, are not computed. In an evaluation containing cross-material covariances, subsections involving the "other" materials will be ignored if IREAD =0 or 1. These subsections can be selectively processed by specifying IREAD=2. Input for a sample problem that illustrates the use of IREAD=2 is given in the next section.

MFCOV... This parameter is used to specify whether covariances of fission v (MFCOV=31) or cross sections (MFCOV=33) are needed. If MFCOV=31, it is necessary to supply multigrouped v values on a GENDF tape (that is, NPEND must be 0); the sample problem below shows the calculational sequence required. MFCOV=32 is reserved for a planned future capability to compute uncertainties in self-shielded cross sections. If MFCOV=33, the contribution of the uncertainty in individual resonance parameters (file 32) is included in the calculated uncertainty in infinite-dilution group cross sections.

Given below is the complete NJOY input for a calculation of a multigroup $\bar{\nu}$ covariance library (^{238}U part only).

0 5 *MODER*/ MOUNT ENDF TAPES 515, 516, and 555 ON UNITS 20, 21, AND 22. 1 - 23*ENDF/B-V NUBAR COVARIANCE MATERIALS*/ 20 1380 20 1381 21 1390 22 1395 22 1398 20 1399 0/ *MODER*/ COPY ENDF FOR USE AS A PENDF. -23 -24 *GROUPR*/ PREPARE GENDF WITH MULTIGROUPED NUBARS. -23 -24 0 25

1380 3 0 3 0 1 1 0 *BIG3 + 2 NUBAR*/ 0. 1.E10 3 452 *TOTAL NUBAR*/ 0/ 1381 3 452 *TOTAL NUBAR*/ 0/ 1390 3 452 *TOTAL NUBAR*/ 0/ 1395 3 452 *TOTAL NUBAR*/ 0/ 1398 3 452 *TOTAL NUBAR*/ 3 455 *DELAYED NUBAR*/ 3 456 *PROMPT NUBAR*/ 0/ 1399 3 452 *TOTAL NUBAR*/ 0/ 0/ *ERROR*/ PREPARE MULTIGROUP NUBAR COVARIANCE LIBRARY. -23 0 25 26/ 1398 19 1 1 2 31 1380 452 1381 452 1390 452 1395 452 1399 452 0/ 1 1.E7 1.7E7 *STOP*

Note the use of the MODER module to prepare a special ENDF tape containing selected evaluations from other ENDF tapes. Because only high-energy \bar{v} data is requested (10-17 MeV), it is unnecessary to use RECONR to prepare a PENDF tape for GROUPR. A copy of the ENDF tape is used instead.

I. DTFR Module

IPTOTL, IPINGP, ITABL, NED... The standard transport table as used in many codes has the following structure for each group.

<u>Position</u>	Contents
1	response function edit
•	cross sections (missing
	if IPTOTL=3)



If IEDIT=1, the first NED positions are printed out as separate tables. If there is not enough room for all the downscatter between IPINGP and ITABL, the array is truncated in such a way as to preserve the scattering cross section; a similar procedure is used for upscatter.

NTHERM,MTI,MTC,NLC... These parameters control the addition of thermal upscatter to the transport table. In the lowest NTHERM groups, the static elastic scattering is replaced by MTI and the total is readjusted appropriately. If MTC is requested, it is added to the in-group of the appropriate tables and the total is increased by MTC.

JPOS,MT,MULT... This system allows the user to specify a response function that is any linear combination of the vector cross sections processed by GROUPR. Before reaction MT is added into JPOS, it is multiplied by MULT. For example, total helium production in ENDF/B-IV ¹²C is $(n,\alpha) + 3 \times (n,n')3\alpha$ and is formed by the cards

with NED=2 and IPTOTL=1+3=4.

Several special values are recognized by DTFR.

<u>MT</u>	Meaning
300	P ₀ flux
470	fission χ (steady-state)
471	delayed χ
455	delayed v

J. CCCCR

NGGRUP,MAXUP... In the current version of the code, these parameters should both be zero. There is no coding for photon files and no provision for upscatter.

ICHIX... Either a vector χ or a matrix χ can be produced by isotope; for vectors, the GROUPR fission matrix can be collapsed with the library flux or an input flux. There is no provision for a set χ vector or χ matrix.

IFOPT,NSBLK... Several options are available for blocking and subblocking matrices. For large group structures, IFOPT=1 and NSBLK=NGROUP are recommended. K. MATXSR

NTYPE,NPART... A MATXS file can include any or all of four "types" of data; neutron cross sections and transfer matrices ("NSCAT"), gamma production matrices ("NGAMA"), photon interaction cross sections and transfer matrices ("GSCAT"), and thermal neutron scattering ("NTHERM"). Depending on which data types are desired, group structures will be required for neutron ("NEUT") or photons ("GAMA") or both.

HTYPE, HPART, HMAT... Data types, particles, materials (and reactions) are identified by Hollerith names rather than integer flags as in the CCCC formats. Data type and particle names are left-justified.

The MATXSR user need not be concerned with conventions for naming reactions, as this is handled automatically.

In a typical MATXS run, the first several cards of input might be these.

-21 -22 23 1 *T2LASL NJOY*/ 1 3 4 2 * */ *TYPICAL MATXS RUN WITH THREE DATA TYPES*/ *BASIC DATA FROM ENDF/B-IV*/ * */ *NEUT * *GAMA */ 30 12 *NSCAT* *NGAMA* *GSCAT */

In this example, three data types are requested for a MATXS file on unit 23. Neutron interaction and gamma production data (in blocked binary) are input from unit 21 and photon interaction data (in blocked binary) are to be found on unit 22. The code expects to find input data averaged over 30 neutron groups and 12 photon groups. Four cards of Hollerith identification are given.

Card input for the first data type ("NSCAT") is given next.

1 1	L	4	3	
1				
1				
*H]	L *	1	1	1269
*FENAT	*	5	2	1192
*PU24() *	1	1	1265

The code will search through unit 21 for MAT=1269, and output data for the first temperature and dilution factor under the name *H 1 *. The output for *FENAT * will include the first five temperatures and the first two dilution factors for MAT=1192. Particle one ("NEUT") is specified as both the incident (IINP) and outgoing (IOUTP) particles.

Input for the final two data types might be as follows.

1	1	4	3	
1				
2				
*H	1 *	1	1	1269
*FEN	AT *	5	2	1192
*PU24	40 *	1	1	1265
1	1	4	3	
2				
2				
* H	*	1	1	1
*FE	*	1	1	26
*PU	*	1	1	94

Note that for photon interaction data (GSCAT), all evaluations are elemental.

IFOPT,NSBLK... These parameters control the size of the matrix records on the MATXS file. If IFOPT=1 and NSBLK=1, then a single record contains all Legendre orders of the entire scattering matrix for a given reaction. The choice of IFOPT=2 separates each Legendre order into a separate record. The NSBLK parameter may be set equal to the number of groups (NOUTG) for the outgoing particle, and each group will be a separate record.

IV. SAMPLE PROBLEMS

The sample problems are designed to demonstrate the major options of NJOY without using too much computer time. For the convenience of the reader, parts of the sample problem output are reproduced in Appendix C. Complete output is available on special request for use in checking an installation of NJOY.

A. Example 1

This run tests group averaging and pointwise file generation for a light isotope (ENDF/B-V carbon). Linearization, Doppler broadening, thinning, heat

production, and thermal scattering are included. ENDF/B tapes 511 and 322 are required.

In RECONR, the addition of 127 points by linearization to the 776 points in the initial union grid gives 903 points, which can be seen in BROADR and in MF3/MT1 on the PENDF tape. In BROADR, only MT2 (elastic) and MT102 (radiative capture) are broadened. Thinning does not remove many points because these cross sections are relatively smooth.

The HEATR run adds both total heat production (MT301) and total damage energy production (MT444) to the PENDF tape, as can be seen by the list of reactions in MF1/MT451. There are two THERMR runs: the first one computes cross sections and matrices for free carbon atoms and stores the results under MT221, and the second run computes both coherent elastic and incoherent inelastic scattering for graphite using MT230 and MT229, respectively. The effective normal temperature shown is the default value. Also note that THERMR uses the 296 K data from tape 322 to satisfy the request for 300 K data.

In the GROUPR run, note that the "LR-flags" on MT52-91 were picked up; these sections provide a "pseudo level" representation of the $(n,n')3\alpha$ reaction. The LR-flag was also processed in HEATR. Note that heating and damage (MT301, MT444) are processed in the same way as any other ENDF/B reaction. The thermal data shown here (MT221,229,230) are supplementary; they are not included in MT=1. Subsequent codes such as DTFR must replace MT2 with MT221 or MT229,230 in the thermal range (groups 1-4) and revise the total accordingly. <u>Caution</u>: GROUPR numbers groups in order of increasing energy.

On the PENDF tape, note that the dictionary and comments are updated. The format used for MF=6 is specially designed for NJOY thermal data. The file uses a "TAB2" record to set up a loop over incident energy (52 points). At each energy (1.E-5, 3.16E-5, etc.), a "LIST" record is given that contains N1 (that is, 640) words organized on a cycle of N2 (that is, 10) words. Each cycle contains a secondary energy, a scattering probability, and 8 equally probable scattering cosines.

B. Example 2

This run processes one isotope for a practical CCCC library. It tests resonance reconstruction, Doppler broadening to several temperatures, unresolved cross sections, self-shielded multigroup cross sections, and CCCC-IV interface files (ISOTXS, BRKOXS, and DLAYXS). Tape 404 is required.

Because this is a resonance material, RECONR includes a table of estimated errors in the resonance integrals due to significant figure reduction and the resonance-integral thinning criteria. In this material, these errors are negligible, but note that 547 energy intervals were affected by the integral check, and 73 energy points were only one significant figure apart for NDIGIT=6. Additional points were removed from the final grid by "back-thinning" to the specified reconstruction tolerance. The BROADR run in this example generates several output temperatures. Each one is written as a separate material on the PENDF tape. Note that the zero-Kelvin cross sections do not appear on tape 23; also note the smoothing effect of Doppler broadening on "POINTS OUT." On the UNRESR listing, effective self-shielded cross sections are given for each energy with σ_0 horizontally and reaction vertically (P_0 total, elastic, fission, capture, P_1 total). The σ_0 list in UNRESR does not have to agree with the σ_0 list in GROUPR.

In GROUPR, it was only necessary to request those reactions with appreciable temperature dependence at the higher temperatures. These normally include MT=1, 2, 18, 19, 102, 301, and 201-250. Also, MT259 is requested to get accurate group-averaged velocities for ISOTXS. In CCCCR, the option to block matrices by group was used. This is the best form for large group structures. The isotope fission chi was produced using the GROUPR flux to collapse the fission matrix. Note also that the BRKOXS file includes shielding factors for elastic removal. XSPO in BRKOXS is simply $4\pi a^2$ using the scattering length from MF2.

C. Example 3

The run demonstrates the generation of photon interaction cross sections * with DTF and MATXS output. The photon interaction tape DLC7E is required.

First, a RECONR run is used to linearize the cross sections. This example illustrates a multiple isotope run. The GAMINR run uses the standard list of reactions. Note how the photon heat production is assembled from the partial reactions and written out as MT=621. In DTFR, the code automatically shifts over to photons when it sees MT501, so read "PHOTON" instead of "NEUTRON" on this listing. Note that one special edit is provided for heating in eV·barns, so the table length is set to 1+3+12=16, and the position of the total is 1+3=4. In these tables, the first 16 numbers are the positions for the first (highest energy) photon group, and so on for the other groups. That is, the group numbering has been restored to the conventional order.

^{*}Available from the Radiation Shielding Information Center at the Oak Ridge National Laboratory, Oak Ridge, Tennessee.

The MATXS listing is fairly self-explanatory. Once again, the groups are numbered in the conventional order. In an isotropic matrix, such as "GPAIR" (gamma pair production), groups are given in the DTF ordering; that is; $10 \leftarrow 9$, $10 \leftarrow 8$, $10 \leftarrow 7$, etc. Also note the "INDEX" at the end of the listing. It provides a concise reference to all the materials and reactions on the library.

D. Example 4

This run demonstrates NJOY's capability to produce cross-section covariances from ENDF/B data. Tape 511 is used. The material processed is the ENDF/B-V "standards" material ²³⁵U.

After converting the mode of the ENDF tape, the RECONR module is used to prepare a PENDF file on unit 22. Note that very coarse tolerances (ERR, ERRMAX, ERRINT) are used. Because only relative covariances are given in this evaluation (that is, there are no sub-subsections with LB=0), the output multigroup relative covariances from the ERRORR module are nearly independent of the cross-section magnitudes; hence, high accuracy is not needed in the resonance reconstruction calculation.

The first ERRORR run operates directly on the PENDF file and produces cross-section covariances (MFCOV=33). The multigroup energy structure includes 1 and 1000 eV, plus any energies in the ENDF file 33 that fall between these limits. This group-structure option (IGN=19) is very useful for test purposes, such as preparing covariance plots of ENDF data using the ERRORR and COVR modules. Because of the use of IGN=19, and because the evaluator has used LB=5, which in a sense is already processed, many of the output numbers here are identical with ENDF entries. (For example, compare the first line of the printed relative covariances with line 10 057 of MAT1395.)

Next, the GROUPR module is used to prepare a GENDF file containing 30-group (IGN=3) fission $\bar{\nu}$ values. This GENDF (on unit 24) is then used as input to a second ERRORR run, in which the ENDF file 31 ($\bar{\nu}$ covariances) is processed (MFCOV=31). In this case, a user-specified 7-group structure is employed. Note that the resulting multigroup $\bar{\nu}$ covariances are added to the cross-section covariances written on unit 23 in the previous ERRORR run, to make a new ERRORR output file on unit 25.

V. PROGRAM DETAILS

The programming details, calculational methods, and theories used by the modules of NJOY are discussed in detail in subsequent volumes of this report. The following sections provide a general overview.

A. Programming Language

NJOY uses ANSI-standard FORTRAN-IV with a few exceptions (for example, CDC overlay commands, mixed-mode arithmetic, expressions as array indices). Machinedependent functions such as timers, input/output, and character manipulation are isolated in special subroutines. The resulting code, with minor modifications, has been compiled using CDC, CRAY, IBM, and UNIVAC compilers. Some loops have been rearranged to facilitate vectorization.

B. Storage Allocation

Variable dimensioning and dynamic storage allocation are used extensively to provide efficient use of the main memory; an easy-to-use system of subroutines called STORAG reserves, releases, retrieves pointers, and repacks stored data when necessary. With this system, most variables are stored in a large container array for each module. In the reference version of NJOY, the container arrays have been chosen for the efficient use of about 150 000₈ words. For many problems, the size of these arrays can be decreased. Increasing the number of words available, if possible, will make BROADR run faster. Dynamic storage allocation leads to a complex set of trade-offs, such as number of groups versus Legendre order in GROUPR. In addition, a few fixed limits remain, such as 620 groups, 10 temperatures, and 10 σ_0 -values in GROUPR. Diagnostic messages are provided when any limits are exceeded. The current allocations have been adequate for all practical problems attempted to date.

C. Running Time

Typical run times are difficult to quote for such a complex system. *
Tables II and III give CRAY-1 times for practical problems.
*
These are centralprocessor times plus an approximately 30% surcharge for memory and I/O.

A few general principles may help the user to guess times for his jobs. The pointwise modules RECONR, BROADR, and HEATR require time proportional to the number of energy points in the cross sections, hence the number of resonances. UNRESR depends mostly on the number of energy points in MF2, MT151. In GROUPR, the time required for vectors depends on the number of energy points, but not on the number of groups, and only weakly on the number of σ_0 values requested. The matrix time does increase with the number of groups. Even more important is the energy range--discrete inelastic reactions require much more time if many groups above 10 MeV are requested, due to the extreme anisotropy found there.

*NJOY runs about 1.6 times faster on the CRAY-1 than on the CDC-7600.

TABLE II

POINTWISE PROCESSING TIMES FOR SEVERAL MATERIALS BY MODULE (approximate CP + I/O in seconds for CRAY-1)

<u>Module</u>	<u>H-1</u>	C-NAT	Fe-NAT	<u>U-235</u>	<u>U-238</u>
RECONR	5	6	160	55	572
BROADR	7	13	190	147	1500
UNRESR		6	<u> </u>	400	210
HEATR	50 J	65	540 ^C	290	633
THERMR	400 ⁰	450 ^e	170	120	208

a. 0.2% tolerance

b. 4 to 7 temperatures

c. With damage

d. In water

e. In graphite

TABLE III

ADDITIONAL TYPICAL PROCESSING TIMES FOR NJOY (approximate CP + I/O in seconds for CRAY-1)

Processing Task	<u>Time</u>
C-NAT GROUPR (80 neutron groups, 24 photon groups, T = 300 K, σ_0 = 1E10)	220
C-NAT GROUPR (30 neutron groups, 12 photon groups, 300 K, infinite dilution)	145
Fe-NAT GROUPR (80 x 24, 6 temper- atures, 6 dilutions)	1960
U-238 GROUPR (80 x 24, 7 temper- atures, 7 dilutions)	2100
Complete 24-group photon inter- action GAMINR run for all elements	2710

This code is extremely "I/O bound." Run times can often be reduced by requesting larger input/output buffers from the system. The blocked-binary mode should be used whenever possible. Also, I/O time in BROADR can be reduced by allocating more storage if available (see /STORE/ and NAMAX in BROADR).

VI. CODE CONVERSION

The reference version of NJOY operates on CDC-7600 equipment at Los Alamos. The code includes comment cards that allow many of the changes required to convert to an IBM system to be made very simply. A typical comment-card example follows.

CCDC INTEGER H(5) CCDC CIBM С REAL*8 H(5)CIBM .

In the (10/81) version there are 146 such blocks of CDC machine-dependent coding and 177 blocks of IBM coding. To convert to IBM form, insert a "C" in column 1 of all card images bracketed with "CCDC" cards, and remove the "C" from column 1 of all cards bracketed by "CIBM" cards. This can be done with a simple preprocessing program such as the one listed in Appendix D.

Additional changes may be required in BANNER (time, date, machine), ERROR (system fatal exit, traceback), character handling for free-form input (FREE, PACK), input/output routines (OPENZ, CLOSEZ, REPOZ, SKIPRZ, REED, RITE), and mathematical subroutines (E1,GAMI). The largest group of system-dependent routines has to do with plotting. COVR uses the DISSPLA software system, which is available at many installations. DTFR uses a local SC-4020 library, which is not widely available; however, the plotting calls have been left in the code as a guide for the conversion to other systems.

As a demonstration of portability, versions of NJOY have been installed on the following machines (operating systems): CDC-7600 (LTSS) and CRAY-1 (CTSS) at Los Alamos National Laboratory, CDC-7600 (LTSS) at the Magnetic Fusion Energy Computing Center at Lawrence Livermore National Laboratory, CDC-7600 (SCOPE) at the Brookhaven National Laboratory, IBM-360/195 (MTD/FTX) at Argonne National Laboratory, UNIVAC-1108 at the Hanford Engineering Development Laboratory of Westinghouse, and the IBM-360/195 at the Oak Ridge National Laboratory.

VII. ACKNOWLEDGEMENTS

The IBM conversion was performed by R. Q. Wright and J. E. Lucius at Oak Ridge National Laboratory and D. Cambra and C. Stenberg at Argonne National Laboratory. The UNIVAC conversion was performed by F. W. Mann at the Hanford Engineering and Development Laboratory. Thanks are also due to Odelli Ozer of the Electric Power Research Institute for help with the RESEND code and the design of the thermal processing capability; to D. E. Cullen of the Lawrence Livermore National Laboratory and J. Hancock of Los Alamos for their work on the Doppler-broadening module; to P. G. Young for help with the heating calculation; to R. E. Seamon, P. D. Soran, W. B. Wilson, D. George, R. B. Kidman, and L. Stewart (all of Los Alamos) for help with testing and validation; to R. J. Barrett of Los Alamos for his extensive work on the CCCC and MATXS systems; and to R. J. LaBauve of Los Alamos, D. R. Harris and M. Becker of Rensselaer Polytechnic Institute, and C. R. Weisbin of Oak Ridge National Laboratory, for moral support, ideas, and constructive criticism.

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APPENDIX A

INPUT INSTRUCTIONS

Los Alamos Identification No. LP-795 NJDY NUCLEAR CRDSS SECTION PROCESSING SYSTEM * VERSION 10/81 ****** NJDY IS A SYSTEM OF PROCESSING MODULES INTENDED TO CONVERT ENDE/B VERS. IV DR V CROSS SECTION DATA INTO FORMS USEFUL FOR PRACTICAL APPLICATIONS. RECONR...RECONSTRUCT POINTWISE CROSS SECTIONS FROM ENDF/B RESONANCE PARAMETERS AND INTERPOLATION SCHEMES. BROADR...ODPPLER BROADEN AND THIN POINTWISE CROSS SECTIONS. UNRESR...COMPUTE EFFECTIVE POINTWISE SELF-SHIELDED CROSS SECTIONS IN THE UNRESOLVED ENERGY RANGE. HEATR...COMPUTE HEAT PRODUCTION CROSS SECTIONS (KERMA) AND DAMAGE ENERGY PRODUCTION. THERMR...GENERATE NEUTRON SCATTERING CROSS SECTIONS AND POINT-TO-POINT SCATTERING KERNELS IN THE THERMAL RANGE FOR FREE OR BOUND ATOMS. GROUPR...GENERATE SELF-SHIELDED MULTIGROUP CROSS SECTIONS AND GROUP-TO-GROUP SCATTERING AND PHOTON PRODUCTION MATRICES. GAMINR...COMPUTE MULTIGROUP PHOTON INTERACTION CROSS SECTIONS, SCATTERING MATRICES, AND HEAT PRODUCTION. ERRORR...CONSTRUCT MULTIGROUP COVARIANCE MATRICES. COVR...PROCESS COVARIANCE DATA FROM ERRORR MODER...CONVERT BETWEEN ENDF/B STANDARD CODED MODE AND THE NJDY BLOCKED BINARY MODE. DTFR...OUTPUT AND PLOT MULTIGROUP DATA FOR DISCRETE ORDINATES TRANSPORT CODES. CCCCR...FORMAT MULTIGROUP DATA INTO THE CCCC STANDARD INTERFACE FILES ISOTXS, BRKOXS, AND DLAYXS. MATXSR...CONVERT MULTIGROUP DATA INTO THE COMPREHENSIVE MATXS CRDSS SECTION INTERFACE FORMAT. ACER... PREPARE LIBRARY FOR THE LOS ALAMOS CONTINUOUS ENERGY MONTE-CARLD CODE MCNP. POWR...CONVERT MULTIGROUP DATA INTO LIBRARIES FOR THE THERMAL POWR REACTOR CODES EPRI-CELL AND EPRI-CPM. EACH MODULE IS A SEPARATE OVERLAY. THE MAIN PROGRAM CONTROLS THE ORDER IN WHICH MODULES ARE USED AND CONTAINS UTILITY SUBROUTINES USED BY ALL MODULES. *---INPUT SPECIFICATIONS (FREE FORMAT)------INPUT OPTION CARD 1 IOPT O FOR CARD INPUT AND FULL DUTPUT. 1 FOR TERMINAL INPUT WITH SHORT OUTPUT ON TERMINAL * * CARD2

IVERF ENDF/B VERSION NUMBER (4 OR 5 ONLY) CARD 3 SIX CHARACTER MODULE NAME DELIMITED WITH *, MOPT E.G., *RECONR* (DNLY FIRST FOUR CHARACTERS ARE USED). REPEAT FOR EACH MODULE DESIRED). USE *STOP* TD TERMINATE PROGRAM. SEE THE COMMENTS AT THE START OF EACH MODULE FOR * ITS SPECIFIC INPUT INSTRUCTIONS. *---CODE CONVERSION------* CONVERSION FROM COC TO IBM OR BACK IS CONTROLLED BY THE COMMENT* * CONVERSION FROM COEMD TO CONVERT FROM COC TO IBM, PUT A C IN * COLUMN 1 OF EVERY LINE BETWEEN TWO CCDC CARDS AND REMOVE THE C * FROM COLUMN 1 OF EVERY LINE BETWEEN TWO CIBM CARDS. A FEW * REMAINING CHANGES MAY BE NEEDED BECAUSE OF SYSTEM DIFFERENCES * BETWEEN INSTALLATIONS PROGRAM RECONR ٠ ***** RECONSTRUCT PDINTWISE CROSS SECTIONS * RESONANCE CRDSS SECTIONS ARE CALCULATED USING THE METHODS OF * RESEND, WITH MODIFICATIONS TO THE METHOD OF GENERATING THE * ENERGY GRID AND THE METHOD OF COMBINING RESONANCE AND * BACKGROUND CROSS SECTIONS. * THIS PROGRAM GENERATES AN ENERGY GRID WHICH IS THE UNION OF * AN INPUT GRID (IF ANY), THE RESONANCE ENERGIES (IF ANY), AND * THE ENERGIES DF CROSS SECTIONS IN MF3 AND MF13 (DR MF23). * THE POINTWISE CROSS SECTIONS ARE THEN COMPUTED ON THIS GRID * AND POINTS ARE ADDED SO THAT THE RESONANCE CROSS SECTIONS AND * ANY CROSS SECTIONS REPRESENTED BY NON-LINEAR INTERPOLATION * ARE REPRODUCED WITHIN A SPECIFIED TOLERANCE BY LINEAR INTER-POLATION. PSI-CHI RECONSTRUCTION CAN BE USED IF DESIRED. SECTIONS WHICH ARE NOT CROSS SECTIONS (MU,NU) AND PHOTON * MULTIPLICITIES (MF12) ARE NOT PROCESSED. REDUNDANT REACTIONS ARE RECONSTRUCTED TO BE THE SUM OF THEIR PARTS. THE PENDF TAPE CONTAINS PDINT CROSS SECTIONS IN MF3 AND MF13 (DR MF23) * * IS CONSTRUCTED WITH THE POTENTIAL SCATTERING LENGTH ADDED. *---INPUT SPECIFICATIONS (FREE FORMAT)------* CARD 1 NENOF UNIT FOR ENOF/B TAPE UNIT FOR PENDE TAPE NPENO CARD 2 66 CHARACTER LABEL FOR NEW PENOF TAPE * LABEL DELIMITED WITH *. ENDED WITH /. CARD 3 MATERIAL TO BE RECONSTRUCTED NUMBER OF CARDS OF DESCRIPTIVE DATA FOR NEW MF1 ΜΔΤ NCARDS (DEFAULT=0.) NGRID NUMBER OF USER ENERGY GRID POINTS TO BE ADDED. ٠ (DEFAULT=0.)

```
CARD 4
٠
     ERR
               FRACTIONAL RECONSTRUCTION TOLERANCE USED WHEN
               RESONANCE-INTEGRAL ERROR CRITERION (SEE ERRINT)
               IS NOT SATISFIED.
               RECONSTRUCTION TEMPERATURE (DEG KELVIN)
     TEMPD
                (DEFAULT=0.)
     NDIGIT
               ND. SIGNIFICANT DIGITS (DEFAULT=6)
     ERRMAX
               FRACTIONAL RECONSTRUCTION TOLERANCE USED WHEN
               RESONANCE-INTEGRAL ERROR CRITERION IS SATISFIED
(ERRMAX.GE.ERR, DEFAULT*20.*ERR)
MAXIMUM RESONANCE-INTEGRAL ERROR (IN BARNS)
     ERRINT
               PER GRID POINT (DEFAULT=ERR/1000D)
 CARD 5
     CARDS
               NCARDS OF DESCRIPTIVE COMMENTS FOR MT451
               EACH CARD DELIMITED WITH *, ENDED WITH /.
  CARD 6
               USERS ENERGY GRID POINTS
     ENODE
        CARDS 3, 4, 5, 6 MUST BE INPUT FOR EACH MATERIAL DESIRED
         MAT=O/ TERMINATES EXECUTION DF RECONR.
PROGRAM BROADR
* ODPPLER BROADEN AND THIN NEUTRON POINT CROSS SECTIONS
 A MODIFIED VERSION OF THE KERNEL BROADENING METHOD DEVELOPED
 FOR SIGMAI (D.E.CULLEN, LLNL) IS USED. CROSS SECTIONS
FOR LOW THRESHOLD REACTIONS ARE UNIONIZED ON THE GRID OF THE
 TOTAL CROSS SECTION, THEN BROADENED AND THINNED IN PARALLEL.
* HIGH THRESHOLD REACTIONS ARE NOT BRDADENED. THE RESULTS ARE
  WRITTEN OUT IN PENDE FORMAT WITH EACH TEMPERATURE REPRESENTED
* AS A DIFFERENT MAT. DICTIONARIES ARE CORRECTED TO REFLECT
* UNIONIZATION AND THINNING. FOR HIGH TEMPERATURES AND LOW
* ENERGIES WHERE THE METHOD OF SIGMAI BREAKS DOWN, A NEW DIRECT
  EXPANSION OF THE OOPPLER INTEGRAL IS USED.
*---INPUT SPECIFICATIONS (FREE FORMAT)-----
  CARD 1
*
                INPUT PENOF TAPE
     NIN
                OUTPUT PENDE TAPE
     NOUT
  CARD 2
                MATERIAL TO BE PROCESSED
     MAT 1
     NTEMP2
                NUMBER OF FINAL TEMPERATURES (MAXIMUM=6)
      ISTART
                RESTART (O ND, 1 YES)
                BOOTSTRAP (O ND, 1 YES)
     ISTRAP
                STARTING TEMPERATURE FROM NIN
     TEMP 1
     FRRTHN
                FRACTIONAL TOLERANCE FOR THINNING
                MAX. ENERGY FOR BRDADENING AND THINNING
      THNMAX
                (DEFAULT=1 MEV)
  CARD 3
     TEMP2
                FINAL TEMPERATURES (DEG KELVIN)
  CARD 4
     MAT 1
                NEXT MAT NUMBER TO BE PROCESSED WITH THESE
                PARAMETERS. TERMINATE WITH MAT1=0.
*---INPUT OPTIONS------
* THE OUTPUT TAPE WILL CONTAIN THE NTEMP2 FINAL TEMPERATURES
  SPECIFIED. IT IS NECESSARY TO HAVE TEMP1.LE.TEMP2(1).
IF TEMP2.EQ.TEMP1, THE OATA WILL BE THINNED ONLY.
```

```
CONTINUE BROADENING AN EXISTING PENDE TAPE.
 RESTART
                                                                  ALL
               TEMPERATURES ARE COPIED THROUGH TEMP1. ADDITIONAL
              FINAL TEMPERATURES ARE ADDED BY STARTING WITH THE
               DATA AT TEMP1.
               IF BOOTSTRAP IS NOT REQUESTED, EACH FINAL TEMPERA-
 BOOTSTRAP
               TURE IS GENERATED BY BROADENING DIRECTLY FROM
               TEMP1 TO TEMP2. IF BOOTSTRAP IS REQUESTED, EACH
FINAL TEMPERATURE IS BROADENED FROM THE PRECEDING
               TEMPERATURE, THIS OPTION IS FASTER DUE TO THE
               THINNING IN THE PREVIOUS STEP. HOWEVER, ERRORS
               ACCUMULATE.
                         PROGRAM UNRESR
                          **************************
  COMPUTE UNRESOLVED RESONANCE CROSS-SECTIONS
*
* THE METHOD OF ETOX IS USED TO COMPUTE SELF-SHIELDED
  UNRESOLVED RESONANCE CROSS-SECTIONS ON THE ENERGY GRID OF
  THE UNRESOLVED PARAMETERS. SUBSEQUENT INTERPOLATION IS
TO BE ON THE CROSS-SECTIONS AND NOT ON THE PARAMETERS.
*
  ADDITIONAL ENERGY GRID POINTS ARE ADDED AT QUARTER LETHARGY
*
  INTERVALS IF ONLY THREE OR FEWER GRID POINTS ARE FOUND.
  THE ACCURATE HWANG QUADRATURE SET IS USED FOR THE INTEGRALS.
*---INPUT SPECIFICATIONS (FREE FORMAT)-----
  CARD 1
    NENDF
              UNIT FOR ENOF/B TAPE
              UNIT FOR INPUT PENOF TAPE
UNIT FOR OUTPUT PENOF TAPE
    NIN
    NOUT
  CARD 2
              MATERIAL TO BE PROCESSED
     MATO
              NO. OF TEMPERATURES (MAX. ND. ALLOWED=9)
ND. OF SIGMA ZEROES (MAX. NO. ALLOWED=8)
PRINT OPTION (O=MIN. 1=MAX) (DEFAULT=0)
     NTEMP
     NSIGZ
     IPRINT
  CARD 3
     TEMP
              TEMPERATURES IN KELVIN (INCLUDING ZERD)
  CARD 4
         .
SIGMA ZERO VALUES (INCLUDING INFINITY)
CARDS 2, 3, 4 MUST BE INPUT FOR EACH MATERIAL DESIRED
MATD=O/ TERMINATES EXECUTION OF UNRESR.
     SIGZ
PROGRAM HEATR
* COMPUTE HEATING KERMA (KINETIC ENERGY RELEASE IN MATERIAL)
   AND RADIATION DAMAGE ENERGY PRODUCTION
  THE PROMPT KERMA IS COMPUTED POINTWISE ON THE GRID OF THE
  TOTAL CROSS SECTION FROM THE INPUT PENDE TAPE AND WRITTEN
 * ONTO THE DUTPUT PENDE TAPE AT INFINITE DILUTION USING THE
  300 SERIES OF MT NUMBERS. ALL TEMPERATURES ON THE INPUT PENDF
 * TAPE FOR THE DESIRED MATERIAL ARE PROCESSED. THE DICTIONARY
                 REACTION Q VALUES ARE OBTAINED FROM THE ENDF/B
 * IS REVISED.
 * TAPE UNLESS THE USER ENTERS HIS DWN VALUE. PARTIAL KERMAS
```

* CAN BE REQUESTED FOR SELF-SHIELDING CALCULATIONS OR OTHER PURPOSES. THE CODE USES THE ENERGY BALANCE METHOD WHERE PHOTON FILES ARE AVAILABLE AND DEPOSITS ALL PHOTON ENERGY LOCALLY WHEN FILES ARE NOT AVAILABLE. THIS ASSURES CONSISTENCY BETWEEN NEUTRON HEATING AND ENERGY DEPOSITION BY SUBSEQUENT PHOTON INTERACTIONS. AN EXCEPTION IS MADE FOR CAPTURE WHERE RECOIL IS COMPUTED BY MOMENTUM CONSERVATION. PHOTON FILES ARE USED TO ESTIMATE THE AVERAGE PHOTON MOMENTUM WHEN AVAILABLE. A DIAGNOSTIC MESSAGE IS PRINTED IF THE ٠ MOMENTUM CALCULATION LEADS TO A SIGNIFICANT ERROR IN ENERGY CONSERVATION. IF DESIRED. THE ENERGY-BALANCE KERMA FACTORS CAN BE COMPARED * WITH CONSERVATIVE KINEMATIC LIMITS (SET IPRINT=2). DAMAGE ENERGY IS COMPUTED USING THE LINDHARD ELECTRONIC SCREENING DAMAGE FUNCTION WITH A 25 EV DISPLACEMENT THRESHOLD. *---INPUT SPECIFICATIONS (FREE FORMAT)------CARD 1 NENDF UNIT FOR ENOF/B TAPE UNIT FOR INPUT PENDE TAPE NIN UNIT FOR OUTPUT PENDE TAPE NOUT CARD 2 MATO MATERIAL TO BE PROCESSED NPK NUMBER OF PARTIAL KERMAS DESIRED (DEFAULT=0) NUMBER OF USER Q VALUES (DEFAULT=0) NOA NUMBER OF TEMPERATURES TO PROCESS NTEMP (DEFAULT=O. MEANING ALL ON PENDF) LOCAL D/1=GAMMA RAYS TRANSPORTED/DEPOSITED LOCALLY (DEFAULT=0) FRINT (O MIN, 1 MAX, 2 CHECK) (DEFAULT=O) FOR NPK GT O DNLY MT NUMBERS FOR PARTIAL KERMAS DESIRED IPRINT CARD 3 MTK TOTAL (MT301) WILL BE PROVIDED AUTOMATICALLY. PARTIAL KERMA FOR REACTION MT IS MT+300 AND MAY NOT BE PROPERLY DEFINED UNLESS A GAMMA FILE FOR MT IS ON ENDF TAPE. SPECIAL VALUES ALLOWED--303 NON-ELASTIC (ALL BUT MT2) INELASTIC (MT51 THRU 91) 304 FISSION (MT18 OR MT19, 20, 21, 38) DISAPPEARANCE (MT102 THRU 120) 318 401 DAMAGE ENERGY PRODUCTION VALUES --TOTAL 444 ELASTIC (MT2) INELASTIC (MT51 THRU 91) 445 446 447 DISAPPEARANCE (MT102 THRU 120) CARDS 4 AND 5 FOR NQA GT 0 DNLY CARD 4 MTA MT NUMBERS FOR USERS Q VALUES CARD 5 OA USER SPECIFIED Q VALUES (EV) PROGRAM THERMR ** ******************************** GENERATE NEUTRON SCATTERING CROSS SECTIONS AND POINT-TO-POINT *

* SCATTERING KERNELS IN THE THERMAL RANGE

ADD POINTWISE SCATTERING CROSS SECTIONS AND SCATTERING MATRICES TO AN EXISTING PENDE TAPE. CROSS SECTIONS ARE ADDED TO MF3 AND MATRICES ARE WRITTEN IN MF6 (USING A MODIFIED FORMAT). BOTH USING MTREF FOR INELASTIC AND MTREF+1 FOR ELASTIC (IF ANY). MULTIPLE SCATTERING TYPES (IE, H FREE AND H IN H2O) CAN BE WRITTEN ON DNE PENDF TAPE BY USING DIFFERENT VALUES OF MTREF FOR EACH THERMR RUN. IF DATA FOR DNE MTREF IS ALREADY ON THE TAPE, IT WILL BE REPLACED WITH THE NEW CROSS SECTIONS. THE ENERGY GRID FOR COHERENT SCATTERING IS DETERMINED ADAPTIVELY SO AS TO REPRESENT THE SHARP BRAGG EDGES TO A SPECIFIED TOLERANCE USING LINEAR INTERPOLATION. THE SECONDARY ENERGY GRID FOR INELASTIC SCATTERING IS ALSO DETER-MINED ADAPTIVELY. ANGULAR DEPENDENCE IS REPRESENTED AS EQUALLY PROBABLE COSINES. THE INITIAL ENERGY GRID IS WIRED IN* (SEE EGRID IN CALCEM). A SPECIAL PROJECTION INTERPOLATION * SCHEME IS USED IN GROUPR TO INTEGRATE THIS RELATIVELY COARSE GRID. CURRENT CAPABILITIES... COMPUTE FREE-GAS SCATTERING MATRICES AND NORMALIZE 1.) TO THE ELASTIC CRDSS SECTION ON THE DLD PENDF TAPE. COMPUTE INCOHERENT MATRICES FROM READ-IN 2.) S(ALPHA, BETA) DATA. COMPUTE COHERENT SCATTERING FROM HEXAGONAL LATTICES. COMPUTE INCOHERENT ELASTIC SCATTERING. 3.) 4.) FUTURE CAPABILITIES. 1.) GENERATE S(ALPHA, BETA) FROM BASIC PHYSICS DATA. *---INPUT SPECIFICATIONS (FREE FORMAT)------CARD 1 NENDF ENDF/B TAPE FOR MF7 DATA OLO PENOF TAPE NEW PENOF TAPE NIN NOUT CARD 2 MATOE MATERIAL DESIRED ON ENDF TAPE MATERIAL DESIRED ON PENDF TAPE MATOP NUMBER OF EQUI-PROBABLE ANGLES NBIN NUMBER OF TEMPERATURES NTEMP INELASTIC OPTIONS IINC NONE 0 COMPUTE AS FREE GAS 1 RESERVED 2 COMPUTE S(A,B) AND MATRIX з READ S(A,B) AND COMPUTE MATRIX ELASTIC OPTIONS ICOH NONE 0 GRAPHITE BERYLLIUM 2 BERYLLIUM OXIDE 3 POLYETHYLENE 11 12 H(ZRH) ZR(ZRH) 13 NATOM NUMBER OF PRINCIPAL ATOMS MT FOR INELASTIC REACTION (201-250 DNLY) MTREF IPRINT PRINT OPTION (O=MINIMUM, 1=MAXIMUM, 2=MAX. NORMAL + INTERMEDIATE RESULTS) (DEFAULT=0) CARD 3 * TEMPR **TEMPERATURES (KELVIN)** CARD 4 FOR IINC=4 ONLY * EFTEMP EFFECTIVE TEMPERATURES FOR SHORT COLLISION TIME * (DEFAULT FOR EACH TEMPERATURE IS STANDARD VALUE *

FROM GENERAL ATOMIC REPORT IF AVAILABLE, OTHERWISE MATERIAL TEMPERATURE) CARD 5 * TOL TOLERANCE MAXIMUM ENERGY FOR THERMAL TREATMENT EMAX PROGRAM GROUPR COMPUTE SELF-SHIELDED GROUP-AVERAGED CRDSS SECTIONS * * PRODUCES SELF-SHIELDED CROSS SECTIONS, NEUTRON SCATTERING MATRICES, AND PHOTON PRODUCTION MATRICES. SCATTERING AND * PHOTON MATRICES MAY BE SELF-SHIELDED IF DESIRED (SEE INIT) * BONDARENKO WEIGHTING IS NORMALLY USED. OPTIONALLY. THE FLUX CAN BE COMPUTED FOR AN INFINITE MIXTURE OF HEAVY ABSORBER * AND LIGHT MODERATOR. DELAYED NEUTRON DATA AND THERMAL * SCATTERING MATRICES ARE HANDLED SPECIALLY. * THE INTEGRATION OVER INITIAL ENERGY IS HANDLED IN THE SAME WAY FOR ALL REACTION TYPES BY USING THE INTEGRAND * * FEED*XSEC*FLUX * FEED IS THE SDURCE INTO FINAL ENERGY GROUP GPRIME AND * LEGENDRE ORDER L FROM INITIAL ENERGY E (SEE GETFF). FOR VECTORS, THE FEED IS 1. OR A YIELD (NUBAR, MUBAR). FOR TWO BDDY SCATTERING, A CENTER-DF-MASS GAUSSIAN INTEGRATION IS USED TO OBTAIN ACCURATE RESULTS EVEN FOR SMALL LEGENDRE COMPONENTS * OF THE GROUP-TD-GROUP SCATTERING. ADDITIONAL INITIAL ENERGY QUADRATURE POINTS ARE ADDED TO INTEGRATE THE KNOWN POLYNOMIAL ORDER DF THIS FEED FUNCTION. FEED FOR TABULATED CONTINUUM REACTIONS IS COMPUTED EXACTLY ON THE ENDF/B GRID POINTS AND * A SPECIAL PROJECTION INTERPOLATION * THEN INTERPOLATED AT E. SCHEME IS USED FOR THERMAL MATRICES (SEE GETAED). THE FEED FOR ANALYTIC CONTINUUM REACTIONS IS EXACT. * *---INPUT SPECIFICATIONS (FREE FORMAT)-----* * CARD 1 NENDF UNIT FOR ENDE/B TAPE UNIT FOR PENOF TAPE UNIT FOR INPUT GOUT TAPE (DEFAULT=0) NPEND NGOUT 1 NGOUT 2 UNIT FOR OUTPUT GOUT TAPE (DEFAULT=O) CARD2 MATERIAL TO BE PROCESSED MATE MATE LT O IS A FLAG TO ADD MTS TO AND/DR REPLACE INDIVIDUAL MTS ON NGOUT1. I GN NEUTRON GROUP STRUCTURE OPTION IGG GAMMA GROUP STRUCTURE OPTION WEIGHT FUNCTION OPTION IWT LEGENDRE DRDER NUMBER OF TEMPERATURES I ORD NTEMP NUMBER OF SIGMA ZERDES NSIGZ LONG PRINT OPTION (0/1=MINIMUM/MAXIMUM) IPRINT (DEFAULT=1) ٠ CARD3 TITLE RUN LABEL (UP TO BO CHARACTERS DELIMITED BY *. ENDED WITH /) (DEFAULT=BLANK) CARD4 TEMP TEMPERATURES IN KELVIN CARD5 SIGMA ZERD VALUES (INCLUDING INFINITY) SIGZ
IF IGN=1, READ NEUTRON GROUP STRUCTURE (6A AND 6B) CARD6A NUMBER OF GROUPS NGN CARD6B NGN+1 GROUP BREAKS (EV) EGN IF IGG=1, READ GAMMA GROUP STRUCTURE (7A AND 7B) CARD7A NGG NUMBER OF GROUPS CAR07B NGG+1 GROUP BREAKS (EV) EGG WEIGHT FUNCTION OPTIONS (8A,8B,8C,8D) CARD8A FLUX CALCULATOR PARAMETERS (IWT.LT.O ONLY) BREAK BETWEEN COMPUTED FLUX AND BONDARENKO FLUX EHI (MUST BE IN RESOLVED RANGE) SIGPOT ESTIMATE OF POTENTIAL SCATTERING CROSS SECTION MAXIMUM NUMBER OF COMPUTED FLUX PDINTS NFLMAX TAPE UNIT FOR NEW FLUX PARAMETERS (DEFAULT=O) NINWT INDEX OF REFERENCE SIGMA ZERD IN SIGZ ARRAY JSIGZ (DEFAULT=O) CAR08B TABULATED (IWT=1 OR -1 ONLY) READ WEIGHT FUNCTION AS TAB1 RECORD. WGHT END WITH A /. CARD8C ANALYTIC FLUX PARAMETERS (IWT=4 DR -4 ONLY) EB THERMAL BREAK (EV) THERMAL TEMPERATURE (EV) ΤВ FISSION BREAK (EV) EC FISSION TEMPERATURE (EV) ŤC INPUT RESONANCE FLUX (IWT=O ONLY) * CAR080 NINWT TAPE UNIT FOR FLUX PARAMETERS CARD9 * MFO FILE TO BE PROCESSED SECTION TO BE PROCESSED DESCRIPTION OF SECTION TO BE PROCESSED MTO MTNAME REPEAT FOR ALL REACTIONS DESIRED MFD≠O/ TERMINATES THIS TEMPERATURE/MATERIAL. CARD 10 * NEXT MAT NUMBER TO BE PROCESSED MATO MATD=O/ TERMINATES GROUPR RUN. *---OPTIONS FOR INPUT VARIABLES-----IGN MEANING ARBITRARY STRUCTURE (READ IN) 1 CSEWG 239 GROUP STRUCTURE 2 LASL 30 GROUP STRUCTURE З ANL 27 GROUP STRUCTURE 4 RRD 50 GROUP STRUCTURE 5 GAM-I 68 GROUP STRUCTURE 6 GAM-II 100 GROUP STRUCTURE LASER-THERMOS 35 GROUP STRUCTURE 7 8 EPRI-CPM 69 GROUP STRUCTURE 9 LASL 187-GROUP STRUCTURE 10 . 11 SAND-II 620-GROUP STRUCTURE 12 13 LASL 80-GROUP STRUCTURE IGG MEANING Ω NONE

ARBITRARY STRUCTURE (LIST RECORD) 1 CSEWG 94 GROUP STRUCTURE LASL 12 GROUP STRUCTURE 2 3 STEINER 21 GROUP GAMMA-RAY STRUCTURE 4 (DRNL-TM-2564) 5 STRAKER 22 GROUP STRUCTURE LASL 48-GROUP STRUCTURE LASL 24-GROUP STRUCTURE 6 7 IWT MEANING 1 READ IN SMOOTH WEIGHT FUNCTION 2 CONSTANT 3 1/E 1/E + FISSION SPECTRUM + THERMAL MAXWELLIAN 4 EPRI-CELL LWR 5 6 (THERMAL) -- (1/E) -- (FISSION + FUSION) 7 FAST REACTOR 8 THERMAL--1/E--FAST REACTOR--FISSION + FUSION COMPUTE FLUX WITH WEIGHT N -N 0 READ IN RESONANCE FLUX FROM NINWT MFO MEANING 3 CROSS SECTION OR YIELD VECTOR FISSION CHI BY SHORT-CUT METHOD 5 6 NEUTRON-NEUTRON MATRIX NEUTRON-GAMMA MATRIX (PHOTON YIELDS GIVEN) NEUTRON-GAMMA MATRIX (PHOTON XSECS GIVEN) 16 * 17 90+LEVEL RADIOACTIVE NUCLIDE PRODUCTION (MF=9) 100+LEVEL RADIDACTIVE NUCLIDE PRODUCTION (MF=10) MTO MEANING -N PROCESS ALL MT NUMBERS FROM THE PREVIOUS ENTRY TO N INCLUSIVE RESERVED FOR THERMAL SCATTERING 221-250 AVERAGE LETHARGY AVERAGE INVERSE VELOCITY (M/SEC) 258 259 ****************** PROGRAM GAMINR * COMPUTE MULTIGROUP PHOTON CROSS SECTIONS PRODUCE MULTIGROUP PHOTON INTERACTION CROSS SECTIONS * AND HEATING KERMA FACTORS USING ENDF/B CROSS SECTIONS AND COHERENT AND INCOHERENT FORM FACTORS. INITIAL ENERGY QUADRATURE TECHIQUES ARE IDENTICAL TO THOSE USED IN GROUPR. * SECONDARY ENERGY-ANGLE QUADRATURE IS PERFORMED USING GAUSSIAN * INTEGRATION * *---INPUT SPECIFICATIONS (FREE FORMAT)------* * CARD 1 NENDE UNIT FOR ENOF/B TAPE NPEND UNIT FOR PENOF TAPE UNIT FOR INPUT NGAM TAPE (DEFAULT=0) UNIT FOR DUTPUT NGAM TAPE (DEFAULT=0) NGAM1 NGAM2 * CARD2 MATR MATERIAL TO BE PROCESSED

INPUT MATERIALS IN ASCENDING ORDER GAMMA GROUP STRUCTURE OPTION IGG WEIGHT FUNCTION OPTION IWT LEGENDRE ORDER LORD IPRINT PRINT OPTION (O/1=MINIMUM/MAXIMUM) (DEFAULT=1) CARD3 TITLE RUN LABEL UP TO 80 CHARACTERS (DELIMITED BY *. ENDED WITH /) (IGG=1 DNLY) CARD4 ٠ NGG NUMBER OF GROUPS NGG+1 GROUP BOUNDS (EV) EGG (IWT=1 DNLY) CAR05 WEIGHT FUNCTION AS TAB1 RECORD WGHT CARD6 MFO FILE TO BE PROCESSED MTO SECTION TO BE PROCESSED DESCRIPTION OF SECTION TO BE PROCESSED MTNAME REPEAT FOR ALL REACTIONS DESIRED MFD=O/ TERMINATES THIS MATERIAL MFD=-1/ IS A FLAG TO PROCESS ALL SECTIONS PRESENT FOR THIS MATERIAL (TERMINATION IS AUTOMATIC) * CARD7 MATO NEXT MAT NUMBER TO BE PROCESSED TERMINATE GAMINE RUN WITH MATD=O. *---OPTIONS FOR INPUT VARIABLES-----I GG MEANING 0 NONE ARBITRARY STRUCTURE (READ IN) CSEWG 94 GROUP STRUCTURE LASL 12 GROUP STRUCTURE 1 2 3 STEINER 21-GROUP GAMMA-RAY STRUCTURE STRAKER 22 GROUP STRUCTURE 4 5 LASL 48-GROUP STRUCTURE LASL 24-GROUP STRUCTURE 6 7 IWT MEANING 1 READ IN CONSTANT 2 1/E + ROLLOFFS 3 PROGRAM ERRORR * PRODUCE CROSS SECTION COVARIANCES FROM ERROR FILES IN ENDF/B FORMAT * FIRST. THE UNION ENERGY GRID OF THE USERS GROUP STRUCTURE * AND THE ENDF COVARIANCE ENERGIES IS DETERMINED. THE ARRA THE ARRAY DF CDEFFICIENTS FOR DERIVED CROSS SECTIONS IS ALSO CONSTRUCTED. * THEN MULTIGROUP CROSS SECTIONS ARE COMPUTED ON THE UNION * GRID (SEE GRPAV), DR THEY ARE READ FROM A MULTIGROUP CROSS * SECTION LIBRARY AND THEN COLLAPSED TO THE UNION GRID. THE * METHODS OF GROUPR ARE USED FOR CROSS SECTION AVERAGING. ENDF COVARIANCES AND THE GROUP CROSS SECTIONS ARE THEN COMBINED * TO GET THE BASIC COVARIANCE MATRICES (SEE COVCAL). FINALLY, * THE BASIC MATRICES ARE COMBINED TO GET COVARIANCES FOR * DERIVED REACTIONS, THE MATRICES ARE COLLAPSED TO THE USER-S FINALLY.

■ GROUP STRUC ■ ONTO AN OUT	TURE, AND THE RESULTS ARE PRINTED AND/OR WRITTEN * PUT GENDF TAPE FOR LATER USE (SEE COVOUT).
INPUT SPE	CIFICATIONS (FREE FORMAT)
CARD 1	•
NENOF	UNIT FOR ENDE APE
NPEND	UNIT FOR PENDE TAPE +
NGOUT	UNIT FOR INPUT GROUP XSEC (GENOF) TAPE *
k	(IF ZERD, GROUP XSECS WILL BE CALCULATED) *
k	(IF MFCOV.EQ.31(SEE BELDW) , NGDUT CANNOT BE O) *
•	(DEFAULT=O) +
NOUT	UNIT FOR DUTPUT COVARIANCE TAPE (DEFAULT=0) +
	UNIT FUR INPUT COVARIANCE TAPE (DEFAULT=0) *
	(NIN AND NUUL MUST BE BUTH CODED OR BOTH BINARY)
	MATERIAL TO RE RROCECCEO
*	(IGN DEFINITION SAME AS GROUPP EXCEPT IGN=19
*	WHICH MEANS READ IN AN ENERGY GRID. AS IN IGN=1.
*	AND SUPPLEMENT THIS WITH THE ENDE COVARIANCE GRID
•	WITHIN THE RANGE OF THE USER-SPECIFIED ENERGIES) +
*	(DEFAULT=1)
IPRINT	PRINT OPTION (O/1=MINIMUM/MAXIMUM) (DEFAULT=1)
IRELCO	COVARIANCE FORM (0/1=ABSOLUTE/RELATIVE) (DEFAULT=1) +
CARD 3	(DMIT IF NGDUT.NE.O) +
	WEIGHT FUNCTION OPTION
* MPRINI * TEMDIN	TEMPEDATURE (DEFAULT-200)
	IEMPERATURE (UEFAULI=300)
FOR ENDF	B VERSION 4 (IVERF=4) ONLY
• CARD 4	
* NEK	NUMBER OF DERIVED XSEC ENERGY RANGES
•	(IF ZERO. ALL XSECS ARE INDEPENDENT)
CARD 5	(DMIT IF NEK=O)
* EK	NEK+1 DERIVED XSEC ENERGY BOUNDS
CARD 6	(DMIT IF NEK=O)
* AKXY	DERIVED CROSS SECTION COEFFICIENTS
- +FOR ENDE	/B VERSION 5 (IVERE=5) ONLY
*	
* CARD 7	
* IREAD	0/1/2=PRDGRAM CALCULATED MTS/INPUT MTS AND EKS/
*	CALCULATED MTS PLUS EXTRA MAT1-MT1 PAIRS FROM INPUT
*	(DEFAULT=O)
* MFCOV	ENDE CUVARIANCE FILE (31, 32, OR 33) TO BE
•	PRUCESSED (DEFAULI#33).
*	COVADIANCES FOOM DECONANCE_DADAMETED INCEDIAINITIES
•	(ME=32) IS INCLUDED WHEN ME=33 IS REQUESTED
÷	(MR 02) 15 INCLOSED WHEN MR -35 15 KEQUESTED:
FOLLOWING	CARDS ONLY IF IREAD.EQ.1
* CARD 8	
* NMT	NO. MTS TO BE EVALUATED
* NEK	ND. DERIVED CROSS SECTION ENERGY RANGES
*	(DEFAULT=1) (NEK=O IS FLAG FOR DEFAULT)
- CARU BA	NMT MTC
	כות וחוי
* EK	NEK+1 DERIVED CROSS SECTION ENERGY ROUNDS
*	(DEFAULT=1.E-5.2.E7)
* CARD 9	· · · · · · · · · · · · · · · · · · ·
* ^KXY	DERIVED CROSS SECTION COFFEICIENTS

```
FOLLOWING CARDS ONLY IF IREAD.EQ.2
*
*
   CARD 10
     MAT 1
     MT 1
               REPEAT FOR ALL MAT1-MT1 PAIRS DESIRED
               TERMINATE WITH MAT1=O.
              CARD 11
               (FDR IGN.EQ.1.DR.IGN.EQ.19)
     NGN
               NUMBER OF GROUPS
   CARD 11A
     EGN
               NGN+1 GROUP BOUNDS (EV)
   CARD 12
               (FOR IWT.EQ.1 ONLY)
               WEIGHT FUNCTION AS A TAB1 RECORD
     WGHT
* *
                  PROGRAM COVR
*********
*
 PROCESS COVARIANCE DATA FROM ERRORR.
* PLOT A MATRIX OF CORRELATION COEFFICIENTS
  AND AN ASSOCIATED PAIR OF STANDARD DEVIATION VECTORS, I.E., A COVARIANCE MATRIX. THE CORRE
                                            THE CORRELATION
  MATRIX IS PLOTTED AS A SHADED CONTDUR PLOT AND THE VECTORS
* ARE PLOTTED AS SEMI-LOG PLOTS, ONE ROTATED BY 90 DEGREES.
* THE LOG ENERGY GRIDS FOR THE VECTOR PLOTS ARE IDENTICAL
* TO THE GRIDS FOR THE MATRIX PLOT.
  NDTE---IF MORE THAN ONE MATERIAL APPEARS ON THE INPUT TAPE,
THE MAT NUMBERS MUST BE IN ASCENDING DRDER.
THIS PROGRAM EMPLOYS THE PROPRIETARY PLOTTING SOFTWARE
  PACKAGE, DISSPLA.
*---INPUT SPECIFICATIONS (FREE FORMAT)------
   CARD 1
       NIN
                         INPUT TAPE UNIT
       NOUT
                         OUTPUT TAPE UNIT
                         (DEFAULT=O=NONE)
   CARD 2
       GAPREF
                         REFERENCE SHADING GAP WIDTH
                         (DEFAULT=.020)
       GAP 1
                         FULL-CORRELATION SHADING GAP WIDTH
                         (DEFAULT=.008)
                         FACTOR FOR NEGATIVE CORRELATION GAP
       XGAP
                         ADJUSTMENT (DEFAULT=1.75)
       NOTE. THE ABOVE DEFAULTS ARE ACTIVATED BY ENTERING ZERDES
       DR JUST A SLASH. A SECOND SET OF DEFAULTS FOR GAPREF AND
GAP1 ARE ACTIVATED BY INPUTTING A VALUE OF GAPREF GREATER
       THAN 0.2. IN THIS CASE, THE VALUES USED ARE 0.030*FACT
AND 0.016*FACT, RESPECTIVELY, WHERE FACT IS THE INPUT
       VALUE OF GAPREF.
*
   CARD 3
       IRELCO
                         RELATIVE COVARIANCE OPTION
                         O/1=ABSOLUTE/RELATIVE COVARIANCES
                         (DEFAULT¤1)
*
*
       NCASE
                          ND. CASES TO BE RUN (MAXIMUM=40)
                          (DEFAULT=1)
       NOLEG
                         PLOT LEGEND OPTION
```

```
-1/0/1=LEGEND FOR FIRST SUBCASE ONLY/
                          LEGEND FOR ALL PLOTS/NO LEGENOS
                          (DEFAULT=O)
       NSTART
                          SEQUENTIAL FIGURE NUMBER
                          O/N=NOT NEEDED/FIRST FIGURE IS FIGURE N.
                          (DEFAULT=0)
   CARD 4
       MAT
                          DESIRED MAT NUMBER
                         DESIRED MT NUMBER
       MT
       MAT 1
                        DESIRED MAT1 NUMBER
       MT 1
                          DESIRED MT1 NUMBER
                          (DEFAULT FOR MT, MAT1 AND MT1 ARE 0,0,0
MEANING PROCESS ALL MTS FOR THIS MAT
                          WITH MAT1=MAT)
             REPEAT CARD 4 NCASE TIMES
                     *****
PROGRAM MODER
   CHANGE THE MODE OF AN ENDF/B TAPE.
   ALSO WORKS FOR PENOF, GENDF AND COVARIANCE TAPES.
 ---INPUT SPECIFICATIONS (FREE FORMAT)-----
  CARD 1
٠
                  UNIT NUMBERS
        NIN
                  INPUT UNIT
                  ABS (NIN) GE 1 AND LE 19 IS A FLAG
TO LOOP OVER DIFFERENT TAPES AND MATERIALS
        NOUT
                  OUTPUT UNIT
   A POSITIVE UNIT IS CODED (MODE 3).
    A NEGATIVE UNIT IS BLOCKED BINARY (NJDY MODE).
        CARDS 2 AND 3 FOR ABS (NIN) GE 1 AND LE 19 ONLY.
                          FOR ENDE /B AND PENDE TAPES
  CARD 2
*
        TPID
                  TAPEID FOR NOUT. 66 CHARACTERS ALLOWED (DELIMITED WITH *, ENDED WITH /)
  CARD 3
                  INPUT UNIT
        NIN
                  TERMINATE MODER BY SETTING NIN=0
        MATO
                  MATERIAL ON THIS TAPE TO ADD TO NOUT
                    ******
PROGRAM OTFR
•
  CONVERT DUTPUT OF GROUPR TO DTF FORMAT.
* PROCESSES NEUTRON AND GAMMA PRODUCTION CROSS SECTIONS AND
* MATRICES. THE NEUTRON TABLES CAN HAVE REDUCED TABLE LENGTH.
* UP-SCATTER IS ALLOWED. THE ABSORPTION REACTION IS COMPUTED
* FROM THE TOTAL CROSS SECTION AND TOTAL SCATTERING. ANY EDIT
* CAN BE PRODUCED WHICH ARE EITHER GIVEN IN THE ENOF/B FILE
                                                                  ANY EDITS
* OR ARE LINEAR COMBINATIONS OF ENOF/B CROSS SECTIONS. THE *
* FISSION NU*SIGF AND CHI ARE COMPUTED FROM THE FISSION MATRICES *
* FOR ALL PARTIAL FISSION REACTIONS. CHI INCLUDES SOURCE
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* WEIGHTING. THE PL TABLES FOR L.GT.O CONTAIN THE PL WEIGHTED * * TOTAL IN THE TOTAL POSITION AND THE PL TRANSPORT CROSS SECTION * * IN THE ABSORPTION POSITION. THE GAMMA TABLES HAVE GAMMA GROUP * * 1 IN POSITION 1, 2 IN POSITION 2, ETC, WITH A TABLE LENGTH * EQUAL TO THE NUMBER OF GAMMA GROUPS. WARNING... THIS PROGRAM IS EXTREMELY MACHINE DEPENDENT BECAUSE OF ITS PLOTTING CAPABILITY. THE CODING IS LEFT AS A GUIDE FOR CONVERSION TO OTHER SYSTEMS. *---INPUT SPECIFICATIONS (FREE FORMAT)------CARD 1 UNITS * INPUT UNIT WITH DATA FROM GROUPR (BINARY). NIN NOUT OUTPUT UNIT CONTAINING OTF TABLES (CODED). (DEFAULT=O=NONE) INPUT UNIT WITH PENDE TAPE FOR POINT PLOTS. NPENO (DEFAULT=O=NDNE) CARD 2 **OPTIONS** IPRINT PRINT CONTROL (O MINIMUM, 1 MAXIMUM) FILM CONTROL (0/1/2=ND/YES WITH 1 PLOT PER FRAME/ IFILM YES WITH 4 PLOTS PER FRAME (DEFAULT=O) EDIT CONTROL (0/1=IN TABLE/SEPARATE) (DEFAULT=O) (SEPARATE ONLY FOR 30 NEUTRON GROUP X 12 PHOTON IEDIT GROUP DATA) CARDS 3 THROUGH 5 ONLY FOR IEDIT=0 CARD 3 NEUTRON TABLES NUMBER OF NEUTRON TABLES DESIRED. NLMAX NUMBER OF NEUTRON GROUPS NG POSITION OF TOTAL CROSS SECTION IPTOTL POSITION OF IN A CROSS SECTION. POSITION OF IN GROUP SCATTERING CROSS SECTION. NEUTRON TABLE LENGTH DESIRED. **IPINGP** ITABL NUMBER OF ENTRIES IN EDIT TABLE (DEFAULT=O). NUMBER OF THERMAL GROUPS (DEFAULT=O). NEO NTHERM CARD 3A ONLY FOR NTHERM NE O CARD 3A THERMAL INCOHERENT AND COHERENT MTS MT FOR THERMAL INCOHERENT DATA MTI MTC MT FOR THERMAL COHERENT DATA (DEFAULT=0) NO. COHERENT LEGENDRE DRDERS (DEFAULT=0) EDIT NAMES NLC CARD 4 SIX CHARACTER HOLLERITH NAMES FOR EDITS FOR AS MANY CARDS AS NEEDED. THERE WILL BE IPTOTL-3 NAMES READ. EACH NAME IS DELIMITED WITH *. EDIT SPECIFICATIONS NED TRIPLETS OF NUMBERS ON AS MANY CARDS AS NEEDED. CARD 5 POSITIONS CAN APPEAR MORE THAN ONCE. REACTION TYPES CAN APPEAR MORE THAN ONCE. POSITION OF EDIT QUANTITY. JPDS ENDF/B REACTION NUMBER. MT MULT MULTIPLICITY TO BE USED WHEN ADDING THIS MT. CARD 6 GAMMA RAY TABLES NPTABL NUMBER OF GAMMA TABLES DESIRED (DEFAULT=O) NUMBER DF GAMMA GROUPS (DEFAULT=0) NGP MATERIAL DESCRIPTION CARD 7 ONE CARD FOR EACH TABLE SET DESIRED. MAT=O/ TERMINATES EXECUTION OF OTFR. HISNAM HOLLERITH ISOTOPE NAME MATERIAL NUMBER AS IN ENDF/B (DEFAULT=O) INDEX NUMBER DF SIGMA-ZERD DESIRED (DEFAULT=1) ΜΔΤ JSIGZ TEMPERATURE DESIRED (DEFAULT=300) OTEMP

PROGRAM CCCCR PRODUCE CCCC-IV FILES FROM NJDY INTERMEDIATE CRDSS SECTION LIBRARY WORKING FROM A GROUPR DUTPUT TAPE, THIS MODULE PRODUCES THE FOLLOWING THREE STANDARD INTERFACE FILES, ISOTXS BRKOXS DLAYXS. AS SPECIFIED BY THE COMMITTEE FOR COMPUTER CODE CODRDINATION (CCCC), TO FACILITATE THE EXCHANGE DF NUCLEAR DATA FOR REACTOR* CALCULATIONS (REFERENCE 1). IN A GIVEN RUN. ALL THREE FILES CAN BE PRODUCED USING THE SAME USER-SPECIFIED LIST OF ISOTOPES. THE CODE WILL IGNORE ISOTOPES WHICH ARE NOT PRESENT ON THE GROUPR TAPE (AND IN THE CASE OF OLAYXS, ISOTOPES WITHOUT DELAYED NEUTRON DATA). THE ISOTXS CODING ALLOWS FOR NSBLK EQUAL TO DNE OR NGROUP.* IN ADDITION, FILES WITH HIGHER ORDER MATRICES CAN BE PRODUCED * WITH A SEPARATE BLOCK FOR EACH L-ORDER (IFOPT=2) OR WITH ALL ORDERS IN ONE BLOCK (IFOPT=1). THIS FLEXIBILITY ACCOMMODATES LARGE GROUP STRUCTURES. FISSION VECTORS OR FISSION MATRICES CAN BE PRODUCED. IN BRKOXS, THE POTENTIAL SCATTERING CROSS SECTION FOR ALL ENERGY GROUPS IS EQUAL TO THE USER-INPUT VALUE (XSPD). THE ELASTIC REMOVAL F-FACTOR IS SUPPLIED AS THE SIXTH REACTION. * 1. R.D.DDELL. STANDARD INTERFACE FILES AND PROCEDURES FOR REACTOR PHYSICS CODES, VERSION IV, LASL REPORT LA-6941-MS (SEPT.77) *---INPUT SPECIFICATIONS (FREE FORMAT)------CCCCR-CARD 1 UNITS * NIN INPUT UNIT FOR DATA FROM GROUPR DUTPUT UNIT FOR ISOTXS (O IF ISOTXS NOT WANTED) DUTPUT UNIT FOR BRKDXS (O IF BRKDXS NOT WANTED) DUTPUT UNIT FOR DLAYXS (O IF DLAYXS NOT WANTED) NISOT NBRKS NOLAY CARD 2 IDENTIFICATION LPRINT PRINT FLAG (0/1=NOT PRINT/PRINTED) FILE VERSION NUMBER (DEFAULT=O) USER IDENTIFICATION (12 CHARACTERS) IVERS HUSE DELIMITED BY *, ENDED BY /. (DEFAULT=BLANK) CARD 3 * HOLLERITH IDENTIFICATION OF SET (12 CHARACTERS) DELIMITED BY *, ENDED BY /. (DEFAULT=BLANK) HSETIO CARD 4 FILE CONTROL NGROUP NUMBER OF NEUTRON ENERGY GROUPS NGGRUP NUMBER OF GAMMA ENERGY GROUPS NUMBER OF ISOTOPES DESIRE NISD MAXIMUM LEGENDRE DRDER MAXORD MATRIX BLOCKING OPTION (1/2=BLOCKING BY IFOPT REACTION/LEGENDRE ORDER) CARD 5 ISOTOPE PARAMETERS (ONE CARD PER ISOTOPE) (FIRST FOUR WORDS ARE HOLLERITH, UP TO SIX CHARACTERS EACH, DELIMITED BY *)

HOLLERITH ISOTOPE LABEL HISNM HABSID HOLLERITH ABSOLUTE ISOTOPE LABEL IDENTIFIER OF DATA SOURCE LIBRARY (ENDF/B) HTOENT HMAT ISOTOPE IDENTIFICATION IMAT NUMERICAL ISOTOPE IDENTIFIER (ENDF/B MAT NUMBER) AVERAGE POTENTIAL SCATTERING CROSS SECT. (BRKDXS) XSPD *-CISDTX- (ONLY IF NISDT.GT.O) * CARD 1 FILE CONTROL NSBLOK SUBBLOCKING OPTION FOR SCATTERING MATRIX (1 DR NGRUP SUB-BLOCKS ALLOWED) MAXIMUM NUMBER OF UPSCATTER GROUPS (ALWAYS ZERD) MAXUP MAXIMUM NUMBER OF DOWNSCATTER GROUPS MAXON FISSION CHI REPRESENTATION ICHIX VECTOR (USING GROUPR FLUX) - 1 NONE 0 VECTOR (USING INPUT FLUX) +1 . GT . 1 MATRIX CARD 2 CHI VECTOR CONTROL (ICHIX=1 ONLY) NGROUP FLUX VALUES USED TO COLLAPSE THE GROUPR SPEC FISSIDN MATRIX INTO A CHI VECTOR CARD 3 CHI MATRIX CONTROL (ICHIX.GT.1 DNLY) * SPEC NGROUP VALUES OF SPEC(I)=K DEFINE THE RANGE OF GROUPS I TO BE AVERAGED TO OBTAIN SPECTRUM K. INDEX K RANGES FROM 1 TO ICHI. THE MODEL FLUX IS USED TO WEIGHT EACH GROUP I. CARD 4 ISOTOPE CONTROL (DNE CARD PER ISOTOPE) * KBR ISOTOPE CLASSIFICATION GRAM ATOMIC WEIGHT AMASS TOTAL THERMAL ENERGY/FISSION EFISS TOTAL THERMAL ENERGY/CAPTURE ECAPT TEMP ISOTOPE TEMPERATURE SIGPOT AVERAGE EFFECTIVE POTENTIAL SCATTERING DENSITY OF ISOTOPE IN MIXTURE ADENS *-CBRKXS- (DNLY IF NBRKS.GT.O) CARD 1 (216) FILE DATA * NTI NUMBER OF TEMPERATURES DESIRED (-N MEANS ACCEPT FIRST N TEMPERATURES) NUMBER OF SIGPO VALUES DESIRE NZI (-N MEANS ACCEPT FIRST N DILUTION FACTORS) * CARD 2 (NOT NEEDED IF NTI.LT.O) ATEM(NTI) VALUES OF DESIRED TEMPERATURES CARD 3 (NOT NEEDED IF NZI.LT.O) ASIG(NZI) VALUES OF DESIRED SIGPD ***-COLAYX-- NO INPUT REQUIRED** PROGRAM MATXSR PRODUCE MATXS INTERFACE FILE FROM NJOY INTERMEDIATE CROSS SECTION LIBRARY THE MATXS FILE IS A GENERALIZED, FLEXIBLE FORMAT SIMILAR TO THE CCCC-ISOTXS FORMAT. WORKING FROM A GROUPR AND/OR GAMINR DUTPUT TAPE, THIS MODULE CAN PROCESS NEUTRON CROSS SECTIONS. GAMMA PRODUCTION DATA AND GAMMA INTERACTION CROSS SECTIONS ONTO A SINGLE DUTPUT FILE. IN ITS PRESENT FORM THIS * MODULE WILL ACCEPT ALL RELEVANT REACTIONS PRESENT ON THE INPUT* TAPE(S) AND PRODUCE AN ARCHIVAL DUTPUT FILE WHICH CAN BE

MANIPULATED BY A SEPARATE CODE CALLED TRANSX. * A MATXS FILE SPECIFICATION MAY BE FOUND FOLLOWING THE INPUT INSTRUCTIONS. *---INPUT SPECIFICATIONS (FREE FORMAT)-----CARD 1 UNITS INPUT UNIT FOR DATA FROM GROUPR INPUT UNIT FOR DATA FROM GAMINR NGEN1 NGFN2 NMATX OUTPUT UNIT FOR MATXS CARD 2 USER IDENTIFICATION LPRINT O/1 MEANS NO PRINT/PRINT FILE VERSION NUMBER (DEFAULT=O) IVERS USER ID (12 CHARACTERS, DELIMITED BY *, ENDED BY /) HUSE (DEFAULT=BLANK) CARD 3 FILE CONTROL * NUMBER OF PARTICLES FOR WHICH GROUP NPART STRUCTURES ARE GIVEN NTYPE NUMBER OF DATA TYPES IN SET NUMBER OF CAROS TO BE READ FOR HOLLERITH NHOLL IDENTIFICATION RECORD. CARD 4 SET HOLLERITH IDENTIFICATION HSETIO HOLLERITH IDENTIFICATION OF SET (A6) (EACH LINE CAN BE UP TO 72 CHARACTERS, DELIMITED WITH *, ENDED BY /) CARD 5 PARTICLE IDENTIFIERS HOLLERITH IDENTIFIERS FOR PARTICLES HPART (UP TO 6 CHARACTERS EACH, DELIMITED BY *) CARD 6 ENERGY GROUPS NGRP NUMBER OF GROUPS FOR EACH PARTICLE CARD 7 DATA TYPE IDENTIFIERS HOLLERITH IDENTIFIERS FOR DATA TYPES HTYPE (UP TO 6 CHARACTERS EACH, DELIMITED BY *) THE FOLLOWING SEQUENCE OF CAROS IS REPEATED FOR EACH DATA TYPE * CARD 11 DATA TYPE CONTROL IFOPT BLOCKING OPTION FOR MATRICES (1/2 MEANS MATRICES NOT BLOCKED/BLOCKED BY L-ORDER) WARNING-IFOPT=2 IS UNDER DEVELOPMENT SUB-BLOCKING PARAMETER (1/NING MEANS THAT A RECORD NSBLK CONTAINS ALL INPUT GROUPS/ONE INPUT GROUP) MAXORO MAXIMUM LEGENDRE ORDER NUMBER OF MAT NAMES TO BE READ NMATN CARD 12 INPUT PARTICLES IINP PARTICLE NUMBERS OF ALL INCIDENT PARTICLES CARD 13 OUTPUT PARTICLES PARTICLE NUMBERS OF ALL OUTGOING PARTICLES TOUTP CARD 14 MATERIAL DATA (DNE CARD PER MATERIAL) HMAT HOLLERITH MATERIAL IDENTIFIER (UP TO 6 CHARACTERS EACH, DELIMITED BY *) MAX NUMBER OF TEMPERATURES MAX NUMBER DF SIGMA ZERD VALUES NTEMP NSIGZ INTEGER MATERIAL IDENTIFIER (IF MATERIAL HAS AN ENOF/B MAT NUMBER. USE THAT) IMAT PROGRAM ACER * PREPARE A DATA LIBRARY FOR MCNP, THE LOS ALAMOS CONTINUOUS

* ENERGY MONTE CARLD CODE. * REACTION CROSS SECTIONS ARE WRITTEN OUT ON THE GRID OF THE TOTAL CROSS SECTION FROM THE INPUT PENOF TAPE (ASSUMED TO BE LINEARIZED AND UNIONIZED). REDUNDANT REACTIONS (EXCEPT MT1 AND MT452) ARE REMOVED. MT18 IS CONSIDERED REDUNDANT IF MT19 IS PRESENT. ANGULAR DISTRIBUTIONS (MF4) AND TABULATED ENERGY DISTRIBUTIONS (MF5, LF1 OR 5) ARE CONVERTED INTO EQUAL PROBABILITY BINS. THE INCIDENT ENERGY GRIDS OF BOTH ARE THINNED FOR LINEAR INTERPOLATION USING THE SPECIFIED TOLERANCE. ANALYTIC SECONDARY ENERGY DISTRIBUTIONS ARE COPIED. ALL PHOTON PRODUCTION CROSS SECTIONS ARE COM-BINED ON THE CROSS SECTION ENERGY GRID AND WRITTEN AS MF13,MT3. MULTIGROUP PHOTON PRODUCTION CROSS SECTIONS ARE OBTAINED FROM THE NGEND INPUT TAPE. THE PHOTON DISTRIBUTIONS ARE SUMMED AND CONVERTED INTO A SET OF EQUALLY PROBABLE MEAN ENERGIES AND WRITTEN AS MF15, MT3LF3 (A SPECIALLY DEFINED LAW). MF14 IS MADE ISOTROPIC. UNRESOLVED REGION PROBABILITY TABLES ARE GENERATED BY A LEAST SQUARES FIT TO THE SELF-SHIELDED CROSS SECTIONS ON MT152 (SEE UNRESR) AND WRITTEN OUT AS MT153 IN A SPECIAL FORMAT. THE DICTIONARY DN THE OUTPUT TAPE IS CORRECTED TO REFLECT THE CHANGES. -- INPUT SPECIFICATIONS (FREE FORMAT)------* * CARD 1 NENDF UNIT FOR INPUT ENDF/B TAPE UNIT FOR INPUT PENDE TAPE UNIT FOR INPUT MULTIGROUP PHOTON DATA NPEND NGEND NACE UNIT FOR OUTPUT ACE TAPE NOUT UNIT FOR OUTPUT ETOPL TAPE CARD2 IOPT TYPE OF ACER RUN OPTION (0/1/2=NORMAL/WRITE THERMAL DATA ONLY/ WRITE PROBABILITY TABLES ONLY) (IOPT=2 NOT YET OPERATIONAL) MATO MATERIAL TO BE PROCESSED TEMPO TEMPERATURE DESIRED (KELVIN) (DEFAULT=300) PRINT CONTROL (O MIN. 1 MAX) (DEFAULT=1) O/NN=ND FICHE OUTPUT/UNIT FOR FICHE IPRINT NPRINT (DEFAULT=O) CARD3 FOR IOPT=O DNLY ERR TOLERANCE FOR THINNING DISTRIBUTIONS NUMBER OF BINS FOR MF4 AND MF5 (DEFAULT=32) NBINA NBINP NUMBER OF PHOTON AVERAGE ENERGIES (DEFAULT=20) CARD4 FOR IOPT=1 DNLY MT FOR THERMAL INCOHERENT DATA MTT NBINT NUMBER OF BINS FOR INCOHERENT SCATTERING MTE MT FOR THERMAL ELASTIC DATA IELAS O/1=COHERENT/INCOHERENT ELASTIC MAXIMUM ENERGY FOR THERMAL TREATMENT (EV) **FMAX** (DEFAULT=1000.=DETERMINED FROM MF3, MTI) IWT WEIGHTING OPTION O/1=VARIABLE/CONSTANT (DEFAULT=VARIABLE) CARD5 FOR IDPT=2 ONLY NO. OF BANDS IN PROBABILITY TABLES (2 OR 3) NBANO (DEFAULT=2) CARD6 FOR NACE NE O AND IOPT=O DNLY TYPE OF THINNING IS DETERMINED BY SIGN OF THIN(1) (+/-=ENERGY SKIP/INTEGRAL FRACTION) E1 ENERGY BELOW WHICH TO USE ALL ENERGIES (EV) THIN(1) OR IWTT WEIGHTING OPTION (1=FLAT,2=1/E) (1/E ACTUALLY HAS WEIGHT=10 WHEN E LT THIN(2) E2 ENERGY ABOVE WHICH TO USE ALL ENERGIES

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OR TARGET NUMBER OF POINTS
                ISKF SKIP FACTOR--USE EVERY ISKF-TH ENERGY
BETWEEN E1 AND E2, OR RSIGZ REFERENCE SIGMA ZERD
     THIN(3)
PROGRAM POWR
   PRODUCE INPUT FOR THE EPRI-CELL CODES GAMTAP (FAST) AND
   LIBRAR (THERMAL), AND THE EPRI-CPM CODE CLIB.
*---INPUT SPECIFICATIONS (FREE FORMAT)-----
  CARD 1
     NGENDF
              UNIT FOR INPUT GOUT TAPE
     NOUT
              UNIT FOR OUTPUT TAPE
  CARD 2
              LIBRARY OPTION (1=FAST, 2=THERMAL, 3=CPM)
     LIB
     IPRINT
              PRINT OPTION (O=MINIMUM, 1=MAXIMUM)
              (DEFAULT=O)
              GROUP COLLAPSING OPTION (O=COLLAPSE FROM 185 GROUP
     ICLAPS
              TO DESIRED GROUP STRUCTURE, 1=ND COLLAPSE)
              (DEFAULT=O)
*---FOR LIB=1-----
  CARD 3
*
     MATO
              MATERIAL TO BE PROCESSED
              IF MATO LT O, READ-IN ABSORPTION DATA ONLY FOR
THIS MATERIAL WITH MAT=ABS(MATO) DIRECTLY FROM
              INPUT DECK (SEE CARD 6)
    FOLLOWING THREE PARAMETERS IRRELEVANT FOR MATO LT O
     RTEMP
              REFERENCE TEMPERATURE (DEGREES KELVIN)
              (DEFAULT=300 K)
     IFF
              F-FACTOR OPTION
               (0/1=D0 NOT CALCULATE F-FACTORS/CALCULATE IF FDUND)
               (DEFAULT=1)
     NSGZ
              ND. OF SIGMA ZERDES TO PROCESS FOR THIS MATERIAL
              (DEFAULT=0=ALL FOUND ON INPUT TAPE)
REF. SIGZERD FOR ELASTIC MATRIX (DEFAULT=1)
     IZREF
  CARDS 4 AND 5 FOR NORMAL RUN ONLY (MATD GT O)
  CARD 4
     WORD
              DESCRIPTION OF NUCLIDE (UP TO 16 CHARACTERS.
              OELIMITED WITH *, ENDED WITH /) (DEFAULT=BLANK)
  CARD 5
     FSN
                TITLE OF FISSION SPECTRUM (UP TO 48 CHARACTERS,
  DELIMITED WITH *, ENDED WITH /) (DEFAULT=BLANK)
CARD 6 FOR READING IN ABSORPTION DATA DNLY
               NGND ABSORPTION VALUES (DEFAUL1 VALUES=0)
     ABS
  REPEAT CARDS 3 THROUGH 6 FOR EACH MATERIAL DESIRED.
TERMINATE WITH MATD=O/ (I.E., A O/ CARD).
CARD 3
     MATO
              MATERIAL TO BE PROCESSED
              TEMPERATURE ID (DEFAULT=300 K)
     IOTEMP
              HOLLERITH NAME OF ISOTOPE (UP TO 10 CHARACTERS,
DELIMITED WITH *, ENDED WITH /) (DEFAULT=BLANK)
     NAME
  CARD 4
              DEFAULT FOR ALL VALUES=0.
     ITRC
               TRANSPORT CORRECTION OPTION (O NO. 1 YES)
              THERMAL INELASTIC MT
     MTT
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THERMAL ELASTIC MT MTC DEFAULT FOR ALL VALUES=O. CARD 5 XI ALPHA MUBAR NU KAPPA FISSION KAPPA CAPTURE LAMBOA * SIGMA S IF O, SET TO SCATTERING CROSS SECTION AT GROUP 35 * * REPEAT CARDS 3 THRU 5 FOR EACH MATERIAL AND TEMPERATURE DESIRED* * (MAXIMUM NUMBER OF TEMPERATURES ALLOWED IS 7.) TERMINATE WITH MATD=O/ (I.E., A O/ CARD). * CARD 3 NLIB NUMBER OF LIBRARY. DATE LIBRARY IS WRITTEN (I FORMAT). IDAT NUMBER OF MATERIALS TO BE ADDED. ADD OPTION (O=MATS WILL BE READ IN, NEWMAT IOPT 1=USE ALL MATS FOUND ON NGENDF). O/1/2=REPLACE ISOTOPE(2) IN CPMLIB/ ADD/CREATE A NEW LIBRARY (DEFAULT=0) MODE FILE5 (BURNUP DATA) OPTION IE5 O/1/2=00 NOT PROCESS FILE5 BURNUP DATA/ PROCESS BURNUP DATA ALONG WITH REST OF DATA/ PROCESS BURNUP DATA DNLY (DEFAULT=0) (DEFAULT=O) FILE4 (CRDSS SECTION DATA) OPTION IF4 O/1=DD NDT PROCESS/PROCESS (DEFAULT=1) CARD 4 FOR IOPT=O DNLY ENDF MAT NUMBER OF ALL DESIRED MATERIALS. MAT FOR MATERIALS NOT ON GENDE TAPE, USE IDENT FOR MAT. IF MAT LT O, ADD 100 TO DUTPUT IDENT (FOR SECOND ISOMER OF AN ISOTOPE) CARD 5 * NINA NINA INDICATOR. 0/1/2/3=NORMAL/ ND FILE2 DATA, CALCULATE ABSORPTION IN FILE4/ NO FILE2 DATA, READ IN ABSORPTION IN FILE4/ READ IN ALL FILE2 AND FILE4 DATA ND. OF TEMPERATURES TO PROCESS FOR THIS MATERIAL (DEFAULT=0=ALL FOUND ON INPUT TAPE) NTEMP NO. OF SIGMA ZERDES TO PROCESS FOR THIS MATERIAL (DEFAULT=O=ALL FOUND ON INPUT TAPE) NSIGZ SGREF REFERENCE SIGMA ZERD FOLLOWING 2 PARAMETERS ARE FOR NINA=0 OR NINA=3. POTENTIAL CROSS SECTION FROM ENDF/B. IRES SIGP FOLLOWING 5 PARAMETERS ARE FOR NTAPEA=O ONLY MTI THERMAL INELASTIC MT MTC THERMAL ELASTIC MT IP10PT 0/1=CALCULATE P1 MATRICES/ CORRECT PO SCATTERING MATRIX INGROUPS. ******IF A P1 MATRIX IS CALCULATED FOR ONE OF THE ISOTOPES HAVING A P1 MATRIX ON THE OLD LIBRARY, FILE 6 ON THE NEW LIBRARY WILL BE COMPLETELY REPLACED. ****** INDRF O/1=INCLUDE RESONANCE FISSION IF FOUND/ DO NOT INCLUDE FOLLOWING TWO PARAMETERS FOR MODE=O ONLY POS POSITION OF THIS ISOTOPE IN CPMLIB POSR (FOR IRES=1) POSITION OF THIS ISOTOPE IN RESONANCE

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TABULATION IN CPMLIB
  REPEAT CARD 5 FOR EACH NUCLIDE.
*
 FOLLOWING THREE CARDS ARE FOR IF5 GT O ONLY
 CARD 6
              ND. TIME-DEPENDENT ISOTOPES
     NTIS
     NFIS
              ND. FISSIONABLE BURNUP ISOTOPES
 CARD 7
     IDENTB IDENT OF EACH OF THE NFIS ISOTOPES
  CARD 8
     IDENTA
             IDENT OF TIME-DEPENDENT ISOTOPE
              DECAY CONSTANT (DEFAULT=0.)
     DECAY
              NFIS YIELDS (DEFAULT=0.)
     YIFLD
   REPEAT CARD 8 FOR EACH OF THE NTIS ISOTOPES.
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  CARD 9 FOR IF5=2 ONLY
     AW
              ATOMIC WEIGHT
     INDFIS FISSION INDICATOR
     NTEMP NO. TEMPERATURES ON OLD LIBRARY
   REPEAT CARD 9 FOR EACH OF THE NTIS ISOTOPES.
  CARD 10
     LAMBDA RESONANCE GROUP GOLDSTEIN LAMBDAS
   *****REMEMBER THAT THE 69-GROUP STRUCTURE HAS 13 RESONANCE
          GROUPS WHILE THE COLLAPSED 185-GROUP STRUCTURE HAS 15.
         USE A SLASH AT END OF EACH LINE OF CARD 10 INPUT.****
   REPEAT CARD 10 FOR EACH NUCLIDE HAVING NINA=O. NINA=3, DR
                        IRES=1.
* CARDS 11 AND 11A FOR NUCLIDES HAVING NINA=3 ONLY.
* CARD 11
               NRG NUS VALUES TO GO WITH THE LAMBOA VALUES
     RESNU
  CARD 11A
   TOT NRG TOTAL XSEC VALUES TO GD WITH THE LAMBDA VALUES
READ CARDS 11 AND 11A FOR EACH NUCLIDE HAVING NINA=3.
  CARDS 12 FOR NINA GT 2 DNLY
     AW
              ATOMIC WEIGHT
     TEMP
              TEMPERATURE
  FPA NGNO ABSORPTION VALUES (DEFAULT=0.)
CARDS 12A, 12B, 12C FDR NUCLIDES HAVING NINA=3 ONLY.
*
  CARD 12A
*
     NUS
               NGND NUS VALUES
               NGND FISSION VALUES
     FIS
               NGND TRANSPORT VALUES
     XTR
*
*
  CARD 12B
                        O MEANS NO SCATTERING FROM THIS GROUP
     IA
                GROUP.
               LOWEST GROUP TO WHICH SCATTERING OCCURS
HIGHEST GROUP TO WHICH SCATTERING OCCURS
     L1
*
     L2
*
  CARD 12C
             FOR IA GT O ONLY
               L2-L1+1 SCATTERING VALUES
     SCAT
   REPEAT CARD 12B AND 12C FOR EACH GROUP
* REPEAT CARDS 12 FOR EACH DF THE NINA GT 2 NUCLIDES
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APPENDIX B

DEFINITION OF ENDF/B REACTION NUMBERS USED BY NJOY

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MT	Description
1	Total cross section (redundant, equal to the sum of all partial cross sections)
2	Elastic scattering cross section
3	Nonelastic cross section (redundant, equal to the sum of all partial cross sections except elastic scattering)
4	Total inelastic cross section (redundant, equal to the sum of MT = 51, 52, 53,, 90, 91)
6	(n,2n) cross section for first excited state (describes first neutron)
7	(n,2n) cross section for first excited state (describes first neutron)
8	(n,2n) cross section for third excited state (describes first neutron)
9	(n,2n) cross section for fourth excited state (describes first neutron)
16	direct (n,2n) cross section [total (n,2n) cross section is sum of MT = 6, 7, 8, 9, and 16]
17	(n,3n) cross section
18	Total fission cross section (sum of MT = 19, 20, 21, 38)
19	(n,f) cross section (first chance fission)
20	(n,n'f) cross section (second chance fission)
21	(n,2nf) cross section (third chance fission)
22	(n,n´α) cross section
23	(n,n´3α) cross section
24	(n,2nα) cross section
25	(n,3nα) cross section
26	(n,2n) isomeric state cross section

28	(n,n´p) cross section
29	(n,n ² α) cross section
30	(n,2n2α) cross section
32	(n,n´d) cross section
33	(n,n´t) cross section
34	(n,n ^{^3} He)
35	(n,n [^] d2α) cross section
36	(n,n´t2α) cross section
37	(n,4n) cross section
38	(n,3nf) cross section (fourth-chance fission)
46	cross section for describing the second neutron from (n,2n) reaction for first excited state
47	cross section for describing the second neutron from (n,2n) reaction for second excited state
48	cross section for describing the second neutron from (n,2n) reaction for third excited state
49	cross section for describing the second neutron from (n,2n) reaction for fourth excited state (Note: MT = 46, 47, 48, and 49 should not be included in the sum for the total (n,2n) cross section)
51	(n,n') to the first excited state
52	(n,n') to the second excited state
•	
•	
90	(n,n') to the 40th excited state
91	(n,n´) to the continuum
102	(n,y) radiative capture cross section
103	(n,p) cross section
104	(n,d) cross section
105	(n,t) cross section

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(n. ³ He)	cross	section
	(n, ³ He)	(n, ³ He) cross

- 107 (n, α) cross section
- 108 (n, 2α) cross section
- 109 (n, 3α) cross section
- 111 (n,2p) cross section
- 112 (n,pα) cross section
- 113 (n,t 2α) cross section
- 114 (n,d 2α) cross section
- 203 Total hydrogen production
- 204 Total deuterium production
- 205 Total tritium production
- 206 Total ³He production
- 207 Total ⁴He production
- 251 $\bar{\mu}_{L}$, the average cosine of the scattering angle (laboratory system) for elastic scattering
- 252 ξ , the average logarithmic energy decrement for elastic scattering
- 253 γ, the average of the square of the logarithmic energy decrement for elastic scattering, divided by twice the average logarithmic decrement for elastic scattering
- 301-450 Energy release rate parameters, E*o, for total and partial cross sections. Subtract 300 from this number to obtain the specific reaction type identification. For example, MT = 302 = (300 + 2) denotes elastic scattering
- 452 v, average total (prompt plus delayed) number of neutrons released per fission event
- 455 Delayed neutrons from fission
- 456 Prompt neutrons from fission
- 501 Total photon interaction cross section
- 502 Photon coherent scattering
- 504 Photon incoherent scattering
- 516 Pair production, nuclear and electron field (that is, pair plus triple production)

602	Photoelectric
700	(n,p ₀) cross section (cross section for leaving the residual nucleus in the ground state)
701	(n,p ₁) cross section for 1st excited state
702	(n,p ₂) cross section for 2nd excited state
703	(n,p ₃) cross section for 3rd excited state
704	(n,p ₄) cross section for 4th excited state
• •	
718	(n,p _c) cross section for continuum excited state
719	(n,p ') cross section for continuum specifically not included in σ total (redundant, used for describing outgoing proton)
720	(n,d ₀) cross section for ground state
721	(n.d.) cross section for 1st excited state
/	· · 1
•	
738	(n,d _c) cross section for continuum
- - - 738 739	<pre>(n,d_c) cross section for continuum (n,d_c') cross section for continuum specifically not included in o_T (redundant, used for describing outgoing deuteron)</pre>
738 739 740	(n,d_c) cross section for continuum (n,d_c') cross section for continuum specifically not included in σ_T (redundant, used for describing outgoing deuteron) (n,t_0) cross section for ground state
738 739 740 741	(n,d_c) cross section for continuum (n,d_c') cross section for continuum specifically not included in σ_T (redundant, used for describing outgoing deuteron) (n,t_0) cross section for ground state (n,t_1) cross section for 1st excited state
738 738 739 740 741	(n,d_c) cross section for continuum (n,d_c') cross section for continuum specifically not included $in \sigma_T$ (redundant, used for describing outgoing deuteron) (n,t_0) cross section for ground state (n,t_1) cross section for 1st excited state
 738 738 739 740 741 . .<!--</td--><td>$(n,d_{c}) \text{ cross section for continuum}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,d_{c}') \text{ cross section for describing outgoing deuteron)}$ $(n,t_{0}) \text{ cross section for ground state}$ $(n,t_{1}) \text{ cross section for 1st excited state}$ $(n,t_{c}) \text{ cross section for continuum}$</td>	$(n,d_{c}) \text{ cross section for continuum}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,d_{c}') \text{ cross section for describing outgoing deuteron)}$ $(n,t_{0}) \text{ cross section for ground state}$ $(n,t_{1}) \text{ cross section for 1st excited state}$ $(n,t_{c}) \text{ cross section for continuum}$
738 739 740 741 758 759	$(n,d_{c}) \text{ cross section for continuum}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,d_{c}') \text{ cross section for describing outgoing deuteron)}$ $(n,t_{0}) \text{ cross section for ground state}$ $(n,t_{1}) \text{ cross section for 1st excited state}$ $(n,t_{c}) \text{ cross section for continuum}$ $(n,t_{c}') \text{ cross section for continuum specifically not included in } \sigma_{T} (redundant, used for describing outgoing triton)$
 738 738 739 740 741 . .<!--</td--><td>$(n,d_{c}) \text{ cross section for continuum}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,t_{0}) \text{ cross section for ground state}$ $(n,t_{1}) \text{ cross section for 1st excited state}$ $(n,t_{c}') \text{ cross section for continuum}$ $(n,t_{c}') \text{ cross section for continuum specifically not included in } \sigma_{T} (redundant, used for describing outgoing triton)}$ $(n,^{3}He_{0}) \text{ cross section for ground state}$</td>	$(n,d_{c}) \text{ cross section for continuum}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,d_{c}') \text{ cross section for continuum specifically not included}$ $(n,t_{0}) \text{ cross section for ground state}$ $(n,t_{1}) \text{ cross section for 1st excited state}$ $(n,t_{c}') \text{ cross section for continuum}$ $(n,t_{c}') \text{ cross section for continuum specifically not included in } \sigma_{T} (redundant, used for describing outgoing triton)}$ $(n,^{3}He_{0}) \text{ cross section for ground state}$

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778	(n, ³ He _c) cross section for continuum
779	(n, ^{3}He ') cross section for continuum specifically not included in σ_{T} (redundant, used for describing outgoing He)
780	(n, α_0) cross section for ground state
781	(n, α_1) cross section for 1st excited state
• •	
798	(n, α_c) cross section for continuum
799	(n, α_1°) cross section for continuum specifically not included in σ_T^{c} (redundant, used to describe outgoing α -particle)

The above MT numbers can also be used as "LR-flags" to indicate the mode of decay of the residual nucleus. For instance, MT68/LR22 denotes a discrete (n,n') scattering event that leaves the residual nucleus in the 18th excited level; the residual nucleus then decays by α emission. The following MT-numbers are used <u>only</u> as LR-flags:

- <u>LR</u> <u>Description</u>
- 31 Indicates that γ-emission is the mode of decay of the residual nucleus formed in the primary reaction.
- 39 Indicates that internal conversion is the mode of decay of the residual nucleus formed in the primary reaction.
- 40 Indicates that electron-positron pair formation is the mode of decay of the residual nucleus formed in the primary reaction.

APPENDIX C

NJOY TEST PROBLEMS

I. EXAMPLE 1, POINTWISE PROCESSING

MOUNT ENDF/B-V TAPE 511 ON UNIT 20. MOUNT ENDF/B-III THERMAL TAPE 322 ON UNIT 26. 0 5 *MODER* 20 -21 *RECONR* -21 -22 *PENDF TAPE FOR C FROM ENDF/B TAPE 511*/ 1306 3 0 .005 0. 6 / *6-C-NAT FROM TAPE 511*/ *PROCESSED BY THE NJOY NUCLEAR DATA PROCESSING SYSTEM*/ *SEE DRIGINAL ENDF/B-V TAPE FOR DETAILS OF EVALUATION*/ 0/ *BRDADR* -22 -23 1306 1 0 0 0. .005 1.E6 300. 0/ *HEATR* -21 -23 -22 1306 1 0 0 0 0 444 *THERMR* 26 - 22 - 24 1065 1306 8 1 1 0 1 221 0 300. .05 4.6 *THERMR* 26 - 24 - 23 1065 1306 8 1 4 1 1 229 0 300. 1 .05 4.6 *GROUPR* -21 -23 0 -24 1306 3 3 3 3 1 1 1 *CARBON IN GRAPHITE*/ 300 1.E10 3 1 +TOTAL +/ 3 2 *ELASTIC*/ 3 4 *INELASTIC*/ 3 51 *DISCRETE INELASTIC*/ 3 -68 *HIGHER LEVELS*/ 3 91 *CONTINUUM INELASTIC*/ 3 102 *N,G*/ 3 103 *(N,P)*/ 3 104 *(N,D)*/ З 107 *(N,A)*/ 3 203 *TOTAL H PRODUCTION*/ 3 204 *TOTAL H2 PRODUCTION*/ 207 *TOTAL HE4 PRODUCTION*/ 3 3 221 *FREE THERMAL SCATTERING*/ 3 229 *GRAPHITE INELASTIC THERMAL SCATTERING*/ 3 230 *GRAPHITE ELASTIC THERMAL SCATTERING*/ 3 251 *MUBAR*/ 3 252 *XI*/ 3 253 *GAMMA*/ 3 301 *TOTAL HEAT PRODUCTION*/

6 2 *ELASTIC*/ 6 51 *DISCRETE INELASTIC*/ 6 -68 *HIGHER LEVELS*/ 6 91 *CONTINUUM INELASTIC*/ 6 221 *FREE THERMAL SCATTERING*/ 6 229 *GRAPHITE INELASTIC THERMAL SCATTERING*/ 6 230 *GRAPHITE ELASTIC THERMAL SCATTERING*/ 17 51 *INELASTIC GAMMA PRODUCTION*/ 16 102 *CAPTURE GAMMA PRODUCTION*/ 0/ 0/ *MODER* -23 25 *STOP* * \$\$ \$\$\$\$\$ * \$\$ \$\$ 55 \$\$ * * VERS. 10/81 \$\$\$ \$\$ \$\$ \$\$ \$\$\$\$\$\$\$ \$\$ * NUCLEAR * RAN AT LANL * \$\$\$\$ \$\$ SS SS SS \$\$\$\$ * CROSS SECTION * ON MACH. U PROCESSING DN 11/09/81 AT 18:46:05 \$\$ \$\$\$\$ \$\$ \$\$ \$\$ \$\$ \$\$ * * * \$\$\$ \$\$\$\$\$\$ \$\$\$\$\$\$\$ \$\$ \$\$ * SYSTEM * * \$\$ \$\$\$\$\$ \$\$ \$\$\$\$\$ \$\$ MODER...CHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE . 6415 INPUT UNIT (+ FOR CODED, - FOR BB) ... DUTPUT UNIT (+ FOR CODED, - FOR BB) ... 20 -21 TAPE LABEL ENDF/B-V TAPE 511 (STANDARDS MATERIALS) 11.0035 RECONR...RECONSTRUCT POINTWISE CROSS SECTIONS IN PENDE FORMAT 11.0525 UNIT FOR ENOF/B TAPE -21 UNIT FOR PENOF TAPE -22 LABEL FOR PENOF TAPE PENOF TAPE FOR C FROM ENDF/B TAPE 511 TAPE LABEL ENDF/B-V TAPE 511 (STANDARDS MATERIALS) MATERIAL TO BE PROCESSED RECONSTRUCTION TOLERANCE RECONSTRUCTION TEMPERATURE NO. SIGNIFICANT FIGURES RESONANCE-INTEGRAL-CHECK TOLERANCE ... 1306 .005 0.K 6 . 100 MAX RESONANCE-INTEGRAL ERROR 5.000E-07 DESCRIPTIVE CARDS FOR PENDE TAPE 6-C-NAT FROM TAPE 511 PROCESSED BY THE NJDY NUCLEAR DATA PROCESSING SYSTEM SEE ORIGINAL ENDF/B-V TAPE FOR DETAILS OF EVALUATION

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PROCESSING MAT 1306

6-C - ORNL EVAL-DEC73 C.Y.FU AND F.G.PEREY MAT HAS NO RESONANCE PARAMETERS PDINTS IN INITIAL UNIONIZED GRID = 776 PDINTS ADDED BY LINEARIZATION = 127 15.8445 20.5505 ******* BROADR...DOPPLER BROADENING OF ENDF/B DATA 20.5865 UNIT FOR INPUT PENOF TAPE -22 UNIT FOR DUTPUT PENOF TAPE -23 MATERIAL TO BE PROCESSED 1306 NUMBER OF FINAL TEMPERATURES 1 BOOTSTRAP (O NO. 1 YES) STARTING MATERIAL TEMPERATURE 0 0 0.K THINNING TOLERANCE 005 MAX. ENERGY 1.00E+06 FINAL TEMPERATURES 3.00E+02 21.6765 BRDADENED MAT1306 FRDM O. TD 3.0000E+02 K PDINTS IN= 903 PDINTS DUT= 866 2 102 MT 22.7435 HEATR...PROMPT KERMA 22.7735 INPUT ENDF/B UNIT -21 INPUT PENDF UNIT -23 - 22 MAT TO BE PROCESSED NO. TEMPERATURES (O=ALL) GAMMA HEAT (O NONLOCAL, 1 LOCAL) PRINT OPTION (O MIN, 1 MORE, 2 CHK) PARTIAL KERMA MT-S DESIRED 1306 0 0 0 444 PROCESSING AT TEMPERATURE= 3.0000E+02 30.3165 ********** THERMR...COMPUTE THERMAL SCATTERING CROSS SECTIONS AND MATRICES 30.3495 UNIT FOR ENOF/B TAPE 26 UNIT FOR INPUT PENDE TAPE -22 UNIT FOR INPUT PENDE TAPE UNIT FOR OUTPUT PENDE TAPE MATERIAL TO BE PROCESSED (ENOF) MATERIAL TO BE PROCESSED (PENDE) NUMBER OF ANGLE BINS NUMBER OF IEMPERATURES INELASTIC OPTION -24 1065 1306 8 1 1

 INELASTIC OPTION
 I

 ELASTIC OPTION
 D

 NUMBER DF PRINCIPAL ATOMS
 1

 REFERENCE MT
 221

 PRINT OPTION (O MIN, 1 MAX)
 0

 TEMPERATURES (KELVIN)
 3.0000E+02

 TDLERANCE
 5.0000E-02

 MAX ENERGY FOR THERMAL TREATMENT
 4.6000E+00

WRDTE THERMAL DATA FOR TEMP= 3.00E+02	44.4515
	44.5155
*****	*****
THERMRCOMPUTE THERMAL SCATTERING CROSS SECTIONS	AND MATRICES 44.516S
UNIT FOR ENDE/R TARE 26	
UNIT FUR UUTPUT PENUF TAPE	
MATERIAL TO BE PROCESSED (ENDF) 1065	
MATERIAL TO BE PROCESSED (PENDF) 1306	
NUMBER OF ANGLE BINS	
NUMBER OF TEMPERATURES	
INFLASTIC OPTION A	
NOMBER OF PRINCIPAL ATOMS	
REFERENCE M1	
PRINT OPTION (O MIN, 1 MAX) O	
TEMPERATURES (KELVIN)	
EFFECTIVE TEMPERATURES 7.1339E+02	
TOLFRANCE	
MAX ENERGY EOR THERMAL TREATMENT & 6000E+00	
MAA ENERGY FOR THERMAL TREATMENT 4.00000000	
SHORT COLLISION TIME USED FOR TRANSFERS GREATER TH	AN 1.00 EV.
DIFFERENCE BETWEEN TEMPERATURES DESIRED AND FOUND	IS 4.00E+00
WRDTE THERMAL DATA FOR TEMP= 3.00E+02	92.400S
*****	*****
GROUPRCOMPUTE SELF-SHIELDED GROUP-AVERAGED CROS	S-SECTIONS 92.2155
UNIT FOR ENDF/B TAPE21	
UNIT FOR PENOF TAPE	
UNIT FOR INPUT GOUT TAPE	
UNIT FOR OUTPUT GOUT TAPE	
MAT TO BE PROCESSED 1306	
NEUTRON GROUP STRUCTURE OPTION 3	
LEGENURE ORDER	
PRINT OPTION (O MIN, 1 MAX) 1	
RUN TITLE	
CARBON IN GRAPHITE	
TEMPERATURES (KELVIN)	
SIGMA ZERDES INFINITY	
1 1 2000 - 01 - 1 5000 - 01	
2 1.5200E-01 - 4.1400E-01	
3 4.1400E-01 - 1.1300E+00	
•	
28 1.2000E+07 - 1.3500E+07	
29 1.3500F+07 - 1.5000F+07	
30 1 5000E+07 - 1 7000E+07	

5.0000E+05 -1.0000E+06 -1.0000E+06 3 4 2.0000E+06 1.0000E+06 -2.0000E+06 -3.0000E+06 -4.0000E+06 -5.0000E+06 -6.0000E+06 -5 3.0000E+06 6 4.000DE+06 7 5.0000E+06 8 6.0000E+06 9 7.0000E+06 7.0000E+06 -10 8.0000E+06 11 8.0000E+06 -9.0000E+06 9.0000E+06 -2.0000E+07 12 WEIGHT FUNCTION.....1/E FOR ALL L PRDCESSING MAT 1306 6-C-NAT FROM TAPE 511 GRDUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 1 TOTAL ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 1 8.108E+00 FLX 7.000E+00 2 4.762E+00 FLX 1.002E+00 4.748E+00 3 FLX 1.004E+00 . . 28 1.361E+00 FLX 1.178E-01 29 1.334E+00 FLX 1.054E-01 1.472E+00 30 FLX 1.252E-01 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 2 ELASTIC ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 8.096E+00 1 4.761E+00 2 3 4.747E+00 . . 28 8.632E-01 29 8.226E-01 30 8.848E-01 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 4 INELASTIC ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION

1.0000E+D4 -

1.0000E+05 -

1.0000E+05

5.0000E+05

1

2

93.4725

93.706S

24	5.168É-02
25	2.465E-01
26	3.311E-01
27	4.290E-01
28	4.016E-01
29	4.300E-01
30	4.932E-01

GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 51 DISCRETE INELASTIC ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 24 5.168E-02 2.465E-01 25 26 3.199E-01 3.115E-01 27 2.155E-01 28 1.850E-01 29 30 1.354E-01 GROUP CONSTANTS AT T=3.000E+02 DFG K FOR MF 3 AND MT 52 CONTINUED LR 23 PARTICLE EMISSION ENRGY GROUP CONSTANTS AT INFINITE DILUTION GROUP 9.781E-03 26 4.607E-02 2.983E-02 27 28 1.261E-02 29 30 7.633E-03 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 53 CONTINUED LR 23 PARTICLE EMISSION ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 5.679E-02 8.611E-02 27 28 6.149E-02 29 30 4.342E-02 ٠ . GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 64 CONTINUED LR 23 PARTICLE EMISSION ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 30 6.363E-04

THRESHOLD IS ABOVE HIGHEST ENERGY BOUND FOR MT 65

94.0235

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94.0935

94.1315

THRESHOLD IS ABOVE HIGHEST ENERGY BOUND FOR MT 66 THRESHOLD IS ABOVE HIGHEST ENERGY BOUND FOR MT 67 THRESHOLD IS ABOVE HIGHEST ENERGY BOUND FOR MT 68 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 91 CONTINUUM INELASTIC LR 23 PARTICLE EMISSION 94.3825 ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 1.369E-03 26 27 1.187E-02 28 1.924E-02 29 2.498E-02 3.645E-02 30 GROUP CONSTANTS AT T=3.000E+02 DEG K 94.4405 FOR MF 3 AND MT 102 N.G ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 1.270E-02 1 2 1.095E-03 3 6.600E-04 . 28 7.142E-05 29 1.258E-04 1.912E-04 30 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT103 (N,P) 94.6695 ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 29 1.599E-04 7.549E-03 30 GROUP CONSTANTS AT T=3.000E+02 DEG K 94.68**0**S FOR MF 3 AND MT104 (N,D) ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 30 1.808E-02 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT107 (N,A) 94.6875 ENRGY GROUP CONSTANTS AT

GROUP	INFINITE OILUTION	
25	1,503E-02	
26	1.670E-01	
27	1.018E-01	
28	9.626E-02	
29	8.114E-02	
30	6.859E-02	
GROUP (CONSTANTS AT T=3.000E+02 DEG K	94.7595
FOR MF	3 AND MT203 TOTAL H PRODUCTION	
ENDOY	CDOUD CONSTANTS AT	
CONID		
GROOF		
29	1.599E-04	
30	7.549E-03	
GROUP (CONSTANTS AT TER COOF+02 DEG K	94 7705
FOR MF	3 AND MT204 TOTAL H2 PRODUCTION	54.7705
ENRGY	GROUP CONSTANTS AT	
GROUP	INFINITE OILUTION	
20	1 9095-02	
30	1.8082-02	
GROUP (CONSTANTS AT T=3.000E+02 DEG K	94.777S
FOR MF	3 AND MT207 TOTAL HE4 PRODUCTION	
ENDOV	OPOUR CONSTANTS AT	
	INFINITE DILUTION	
25	1.503E-02	
26	2.004E-01	
27	4.545E-01	
28	6.544E-01	
29	8.160E-01	
30	1.142E+00	
GROUP	CONSTANTS AT T=3.00DE+02 DEG K	94.8365
FOR MF	3 AND MT221 FREE THERMAL SCATTERING	
ENDO V		
ENRGY	GRUUP CUNSIANIS AI	
GRUUP	INFINITE DILOTION	
1	8.096E+00	
2	4.761E+00	
3	4.747E+00	
4	4.742E+00	
GROUP	CONSTANTS AT T=3.000E+02 DEG K	94.946S
FOR MF	3 AND MT229 GRAPHITE INELASTIC THERMAL SCATTERING	
	GROUP CUNSTANTS AT	
GROUP		
1	6.399E-01	
2	2.483E+00	
3	3 7605+00	

GRDUP FOR MF	CONSTANTS AT T=3.000E+02 DEG K 3 AND MT230 GRAPHITE ELASTIC THERMAL SCATTERING	95.1485
ENRGY GROUP	GROUP CONSTANTS AT INFINITE DILUTION	
1 2 3 4	2.714E+00 2.175E+00 9.166E-01 3.391E-01	
GROUP FOR MF	CONSTANTS AT T=3.000E+02 DEG K 3 AND MT251 MUBAR	95.3375
ENRGY GROUP	GROUP CONSTANTS AT INFINITE DILUTION	
1 2 3	5.6C0E-02 5.604E-02 5.604E-02	
28 29 30	5.793E-01 6.259E-01 6.848E-01	
GROUP FDR MF	CONSTANTS AT T=3.000E+02 DEG K 3 AND MT252 XI	102.4635
ENRGY GROUP	GROUP CONSTANTS AT INFINITE DILUTION	
2 3	1.673E-01 1.592E-01	
28 29 30	4.616E-02 3.970E-02 3.951E-02	
GROUP FDR MF	CONSTANTS AT T=3.000E+02 DEG K 3 AND MT253 GAMMA	107.2375
ENRGY GROUP	GROUP CONSTANTS AT INFINITE DILUTION	
2 3	1.671E-01 1.196E-01	
28 29 30	4 . 407E - 02 3 . 880E - 02 4 . 262E - 02	

4 4.260E+00

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GROUP CONSTANTS AT T=3.000E+02 DEG K 112.1435 + FOR MF 3 AND MT301 TOTAL HEAT PRODUCTION ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 1.128E+01 1 2 1.149E+00 З 1.072E+00 . 28 2.308E+06 2.840E+06 29 3.873E+06 30 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 6 AND MT 2 ELASTIC 112.5495 INITL FINAL GROUP CONSTANTS VS LEGENDRE ORDER GROUP GROUP O 1 8.091E+00 4.534E-01 1.144E-02 4.182E-05 7.572E-01 -2.232E-01 -1.689E-')2 -4.359E-04 4.004E+00 4.900E-01 2.363E-02 4.539E-04 1 1 2 1 2 2 . . 6.865E-02 -1.719E-02 -1.846E-02 1.031E-02 30 28 1.834E-01 1.016E-01 1.937E-02 -7.484E-03 6.055E-01 5.428E-01 4.372E-01 3.182E-01 29 30 30 30 GROUP CONSTANTS AT T=3.000E+02 DEG K 123.4495 FOR MF 6 AND MT 51 DISCRETE INELASTIC INITL FINAL GROUP CONSTANTS VS LEGENDRE ORDER GROUP GROUP O 3 1 2 2.650E-09 -1.311E-09 -8.178E-12 3.171E-10 1.222E-08 -5.999E-09 -6.950E-11 1.420E-09 5.751E-08 -2.796E-08 -4.064E-10 6.263E-09 24 7 24 8 24 q • . 9.845E-03 -8.451E-03 6.152E-03 -3.686E-03 4.082E-02 -7.808E-03 -3.619E-03 -1.275E-03 7.873E-02 5.860E-02 3.053E-02 8.245E-03 5.959E-03 5.566E-03 4.841E-03 3.891E-03 25 30 30 26 30 27 30 28 GRDUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 6 AND MT 52 CONTINUED 128.3865 LR 23 PARTICLE EMISSION FINAL GROUP CONSTANTS VS LEGENDRE ORDER GROUP O 1 2 3 INITL GROUP 2 1.555E-09 -7.688E-10 -3.943E-12 1.834E-10 7.217E-09 -3.534E-09 -4.068E-11 8.205E-10 3.421E-08 -1.650E-08 -3.360E-10 3.636E-09 26 8 26 9 26 10

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302430253026	2.152E-03 -1.461E-03 7.333E-04 -3.335E-04 3.424E-03 1.737E-03 3.777E-04 2.235E-04 2.057E-03 1.848E-03 1.510E-03 1.150E-03	
GROUP CONSTAN FOR MF 6 AND	NTS AT T=3.000E+02 DEG K MT 53 CONTINUED LR 23 PARTICLE EMISSION	131.5615
INITL FINAL GROUP GROUP	GROUP CONSTANTS VS LEGENDRE DRDER O 1 2' 3	
27 7 27 8 27 9	2.360E-09 -1.174E-09 -1.905E-12 2.856E-10 1.078E-08 -5.320E-09 -4.813E-11 1.293E-09 4.958E-08 -2.419E-08 -4.647E-10 5.805E-09	
30 23 30 24 30 25	2.763E-03 -1.558E-03 1.204E-04 5.831E-04 3.533E-02 1.038E-02 -2.794E-03 -1.265E-03 5.313E-03 4.335E-03 2.805E-03 1.320E-03	
GROUP CONSTAI FOR MF 6 AND	NTS AT T=3.000E+02 DEG K MT 64 CONTINUED LR 23 PARTICLE EMISSION	138.1165
INITL FINAL GROUP GROUP	GROUP CONSTANTS VS EGENDRE DRDER O 1 2 3	
30 8 30 9 30 10 30 11 30 12 30 13 30 14 30 15 30 16 30 17 30 18 30 19	1.481E-09 -7.357E-10 -1.898E-12 1.783E-10 6.805E-09 -3.351E-09 -3.229E-11 8.018E-10 3.171E-08 -1.541E-08 -3.062E-10 3.594E-09 1.501E-07 -7.137E-08 -2.527E-09 1.586E-08 7.416E-07 -3.399E-07 -2.039E-08 6.878E-08 3.848E-06 -1.655E-06 -1.628E-07 2.743E-07 2.167E-05 -8.291E-06 -1.302E-06 8.066E-07 1.040E-04 -1.647E-05 -2.358E-05 4.000E-06 1.099E-04 2.765E-05 -3.111E-05 -1.994E-05 1.763E-04 1.009E-04 1.164E-05 -3.152E-05 1.710E-04 1.396E-04 8.962E-05 3.949E-05 8.226E-06 7.975E-06 7.491E-06 6.807E-06	
THRESHOLD IS	ABOVE HIGHEST ENERGY BOUND FOR MT 65	
THRESHOLD IS	ABOVE HIGHEST ENERGY BOUND FOR MT 66	
THRESHOLD IS	ABOVE HIGHEST ENERGY BOUND FOR MT 67	
THRESHOLD IS	ABOVE HIGHEST ENERGY BOUND FOR MT 68	

GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 6 AND MT 91 CONTINUUM INELASTIC LR 23 PARTICLE EMISSION

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INITL	FINAL	ISOTROPIC	MATRIX V.S F	INAL GROUP			
GROUP	GROUP	+0	+1	+2	+3		
26	9	2.693E-09	1.989E-08	1.216E-07	8.810E-07	5.537E-06	3.511E-05
26	15	1.827E-04	2.303E-04	3.441E-04	3.511E-04	1.903E-04	2.577E-05
26	21	3.463E-06					
27	8	1.591E-09	1.175E-08	8.680E-08	6.348E-07	4.644E-06	3.304E-05
27	14	2.215E-04	1.241E-03	1.682E-03	2.714E-03	3.122E-03	2.142E-03
27	20	4.740E-04	1.868E-04	4.606E-05	5.244E-06	1.042E-07	
28	8	2.577E-09	1.903E-08	1.4068-07	1.028E-06	7.522E-06	5.352E-05
28	14	3.587E-04	2.010E-03	2.7252-03	4.395E-03	5.056E-03	3.469E-03
28	20	7.677E-04	3.030E-04	8.092E-05	1.325E-05	1.132E-06	
29	8	3.345E-09	2.470E-08	1.825E-07	1.335E-06	9.764E-D6	6.947E-05
29	14	4.656E-04	2.608E-03	3.537E-03	5.705E-03	6.563E-03	4.503E-03
29	20	9.964E-04	3.933E-04	1.050E-04	1.720E-05	1.558E-06	
30	8	4.882E-09	3.605E-08 .	2.663E-07	1.948E-06	1.425E-05	1.014E-04
30	14	6.795E-04	3.806E-03	5.161E-03	8.325E-03	9.577E-03	6.571E-03
30	20	1.454E-03	5.740E-04	1.533E-04	2.510E-05	2.274E-06	1.254E-09

GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 6 AND MT221 FREE THERMAL SCATTERING

138.7115

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INITL	FINAL	GROUP CONS	TANTS VS LE	GENORE ORDE	R
GRUUP	GRUUP	0	1	2	3
1	1	8.063E+00	2.232E-01	3.138E-02	7.594E-02
1	2	3.241E-02	-1.425E-03	-1.588E-03	-3.647E-04
1	3	1.677E-08	-1.282E-08	7.246E-09	-2.772E-09
2	1	7.423E-01	-1.557E-01	-2.514E-02	-2.682E-03
2	2	3.962E+00	4.477E-01	2.841E-02	7.789E-02
2	3	5.632E-02	8.940E-03	-1.716E-03	-8.155E-04
3	1	8.166E-05	-6.612E-05	4.254E-05	-2.083E-05
3	2	7.281E-01	-1.809E-01	-1.947E-02	2.330E-03
3	3	4.007E+00	4.726E-01	1.381E-02	9.593E-02
3	4	1.153E-02	6.297E-03	2.210E-03	9.408E-04
4	3	7.408E-01	-1.995E-01	-1.954E-02	1.771E-03
4	4	3.999E+DO	4.982E-01	2.364E-02	7.423E-02
4	5	1.841E-03	1.514E-03	1.030E-03	5.977E-04

---WARNING FROM BINA---DISC= -3.0170E-14. SET TO ABS VALUE AND CONTINUE.

GROUP	CD	NSTA	NTS AT	T=3.000E+	HO2 DEG K		
FOR M	F 6	AND	MT229	GRAPHITE	INELASTIC	THERMAL	SCATTERING

INITL GROUP	F I NAL GROUP	GROUP CONS	TANTS VS LE	GENORE DROI	ER 3
1	1	6.259E-01	-9.509E-02	-1.110E-02	-3.864E-03
1	2	1.402E-02	-3.024E-03	-5.689E-04	-1.220E-04
1	3	1.159E-06	-3.364E-07	2.09DE-08	-8.742E-09
2	1	4.251E-01	-1.161E-01	-1.290E-02	-4.055E-03
2	2	2.023E+00	-3.631E-01	-1.554E-01	-3.522E-02
2	3	3.446E-02	-3.703E-03	-3.396E-03	-9.096E-04
3	1	1.049E-02	-4.201E-03	6.778E-04	-1.779E-D4
3	2	6.057E-01	-1.672E-01	-2.143E-02	-5.394E-03
3	3	3.135E+00	-1.797E-01	-2.928E-01	-1.046E-01
3	4	9.094E-03	2.229E-03	-7.498E-04	-6.540E-04
4	1	3.527E-08	-3.201E-08	2.631E-08	-1.948E-08
4	2	6.161E-05	-5.000E-05	3.255E-05	-1.638E-05
4	3	6.864E-01	-1.951E-01	-2.065E-02	-3.238E-03
4	4	3.571E+00	1.648E-01	-2.419E-01	-1.212E-01
4	5	1.822E-03	1.175E-03	4.590E-04	6.192E-05

INITL FINAL GROUP CONSTANTS VS LEGENDRE DRDER GROUP GROUP 0 1 2 3 1 1 2.714E+00 -1.774E+00 -1.158E-01 -3.057E+01 2 2 2.175E+00 1.537E+00 2.702E+01 2.2418E+02 3 3 9.166E+01 5.653E+01 2.540E+01 8.639E+02 4 4 3.391E+01 1.064E+01 7.835E+02 5.067E+02 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF17 ANO MT 51 INELASTIC GAMMA PRODUCTION INITL FINAL GROUP CONSTANTS VS LEGENDRE DRDER GROUP GROUP 0 1 2 3 24 7 5.168E+02 0. 1.338E+02 0. 25 7 2.465E+01 0. 2.272E+02 0. 25 7 2.465E+01 0. 3.345E+02 0. 27 7 3.115E+01 0. 5.306E+02 0. 28 7 1.850E+01 0. 2.532E+02 0. 29 7 1.850E+01 0. 2.532E+02 0. 30 7 1.354E+01 0. 1.647E+02 0. GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF16 AND MT102 CAPTURE GAMMA PRODUCTION INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP GROUP GROUP +0 +1 +2 +3 1 4 4.065E+03 0. 4.065E+03 8.638E+03 2 4 3.504E+04 0. 3.504E+04 4.509E+04 3 4 2.122E+04 0. 2.122E+04 4.509E+04 3 4 2.122E+04 0. 2.122E+04 4.509E+04 3 4 2.122E+04 0. 2.122E+04 4.509E+04 3 6 6.118E+05 0. 0. 0. 0. 0. 30 12 1.452E+05 0. 0. 0. 0. 0. 180.7235 MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY OUTPUT TAPE 180.7495 TAPE LABEL PENOF TAPE FOR C FROM ENDF/B TAPE 511 10 10 0 0 0 0 0 01306 1451 1 PENOF TAPE FOR C FROM ENDF/B TAPE 511 10 0 0 0.00000+ 0.000000+ 0 0 0 0 0 01306 1451 1 10.0000E+0 0.00000+0 0 0 0 0 0 01306 1451 1 10.0000E+0 0.00000+0 0 0 0 0 0 0 01306 1451 1 10.0000E+0 0.00000+0 0 0 0 0 0 01306 1451 1 10.0000E+0 0.00000+0 0 0 0 0 0 0 01306 1451 1 10.0000E+0 0 0.00000+0 0 0 0 0 0 0 0 0 0 01306 1451 1 100000E+0 0.00000+0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	GROUP	CONSTAN 6 AND	ITS AT T=3.0 MT230 GRAPH	DOE+O2 DEG ITE ELASTIC	K THERMAL SC	ATTERING		175.1285	
1 1 2.714E+00 -1.774E+00 2.702E-01 -2.418E-02 3 3 9.166E-01 5.632E-01 2.418E-02 4 4 3.391E-01 1.064E-01 7.853E-02 5.067E-02 GROUP CONSTANTS AT T=3.000E+02 DEG K 178.9365 178.9365 INITL FINAL GROUP CONSTANTS VS LEGENDRE DRDER GROUP GROUP 0 2 3 24 7 5.168E-02 0. 1.38E-02 0. 25 7 2.468E-01 0. 2.122E-02 0. 26 7 2.199E-01 0. 3.345E-02 0. 27 7 3.195E-01 0. 3.345E-02 0. 29 7 1.85E-01 0. 2.53E-02 0. 29 7 1.85E-01 0. 1.647E-02 0. 29 7 1.85E-01 0. 1.647E-02 0. 1011L FINAL GROUP 400 +0 +1 +2 +3 11 4 4.065E-03 0. 0. 0. 0. 11 4 4.052E-05 0.<	INITL GROUP	F INAL GROUP	GROUP CONS	TANTS VS LE 1	GENDRE ORDE 2	R 3			
GROUP CONSTANTS AT T=3.000E+02 DEG K FOR WE17 AND WT 51 INELASTIC GAMMA PRODUCTION 178.9365 INITL FINAL GROUP GROUP 0 1 2 3 24 7 5.168E-02 0. 2.372E-02 0. 25 7 2.465E-01 0. 2.272E-02 0. 27 7 3.115E-01 0. 5.306E-02 0. 28 7 2.155E-01 0. 2.532E-02 0. 29 7 1.354E-01 0. 1.647E-02 0. 30 7 1.354E-01 0. 1.647E-02 0. GROUP CONSTANTS AT T=3.000E+02 DEG K 179.4055 179.4055 FOR MF16 AND MT 102 CAPTURE GAMMA PRODUCTION 179.4055 INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP GROUP 40 +1 +2 +3 1 4 4.055E-03 0. 0. 0. 0. 29 6 4.025E-05 0. 0. 0. 0. 0. 29 1 2.58E-04 0. 0. 0. 0. 0. 0. 30 1 1.258E-05	1 2 3 4	1 2 3 4	2.714E+00 2.175E+00 9.166E-01 3.391E-01	-1.774E+00 1.537E+00 5.653E-01 1.064E-01	-1.156E-01 2.702E-01 2.540E-01 7.853E-02	-3.067E-01 -2.418E-02 8.639E-02 5.067E-02			
INITL FINAL GROUP CONSTANTS VS LEGENDRE DROER GROUP GROUP 0 1 2 3 24 7 5.168E-02 0. 1.388E-02 0. 25 7 2.468E-01 0. 4.109E-02 0. 26 7 3.119E-01 0. 5.306E-02 0. 27 7 3.115E-01 0. 5.306E-02 0. 28 7 2.155E-01 0. 3.345E-02 0. 29 7 1.850E-01 0. 2.532E-02 0. 30 7 1.354E-01 0. 1.647E-02 0. GROUP CONSTANTS AT T=3.000E+02 DEG K 179.405S 179.405S FOR MFI6 AND MT102 CAPTURE GAMMA PRODUCTION 179.405S INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP 179.405S GROUP GROUP +0 +1 +2 +3 1 4 .065E-03 0.638E-03 0. 2 4 .504E-04 0. 2.122E-04 0. 30 6 6.118E-05 0. 0. 0. 0. 10 6 6.118	GROUP FOR MF	CONSTAN 17 AND	ITS AT T=3.0 MT 51 INELA	OOE+O2 DEG STIC GAMMA	K PRODUCTION			178.9365	
24 7 5.168E-02 0. 1.388E-02 0. 25 7 2.465E-01 0. 2.272E-02 0. 26 7 3.195E-01 0. 4.109E-02 0. 27 7 3.115E-01 0. 5.306E-02 0. 28 7 1.850E-01 0. 2.523E-02 0. 29 7 1.850E-01 0. 2.523E-02 0. 300 7 1.354E-01 0. 1.647E-02 0. INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP GROUP GROUP +0 +1 +2 +3 1 4 4.065E-03 0. 4.065E-03 8.638E-03 2 4 3.504E-04 0. 2.122E-04 4.509E-04 3 4 2.122E-04 0. 0. 0. 0. 29 6 4.025E-05 0. 0. 0. 0. 0. 30 12 1.452E-04 0. 0. 0. 0. 0. 10012 1.452E-04 0. 0. 0.	INITL GROUP	FINAL GROUP	GRDUP CONS	TANTS VS LE 1	GENDRE DRDE 2	R 3			
GRDUP CONSTANTS AT T=3.000E+02 DEG K FOR MF16 AND MT102 CAPTURE GAMMA PRODUCTION 179.405S INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP GROUP GROUP +0 +1 +2 +3 +3 1 4 4.065E-03 0. 4.065E-03 8.638E-03 2 4 3.504E-04 0. 3.504E-04 7.446E-04 3 4 2.122E-04 0. 2.122E-04 4.509E-04 -3 29 6 4.025E-05 0. 0. 0. 0. 0. 0. 0. 29 12 1.258E-04 30 6 6.118E-05 0. 0. 0. 0. 0. 0. 30 12 1.452E-04 0. 0. 0. 180.7235 MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE DUTPUT UNIT (+ FOR CODED FOR BB)23 DUTPUT UNIT (+ FOR CODED FOR BB) 25 -23 74PE LABEL 	24 25 26 27 28 29 30	7 7 7 7 7 7 7	5.168E-02 2.465E-01 3.199E-01 3.115E-01 2.155E-01 1.850E-01 1.354E-01	0. 0. 0. 0. 0. 0. 0.	1.388E-02 2.272E-02 4.109E-02 5.306E-02 3.345E-02 2.523E-02 1.647E-02	0. 0. 0. 0. 0. 0.			
INITL FINAL ISOTROPIC MATRIX VS FINAL GROUP GROUP GROUP +0 +1 +2 +3 1 4 4.065E-03 0. 4.065E-03 8.638E-03 2 4 3.504E-04 0. 3.504E-04 7.446E-04 3 4 2.122E-04 0. 2.122E-04 4.509E-04 29 6 4.025E-05 0. 0. 0. 0. 0. 0. 29 12 1.258E-04 30 6 6.118E-05 0. 0. 0. 0. 0. 0. 30 12 1.452E-04 MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE 180.749S INPUT UNIT (+ FOR CODED FOR BB)23 DUTPUT UNIT (+ FOR CODED FOR BB) 25 TAPE LABEL 	GROUP FOR MF	CONSTAN 16 AND	ITS AT T=3.0 MT102 CAPTU	OOE+O2 DEG Re gamma Pr	K DDUCTION			179.4055	
1 4 4.065E-03 0. 4.065E-03 8.638E-03 2 4 3.504E-04 0. 3.504E-04 7.446E-04 3 4 2.122E-04 0. 2.122E-04 4.509E-04 	INITL GROUP	F INAL GROUP	ISOTROPIC +0	MATRIX VS F +1	INAL GROUP +2	+3			
29 6 4.025E-05 0. 0. 0. 0. 0. 29 12 1.258E-04 0. 0. 0. 0. 30 6 6.118E-05 0. 0. 0. 0. 0. 30 12 1.452E-04 0. 0. 0. 0. 0. MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE 180.723S INPUT UNIT (+ FOR CODED, - FOR BB) -23 DUTPUT UNIT (+ FOR CODED, - FOR BB) 25 25 TAPE LABEL PENOF TAPE FOR C FROM ENDF/B TAPE 511 190.553S PENOF TAPE FOR C FROM ENDF/B TAPE 511 10 0 0 6 0 0 0 0 0 0 100 0 0 0 0 0 0	1 2 3	4 4 4	4.065E-03 3.504E-04 2.122E-04	0. 0. 0.	4.065E-03 3.504E-04 2.122E-04	8.638E-03 7.446E-04 4.509E-04			
30 6 6.118E-05 D. O. O. O. O. 30 12 1.452E-04 180.7235 MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE 180.7235 INPUT UNIT (+ FOR CODED, - FOR BB) -23 -23 DUTPUT UNIT (+ FOR CODED, - FOR BB) 25 -25 TAPE LABEL -25 190.5535 PENOF TAPE FOR C FROM ENDF/B TAPE 511 190.5535 PENOF TAPE FOR C FROM ENDF/B TAPE 511 10 0 6.00000+ 3 1.18969+ 1 0 0 01306 1451 1	29 29	6 12	4.025E-05 1.258E-04	0.	0.	0.	0.	0.	
180.7235 MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE 180.7495 INPUT UNIT (+ FOR CODED, - FOR BB) -23 DUTPUT UNIT (+ FOR CODED, - FOR BB) 25 TAPE LABEL -25 PENDF TAPE FOR C FROM ENDF/B TAPE 511 190.5535 PENDF TAPE FOR C FROM ENDF/B TAPE 511 10 0 0 6.00000+ 3 1.18969+ 1 0 0 01306 1451 1 0.00000+ 0 0.00000+ 0 0 0 0 01306 1451 1	30 30	6 12	6.118E-05 1.452E-04	D.	0.	0.	Ο.	0.	
MODERCHANGE THE MODE OF AN ENDF/B TAPE OR NJDY DUTPUT TAPE 180.749S INPUT UNIT (+ FOR CODED, - FOR BB) -23 DUTPUT UNIT (+ FOR CODED, - FOR BB) 25 TAPE LABEL	*****	******	*****	********	********	*********	* * * * * * * * * * * * *	180.7235	
INPUT UNIT (+ FOR CODED, - FOR BB)23 OUTPUT UNIT (+ FOR CODED, - FOR BB) 25 TAPE LABEL PENDF TAPE FOR C FROM ENDF/B TAPE 511 PENDF TAPE FOR C FROM ENDF/B TAPE 511 10 0 0 6.00000+ 3 1.18969+ 1 0 0 01306 1451 1 0.00000+ 0 0.00000+ 0 0 0 0 01306 1451 2	MODER.	CHANG	E THE MODE	OF AN ENDF/	B TAPE OR N	NJOY OUTPUT	ΤΑΡΕ	180.7495	
TAPE LABEL PENDF TAPE FOR C FROM ENDF/B TAPE 511 190.5535 ************************************	INPUT OUTPUT	UNIT (+ UNIT (FOR CODED, + FOR CODED	- FOR BB) FOR BB)	•••	-23 25			
PENDF TAPE FOR C FROM ENDF/B TAPE 511 190.5535 ************************************									
PENDF TAPE FOR C FROM ENDF/B TAPE 511 1 0 0 0 6.00000+ 3 1.18969+ 1 0 0 01306 1451 1 0.00000+ 0 0.00000+ 0 0 0 01306 1451 2	*****	******	*****	NUF/B TAPE	*********	*********	*********	190.553S	
	PENDF T 6.0000 0.0000	APE FOR 0+31. 0+00.	R C FROM END 18969+ 1 .00000+ 0	F/B TAPE 51 D O	1 0 0	0	0130 0130	1 0 0 6 1451 6 1451	0 1 2

.

PENDF TAP	PE FC	R C FROM	E١	IDF/B TAPE 5	511		1 () ()	0
6.00000+	· 3 1	. 18969+	1	0	0	0	01306	1451	1
0.0000+	· o c	.00000+ (D	0	0	0	01306	1451	2
3.00000+	- 25	i.00000- 3	3	0	0	3	391306	1451	3
6-C-NAT F	ROM	TAPE 511					1306	1451	4
PROCESSEC) BY	THE NJOY	NU	JCLEAR DATA	PROCESSING	SYSTEM	1306	1451	5
SEE DRIGI	NAL	ENOF/B-V	ΤI	APE FOR DETA	ILS OF EVAL	LUATION	1306	1451	6
				1	451	43	01306	1451	7
				2	151	4	01306	1451	8
				3	1	292	01306	1451	9
				3	2	292	01306	1451	10
				3	4	139	01306	1451	11
				3	51	139	D1306	1451	12
				3	52	82	01306	1451	13
				3	53	56	01306	1451	14
				3	54	49	01306	1451	15
				3	55	46	01306	1451	16
				3	56	42	01306	1451	17
				3	57	39	01306	1451	18
				3	58	36	01306	1451	19
				3	59	33	01306	1451	20
				3	60	30	01306	1451	21
				3	61	25	01306	1451	22
				3	62	21	01306	1451	23
				3	63	19	01306	1451	24
				3	64	17	01306	1451	25
				3	65	14	01306	1451	26
				3	66	12	01306	1451	27
				3	67	10	01306	1451	28
				3	68	8	01306	1451	29
				3	91	91	01306	1451	30
				3	102	292	01306	1451	31
				3	103	30	01306	1451	32
				3	104	21	01306	1451	33
				3	107	118	01306	1451	34
				3	203	30	01306	1451	35
				3	204	21	01306	1451	36
				3	207	118	01306	1451	37
				3	221	32	01306	1451	38
				3	229	209	01306	1451	39
				3	230	209	01306	1451	40
				3	301	292	0 1306	1451	41
				3	444	292	01306	1451	42
				6	221	4010	01306	1451	43
				6	229	4644	01306	1451	44
				6	230	2	01306	1451	45
					_		1306	1 0	46
							1306	0 0	47
6.00000)+ 3	1,18969+	1	0	0	1	01306	2151	48
6.00000)+ 3	0.00000+	0	Ō	0	1	01306	2151	49
1.00000)- 5	2.00000+	7	0	0	0	01306	2151	50
0.00000)+ 0	6.14112-	1	Ō	0	0	01306	2151	51
							1306	2 0	52
							1306	0 0	53
6.00000)+ 3	1.18969+	1	0	99	0	01306	3 1	54
3.00000)+ 2	0.00000+	Ó	ō	0	1	8661306	3 1	55
2.30000	866		2	Ū	-		1306	3 1	56
1.00000	0-5	7,911976	+1	1.193350-5	7.244845+1	1.386710-5	6.722771+11306	3 1	57
1.58007	70-5	6.299862	+1	1.773430-5	5.948262+1	2.160150-5	5.392759+11306	3 1	58
2.54687	70-5	4.969414	+1	2.933590-5	4.633020+1	3.320310-5	4.357423+11306	3 1	59
4.09375	50-5	3.928872	+1	4.867180-5	3.607434+1	5.640620-5	3.354913+11306	3 1	60
•	-								
•									
6.0000) + 3	1.18969+	1	0	0	0	01306	3229	2238
3.0000)+ 2	0.00000+	0	0	0	1	6181306	3229	2239
	618		2				1306	3229	2240

1.000000-5 3.051823+0 4.555443-4 4."33945-1 4.555535-4 4.733903-11306 3229 2241 1.822177-3 2.423495-1 1.822214-3 2.423479-1 2.391635-3 2.228712-11306 3229 2242 2.961056-3 2.107348-1 4.099899-3 1.979619-1 4.099981-3 1.979613-11306 3229 2243 4.515467-3 1.957387-1 4.515557-3 1.957384-1 4.971011-3 1.941680-11306 3229 2244 6.00000+ 3 1.18969+ 1 0 0 01306 3230 2448 3.00000+ 2 0.00000+ 0 0 O 6181306 3230 2449 618 2 1306 3230 2450 1.000000-5 0.000000+0 4.555443-4 0.000000+0 4.555535-4 0.000000+01306 3230 2451 1.822177-3 0.000000+0 1.822214-3 7.392451+0 2.391635-3 5.632392+01306 3230 2452 2.961056-3 4.549263+0 4.099899-3 3.285599+0 4.099981-3 3.285534+01306 3230 2453 4.515467-3 2.983219+0 4.515557-3 3.330007+0 4.971011-3 3.024905+01306 3230 2454 4.971110-3 4.819179+0 6.337644-3 3.780059+0 6.337771-3 4.190413+01306 3230 2455 6.00000+ 3 1.18969+ 1 01306 3301 2658 0 0 0 3.00000+ 2 0.00000+ 0 8661306 3301 2659 0 0 866 1306 3301 2660 2 1.000010-5 1.513444+2 1.193350-5 1.385512+2 1.386710-5 1.285330+21306 3301 2661 1.580070-5 1.204165+2 1.773430-5 1.136650+2 2.160150-5 1.029944+21306 3301 2662 2.546870-5 9.485684+1 2.933590-5 8.838563+1 3.320310-5 8.308118+11306 3301 2663 4.093750-5 7.482474+1 4.867180-5 6.862420+1 5.640620-5 6.374702+11306 3301 2664 6.414060-5 5.978092+1 7.187500-5 5.647769+1 8.734350-5 5.123677+11306 3301 2665 6.00000+ 3 1.18969+ 1 01306 3444 2951 0 0 0 3.00000+ 2 0.00000+ 0 0 8661306 3444 2952 O 1306 3444 2953 866 2 1.000010-5 1.142386+2 1.193350-5 1.045362+2 1.386710-5 9.697747+11306 3444 2954 1.580070-5 9.085359+1 1.773430-5 8.575963+1 2.160150-5 7.770867+11306 3444 2955 2.546870-5 7.156891+1 2.933590-5 6.668641+1 3.320310-5 6.268422+11306 3444 2956 4.093750-5 5.645475+1 4.867180-5 5.177646+1 5.640620-5 4.809663+11306 3444 2957 6.00000+ 3 1.18969+ 1 O 5 0 01306 6229 7256 3.00000+ 2 0.00000+ 0 0 0 521306 6229 7257 1 52 1306 6229 7258 2 0.00000+ 0 1.00000- 5 0 450 101306 6229 7259 0 0.000000+0 0.000000+0 0.000000+0 0.000000+0 0.000000+0 0.000000+01306 6229 7260 0.00000+0 0.00000+0 0.00000+0 0.00000+0 1.081430-4 0.00000+01306 6229 7261 0.000000+0 0.000000+0 0.000000+0 0.000000+0 0.000000+0 0.000000+01306 6229 7262 0.000000+0 0.000000+0 2.091463-4 1.114684-1-9.050929-1-7.089504-11306 6229 7263 -5.020376-1-2.821791-1-4.633196-2 2.100009-1 4.940009-1 8.185834-11306 6229 7264 4.111530-4 2.668334-1-8.984247-1-6.896068-1-4.713482-1-2.421188-11306 6229 7265 6.00000+ 3 1.18969+ 1 01306 623011901 O 6.00000+ 3 1.18969+ 1 0 0 0 3141306 623011902 1306 6 011903

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II. EXAMPLE 2, CCCC CRDSS SECTION LIBRARY MOUNT ENDF/B-IV TAPE 404 ON UNIT 20. 0 4 *MODER* 20 -21 *RECONR* -21 -22 *PENDF TAPE FOR PU-238 FROM ENDF/B-IV TAPE 404*/ 1050 3 0 .005 0 6 / *94-PU-238 FROM ENDF/B TAPE T404*/ *PROCESSED BY THE NJOY NUCLEAR DATA PROCESSING SYSTEM*/ *SEE ORIGINAL ENDF/B-IV TAPE FOR DETAILS OF EVALUATION*/ 0/ *BRDADR* -22 -23 1050 3 0 1 0 .002 1.E6 300. 900. 2100. 0/ *UNRESR* -21 -23 -22 1050 3 7 1 300 900 2100 1.E10 1.E5 1.E4 1000. 100. 10. 1 0/ *GROUPR* -21 -22 0 -25 1050 5 0 4 3 3 7 1 *94-PU-238*/ 300. 900. 2100. 1.E10 1.E5 1.E4 1000. 100. 10. 1 .1 0.025 0.8208E06 1.4E06 3 1 *TOTAL*/ 3 2 *ELASTIC*/ 3 16 *N2N*/ 3 17 *N3N*/ 3 18 *FISSIDN*/ 3 102 *CAPTURE*/ 3 251 *MUBAR*/ 3 252 *XI*/ 3 253 *GAMMA*/ 3 259 *1/V*/ 6 2 *ELASTIC*/ 6 16 *N2N*/ 6 17 *N,3N*/ 6 18 *FISSION*/ 6 51 *DISCRETE INELASTIC*/ 6 -59 *CONTINUED*/ 6 91 *CONTINUUM INELASTIC*/ 0/ 3 1 *TDTAL*/ 3 2 *ELASTIC*/ 3 18 *FISSION*/ 3 102 *CAPTURE*/ 6 2 *ELASTIC*/ 0/ 3 1 *TOTAL*/ 3 2 *ELASTIC*/ 3 18 *FISSION*/ 3 102 *CAPTURE*/ 6 2 *ELASTIC*/ 0/ ō/ *CCCCR* -25 21 22 0 1 1 *T2LANL NJOY*/

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CCCCR TESTS DCT, 1981/ 50 0 1 4 *PU238* *PU238* *ENDFB4* * 1050 * 1050 10.89 1 0 50 -1 0 2.3821E02 3.3003E-11 1.7461E-12 0. 1.E10 0.0 3 6 300 900 2100 1.65 1.64 1000. 100. 10. 1 *STOP* 1 * * * * * * * * * * * \$\$ \$\$ \$\$ \$\$\$\$\$ \$\$ 22 * VERS. 10/81 * \$\$\$ \$\$ 55 \$5 \$\$\$\$\$\$\$ NUCLEAR \$\$ * * RAN AT LANL * \$\$ \$\$\$\$ \$\$ \$\$\$\$ \$\$ \$\$ * CRDSS SECTION * DN MACH, U \$\$ * DN 11/06/81 * AT 18:33:09 \$\$ \$\$\$\$ \$\$ \$\$ \$\$ \$\$ * PRDCESSING * \$\$\$\$\$\$ \$\$ \$\$\$ \$\$\$\$\$\$\$ \$5 * SYSTEM \$\$ \$£ 22222 222222 \$\$ * * * **** ******** MODER...CHANGE THE MODE OF AN ENDF/B TAPE OR NJOY DUTPUT TAPE .6295 INPUT UNIT (+ FOR CODED, - FOR BB) ... DUTPUT UNIT (+ FOR CODED, - FOR EB) ... 20 -21 TAPE LABEL ENDF/B-IV TAPE 404 (REV. 3) 1-SEPT-76 7.9625 *********** RECONR...RECONSTRUCT POINTWISE CROSS SECTIONS IN PENDE FORMAT 8.0025 UNIT FOR ENDF/B TAPE -21 UNIT FOR PENOF TAPE - 22 LABEL FOR PENOF TAPE PENDE TAPE FOR PU-238 FROM ENDE/B-IV TAPE 404 TAPE LABEL ENDF/B-IV TAPE 404 (REV. 3) 1-SEPT-76 MATERIAL TO BE PROCESSED 1050 .005 0.K NO. SIGNIFICANT FIGURES 6 RESONANCE-INTEGRAL-CHECK TOLERANCE 100 MAX RESONANCE-INTEGRAL ERROR 5.000E-07 DESCRIPTIVE CARDS FOR PENDE TAPE 94-PU-238 FROM ENDF/B TAPE T404 PROCESSED BY THE NJDY NUCLEAR DATA PROCESSING SYSTEM SEE ORIGINAL ENDF/B-IV TAPE FOR DETAILS OF EVALUATION PRDCESSING MAT 1050 94-PU-238 AI EVAL-MAY67 ALTER AND DUNFORD

PDINTS IN INITIAL UNIONIZED GRID = 111
PDINTS ADDED BY LINEARIZATION = 338	10.002\$
ESTIMATED MAXIMUM ERROR DUE TO RESONANCE INTEGRAL CHECK (ERRMAX,ERRINT) AND SIGNIFICANT FIGURE TRUNCATION (NDIGIT)	
UPPER ELASTIC PERCENT ERROR CAPTURE PERCENT ERROF ENERGY INTEGRAL RES-INT SIG-FIG INTEGRAL RES-INT SIG-FI 1.00E+00	R LG
3.17E+00 1.11E+01 0.000 0.000 3.62E+01 0.000 0.000 1.04E+01 1.16E+01 0.000 0.000 7.49E+00 0.000 0.000 3.92E+01 1.65E+01 .000 .000 3.68E+01 .000 .000 1.24E+02 2.23E+01 .000 .000 2.19E+01 .000 .000 2.00E+02 1.61E+01 .000 .000 7.93E+00 .000 .000 1.00E+04 1.07E+02 0.000 .000 1.96E+01 0.000 .000	
POINTS ADDED BY RESONANCE RECONSTRUCTION = 2703 POINTS AFFECTED BY RESONANCE INTEGRAL CHECK = 547 POINTS AFFECTED BY SIGNIFICANT FIGURE REDUCTION = 73 POINTS REMOVED BY BACKTHINNING = 569 FINAL NUMBER OF RESONANCE POINTS = 2753	20, 4545
****	****
BROADRDDPPLER BROADENING OF ENDF/B DATA	20.4935
UNIT FOR INPUT PENDF TAPE-22UNIT FOR OUTPUT PENDF TAPE-23MATERIAL TO BE PROCESSED1050NUMBER OF FINAL TEMPERATURES3RESTART (O ND. 1 YES)0BODTSTRAP (O ND. 1 YES)1STARTING MATERIAL TEMPERATURE0.KTHINNING TOLERANCE.002MAX. ENERGY1.00E+06FINAL TEMPERATURES3.00E+029.00E+022.10E+03	22.9685
BRDADENED MAT1050 FRDM O. TD 3.0000E+02 K POINTS IN≖ 3136 PDINTS JUT= 2323 MT 2 18 102	26 1965
BRDADENED MAT1050 FROM. 3.0000E+02 TO 9.0000E+02 K PDINTS IN= 2323 POINTS OUT= 2067 MT 2 18 102	50.1505
BROADENED MAT1050 FRDM 9.0000E+02 TD 2.1000E+03 K PDINTS IN= 2067 PDINTS OUT= 1822 MT 2 18 102	41.7725
M1 Z 10 VZ	46.4235
UNRESRCALCULATION OF UNRESOLVED RESONANCE CROSS SECTIONS	46.4605
UNIT FOR INPUT ENDF/B TAPE	

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		1.00 1.00 1.00	E+03 E+02 E+01		
PRINT OPTION (O MIN	I MAX.)		1		
MAT = 1050 TEMP = 3	3.00E+02				46.9535
6.538E+01 6.442E+01	5.780E+01	3.715E+01	2.018E+01	1.492E+01	1.402E+01
4.484E+01 4.414E+01	3.937E+01	2.559E+01	1.569E+01	1.292E+01	1.246E+01
3.582E+00 3.537E+00	3.211E+00	2.004E+00	7.677E-01	3.314E-01	2.553E-01
1.687E+01 1.666E+01	1.513E+01	9.466E+OO	3.633E+00	1.570E+00	1.211E+00
6.538E+01 6.349E+01	5.185E+O1	2.683E+01	1.520E+01	1.284E+01	1.248E+01
ENERGY = 5.0000E+02					
4.5412+01 4.5162+01	4.318E+01	3.351E+01	2.037E+01	1.506E+01	1.409E+01
3.4302+01 3.4112+01	3.2561+01	2.5301+01	1.6361+01	1.314E+01	1.2581+01
7 9525400 7 9155400	2.9/30+00	2.2852+00	1.0885+00	4.9036-01	3.8112-01
A 541E+01 A 492E+01	4 1195+01	2 7135+01	1 5625+01	1 2845+01	1 2435+01
FNFRGY = 6.4000F+02	4.1152.01	2.7102.01	1.5022.01	1.2042.01	1.2452.01
4.143E+01 4.126E+01	3.985E+01	3.226E+01	2.037E+01	1.511E+01	1.413E+01
3.210E+01 3.196E+01	3.084E+01	2.497E+01	1.654E+01	1.322E+01	1.263E+01
2.734E+00 2.723E+00	2.638E+00	2.119E+00	1.079E+00	5.065E-01	3.899E-01
6.434E+00 6.410E+00	6.214E+00	5.012E+00	2.584E+00	1.229E+00	9.509E-01
4.143E+01 4.109E+01	3.841E+01	2.690E+01	1.577E+01	1.286E+01	1.243E+01
ENERGY = 8.1920E+02					
3.791E+01 3.779E+01	3.680E+01	3.094E+01	2.034E+01	1.518E+01	1.418E+01
3.004E+01 2.995E+01	2.914E+01	2.449E+01	1.670E+01	1.330E+01	1.268E+01
2.4232+00 2.4162+00	2.35/1400	1.9/02+00	1.0/51+00	5.2211-01	4.040E-01
3.2012+00 5.2402+00	3.1226+00	4.2986+00	2.3/96+00	1.1762+00	a 1285-01
3./912+0: 3./0/2+01	3.3772401	2.0526+01	1.3546+01	1.2906+01	1.2446701
•					
ENERGY = 1.0000E+04					
1.893E+01 1.893E+01	1.891E+01	1.871E+01	1.749E+01	1.546E+01	1.474E+01
1.623E+01 1.623E+01	1.621E+01	1.603E+01	1.498E+01	1.330E+01	1.274E+01
1.364E+00 1.364E+00	1.362E+00	1.3458+00	1.232E+00	9.902E-01	8.829E-01
7.047E-01 7.047E-01	7.039E-01	6.9679-01	6.481E-01	5.404E-01	4.901E-01
1.893E+01 1.893E+01	1.888E+01	1.848E+01	1.6421+01	1.381E+01	1.304E+01
GENERATED CRUSS SECTIO	NS AT 15 PU	INIS			50.4345
MAT = 1050 TEMP =	9.00E+02				56.4345
ENERGY = 2.0000E+02					
6.538E+01 6.472E+01	5.986E+01	4.128E+01	2.214E+01	1.557E+01	1.445E+01
4.484E+01 4.435E+01	4.079E+01	2.805E+01	1.660E+01	1.317E+01	1.261E+01
3.5822+00 3.5522+00	3.323E+00	2.2941+00	9.4772-01	4.000E-01	3.019E-01
	1.3000+01	1.0851+01	4.5032+00	1.9065+00	1.4402+00
0.5382+01 0.4082+01	5.5252+01	3.0982+01	1.5972401	1.2902+01	1.2546+01
•					
ENERGY = 1.0000E+04					
1.893E+01 1.893E+01	1.892E+01	1.379E+01	1.798E+01	1.636E+01	1.570E+01
1.623E+01 1.623E+01	1.622E+01	1.611E+01	1.539E+01	1.399E+01	1.345E+01
1.364E+00 1.364E+00	1.363E+00	1.354E+00	1.292E+00	1.135E+00	1.055E+00
7.047E-01 7.047E-01	7.043E-01	7.008E-01	6.749E-01	6.077E-01	5.714E-01
1.893E+01 1.893E+01	1.890E+01	1.865E+01	1.718E+01	1.476E+01	1.386E+01
GENERATED CRUSS SECTIO	INS AL 15 PC	11012			64.4515
MAT = 1050 TEMP =	2.10E+03				64.4515
65285+01 - 64045+02	6 1205-04	4 5005104	3 4405104	1 6405104	
4.484E+01 4.449E+01	4.180E+01	4.503E+01 3.041E+01	2.448E+01 1.777E+01	1.355E+01	1.287E+01

3.582E+00 3.561E+00 3.396E+00 2.537E+00 1.150E+00 4.925E-01 3.683E-01 1.687E+01 1.677E+01 1.600E+01 1.200E+01 5.473E+00 2.354E+00 1.763E+00 6.538E+01 6.445E+01 5.771E+01 3.469E+01 1.713E+01 1.320E+01 1.268E+01 ENERGY = 1.0000E+041.893E+01 1.893E+01 1.892E+01 1.885E+01 1.833E+01 1.719E+01 1.671E+01 1.623E+01 1.623E+01 1.623E+01 1.616E+01 1.568E+01 1.464E+01 1.421E+01 1.364E+00 1.364E+00 1.364E+00 1.360E+00 1.331E+00 1.258E+00 1.220E+00 7.047E-01 7.048E-01 7.046E-01 7.031E-01 6.923E-01 6.633E-01 6.469E-01 1.893E+01 1.893E+01 1.891E+01 3.876E+01 1.777E+01 1.583E+01 1.497E+01 GENERATED CROSS SECTIONS AT 15 POINTS 72.2885 GROUPR...COMPUTE SELF-SHIELDED GROUP-AVERAGED CROSS-SECTIONS 72.3455 UNIT FOR ENOF/B TAPE -21 UNIT FOR PENDF TAPE UNIT FOR INPUT GOUT TAPE -22 0 UNIT FOR DUTPUT GOUT TAPE - 25 MAT TO BE PROCESSED 1050 GAMMA GROUP DETION 5 0 WEIGHT FUNCTION OPTION 4 LEGENDRE DRDER .. З
 LEGENORE ORDER

 PRINT OPTION (O MIN, 1 MAX)
 1 RUN TITLE -----94-PU-238 TEMPERATURES (KELVIN) 3.00E+02 9.00E+02 2.10E+03 SIGMA ZERDES INFINITY 1.00E+05 1.00E+04 1.00E+03 1.00E+02 1.00E+01 1.00E+00 NEUTRON GROUP STRUCTURE.....RRD 50 GR/JUP 1. 1.0000E-05 - 6.8256E-01 2 6.8256E-01 - 1.1254E+00 3 1.1254E+00 -1.8554E+00 . 48 3.6788E+D6 -6.0653E+06 6.0653E+06 -49 1.0000E+07 1.0000E+07 -1.9971E+07 50 WEIGHT FUNCTION.....THERMAL + 1/E + FISSIDN THERMAL BREAKPOINT AND TEMPERATURE 1.0000E-01 2.5000E-02 FISSION BREAKPOINT AND TEMPERATURE 8.2080E+05 1.4000E+06 PROCESSING MAT 1050 _____ 94-PU-238 FROM ENDF/B TAPE T404 76.2695 GROUP CONSTANTS AT T=3.000E+02 DEG K

1 3.688E+02 3.677E+02 3.594E+02 3.125E+02 2.184E+02 1.663E+02 1.570E+02 0 FLUX O 5.021E+00 5.003E+00 4.847E+00 3.826E+00 1.577E+00 2.848E-01 3.179E-02 7.300E+01 3.688E+02 3.667E+02 3.506E+02 2.686E+02 1.318E+02 8.010E+01 1 FLUX 1 5.021E+00 3.016E+00 6.803E-01 3.161E-02 4.985E+004.683E+00 4.295E-04 2.472E+01 2 0 2.476E+01 2.476E+01 2.475E+01 2.440E+01 2.330E+01 2.265E+01 FLUX O 5.002E-01 5.000E-01 4.989E-01 4.881E-01 4.021E-01 1.502E-01 2.116E-02 2 2.476E+01 2.476E+01 2.475E+01 2.467E+01 2.402E+01 2.172E+01 2.035E+01 FLUX 1 5.002E-01 4.999E-01 4.977E-01 4.763E-01 3.242E-01 4.735E-02 9.912E-04 . 0 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.879E+00 50 FLUX O 9.012E-03 9.012E-03 9.007E-03 8.959E-03 8.512E-03 5.675E-03 1.310E-03 50 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.879E+00 FLUX 1 9.012E-03 9.011E-03 9.002E-03 8.907E-03 8.039E-03 3.574E-03 1.904E-04 GROUP CONSTANTS AT T=3.000E+02 DEG K 79.9235 FOR MF 3 AND MT 2 ELASTIC GROUP CONSTANTS VS SIGMA ZERD ENDGY GROUP INFINITY 1.000E+05 1.000E+04 1.000E+03 1.000E+02 1.000E+01 1.000E+00 1.748E+01 1.918E+01 1.893E+01 1.817E+01 1.760E+01 1.922E+01 1.921E+01 1.377E+01 1.369E+01 1.325E+01 1.378E+01 1.378E+01 1.342E+01 1.378E+01 2 9.597E+00 9.597E+00 9.597E+00 9.597E+00 9.597E+00 9.595E+00 3 9.594E+00 3.654E+00 3.654E+00 3.654E+00 3.654E+00 3.654E+00 3.654E+00 48 3.653E+00 49 3.115E+00 3.115E+00 3.115E+00 3.115E+00 3.114E+00 3.111E+00 3.106E+00 50 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT 16 N2N 81.8835 ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 49 2.279E-02 1.109E-01 50 GROUP CONSTANTS AT T=3.000E+02 DEG K 82.7595 FOR MF 3 AND MT 17 N3N ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 50 2.902E-03 GROUP CONSTANTS AT T=3.000E+02 DEG K 83.3745 FOR MF 3 AND MT 18 FISSION ENRGY GROUP CONSTANTS VS SIGMA ZERO GROUP INFINITY 1.000E+05 1.000E+04 1.000E+03 1.000E+02 1.000E+01 1.000E+00

1.000E+05 1.000E+04 1.000E+03 1.000E+02 1.000E+01 1.000E+00

FOR MF 3 AND MT 1 TOTAL

INFINITY

GROUP CONSTANTS VS SIGMA ZERD

ENRGY LGEND

ORDER

GROUP

 1.010E+01
 1.007E+01
 9.830E+00
 8.478E+00
 5.772E+00
 4.277E+00
 4.009E+00

 2.991E-01
 2.991E-01
 2.992E-01
 2.917E-01
 2.698E-01
 2.568E-01

 2.819E-02
 2.819E-02
 2.819E-02
 2.819E-02
 2.819E-02
 2.817E-02
 2.810E-02

 1 2 З . . 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.612E+00 2.602E+00 2.605E+00 2.605E+00 2.604E+00 2.604E+00 2.602E+00 2.470E+00 2.470E 48 49 50 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT102 CAPTURE 85.2135 ENRGY GROUP CONSTANTS VS SIGMA ZERO GROUP INFINITY 1.000E+05 1.000E+04 1.000E+03 1.000E+02 1.000E+01 1.000E+00 3.394E+02 3.384E+02 3.304E+02 2.851E+02 1.945E+02 1.445E+02 1.355E+02 1.068E+01 1.068E+01 1.068E+01 1.065E+01 1.042E+01 9.615E+00 9.139E+00 1 1.068E+01 1.068E+01 1.068E+01 1.065E+01 1.042E+01 9.615E+00 9.139E+00 6.354E-01 6.353E-01 6.353E-01 6.353E-01 6.347E-01 6.320E-01 6.296E-01 2 3 . . 1.087E-02 1.087E-02 1.087E-02 1.087E-02 1.087E-02 1.086E-02 1.085E-02 7.328E-03 7.328E-03 7.328E-03 7.328E-03 7.327E-03 7.317E-03 7.303E-03 48 49 5.584E-03 5.584E-03 5.584E-03 5.584E-03 5.584E-03 5.584E-03 5.585E-03 50 GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 3 AND MT251 MUBAR 87, 1955 ENRGY GROUP CONSTANTS AT INFINITE DILUTION GROUP 4 2.823E-03 2 2.823E-03 3 2.823E-03 . 48 8.522E-01 49 8.829E-01 8.714E-01 50 GROUP CONSTANTS AT T=3.000E+02 DEG K 91.1535 FOR MF 3 AND MT252 XI ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 2 1.391E-02 8.964E-03 3 48 2.278E-03 3.907E-03 49 50 6.814E-04 GROUP CONSTANTS AT T∓3.000E+02 DEG K FOR MF 3 AND MT253 GAMMA 94.3205

ENRGY GROUP	GROUP	CONSTAN TE DILL	NTS AT JTION						
2 3	2.644E 3.961E	-01 -02							
48 49 50	5.238E 2.461E 1.066E	-02 -01 -02							
GROUP FOR MF	CDNSTAN 3 AND	TS AT ⁻ MT259	r=3.000E+0 1/V	2 DEG K			97.08	35	
ENRGY GROUP	GROUP INFINI	CONSTANTE DIL	NTS AT JTION						
1 2 3	3.184E 7.731E 6.021E	-04 -05 -05							
48 49 50	3.393E 2.693E 2.142E	-08 -08 -08							
GROUP FOR MF	CONSTAN 6 AND	TS AT MT 2	T=3.000E+0 ELASTIC	DEG K			98.18	35	
INITL GROUP	FINAL GROUP	LGEND ORDER	GRDUP CO INFINITY	NSTANTS VS S 1.000E+05	IGMA ZERD 1.000E+04	1.000E+03	1.000E+02	1.000E+01	1.000E+00
1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	1 1 1 1 1 1 2 2	0 1 2 3 0 1 2 3 0 1	1.922E+0 5.425E-0 6.891E-0 0. 2.565E-0 -3.068E-0 -1.296E-0 1.352E+0 1.239E-0	1.921E+01 2.5.423E-02 5.6.887E-05 0. 1.2.565E-01 0.2.8.498E-02 0.43.066E-04 0.61.291E-06 0.1.352E+01 0.1.239E-01	1.918E+01 5.403E-02 6.849E-05 0. 2.563E-01 -8.484E-02 -3.047E-04 1.241E-06 1.352E+01 2.37E-01	1.893E+01 5.265E-02 6.597E-05 0. 2.544E-01 -8.358E-02 -2.867E-04 -7.816E-07 1.351E+01 1.224F-01	1.817E+01 4.889E-02 6.015E-05 0. 2.392E-01 -7.379E-02 -1.666E-04 1.434E-06 1.345E+01 1.122E-01	1.760E+01 4.666E-02 5.762E-05 0. 1.969E-01 -4.788E-02 3.438E-06 1.144E-06 1.322E+01 8.462E-02	1.748E+01 4.627E-02 5.723E-05 0. 1.763E-01 -3.646E-02 2.300E-05 4.647E-07 1.307E+01 7.219E-02
2 2	2	2 3	3.562E-0 1.296E-0	04 3.560E-04 06 1.291E-06	3.541E-04 1.241E-06	3.360E-04 7.816E-07	2.150E-04 -1.434E-06	4.177E-05 -1.144E-06	2.038E-05 -4.647E-07
50 50 50 50 50 50 50 50	49 49 49 50 50 50 50	0 1 2 3 0 1 2 3	9.862E-C 8.107E-C 7.195E-C 2.49E-C 2.621E+C 2.288E+C 1.983E+C 1.721E+C	9.862E-02 8.107E-02 9.862E-02 9.8107E-02 9.8107E-02 9.8249E-02 9.8249E-02	 9.862E-02 8.107E-02 7.195E-02 6.249E-02 2.621E+00 2.288E+00 1.983E+00 1.721E+00 	9.861E-02 8.107E-02 7.194E-02 6.249E-02 2.621E+00 2.88E+00 1.983E+00 1.721E+00	9.860E-02 8.104E-02 7.190E-02 6.244E-02 2.621E+00 2.288E+00 1.983E+00 1.721E+00	9.848E-02 8.085E-02 7.166E-02 6.216E-02 2.620E+00 2.288E+00 1.983E+00 1.721E+00	9.831E-02 8.056E-02 7.127E-02 6.170E-02 2.620E+00 2.288E+00 1.982E+00 1.720E+00

GROUP CONSTANTS AT T=3.000E+02 DEG K FOR MF 6 AND MT 16 N2N

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106.5305

INITL GRDUP	FINAL GROUP	ISDTROPIC +0	MATRIX VS F +1	INAL GRDUP +2	+3		
49 49 49 49 49 49 49	10 16 22 28 34 40 46	1.393E-09 7.749E-08 1.516E-06 2.744E-05 4.056E-04 3.610E-03 1.071E-03	3.784E-05 1.275E-07 2.475E-06 4.377E-05 6.185E-C4 4.566E-03 3.635E-05	1.028E-08 2.096E-07 4.033E-06 6.942E-05 9.266E-04 5.411E-03	2.790E-08 3.443E-07 6.553E-06 1.094E-04 1.362E-03 5.900E-03	2.860E-08 5.650E-07 1.061E-05 1.703E-04 1.952E-03 1.075E-02	4.708E-08 9.261E-07 1.710E-05 2.623E-04 2.709E-03 5.532E-03
50 50 50 50 50 50 50	10 16 22 28 34 40 46	1.995E-09 1.117E-07 2.231E-06 4.380E-05 7.944E-04 1.012E-02 2.329E-02	5.423E-09 1.840E-07 3.672E-06 7.162E-05 1.262E-03 1.413E-02 4.350E-03	1.474E-08 3.032E-07 6.039E-06 1.168E-04 1.983E-03 1.885E-02 2.214±-04	4.006E-08 4.996E-07 9.927E-06 1.900E-04 3.075E-03 2.370E-02 2.778E-06	4.110E-08 8.229E-07 1.630E-05 3.077E-04 4.685E-03 5.701E-02 6.290E-09	6.774E-08 1.355E-06 2.674E-05 4.960E-04 6.983E-03 5.008E-02
GROUP FOR MF	CONSTAN 6 AND	ITS AT T=3.0 MT 17 N,3N	000E+02 DEG	к			107.3325
INITL GROUP	F INAL GROUP	ISOTROPIC +0	MATRIX VS F +1	INAL GROUP +2	+3		
50 50 50 50 50 50 50	12 18 24 30 36 42 48	1.249E-09 2.566E-08 5.084E-07 9.609E-06 1.470E-04 9.852E-04 1.227E-06	3.394E-09 4.226E-08 8.340E-07 1.550E-05 2.210E-04 1.105E-03 2.685E-09	3.48 E-09 6.957E-08 1.366E-06 2.474E-05 3.239E-04 2.127E-03	5.737E-09 1.145E-07 2.234E-06 3.899E-05 4.594E-04 1.253E-03	9.454E-09 1.883E-07 3.645E-06 6.151E-05 6.261E-04 3.468E-04	1.558E-08 3.095E-07 5.929E-06 9.583E-05 8.110E-04 3.811E-05
GROUP For MF	CONSTAN 6 AND	ITS AT T≖3.(MT 18 FISS]	000E+02 DEG	к			107.9485
INITL GROUP	F I NAL GROUP	ISOTROPIC +0	MATRIX VS F +1	INAL GROUP +2	+3		
1 1 1 1 1 1 1 1	1 7 13 25 31 37 43 49	1.502E-07 4.942E-07 3.313E-05 1.841E-04 1.739E-03 1.629E-02 1.459E-01 1.069E+00 7.278E-01	4.860E-03 9.327E-07 2.843E-05 2.676E-04 2.528E-03 2.359E-02 2.079E-01 3.240E+00 5.046E-02	6.913E-08 1.831E-06 4.127E-05 3.392E-04 3.673E-03 3.412E-02 2.944E-01 4.981E+00	1.032E-07 3.697E-06 5.995E-05 5.659E-04 5.335E-03 4.926E-02 4.137E-01 6.252E+00	1.634E-07 7.600E-06 8.711E-05 8.229E-04 7.746E-03 7.097E-02 5.756E-01 5.656E+00	2.756E-07 1.581E-05 1.266E-04 1.196E-03 1.124E-02 1.019E-01 7.909E-01 3.043E+00
50 50 50 50 50 50 50 50 50 50	1 7 13 25 31 37 43 49	5.479E-08 1.802E-07 1.208E-05 6.713E-05 6.343E-04 5.941E-03 5.321E-02 3.899E-01 2.654E-01	1.772E-08 3.402E-07 1.037E-05 9.760E-05 9.219E-04 8.604E-03 7.581E-02 1.182E+00 1.840E-02	2.521E-08 6.678E-07 1.505E-05 1.419E-04 1.340E-03 1.244E-02 1.074E-01 1.817E+00	3.764E-08 1.348E-06 2.186E-05 2.064E-04 1.946E-03 1.797E-02 1.509E-01 2.280E+00	5.958E-08 2.772E-06 3.177E-05 3.001E-04 2.825E-03 2.588E-02 2.099E-01 2.063E+00	1.005E-07 5.765E-06 4.617E-05 4.363E-04 4.098E-03 3.718E-02 2.884E-01 1.110E+00

GROUP CONSTANTS AT T=3.000E+02 DEG K

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FOR MF 6 AND MT 51 DISCRETE INELASTIC

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INITL	F I NAL	GRDUP CONSTANTS VS LEGENORE ORDER	
GROUP	GROUP	O 1 2 3	
34 34 34 34 34 34 34	7 8 9 10 11 12	6.049E-09 -4.087E-09 1.650E-09 -1.675E-10 6.705E-08 -2.319E-08 -1.058E-10 6.584E-10 3.394E-07 -4.658E-08 -2.401E-09 -1.767E-10 1.198E-06 -8.683E-08 -3.170E-09 -2.305E-10 3.712E-06 -1.681E-07 -4.294E-09 -2.274E-10 1.083E-05 -3.358E-07 -6.115E-09 -2.057E-10	
49 50 50	49 49 50	6.838E-01 1.102E-02 8.335E-05 3.840E-08 4.998E-02 -7.339E-03 -1.952E-03 1.240E-03 5.148E-01 8.936E-03 1.954E-03 -1.240E-03	
GROUP	CONSTAN	TS AT T=3.000E+02 DEG K	112.6705
FOR MF	6 AND	MT 52 CONTINUED	
INITL	F I NAL	GRDUP CONSTANTS VS LEGENDRE ORDER	
GROUP	GROUP	O 1 2 3	
39 39 39 39 39 39 39	23 24 25 26 27 28	6.203E-08 -1.156E-08 5.300E-10 1.642E-11 2.830E-07 -8.121E-09 -6.178E-10 -1.351E-11 5.311E-07 -7.251E-09 -4.973E-11 -5.492E-13 9.425E-07 -9.902E-09 -5.765E-11 -5.285E-13 1.630E-06 -1.364E-08 -1.660E-11 -3.910E-11 3.371E-06 -7.323E-08 1.004E-09 5.536E-11	
47	47	1.420E-02 1.654E-04 7.193E-07 -1.605E-10	
48	47	5.958E-04 -1.926E-05 -1.048E-06 -8.418E-11	
48	48	1.401E-04 2.138E-05 1.050E-06 8.418E-11	
GROUP	CONSTAN	ITS AT T=3.000E+02 DEG K	121.0785
FOR MF	6 AND	MT 59 CONTINUED	
INITL	FINAL	GRDUP CONSTANTS VS LEGENDRE DRDER	
GROUP	GROUP	O 1 2 3	
45 45 45	31 32 33	4.304E-09 -3.570E-09 2.402E-09 -1.230E-D9 1.957E-06 -2.728E-07 6.661E-09 1.462E-09 8.935E-06 -3.852E-07 -3.297E-09 -4.346E-11	
47	46	9.251E-03 5.328E-05 -2.240E-07 -3.366E-10	
47	47	1.441E-03 5.122E-05 5.729E-07 6.328E-10	
48	47	6.231E-04 2.082E-06 3.130E-09 2.416E-15	
GROUP	CONSTAN	NTS AT T=3.000E+02 DEG K	122.6305
FOR MF	6 AND	MT 91 CONTINUUM INELASTIC	
INITL	F INAL	GROUP CONSTANTS VS LEGENDRE DRDER	
GROUP	GROUP	O 1 2 3	

45 12 2.470E-09 6.972E-12 7 4115-15 0. 6.713E-09 1.894E-11 2.014E-14 45 13 0. 2.066E-14 45 14 6.886E-09 1.943E-11 Ο. 15 1.135E-08 3.203E-11 3.405E-14 45 Ο. 1.870E-08 5.278E-11 5.611E-14 45 16 ο. 9.245E-14 45 17 3.082E-08 8.696E-11 Э. . . 50 48 1.018E-04 2.872E-07 3.054E-10 0. ο. 50 40 3.252E-06 9.176E-09 9.755E-12 50 50 5.733E-09 1.618E-11 1.720E-14 0. GROUP CONSTANTS AT T=9.000E+02 DEG K 127.7375 FOR MF 3 AND MT 1 TOTAL FNRGY LGEND GROUP CONSTANTS VS SIGMA ZERD INFINITY 1.000E+05 1.000E+04 GROUP ORDER 1.000E+03 1.000E+02 1 000E+01 1.000E+00 3.690E+02 3.679E+02 3.596E+02 3.127E+02 2.186E+02 1.665E+02 1.571E+02 1 0 FLUX O 5.021E+00 5.003E+00 4.847E+00 3.825E+00 1.576E+00 2.845E-01 3.176E-02 3.669E+02 8.019E+01 7.310E+01 1 3.690F+02 3.508E+02 2.687E+02 1.319E+02 FLUX 1 4.683E+00 5.021E+00 4.985E+00 3.015E+00 6.797E-01 3.155E-02 4.286F-04 2 0 2.485E+01 2.485E+01 2.485E+01 2.481E+01 2.451E+01 2.352E+01 2.294E+01 FLUX O 5.002E-01 5.000E-01 4.989E-01 4.881E-01 4.017E-01 1.492E-01 2.089E-02 2 2.485E+01 2.485E+01 2.484E+01 2.477E+01 2.417E+01 2.210E+01 2.088E+01 FLUX 1 5.002E-01 4.999E-01 4.977E-01 3.235E-01 4.649E-02 4.763E-01 9.552E-04 3 1.032E+01 1.032E+01 1.032E+01 1.031E+01 1.031E+01 0 1.032E+01 1.030F+01 FLUX O 5.001E-01 5.001E-01 4.996E-01 4.950E-01 4.534E-01 2.463E-01 4.426E-02 1.032E+01 1.029E+01 1.032E+01 1.032E+01 1.032E+01 1.031E+01 1.030E+01 3 FLUX 1 5.001E-01 5.000E-01 4.991E-01 4.900E-01 4.110E-01 1.213E-01 3.922E-03 50 0 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.880E+00 5.879E+00 FLUX O 9.012E-03 9.012E-03 9.007E-03 8.959E-03 8.512E-03 5.675F-03 1.310E-03 5.880E+00 5.880E+00 5.880E+00 5.880F+00 5 880E+00 50 5.880F+00 5.879F+00 1 FLUX 1 9.012E-03 9.011E-03 9.002E-03 8.907E-03 8.039E-03 3.574E-03 1.904E-04 GROUP CONSTANTS AT T=9.000E+02 DEG K FOR MF 3 AND MT 2 ELASTIC 131.0125 ENRGY GROUP CONSTANTS VS SIGMA ZERO INFINITY 1.000E+05 1.000E+04 1.000E+03 1.000E+02 GROUP 1.000E+01 1.000E+00 1.820E+01 1.762E+01 1.750E+01 1 1.927E+01 1.927E+01 1.923E+01 1.897E+01 2 1.380E+01 1.380E+01 1.379E+01 1.379E+01 1.372E+01 1.347E+01 1.333E+01 3 9.615E+00 9.615E+00 9.615E+00 9.615E+00 9.614E+00 9.612E+00 9.610E+00 . 48 3.654E+00 3.654E+00 3.654E+00 3.654E+00 3.654E+00 3.654E+00 3.653E+00 49 3.114E+00 3.115E+00 3.115E+00 3.115E+00 3.115E+00 3.111E+00 3.106E+00 50 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 2.719E+00 GRDUP CONSTANTS AT T=9.000E+02 DEG K 133.4135 FOR MF 3 AND MT 18 FISSION ENRGY GROUP CONSTANTS VS SIGMA ZERO 1.000E+05 1.000E+04 1.000E+03 1.000E+02 1.000E+01 1.000E+00 GROUP INFINITY

1 2 3	1.011E4 3.010E 2.916E	01 1.0 -01 3.0 -02 2.9	008E+01 010E-01 916E-02	9.834E+(3.009E-(2.916E-(00 8.482E+0 01 3.002E-0 02 2.916E-0	0 5.775E+00 1 2.942E-01 2 2.913E-02	4.281E+00 2.742E-01 2.902E-02	4.013E+00 2.626E-01 2.892E-02	
48 49 50	2.6!2E 2.605E 2.470E	HOO 2.0 HOO 2.0 HOO 2.0	612E+00 605E+00 470E+00	2.612E+(2.605E+(2.470E+(00 2.612E+0 00 2.605E+0 00 2.470E+0	0 2.612E+00 0 2.604E+00 0 2.470E+00	2.612E+00 2.604E+00 2.470E+00	2.612E+00 2.602E+00 2.470E+00	
GROUP FOR MF	CONSTAN 3 AND 1	TS AT T MT102 C	=9.000E+ APTURE	O2 DEG K			135	. 595S	
ENRGY GROUP	GROUP (INFINI	CONSTAN TY 1.0	TS VS SI 000E+05	GMA ZERD	04 1.000E+0	3 1.000E+02	1.000E+01	1.000E+00	
1 2 3	3.396E 1.075E 6.717E	+02 3.: +01 1.0 -01 6.	386E+02 075E+01 717E-01	3.305E+(1.075E+(6.717E-(02 2.852E+0 01 1.072E+0 01 6.716E-0	2 1.946E+02 1 1.050E+01 1 6.707E-01	1.446E+02 9.775E+00 6.664E-01	1.356E+02 9.352E+00 6.627E-01	
48 49 50	1.087E 7.328E 5.584E	-02 1.0 -03 7.3 -03 5.3	087E-02 328E-03 584E-03	1.087E- 7.328E- 5.584E-	02 1.087E-0 03 7.328E-0 03 5.584E-0	2 1.087E-02 3 7.327E-03 3 5.584E-03	1.086E-02 7.317E-03 5.584E-03	1.085E-02 7.303E-03 5.585E-03	
GROUP FOR MF	CONSTAN 6 AND 1	TSATT MT2E	=9.000E+ LASTIC	02 DEG K			137	.8785	
INITL GROUP	F I NAL GROUP	LGEND ORDER	GROUP C	CONSTANTS	VS SIGMA ZE 08+05 1.000	RD E+04 1.000E	+03 1.000E+	02 1.000E+01	1.000E+00
1 1 1	1 1 1	0 1 2	1.927E+ 5.441E- 6.912E-	01 1.92 02 5.43 05 6.90	7E+O; 1.923 8E-O2 5.41 7E-O5 6.863	8E+01 1.897E 7E-02 5.274E 7E-05 6.606E	+01 1.820E+ -02 4.892E- -05 6.017E-	01 1.762E+01 02 4.668E-02 05 5.763E-05	1.750E+01 4.629E-02 5.724E-05
50 50 50 50	50 50 50 50	0 1 2 3	2.621E+ 2.288E+ 1.983E+ 1.721E+	+00 2.62 +00 2.28 +00 1.98 +00 1.72	fE+00 2.62 &E+00 2.28 SE+D0 1.98 1E+00 1.72	E+00 2.621E E+00 2.288E E+00 1.983E E+00 1.721E	+00 2.621E+ +00 2.288E+ +00 1.983E+ +00 1.721E+	00 2.620E+00 00 2.288E+00 00 1.983E+00 00 1.721E+00	2.620E+00 2.288E+00 1.982E+00 1.720E+00
GROUP FDR MF	CONSTAN 3 AND	TS AT T MT 1 T	=2.100E- OTAL	HO3 DEG K			149	.3345	
ENRGY GROUP	LGENO DRDER	GROUP INFINI	CONSTAN TY 1.0	TS VS SIG DOOE+05	MA ZERD 1.000E+04	I.000E+03 1.	000E+02 1.C	000E+01 1.000E	+00
1 FL 1 FL	0 .UX 0 1 .UX 1	3.688E 5.021E 3.688E 5.021E	+02 3.0 +00 5.0 +02 3.0 +00 4.9	678E+02 D03E+00 667E+02 985E+00	3.59,4E+02 4.847E+00 3.507E+02 4.683E+00	3.126E+02 2. 3.826E+00 1. 2.687E+02 1. 3.015E+00 6.	186E+02 1.6 576E+00 2.8 320E+02 8.0 793E-01 3.1	66E+02 1.573E 43E-01 3.173E 031E+01 7.321E 148E-02 4.276E	+02 -02 +01 -04
50 FL 50 FL	0 _UX 0 _1 _UX 1	5.880E 9.012E 5.880E 9.012E	+00 5. -03 9. +00 5. -03 9.	880E+00 012E-03 880E+00 011E-03	5.880E+00 9.007E-03 5.8802+00 9.002E-03	5.880E+00 5. 3.959E-03 8. 5.880E-00 5. 8.907E-03 8.	880E+00 5.8 512E-03 5.6 880E+00 5.8 039E-03 3.5	880E+00 5.879E 575E-03 1.310E 880E+00 5.879E 574E-03 1.904F	+00 -03 +00 -04

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GROUP FOR MF	CONSTAN 6 AND	NTS AT 1 MT 2 E	=2.100E+03 ELASTIC	DEG K			157.48	35	
INITL GROUP	FINAL GROUP	LGEND ORDER	GROUP CONS INFINITY	TANTS VS SI 1.000E+05	GMA ZERD 1.000E+04	1.000E+03	1.000E+02	1.000E+01	1.000E+00
1 1 1	1 1 1	0 1 2	1.939E+01 5.473E-02 6.952E-05	1.938E+01 5.470E-02 6.946E-05	1.934E+01 5.445E-02 6.901E-05	1.905E+01 5.292E-02 6.624E-05	1.825E+01 4.899E-02 6.022E-05	1.765E+01 4.671E-02 5.765E-05	1.753E+01 4.632E-02 5.727E-05
50 50 50	50 50 50	1 2 3	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.983E+00 1.721E+00	2.288E+00 1.982E+00 1.720E+00
*****	*****	******	*********	**********	*********	*********	165.15 *****	7S **	
CCCCR.	PR00	UCE CCC	C FORMAT OUT	PUT FILES			165.19	45	
INPUT OUTPUT OUTPUT OUTPUT	GENDF ISDTX BRKOX DLAYX	UNIT S UNIT S UNIT S UNIT S UNIT	· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	-25 21 22 0				
FIL **USER	E ISOT IDENT	XS V Ificati	ERSIDN 1 DN**T2LANL	UNIT 21 NJOY					
FILE C	ONTROL	PARAME	TER5						
NGROU NISO Maxuf Maxon Maxof Ichis	JP N N M M RD M ST S	UMBER D UMBER O AXIMUM AXIMUM AXIMUM ET FISS ICHIST	F ENERGY GRC F ISOTOPES I NUMBER OF UF NUMBER OF OC SCATTERING C ION SPECTRUN =1 SET =NGROUP. SET	DUPS IN SET SCATTER GRO DWNSCATTER G DRDER 4 FLAG 7 VECTDR 7 MATRIX	NUPS ROUPS	50 1 50 4 0			
NSCMA	X M	AXIMUM SCATTE	NUMBER OF BL RING DATA	DCKS OF		4			
NSBLO		LDCKING	CONTROL FOR	R SCATTERING	G DATA	1			
ISC	DTOPE	NAME PU238							
GROUP	NE	UTRON V (CM/SEC)	UPPER I	ENERGY (EV)				
1 2 3		4.667 3.712 2.947	7684E+09 734E+09 7111E+09	1.99710 1.0000 6.06530	08E+07 00E+07 07E+06				
48		1.660	9850E+06	1.8553	91E+00				

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49	1.293479E+06	1.125352E+00
50	3.140241E+05	6.825603E-01
		1.000021E-05

NUMBER OF RECORDS TO BE SKIPPED

ISOTOPE NUMBER 1 O

ISOTOPE 1

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ISOTOPE CONTROL PARAMETERS

HIDENT LIBRARY IDENTIFIER ENDFE HMAT ISOTOPE IDENTIFICATION 1050 AMASS GRAM ATOMIC WEIGHT 2.38210E+0	34))2 1 2
HMAT ISOTOPE IDENTIFICATION 1050 AMASS GRAM ATOMIC WEIGHT 2.38210E+0))2 1 2
AMASS GRAM ATOMIC WEIGHT 2.38210E+C	2
	11
LFISS IMERMAL ENERGY/FISSION (W*SEC/FISS) 3.30030E-1	2
ECAPT THERMAL ENERGY/CAPTURE (W*SEC/CAPT) 1.74610E-	
TEMP ISOTOPE TEMPERATURE (DEG K) O.	0
SIGPOT AVE. POTENTIAL SCATTERING (BARNS/ATOM) 1.00000E+	0
ADENS REFERENCE ATOM DENSITY (A/B+CM) O.	-
KBR ISOTOPE CLASSIFICATION	0
ICHI FISSION SPECTRUM FLAG (0/1/N=SET CHI/VECTOR/MATRIX)	1
IFIS (N.F) X-SEC FLAG (0/1=ND/YES)	1
IALF (N.A) X-SEC FLAG (O/1=NO/YES)	0
INP (N.P) X-SEC FLAG (O/1=ND/YES)	õ
IN2N (N.2N) X-SEC FLAG (O/1=ND/YES)	1
IND (N,D) X-SEC FLAG (O/1=ND/YES)	0
INT (N.T) X-SEC FLAG (O/1=ND/YES)	ō
LTOT NUMBER OF TOTAL X-SEC MOMENTS	1
LTRN NUMBER OF TRANSPORT X-SEC MOMENTS	1
ISTRPD NUMBER OF TRANSPORT X-SEC DIRECTIONS	0

BLOCK	TYPE	IDENT	ORDERS
4	TNELAC	100	

1	INELAS	200	- 4
2	ELASTC	100	4
3	N2N	300	1
4	TOTAL	0	4

SCATTERING BANDWIDTH AND IN-GROUP SCATTERING POSITION

GROUP/BLOCK	1	2	3	4	1	2	3	4
1	1	1	1	1	1	1	1	i
2	2	2	2	2	1	1	1	i
3	3	2	3	3	1	1	i	1
• •								
•								
•								
48	0	2	0	2	1	1	1	ſ
49	0	2	Ō	2	1	1	1	1
50	0	2	Ō	2	1	1	1	i.

PRINCIPAL CROSS-SECTIONS

GROUP 1 2 3	STRPL 3.508785E+00 3.835912E+00 4.199807E+00	STOTPL 5.879757E+00 6.588224E+00 7.316763E+00	SNGAM 5.583553E-03 7.328427E-03 1.036872E-02	SFIS 2.469865E+00 2.604587E+00 2.611864E+00	SNUTOT 4.101729E+00 3.605884E+00 3.292566E+00	CHISO 1.816369E-03 2.619729E-02 1.095327E-01	SN2N 1.109183E-01 2.278995E-02 0.
•							

 48
 1.023363E+01
 1.026073E+01
 6.353501E-01
 2.818899E-02
 2.750000E+00
 2.486807E-09
 0.

 49
 2.471991E+01
 2.475880E+01
 1.068274E+&1
 2.990718E-01
 2.750000E+00
 1.748206E-09
 0.

 50
 3.687034E+02
 3.687577E+02
 3.394378E+02
 1.010177E+01
 2.750000E+00
 5.404519E-09
 0.

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MATRIX ** INELAS **

FINAL	INITL	XSEC VS LE	GENORE ORDER		
GROUP	GROUP	ORDER O	ORDER 1	ORDER 2	ORDER 3
		5 1485-01	9 936F-03	1 9545-02	-1 2405-02
	2	6 9395-01	1 1025-02	9 2355-05	3 9405-09
2		4 0095-07	-7 2205-02	-1 0525-02	1 2405-02
2	2	4.9900-02	-7.3392-03	-1.9520-05	7.0465-00
3	3	7.8302-01	8.7732-03	3.4102-05	7.0162-09
	2	5.210E-02	-8.934E-03	-8.070E-05	-3.840E-08
	1	1.030E-04	2.872E-07	3.054E-10	0.
•					
•					
41	17	1.198E-06	-8.683E-08	-3.170E-09	-2.305E-10
	5	2.396E-09	6.762E-12	7.188E-15	0.
	4	1.693E-09	4.779E-12	5.080E-15	0.
	3	1.763E-09	4.974E-12	5.288E-15	0.
42	17	3.394E-07	-4.658E-08	-2.401E-09	-1.767E-10
43	17	6.705E-08	-2.319E-08	-1.058E-10	6.584E-10
44	17	6.049E-09	-4.087E-09	1.650E-09	-1.675E-10

MATRIX ** ELASTC **

F I NAL GROUP	INITL GROUP	XSEC VS LEC ORDER O	GENORE ORDER ORDEF: 1	ORDER 2	ÖRDER 3	
1	1	2.621E+00	2.288E+00	1.983E+00	1.721E+00	
2	2	3.101E+00	2.747E+00	2.440E+00	2.089E+00	
	1	9.862E-02	8.107E-02	7.195E-02	6.249E-02	
3	3	3.638E+00	3.111E+00	2.685E+00	2.185E+00	
	2	1.389E-02	3.195E-03	4.247E-03	1.441E-03	
•						
50	50	1.922E+01	5.425E-02	6.891E-05	0.	
-	49	2.565E-01	-8.499E-02	-3.068E-04	-1.296E-06	
MATRIX	** N2N	**				
FINAL	INITL	XSEC VS IN	ITL GROUP			
GROUP	GROUP	-0	- 1	-2	-3	
1	!	6.290E-09				
2	-	2.7785-00				
	2	3 6355-05	4 350F-03			
5	2	1.071E-03	2.329E-02			
	-					
39	2	1.028E-08	1.474E-08			
40	2	3.784E-09	5.423E-09			
41	2	1.393E-09	1.995E-09			
MATRIX	** TOTAL	**				
FINAL	INITL	XSEC VS LE	GENORE ORDER			
GROUP	GROUP	ORDER O	ORDER 1	ORDER 2	ORDER 3	

1 2 3	1 2 1 3 2 1	3.135E+00 3.785E+00 1.486E-01 4.421E+00 6.599E-02 3.244E-04	2.297E+00 2.758E+00 7.373E-02 3.120E+00 -5.739E-03 2.872E-07	1.985E+00 2.440E+00 7.000E-02 2.685E+00 4.36EE-03 3.054E-10	1.720E 2.089E 6.373E 2.185E 1.440E 0.	+00 +00 -02 +00 -03		
49 50	49 48 50 49	1.352E+01 1.720E-01 1.922E+01 2.565E-01	1.239E-01 -5.678E-02 5.425E-02 -8.499E-02	3,562E-04 -3.417E-04 6.891E-05 -3.068E-04	1.296E 1.393E 0. -1.296E	-06 -07 -06		
1***FILE **USER I	BRKOXS DENTIFI	VERSION CATION**T2LA	1 UNIT 22 NL NJOY	***				
FILE CON	TROL PA	RAMETERS						
NGROUP	NUME	BER DF ENERGY	GROUPS IN S	ET		50		
NISUSH		IELDING FACT	ORS	-		1		
NSIGPI	WH	ICH ARE GIVE	N. NSIGPT I	S EQUAL TO		-		
NTEMPT	TOT A	L SUM FROM 1	VALUES OF VA	RIABLE TB		6		
	THE	SUM FROM 1 T	O NISOSH OF	NTABY(I)		3		
IBLK	BLOC	CKING OPTION	UNS			6 0		
I SOTO 1	PE N F	NAME PU238						
LN(SIGPO ISOT)/LN(10 OPE 1 5.	D) VALUES FOR 1ST VALUE .00000E+00	ALL ISDTOPE 2ND VAL 4.00000E+00	:S .UE) 3.00000	 E+00	2.00000E+00	1.00000E+00	0.
TEMPERAT	URES (C	DEG C) FOR AL	L ISOTOPES					
ISOTOPE	1 2.	ST VALUE . 68400E+01	2ND VALUE 6.26840E+02	2 1.82684	E+03			
MAXIMUM	ENERGY	BOUND						
GROUP	1 1.99							
	2 1.00 3 6.06	9711E+07 DOODE+07 6531E+06						
	2 1.00	9711E+07 2000E+07 3531E+06						
4 4 5	2 1.00 3 6.00	9711E+07 0000E+07 5531E+06 5539E+00 2535E+00 2560E-01						
4 4 5 Minimum	2 1.00 3 6.06	9711E+07 0000E+07 5531E+06 5539E+00 2535E+00 2560E-01 BDUND OF SE1	r					
4 4 5 MINIMUM	2 1.00 3 6.00	9711E+07 0000E+07 5531E+06 2535E+00 2535E+00 2560E-01 BDUND OF SE1 1.00002E-05	r 5					
4 4 5 MINIMUM F-FACTOR AND NUME	2 1.00 3 6.00	9711E+07 0000E+07 5531E+06 2535E+00 2560E-01 BDUND OF SET 1.00002E-05 AND STDP GRO SIGO AND TEMP	5 DUPS PERATURE VALU	JES				

TOTAL SELF-SHIELDING FACTORS ISOTOPE 1

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GROUP	1		
SIGD	TEMP 1	TEMP 2	TEMP 3
1	1.00000E+00	1.00000E+00	1.00000E+00
2	1.00000E+00	1.00000E+00	1.00000E+00
3	9.99999E-01	9.99999E-01	9.99999E-01
4	9.99994E-01	9.99994E-01	9.99994E-01
5	9.99957E-01	9.99957E-01	9.99957E-01
6	9.99903E-01	9.99903E-01	9.99903E-01
GROUP	2		
SIGD	TEMP 1	TEMP 2	TEMP 3
1	1.00000E+00	1.00000E+00	1.00000E+00
2	9.99997E-01	9.99997E-01	9.99997E-01
3	9.99969E-01	9.99969E-01	9.99969E-01
4	9.99702E-01	9.99702E-01	9.99702E-01
5	9.98061E-01	9.98061E-01	9.98061E-01
6	9.95676F-01	9.95676F-01	9.95676F-01
0	5.5557.62 01	5.555.52 0.	5.555.52 0,
•			
•			
CROUR	50		
SLCO			TEMD 2
3100			0.045455-01
1	9.943922-01	9.949346-01	9.943436-01
2	9.50/292-01	9.512536-01	9.309082-01
3	7.28257E-01	7.28645E-01	7.28597E-01
4	3.5/339E-01	3.576312-01	3.5/933E-01
5	2.17207E-01	2.17473E-01	2.17792E-01
6	1.97971E-01	1.98227E-01	1.98536E-01
CAPTURE SELF-	SHIELDING FACTO	RS I	SOTOPE 1
GROUP	1		
SIGD	TEMP 1	TEMP 2	TEMP 3
1	1.00000E+00	1.00000E+00	1.00000E+00
2	1.00000E+00	1.00000E+00	1.00000E+00
3	1.00000E+00	1.00000E+00	1.00000E+00
4	1.00001E+00	1.00001E+00	1.00001E+00
5	1.00008E+00	1.00008E+00	1.00008E+00
6	1.00017E+00	1.00017E+00	1.00017E+00
GRDUP	2		
SIGD	TEMP 1	TEMP 2	TEMP 3
1	1.00000E+00	1.00000E+00	1.00000E+00
2	9.99997E-01	9.99997E-01	9.99997E-01
3	9.99974E-01	9.99974E-01	9.99974E-01
4	9.99753E-01	9.99753E-01	9.99753E-01
5	9.98403E-01	9.98403E-01	9.98403E-01
6	9.96466E-01	9.96466E-01	9.96466E-01
•			
•			
	50		
SICO	TEMP 1	TEMP 2	TEMP 3
5100	9 97040F-01	9.974755-01	9.96721F-01
,	9 732415-01	9 736625-01	9 729575-01
2	8 39920F-01	8 40290E-01	8 39841E-01
Д	5 730075-01	5 733545-01	5 733365-01
* E	A 25570E-01	A 25969E-01	A 261885-01
5 6	3 001315-01	3 005245-01	3 007005-01
Ū	3.331210-01	J. 55534E-UI	3.33/302-01
EISSIAN SELE-		100 1	
	4		SUIVEL
GRUUP			TEMD 2
2100		1 CMP 2	1 CMP 3
		1.0000000000000000000000000000000000000	1.000000000000
2	1.000001+00	1.000000000000	1.00000E+00
3	1.00000E+00	1.00000E+00	1.00000E+00

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	4 5 6	1.00000E+00 1.00000E+00 1.00001E+00	1.00000E+00 1.00000E+00 1.00001E+00	1.00000E+00 1.00000E+00 1.00001E+00
	GROUP	2		
	5160	1 00000E+00	1 00000E+00	1 00000E+00
	2	9.999998-01	9.999998-01	9,999998-01
	3	9.99993E-01	9.99993E-01	9.99993E-01
	4	9.99937E-01	9.99937E-01	9.99937E-01
	5	9.99589E-01	9.99589E-01	9.99589E-01
	0	9.990882-01	9.990882-01	9.990882-01
	•			
	GROUP 5	50		
	SIGO	TEMP 1	TEMP 2	TEMP 3
	1	9.97027E-01	9.974518-01	9.96669E-01
	23	8.39276E-01	8.39638E-01	8.39169E-01
	4	5.71351E-01	5.71694E-01	5.71665E-01
	5	4.23412E-01	4.23808E-01	4.24021E-01
	6	3.96881E-01	3.97292E-01	3.97550E-01
TRANSP	ORT SELF-S	SHIELDING FAC	TORS	ISOTOPE 1
	SIGD	TEMP 1	TEMP 2	TEMP 3
	1	1.00000E+00	1.00000E+00	1.00000E+00
	2	1.00000E+00	1.00000E+00	1.00000E+00
	3	1.00000E+00	1.00000E+00	1 00001E+00
	5	1.00005E+00	1.00005E+00	1.00005E+00
	6	1.00010E+00	1.00010E+00	1.00010E+00
	GROUP	2	75WD 0	75WD 0
	SIGU	1 00000E+00	1 00000E+00	1 00000E+00
	2	9,99998E-01	9,999988-01	9.99998E-01
	3	9.99979E-01	9.99979E-01	9.99979E-01
	4	9.99798E-01	9.99798E-01	9.99798E-01
	5	9.986861-01	9.986861-01	9.986861-01
		5.570782 01	5.570782 01	5.570782 01
	•			
	GROUP	50		
	SIGO	TEMP 1	TEMP 2	TEMP 3
	2	9.94391E-01 9.50723E-01	9.51246E-01	9.50899F-01
	3	7.28222E-01	7.28609E-01	7.28561E-01
	4	3.57259E-01	3.57551E-01	3.57853E-01
	5	2.17112E-01	2.17378E-01	2.17698E-01
	6	1.978752-01	1.98131E-01	1.98439E-01
ELASTI	C SELF-SH GRDUP	IELDING FACTO	DRS I	SOTOPE 1
	SIGO	TEMP 1	TEMP 2	TEMP 3
	1	1.00000E+00	1.00000E+00	1.00000E+00
	2	1.00000E+00	1.00000E+00	1.00000E+00
	4	9.99990E-01	9.99990E-01	9.99990E-01
	5	9.99932E-01	9.99932E-01	9.99932E-01
	6	9.99846E-01	9.99846E-01	9.99846E-01
	SIGO	Z TEMP 1	TEMP 2	TEMP 2
	1	1.00000E+00	1.00000E+00	1.00000E+00

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	2	9.99998E-0	1 9.99998E-01	9.99998E-01	
	3	9.99979E-0	1 9.99979E-01	9 99979E-01	
	4 5	9.998021-0	1 9.99802E-01	9.998021-01	
	6	9.97146F-0	1 9.97146F-01	9.98713E-01	
		5.57 402 0	. 5.571462 01	5.571462 01	
	•				
	ondun	50			
	SIGO			TEND 2	
	1	9 997815-0	1 1 00271F+00	1 008515+00	
	2	9 97927E-0	1 1 000715+00	1 006235+00	
	3	9.84846E-O	1 9.86992E-01	9.91264E-01	
	4	9.45681E-0	1 9.46953E-01	9.49488E-01	
	5	9.15673E-0	1 9.16588E-01	9.18391E-01	
	6	9.09501E-0	1 9.10359E-01	9.12043E-01	
DE 101/4					
REMUVA	CPOUP	IELDING FAC	IURS I	SOTOPE 1	
	SIGD		TEMP 2	TEMD 3	
	1	1.00000E+0	0 1.00000E+00	1.00000E+00	
	2	9.99998E-0	1 9.99998E-01	9,99998E-01	
	3	9.99979E-0	1 9.99979E-01	9.99979E-01	
	4	9.99800E-0	1 9.99800E-01	9.99800E-01	
	5	9.98663E-D	1 9.98663E-01	9.98663E-01	
	6	9.96896E-0	1 9.96896E-01	9.96896E-01	
	GROUP	2			
	SIGU	IEMP 1	1 EMP 2	TEMP 3	
	2	9.999961-0	1 9.999966-01	9.999965-01	
	23	9.999582-0	1 9 99 9 38 - 01	9.999586-01	
	4	9.96018E-0	1 9 96018E-01	9 960185-01	
	5	9.74638E-0	1 9.74638E-01	9.74638E-01	
	6	9.45192E-0	1 9.45192E-01	9.451928-01	
	•				
	•				
	- CROUR	50			
	SIGO			TEMD 2	
	1	1 00000F+0	0 1 00000E+00	1 00000E+00	
	2	1.00000E+0	0 1.00000E+00	1.00000E+00	
	3	1.00000E+0	0 1.00000E+00	1.00000E+00	
	4	1.00000E+0	0 1.00000E+00	1.00000E+00	
	5	1.00000E+0	0 1.00000E+00	1.00000E+00	
	6	1.00000E+0	0 1.00000E+00	1.00000E+00	
CROUR	,		VETN	VCF VCMI	
1	1 089000	5PU 5+01 5 80031	ASIN 65-01 2 7101575	ASE ASMU	
2	1.089000	E+01 8 41303	9E-01 3 114775	+00 8.713364E-01	1 3888185-02 3 9071215-03
3	1.0890001	E+01 1.03980	4F+00 3.654240E	+00 8.521630E-01	1.664873E-02 2.278001E-03
4	1.089000	E+01 9.59599	2E-01 3.767679E	+00 7.600857E-01	2.036892E-02 2.703112E-03
5	1.089000	E+01 8.05369	9E-01 3.9506368	+00 6.621020E-01	2.501423E-02 3.165847E-03
6	1.089000	E+O1 6.94334	9E-01 4.4127618	+00 5.619092E-01	3.262864E-02 3.697078E-03
•					
•					
49	1 089000	E+01 0	9 597100		1 7204895-01 9 0625825-02
49	1.089000	E+01 0	1 3776945	+01 2 822801E-03	2 565092F-01 1 2910995-02
50	1.089000	E+01 0.	1.9218125	+01 2.822861E-03	0. 0.
					169.7525
*****	********	*********	**********	**************	* * * * * * * * * * * * * * * * * * *

III. EXAMPLE 3. PHOTON INTERACTION PROCESSING MOUNT FILE 23 PART OF DLC7E ON UNIT 20. MOUNT FILE 27 PART OF DLC7E DN UNIT 22. 0 4 *RECONR* 20 21 *PENDE TAPE FOR PHOTON INTERACTION CROSS SECTIONS FROM OLC7E*/ 1 1 0 .001 0 6 / *1-HYDROGEN*/ 92 1 0 .001 0. 6 / *92-URANIUM*/ 0/ *GAMINR* 22 21 0 23 1 3 3 4 1 *12 GROUP PHOTON INTERACTION LIBRARY*/ -1 0/ 92 -1 0/ *DTFR* 23 24 0 100 5 12 4 5 16 1 0 *PHEAT * 1 621 1 0/ *H* 1 1 0./ *U* 92 1 0./ 0/ *MATXSR* 0 23 25 1 1 *T2LASL NJDY*/ 1 1 1 *12-GROUP FHOTON INTERACTION LIBRARY*/ *GAMA* 12 *GSCAT*/ 1 1 3 2 4 *H* 1 1 1 *U* 1 1 92 *STOP* * * * \$\$ \$\$ \$\$\$\$\$ \$\$ * * VERS. 10/81 * \$\$ \$\$ \$\$ \$\$ \$\$ \$\$\$\$\$\$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$ \$\$\$ \$\$ NUCLEAR RAN AT LANL * * * \$\$\$\$ \$\$ \$\$ \$\$\$\$ CROSS SECTION * PROCESSING * * ON MACH. U * DN 11/06/81 \$\$\$\$ * * * - \$\$ * * \$\$ * AT 18:59:14 * \$\$ \$\$\$ \$\$\$\$\$\$\$ \$\$\$\$\$\$\$ \$\$ * SYSTEM * * * \$\$ \$\$ \$\$\$\$\$ \$\$\$\$\$ \$\$ * * * ************ RECONR...RECONSTRUCT POINTWISE CROSS SECTIONS IN PENDE FORMAT .7685 UNIT FOR ENDF/B TAPE 20 UNIT FOR PENDE TAPE 21 LABEL FOR PENOF TAPE _____

PENDE TAPE FOR PHOTON INTERACTION CROSS SECTIONS FROM DLC7E TAPE LABEL OLC-7E PHOTON INTERACTION LIBRARY IN ENDF FORMAT - DATAFOR FILE 23

 MATERIAL TO BE PROCESSED

 RECONSTRUCTION TOLERANCE

 RECONSTRUCTION TEMPERATURE

 NO. SIGNIFICANT FIGURES

 RESONANCE-INTEGRAL-CHECK TOLERANCE

 .001 0.K 6 RESONANCE-INTEGRAL-CHECK TOLERANCE020 MAX RESONANCE-INTEGRAL ERROR 1.000E-07 .020 DESCRIPTIVE CARDS FOR PENDE TAPE 1-HYDRDGEN PRDCESSING MAT 1 PHOTON INTERACTION DATA FOR 1 H MAT HAS NO FILE 2. MAT HAS NO RESONANCE PARAMETERS PDINTS IN INITIAL UNIONIZED GRID = 40 2.0285 POINTS ADDED BY LINEARIZATION = 263 MATERIAL TO BE PROCESSED 92 .001 RECONSTRUCTION TOLERANCE RECONSTRUCTION TEMPERATURE 0.К 6 .020 DESCRIPTIVE CARDS FOR PENDE TAPE _ _ _ _ _ _ _ 92-URANIUM PROCESSING MAT 92 ------PHOTON INTERACTION DATA FOR 92 U MAT HAS NO FILE 2. MAT HAS NO RESONANCE PARAMETERS POINTS IN INITIAL UNIONIZED GRID = 60 POINTS ADDED BY LINEARIZATION = 493 7.0415 9.1385 GAMINR... PRODUCE PHOTON INTERACTION CROSS SECTIONS 9.1695 UNIT FOR ENDF/B TAPE 22 UNIT FOR PENOF TAPE UNIT FOR INPUT GAMOUT TAPE UNIT FOR OUTPUT GAMOUT TAPE 21 0 23 MAT TO BE PROCESSED 1 GAMMA GROUP OPTION 3

WEIGHT FUNCTION OPTION LEGENORE DRDER PRINT OPTION (O MIN, 1 MAX)
RUN TITLE
12 GROUP PHOTON INTERACTION LIBRARY
GAMMA GROUP STRUCTURELASL 12 GROUP 1 1.0000E+04 - 1.0000E+05 2 1.0000E+05 - 5.0000E+05 3 5.0000E+05 - 1.0000E+06 4 1.0000E+06 - 2.0000E+06 5 2.0000E+06 - 3.0000E+06 6 3.000E+06 - 4.0000E+06 7 4.0000E+06 - 5.0000E+06 8 5.0000E+06 - 6.0000E+06 9 6.0000E+06 - 7.0000E+06 10 7.0000E+06 - 8.0000E+06 11 8.0000E+06 - 9.0000E+06 12 9.0000E+06 - 2.0000E+07
WEIGHT FUNCTION1/E WITH ROLLOFFS
PROCESSING MAT 1
1-HYDRDGEN
GROUP CONSTANTS FOR MF23 AND MT501 TOTL
GAMMA SIGMA GROUP (BARNS) 1 5.256E-01 2 3.914E-01 3 2.493E-01 4 1.77E-01 5 1.307E-01 6 1.063E-01 7 9.072E-02 8 7.978E-02 9 7.160E-02 10 6.532E-02 11 6.032E-02 12 4.996E-02
GROUP CONSTANTS FDR MF23 AND MT502 CDHT
GAMMA SIGMA GRDUP (BARNS) 1 1.235E-03 2 1.386E-04 3 1.004E-05 4 2.510E-06 5 7.934E-07 6 3.919E-07 7 2.337E-07 8 1.553E-07 9 1.109E-07 10 8.311E-08

3 4 1

10 8.311E-08 11 6.451E-08 12 3.677E-08

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9.2945

9.3845

INITL GROUP 1 3 4 5 6 7 8 9 10	F INAL GROUP 1 2 3 4 5 6 7 8 9 10	CRDSS SECT LEGENDRE OI 1.235E-03 1.386E-04 1.004E-05 2.510E-06 7.934E-07 3.919E-07 2.337E-07 1.553E-07 1.109E-07 8.311E-08	IONS VS RDER 1.220E-03 1.386E-04 1.004E-05 2.510E-06 7.934E-07 3.919E-07 2.337E-07 1.553E-07 1.109E-07 8.311E-08	1.193E-03 1.384E-04 1.004E-05 2.510E-06 7.934E-07 3.919E-07 2.337E-07 1.553E-07 1.109E-07 8.311E-08	1.158E-03 1.381E-04 1.004E-05 2.510E-06 7.934E-07 3.919E-07 2.337E-07 1.553E-07 1.109E-07 8.311E-08	1.117E-03 1.377E-04 1.004E-05 2.510E-06 7.934E-07 3.919E-07 2.337E-07 1.553E-07 1.109E-07 8.311E-08	
11 12	11 12	6.451E-08 3.677E-08	6.451E-08 3.677E-08	6.451E-08 3.677E-08	6.451E-08 3.677E-08	6.451E-08 3.677E-08	
GROUP FOR MF	CONSTAN 23 AND	TS MT5D4 INCH					12.7785
GAMMA GROUP 1 2 3 4 5 6 7 8 9 10 10 11	SIGMA (BARNS 5.243E 3.912E 2.493E 1.777E 1.304E 8.952E 7.818E 6.965E 6.300E 5.762E 4.622F) -01 -01 -01 -01 -01 -02 -02 -02 -02 -02 -02 -02 -02					
GROUP FOR MF	CONSTAN	TS MT504 INCH					12.8735
INITL GROUP 1 1 2 2 2 2	FINAL GROUP 1 XSEC HEAT 1 2 XSEC	CRDSS SECT LEGENDRE O 5.243E-O1 4.488E+O3 5.769E-O2 3.335E-O1 3.912E-O1	IONS VS RDER 4.719E-02 -1.966E-02 9.507E-02	5.465E-02 3.794E-03 4.679E-02	4.790E-03 -2.102E-03 1.466E-02	-6.011E-04 -3.024E-04 3.093E-03	
12 12 12 12 12	10 11 12 XSEC HEAT	2.167E-03 2.101E-03 4.112E-03 4.622E-02 3.711E+05	2.121E-03 2.073E-03 4.082E-03	2.031E-03 2.019E-03 4.023E-03	1.899E-03 1.940E-03 3.935E-03	1.731E-03 1.837E-03 3.821E-03	
GROUP FOR M	CONSTAN F23 AND	NTS MT516 PAIR					24.7305
GAMMA GRDUP 4 5	SIGMA (BARNS 4.6411 3.3081	5) -05 -04					

9.4725

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6 7 8 9 10 11 12	7.523E 1.201E 1.558E 1.950E 2.318E 2.704E 3.740E	-04 -03 -03 -03 -03 -03 -03			
GROUP FOR MF	CONSTAN 26 AND	ITS MT516	5 PAI	R	
INITL GROUP 4 5 5	FINAL GROUP 3 HEAT 3 HEAT	CRDS LEGE 9.28 3.32 6.6 5.08	SS SE ENDRE 32E-C 29E+C 15E-C 89E+C	CTIDNS ORDER 05 01 04 02	vs
12 12	3 Heat	7.48 4.2	B1E-0 13E+0)3)4	
GROUP FOR MI	CONSTAN 723 AND	ITS MT60:	2 ABS	ST	
GAMMA GRÒUP 1 2 3 4 5 6 7 8 9 10 11 11 12	SIGMA (BARNS 1.926E 3.243E 3.915E 6.579E 1.590E 1.590E 1.152E 9.006E 7.424E 5.445E 3.937E	-) -05 -07 -10 -10 -10 -11 -11 -11 -11 -11	HEA1 (EV- 6.33 4.56 8.84 5.45 5.13 1.93 4.80 4.70 4.6 4.41	F I NG - BARNS) 31E - 01 59E - 02 07E - 03 47E - 04 34E - 04 34E - 04 75E - 04 31E - 04 20E - 04 03E - 04 03E - 04 BOE - 04	
GROUP FOR M	CONSTAN F23 AND	NTS MT62	1 HE	A T	
GAMMA GRDUP 1 2 3 4 5 6 7 8 9 10 10 11 12	HEATIN (EV-B/ 4.4898 2.2828 7.0508 1.2408 1.7848 2.1900 2.5191 2.7998 3.0468 3.2741 3.4888 4.1331	NG ARNS) +03 +04 +04 +05 +05 +05 +05 +05 +05 +05 +05 +05 +05			
PROCE	SSING M	AT	92 		
GROUP	CONSTA	NTS			

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1

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24.9255

24.7645

24-8175

24.9075

92

2

3

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5

6

953E+01 623E+00 6.803E-01 4 4 5 5 2.188E-01 6 6 1.085E-01 7 7 6.483E-02 4.313E-02 3.079E-02 8 8 9 9 10 10 2.306E-02 11 11 1.793E-02 1.021E-02 12 12 GROUP CONSTANTS FOR MF23 AND MT504 INCH GAMMA SIGMA GROUP (BARNS) 3.676E+01 1

3.280E+01

2.252E+01 1.623E+01

1.196E+01

9.688E+00

FOR MF	26 AND	MT502 COHT	
INITL	FINAL	CROSS SECT	IONS VS
GROUP	GROUP	LEGENDRE D	RDER
1	1	1.844E+O2	1.292E
2	2	2.953E+01	2.541E
3	3	2.623E+00	2.495E

1.800E+01 q 10 1.849E+01 1.909E+01 11 2.103E+01 12 GROUP CONSTANTS FOR MF23 AND MT502 CDHT GAMMA SIGMA G

ROUP	(BARNS)
1	1.844E+02
2	2.953E+01
3	2.623E+00
4	6.803E-01
5	2.188E-01
6	1.085E-01
7	6.483E-02
8	4,313E-02
9	3.079E-02
10	2 306F-02

1.021E-02 12

2.306E-02 1.793E-02 10 11

2.541E+01

2.495E+00

6.664E-01

2.170E-01

1.080E-01

6.466E-02

4.305E-02

3.075E-02

2.304E-02

1.792E-02

1.021E-02

1.292E+02 9.996E+01

2.232E+01

2.374E+00

6.477E-01

2.142E-01

1.072E-01

6.435E-02

4.291E-02

3.067E-02

2.300E-02

1.789E-02

1.0208-02

7.350E+01

1.922E+01

2.255E+00

6.275E-01

2.107E-01

1.062E-01

6.393E-02

4.271E-02

3.057E-02

2.293E-02

1.785E-02

1.019E-02

GROUP CONSTANTS

02 COHT

35.8795

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25.2325

25.080S

1 TOTL

FOR	MF2	3	AND	MT 50
GAM	AA JP	SI (B		5)

1 2

3

4

5 6

7

8

MF2	3	AND	MT 50
A	ŞI	GMA	_ 、

2.768E+03

5.452E+02

4.882E+01

2.345E+01 1.781E+01

1.718E+01

1.735E+01

1.766E+01

5.558E+01

1.664E+01

2.151E+00

6.081E-01

2.069E-01

1.049E-01

6.343E-02

4.247E-02

3.044E-02

2.286E-02

1.781E-02

1.017E-02

8.222E+00 7.183E+00 8 9 6.401E+00 10 5.788E+00 5.296E+00 11 4.247E+00 12 GROUP CONSTANTS 36.0375 FOR MF26 AND MT504 INCH FINAL CROSS SECTIONS VS INITL LEGENDRE ORDER GROUP GROUP 3.676E+01 -3.149E-02 2.240E+00 -8.225E-01 -6.794E-01 1 1 3.676E+01 XSEC 1 3.561E+05 1 HEAT 4.864E+00 -1.777E+00 3.398E-01 -1.769E-01 -1.700E-02 2.794E+01 6.794E+00 2.958E+00 4.526E-01 -3.175E-01 2 1 2 2 2 XSEC 3.280E+01 . . 12 12 3.872E-01 3.845E-C1 3.790E-01 3.710E-01 3.605E-01 4.247E+00 XSEC 12 12 HEAT 3.400E+07 GROUP CONSTANTS 71.4975 FOR MF23 AND MT516 PAIR GAMMA SIGMA GROUP (BARNS) . 6.051E-01 4 3.491E+00 5 6 6.143E+00 7 8.217E+00 8 9.796E+00 1.105E+01 q 10 1.225E+01 11 1.341E+01 1.652E+01 12 GROUP CONSTANTS 71.5695 FOR MF26 AND MT516 PAIR FINAL CROSS SECTIONS VS INITL GROUP LEGENDRE ORDER GROUP 1.210E+00 4 З 4 HEAT 4.503E+05 5 3 6.982E+00 5 HEAT 5.276E+06 . . 3.303E+01 12 З HEAT 1.842E+08 12 GROUP CONSTANTS 71.6745 FOR MF23 AND MT602 ABST GAMMA SIGMA HEATING GROUP (BARNS) (EV-BARNS) 2.547E+03 1 1.289E+08 2 4.829E+02 8.101E+07 3 2.368E+01 1.570E+07 5.937E+00 7.955E+06 4

7

5 6 7 8 9 10 11 12	2.140E+0 1.244E+0 8.503E-0 6.398E-0 5.150E-0 4.276E-0 3.637E-0 2.563E-0	XX 5.164E+C XX 4.278E+C XX 3.786E+C XX 3.496E+C XX 3.331E+C XX 1.96E+C XX 3.083E+C XX 2.916E+C	96 96 96 96 96 96 96 96 96			
GROUP (FOR MF2	CONSTANTS	621 HEAT				71.8265
GAMMA GRDUP 1 2 3 4 5 6 7 8 9 9	HEATING (EV-BARN 1.293E+C 8.305E+C 2.208E+C 1.969E+C 2.673E+C 3.945E+C 5.509E+C 7.236E+C 9.081E+C 9.081E+C	IS) 98 97 97 97 97 97 97 97 97				
10	1.112E+(1.335E+()8)8				
12	2.211E+C	08				
						71.7485
	•••••					****
DTFR	. PRODUCE	OTF FORMAT	FROM GROUPR	OUTPUT		71.7915
INPUT	GENDE UNI	T		2:	3	
OUTPUT	UNIT			. 24	4	
INPUT I	PENDF UNI	IT		. ()	
PRINT	OPTION ((D MIN. 1 MAX	<)	••	1	
FILM D	PTION (NO	DNE, 1/PLOT		(2	
	PIION (O	TABLE, 1 SI	ΕΡ.)	(
NUMBER	OF NEUT	ON CONIDS	• • • • • • • • • • • • •		כ ת	
POSITI	ON OF TO		• • • • • • • • • • • • •		4	
POSITI	ON OF IN	GROUP			5	
TABLE	LENGTH			. 10	6	
EDIT C	ROSS SECT	TIONS		. NAME PO	DSITION REACT	ION MULTIPLICITY
NUMBED		IN TARIES		PHEAT	1 621	1
NOMBER		IN TABLES .			5	
MAT= IL= 1	1 ISD= TABLE 12	H SIG	ZERD ND= 1 MAT= 13	TEMP= O.	0.	
4.13	26E+05 -:	3.7403E-03	0.	4.9959E-02	4.1123E-03	0.
0.		5.	0.	0.	0.	0.
0.		J. 5 03315-03	0.	0.	3.48841+05	-2.7037E-03
0. 0.		D.0321E-02	0	2.10092-03	0.	0.
0.		D.	3.2740E+05	-2.3184E-03	0.	6.5317E-02
2.06	63E-03	3.5675E-03	2.1674E-03	0.	0.	0.
0.		D.	0.	0.	0.	0.
3.04	012+05 -	1.9501E-03	0.	7.1605E-02	2.7252E-03	4.5910E-03,
0	112-03	2.2030E-U3 N	0.	0.	U. 2 79015±05	U. -1 5979F-03
ŏ.		7.9779E-02	3.7626E-03	6.1295E-03	4.7009E-03	3.7957E-03
2.47	78E-03	D.	0.	0.	0.	0.
Ο.		0.	2.5188E+05	-1.2009E-03	0.	9.0717E-02

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6.3377E-03 5.5276E-03 8.6030E-03 5.0001E-03 4.1302E-03 2.8022E-03 ο. ο. Ο. Ο. Ο. Ο. 2.1901E+05 -7.5223E-04 ο. 1.0629E-01 8.9122E-03 1.2960E-02 9.0525E-03 6.9481E-03 5.6558E-03 4.7847E-03 3.3717E-03 ο. ο. O., 0. Ο. 1.7844E+05 -3.3080E-04 1.3071E-01 1.8756E-02 2.1796E-02 0. 1.4155E-02 1.0549E-02 8.4713E-03 7.1196E-03 6.1633E-03 4.4949E-03 0. ο. о. 1.2397E+05 -4.6426E-05 1.7774E-01 Ο. ο. 4.3226E-02 4.4858E-02 1.5488E-02 2.6576E-02 1.9396E-02 1.2980F-02 1.1216E-02 9.8948E-03 7.4090E-03 ο. Ο. ο. 7.0497E+04 -7.3820E-08 Ο. 2.4930E-01 8.1577E-02 6.7262E-02 3.2719E-02 2.4173E-02 2.0211E-02 1.7934E-02 1.6494E-02 1.5644E-02 1.4867E-02 3.9136E-01 1.5187E-02 ο. Ο. 2.2817F+04 3.2150E-07 3.3367E-Q1 ο. 1.6772E-01 6.7295E-02 3.6706E-02 2.5581E-02 1.5988E-02 1.3469E-02 1.9669E-02 1.1641E-02 1.0250E-02 7.6122E-03 ο. 4.4889E+03 5.8176E-05 Ο. 5.2555E-01 5-2550E-01 5.7690E-02 Ο. 0. ο. Ο. Ο. ο. 0. Ο. Ο. Ο. IL= 2 TABLE 12 GP 16 PDS, MAT= 1 IZ= 1 TEMP= 0. ο. Ο. 0. ο. 4.0823E-03 ο. о. Ο. ο. ο. ο. ο. ō. Ο. ο. ο. 0. Ο. -9.7256E-03 -7.8220E-03 -6.5324E-03 -5.6042E-03 -4.9071E-03 -4.3633E-03 -3.2954E-03 O. Ο. Ο. 0. ο. 4.8415E-02 -1.9662E-02 ο. IL= 3 TABLE 12 GP 16 PDS, MAT= 1 IZ= 1 TEMP= 0. Ο. ο. ο. c. 4.0229E-03 ο. Ο. Ο. ο. е. Ο. Ο. ο. Ο. 0, ο. ο. ο. -6.7256E-04 0. ο. ο. Ο. ο. 5.5845E-02 3.7941E-03 Ο. Ο. Ο. Ο. Ο. ο. Ο. Ο. Ο. 0. IL= 4 TABLE 12 GP 16 POS, MAT= 1 IZ= 1 TEMP= O. ο. ο. Ο. ο. 3.9352E-03 0. 0. Ο. Ο. о. Ο. ο. Ο. ο. ο. ο. ο. ο. 1.1390E-03 0. ο. Ο. ο. ο. 5.9479E-03 -2.1019E-03 о. о. ο. ο. Ο. ο. ο. Ο. Ο. Ο. IL= 5 TABLE 12 GP 16 PDS, MAT= 1 IZ= 1 TEMP= 0. ο. ο. ο. ο. 3.8207E-03 Ο. 2.4938E-04 O. ο. Ο. Ο. ο. 5.1587E-04 -3.0239E-04 ο. Ο. Ο. 0. ο. ο. Ο. Ο. ο. 0. MAT = 92 iSO=U SIGZERO NO= 1 TEMP= 0. IL= 1 TABLE 12 GP 16 POS, MAT= 92 IZ= 1 TEMP= 0. 2.2110E+08 -1.6259E+01 0. 2.1029E+01 3.9740E-01 0.

0. 0. 1.3350E+08 -1.3044E+01 ο. 0. 0. 0. 0. 0. 7). Ο. . . 6.9779E-01 0. 1.2926E+08 2.5467E+03 0. 2.7679E+03 2.2116E+02 4.8636E+00 0. Ο. Ο. Ο. ο. ο. o. Ο. ο. ο. IL= 5 TABLE 12 GP 16 PDS. MAT= 92 IZ= 1 TEMP= 0. ο. ο. Ο. ο. 3.7071E-01 ο. 0. 0. ο. Ο. Ο. 0. 0. 0. Ο. 0. Ο. Ο. . 2.2866E-02 0. 5.4900E+01 -1.6996E-02 ο. 75.0885 MATXSR... PRODUCE A MATXS FORMAT DUTPUT FILE 75.1285 INPUT GENOF UNIT INPUT GAMOUT UNIT OUTPUT MATXS UNIT 0 23 25
 PRINT OPTION (O MIN, 1 MAX)
 1

 FILE VERSION NUMBER
 1

 USER IO
 1

 NPART
 1

 NTYPE
 1
 1 SET IDENTIFICATION 12-GROUP PHOTON INTERACTION LIBRARY NAMES OF DATA TYPES GSCAT FOR DATA TYPE 1 IFOPT 1 NSBLK MAX. LEGENDRE ORDER ND. MATERIALS 1 3 2 I INP I OUTP 1 MATERIAL MAX. MAX. NAME TEMPS SIGZ IMAT - - - - - - - -- - - - -----____ н 1 1 1 U 1 1 92 1

76.4995

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*	*	*		F	I	LI	Ε		M	A	T)	κs	;							*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
*	*	*		V	Ξ١	R	5				1									*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
*	*	*	ł	U:	51	EF	2		т	2	L	١S	iL	1	Ν	J	D	4		*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
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12-GROUP PHOTON INTERACTION LIBRARY

NAME

- - - -

GAMA

NAME

- - - -

GSCAT

NUMBER OF PARTICLES FOR WHICH GROUP STRUCTURES ARE GIVEN

NUMBER OF DATA TYPES PRESENT IN SET NUMBER OF WORDS IN SET HOLLERITH IDENTIFICATION RECORD

NGRP

- - - -

NMAT

- - - -

MAXIMUM ENERGY 2.00000E+07

9.00000E+06 8.00000E+06

7.00000E+06

6.00000E+06

5.00000E+06

4.00000E+06

3.00000E+06

2.00000E+06

1.00000E+06 5.00000E+05

1.00000E+05

1.00000E+04

TEMP

- - - -

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Ο.

SIGZ

1.00E+10

1.00E+10

HMATNM

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INCIDENT PARTICLES

DUTGOING PARTICLES

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NINP

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12

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FILE CONTROL PARAMETERS

NPART

NTYPE

NHOLL

FILE DATA PARTICLE

- - - - - - - -

DATA TYPE

1

1

PARTICLE 1

GROUP

1 2

3 4

5

6

7

8

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10

11

12 EMIN

MATERIAL

1

2

NSBLK

DATA TYPE CONTROL

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GROUP STRUCTURES

FILE DESCRIPTION

MATERIAL CONTROL

HMAT	н
AMASS	9.992E-01
TEMP	Ο.
SIGZ	1.000E+10
IMAC	1
N1DR	7
N10B	1
N20B	3

VECTOR CONTROL

REACTION	REACTION ID	BLOCK	FIRST GROUP	LAST GROUP
1	GWTO	1	1	12
2	GTOTO	1	1	12
3	GCDH	1	9	:2
4	GINCH	1	1	12
5	GPAIR	1	1	9
6	GABS	1	12	12
7	GHEAT	1	1	12

VECTOR BLOCK 1

GROUP	GWTO	GTOTO	GCDH	GINCH	GPAIR	GABS	GHEAT
1	3.855E+04	4.996E-02		4.622E-02	3.740E-03		4.133E+05
2	1.178E+04	6.032E-02		5.762E-02	2.704E-03		3.488E+05
3	1.336E+04	6.532E-02		6.300E-02	2.318E-03		3.274E+05
4	1.542E+04	7.160E-02		£.965E-02	1.950E-03		3.046E+05
5	1.824E+04	7.978E-02		7.8 8E - 02	1.598E-03		2.799E+05
6	2.232E+04	9.072E-02		8.952E-02	1.201E-03		2.519E+05
7	2_878E+04	1.063E-01		1.055E-01	7.523E-04		2.190E+05
8	4.056E+04	1.307E-01		1.304E-01	3.308E-04		1.784E+05
9	6.934E+04	1.777E-01	2.510E-06	1.777E-01	4.641E-05		1.240E+05
10	6.934E+04	2.493E-01	1.004E-05	2.493E-01			7.050E+04
11	1.610E+05	3.914E-01	1.386E-04	3.912E-01			2.282E+04
12	3.331E+04	5.256E-01	1.235E-03	5.243E-01		1.926E-05	4.489E+03

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MATRIX ** GCDH **

FINAL		XSEC VS LEGE	NORE ORDER	00053 0	00050 3
		URUER U	URUER 1	URDER 2	URDER 3
9	9	2.510E-06	2.510E-06	2.510E-06	2.510E-06
10	10	1.004E-05	1.004E-05	1.0046-05	1.004E-05
11	11	1.386E-04	1.386E-04	1.384E-04	1.381E-04
12	12	1.235E-03	1.220E-03	1.193E-03	1.158E-03

MATRIX ** GINCH **

FINAL	INITL	XSEC VS LEG	ENDRE ORDER		
GROUP	GROUP	ORDER O	ORDER 1	ORDER 2	ORDER 3
				`	
1	1	4.112E-D3	4.082E-03	4.023E-03	3.935E-03
2	2	1.620E-03	1.616E-03	1.008E-03	1.597E-03
	1	2.101E-03	2.073E-03	2.019E-03	1.940E-03
3	3	2.066E-03	2.060E-03	2.047E-03	2.029E-03
	2	3.568E-03	3.540E-03	3.484E-D3	3.401E-03
	1	2.167E-03	2.121E-03	2.031E-03	1.899E-03

•						
12	1 12 11	7.612E-03 5.243E-01 5.769E-02	-3.295E-03 4.719E-02 -1.966E-02	-6.726E-04 5.465E-02 3.794E-03	1.139E-03 4.790E-03 -2.102E-03	
MATRIX	** GPAIR	**				
F I NAL GROUP	INITL GROUP	XSEC VS IN -0	ITL GROUP -1	-2	-3	-4
10	9 4	9.282E-05 3.900E-03	6.615E-04 4.637E-03	1.505E-03 5.407E-03	2.402E-03 7.461E-03	3.196E-03
*** GS(4 CAT *** M	3.900E-03	4.637E-03	5.407E-03	7.481E-03	

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MATERIAL CONTROL

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HMAT	U
AMASS	2.360E+02
TEMP	0.
SIGZ	1.000E+10
IMAC	1
N10R	7
N10B	1
N20B	3

VECTOR CONTROL

REACTION	REACTION ID	BLOCK	FIRST GROUP	LAST GROUP
1	GWTO	1	1	12
2	GTDTO	1	1	12
3	GCOH	1	1	12
4	GINCH	1	1	12
5	GPAIR	1	1	9
6	GABS	1	1	12
7	GHEAT	1	1	12

VECTOR BLOCK 1

GROUP	GWTO	GTOTO	GCOH	GINCH	GPAIR	GABS	GHEAT
1	3.848E+O4	2.103E+01	1.021E-02	4.247E+00	1.652E+01	2.563E-01	2.211E+08
2	1.178E+04	1.909E+01	1.793E-02	5.296E+00	1.341E+01	3.637E-01	1.335E+08
3	1.335E+04	1.849E+01	2.306E-02	5.788E+00	1.225E+01	4.276E-01	1.112E+08
4	1.542E+04	1.800E+01	3.079E-02	6.401E+00	1.105E+01	5.150E-01	9.081E+07
5	1.823E+04	1.766E+01	4.313E-02	7.183E+00	9.796E+00	6.398E-01	7.236E+07
6	2.232E+04	1.735E+01	6.483E-D2	8.222E+00	8.217E+00	8.503E-01	5.509E+07
7	2.877E+04	1.718E+01	1.085E-01	9.688E+00	6.143E+00	1.244E+00	3.945E+07
8	4.055E+04	1.781E+01	2.188E-01	1.196E+01	3.491E+00	2.140E+00	2.673E+07
9	6.932E+04	2.345E+01	6.803E-01	1.623E+01	6.051E-01	5.937E+00	1.969E+07
10	6.932E+04	4.882E+01	2.623E+00	2.252E+01		2.368E+01	2.208E+07
11	1.610E+05	5.452E+O2	2.953E+01	3.280E+01		4.829E+02	8.305E+07
12	3.330E+04	2.768E+03	1.844E+O2	3.676E+01		2.547E+03	1.293E+08

MATRIX ** GCOH **

FINAL	INITL	XSEC VS LEG	ENDRE ORDER		
GROUP	GROUP	ORDER O	DRDER 1	ORDER 2	ORDER 3

1	1	1.021E-02	1.021E-02	1.020E-02	1.019E-02
2	2	1.793E-02	1.792E-02	1.789E-02	1.785E-02
3	3	2.306E-02	2.304E-02	2.300E-02	2.293E-02
4	4	3.Q79E-02	3.075E-02	3.067E-02	3.057E-02
5	5	4.313E-02	4.305E-02	4.291E-02	4.271E-02
6	6	6.483E-02	6.466E-02	6.435E-02	6.393E-02
7	7	1.0858-01	1.080E-01	1.072E-01	1.062E-01
8	8	2.188E-01	2.170E-01	2.142E-01	2.107E-01
9	9	6.803E-01	6.664E-01	6.477E-01	6.275E-01
10	10	2.623E+00	2.495E+00	2.374E+00	2.255E+00
11	11	2.953E+01	2.541E+01	2.232E+01	1.922E+D1
12	12	1.844E+O2	1.292E+02	9.996E+01	7.350E+01

MATRIX ** GINCH **

FINAL	INITL	XSEC VS LE	GENORE ORDER		
GROUP	GROUP	ORDER O	ORDER 1	ORDER 2	ORDER 3
1	1	3.872E-01	3.845E-01	3.790E-01	3.710E-01
2	2	1.511E-01	1.507E-01	1.500E-01	1.490E-01
	1	1.926E-01	1.901E-01	1.851E-01	1.778E-01
3	3	1.926E-01	1.920E-01	1.909E-01	1.892E-01
	2	3.274E-01	3.248E-01	3.197E-01	3.122E-01
	1	1.987E-01	1.945E-01	1.862E-01	1.742E-01
•					
•					
•					
	1	6.978E-01	-3.021E-01	-6.166E-02	1.044E-01
12	12	3.676E+01	-3.149E-02	2.240E+00	-8.225E-01
	11	4.864E+00	-1.777E+00	3.398E-01	-1.769E-01

MATRIX ** GPAIR **

FINAL	INITL	XSEC VS IN	ITL GROUP			
GROUP	GROUP	-0	- 1	-2	-3	-4
10	9	1.210E+00	6.982E+00	1.229E+01	1.643E+01	1.959E+01
	4	2.210E+01	2.450E+01	2.682E+01	3.303E+01	

INDEX OF MATXS FILE

76.6025

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FILE DESCRIPTION

12-GROUP PHOTON INTERACTION LIBRARY

DATA TYPES ON FILE

	NAME	LDCT
1	GSCAT	0

**** DATA TYPE 1 ****** GSCAT *******

MATERIALS ON FILE FOR THIS DATA TYPE

	NAM	Ε	TEMP	SIGZ	LOCA
		-			
1	н	Ο.		1.00E+10	0
2	U	0.		1.00E+10	7

VECTOR REACTION TYPES ON FILE BY MATERIAL

	1	2
1	GWTO	GWTO
2	GTDTO	GTDTO
3	GCDH	GCDH
4	GINCH	GINCH
5	GPAIR	GPAIR
6	GABS	GABS
7	GHEAT	GHEAT

MATRIX REACTION TYPES ON FILE BY MATERIAL

	1	2
1	GCOH	GCDH
2	GINCH	GINCH
3	GPAIR	GPAIR

INDEX COMPLETE

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76.6355 76.671S ********* .

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IV. EXAMPLE 4, TEST OF COVARIANCE PROCESSING.

MOUNT ENDE V TAPE 511 ON UNIT 20.

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0
5
*MODER*
20 -21
*RECONR*
-21 -22
+U-235 10% PENDF FOR ERRORR TEST PRUBLEM FROM T511+/
1395 0 0
.10 0. 6 0 .1
0/____
*ERRDRR*
-21 -22 0 23 0
1395 19 1 1
3 0 0 0 33
1
.
1.EO 1.E3
*GROUPR*
*GROUPR*
-21 -22 0 24
1395 3 0 3 0 1 1 1
*U-235 MULTIGROUP NUBAR CALCULATION*/
ο.
1.E10
3 452 *TOTAL NUBAR*/
0/
0/
*ERRORR*
-21 0 24 25 23
1395 1 1 1
```

0 31 1.EO 1.E1 1.E2 1.E3 1.E4 1.E5 1.E6 1.E7 *STOP* 1 * * * ************************ * * * \$\$ \$\$ \$\$ \$\$\$\$\$ \$\$ \$\$ * VERS. 10/81 * * ÷ - 55 \$\$ \$\$ \$\$\$ \$\$ \$\$\$\$\$\$\$ * NUCLEAR * RAN AT LANL CRDSS SECTION * PROCESSING * \$\$\$\$ \$\$ \$\$ \$\$ - \$\$ \$\$\$\$ * * ON MACH. T \$\$ \$\$ \$\$ \$\$\$\$ \$\$ ON 11/06/81 \$\$ \$\$ \$\$ * * \$\$\$\$\$\$\$ \$\$\$\$\$\$\$ 222 SYSTEM * 22 * * AT 19:25:12 * * 22 22 22222 22222 \$\$ * * ********************** MODER...CHANGE THE MODE OF IN ENDE BAPE OR NUOY OUTPUT TAPE .6335 INPUT UNIT (+ FOR CODED, - FOR BB) ... DUTPUT UNIT (+ FOR CODED, - FOR BB) ... 20 -21 TAPE LABEL -----ENDF/B-V TAPE 511 (STANDARDS MATERIALS) 11.4565 **** RECONR...RECONSTRUCT POINTWISE CROSS SECTIONS IN PENDE FORMAT 11.5075 UNIT FOR ENDF/B TAPE -21 UNIT FOR PENDE TAPE - 22 LABEL FOR PENOF TAPE U-235 10% PENDE FOR ERRORE TEST PROBLEM FROM T511 TAPE LABEL ENDF/B-V TAPE 511 (STANDARDS MATERIALS) MATERIAL TO BE PROCESSED 1395 RECONSTRUCTION TOLERANCE . 100 0.K 6 2.000 MAX RESONANCE-INTEGRAL ERROR 1.000E-01 DESCRIPTIVE CARDS FOR PENDE TAPE PENDE TAPE PREPARED BY THE NUOY NUCLEAR DATA PROCESSING SYSTEM PROCESSING MAT 1395 92-U -2350BNL EVAL-APR77 M.R.BHAT PDINTS IN INITIAL UNIONIZED GRID = 1662 POINTS ADDED BY LINEARIZATION = 15 16.6185 ESTIMATED MAXIMUM ERROR DUE TO RESONANCE INTEGRAL CHECK (ERRMAX.ERRINT) AND SIGNIFICANT FIGURE TRUNCATION (NDIGIT) UPPER ELASTIC PERCENT ERROR CAPTURE PERCENT ERROR

3

ENERGY INTEGRAL RES-INT SIG-FIG INTEGRAL RES-INT SIG-FIG 1.00E+00 1.48E+01 .002 3.18E+00 0.000 . 428 1.12F+01 0.000 1.29E+01 1.01E+01 .337 0.000 3.25E+01 1.315 0.000 3.20E+01 1.38E+01 1.008 0.000 4.23E+01 3.409 0.000 1.925 8.20E+01 1.26E+01 0:000 2.31E+01 8.113 0.000 2.50E+04 6.90E+01 .002 0.000 2.69F+01 .029 0.000 POINTS ADDED BY RESONANCE RECONSTRUCTION = 149 POINTS AFFECTED BY RESONANCE INTEGRAL CHECK = 347 POINTS AFFECTED BY SIGNIFICANT FIGURE REDUCTION = POINTS REMOVED BY BACKTHINNING = 48 o 48 FINAL NUMBER OF RESONANCE PDINTS = 941 33.0815 ERRORR... PRODUCE CROSS SECTION COVARIANCES. 33.1155 UNIT FOR ENOF/B TAPE -21 UNIT FOR PENDF TAPE UNIT FOR INPUT GENDF TAPE -22 0 UNIT FOR OUTPUT COVARIANCE TAPE 23 UNIT FOR INPUT COVARIANCE TAPE 0 MATERIAL TO BE PROCESSED 1395 19 1 3 0 TEMPERATURE TEMPERATURE READ OPTION (O CALC, 1 READ 2 COMBD) ENOF COVARIANCE FILE TO BE PROCESSED ο. 0 33 NDTE TO USER. COVARIANCES OF THE STANDARD REACTION MTS= 18 WITH THE REACTION (MAT=1342.MT= 18) WILL NOT BE CALCULATED. LTY=3. NDTE TO USER. COVARIANCES OF THE STANDARD REACTION MTS= 18 WITH THE REACTION (MAT= 1361, MT= 18) WILL NOT BE CALCULATED. LTY=3. NOTE TO USER. COVARIANCES OF THE STANDARD REACTION MTS= 18 WITH THE REACTION (MAT=1399.MT= 18) WILL NOT BE CALCULATED. LTY=3. NDTE TO USER. COVARIANCES OF THE STANDARD REACTION MTS= 18 WITH THE REACTION (MAT=1399, MT=102) WILL NOT BE CALCULATED. LTY=3. COMPUTING MULTIGROUP CROSS SECTIONS 35.0445 REPACKING UNION STRUCTURE (= USER STRUCTURE) HAS 9 GROUPS 1 1.00000E+00 - 1.80000E+00 1.8000CE+00 - 5.00000E+00 2 5.00000E+00 - 1.00000E+01 1.000D0E+01 - 2.00000E+01 З 4 2.00000E+01 - 4.0000E+01 4.00000E+01 - 8.00000E+01 5 6 4.00000E+01 - 2.00000E+02 2.00000E+02 - 4.00000E+02 4.00000E+02 - 1.00000E+03 7 8 WEIGHT FUNCTION..... 1/E FOR ALL L GROUP AVERAGING COMPLETED 36.0575

REPACKING COVARIANCES CALCULATED FOR 2 REACTIONS AND 9 GROUPS 36.8945 TABLE OF MULTIGROUP CROSS SECTIONS GROUP CRDSS SECTION NO. ENERGY MT 18 MT 102 - - - - -1.0000E+00 4.1646E+01 1.0751E+01 1 1.8000E+00 1.6744E+01 1.3156E+01 2 3 5.0000E+00 5.2733E+01 3.5074E+01 4 1.0000E+01 5.2362E+01 4.4286E+01 2.0000E+01 5 4.9641E+01 3.0819E+01 4.0000E+01 4.1728E+01 1.9486F+01 6 7 8.0000E+01 2.1709E+01 1.2707E+01 2.0000E+02 1.7192E+01 8.1314E+00 8 4.0000E+02 1.1587E+01 4.7376E+00 q RELATIVE COVARIANCE (MT 18 , IG , MT 18 , IGP) 37.4165 IGP IG +0 + 1 +2 - - -- - - -- - - -- - - -1 5.000E-04 4.000E-04 4.001E-04 4.001E-04 3.999E-04 4.000E-04 4.000E-04 4.000E-04 4.000E-04 7 1 4.000E-04 5.000E-04 4.001E-04 4.001E-04 3.999E-04 4.000E-04 2 1 2 7 4.000E-04 4.000E-04 4.000E-04 3 4.001E-04 4.001E-04 7.001E-04 6.001E-04 5.998E-04 5.998E-04 3 5.998E-04 5.998E-04 5.998E-04 7 4 4.001E-04 4.001E-04 6.001E-04 7.001E-04 5.998E-04 5.998E-04 1 5.998E-04 5.998E-04 5.998E-04 4 7 5 3.999E-04 3.999E-04 5.998E-04 5.998E-04 7.998E-04 7.001E-04 5 7 7.001E-04 7.001E-04 7.001E-04 4.000E-04 4.000E-04 5.998E-04 6 5.998E-04 7.001E-04 9.998E-04 1 8.999E-04 6 8.999E-04 8.999E-04 7 4.000E-04 5.998E-04 5.998E-04 7.001E-04 8.999E-04 7 1 4.000E-04 8.999E-04 7 7 9.998E-04 8.999E-04 8 1 4.000E-04 4.000E-04 5.998E-04 5.998E-04 7.001E-04 8.999E-04 8.999E-04 9.998E-04 3,999E-04 7 8 4.000E-04 5.998E-04 7.001E-04 8.999E-04 9 1 4.000E-04 5.998E-04 9 7 8.999E-04 8.999E-04 9.998E-04 RELATIVE COVARIANCE (MT 18 , 3G , MT102 . IGP) 37.4415 ZERD RELATIVE COVARIANCE (MT102 . IG . MT102 , IGP) 37.4505 IG IGP +0 +2 ÷1 - - -- - -- - - -- - - -1.440E-02 1.440E-02 1.440E-02 1 1.690E-02 1.440E-02 1.440E-02 1 7 2.500E-03 2.500E-03 2.500E-03 1 1.440E-02 1.690E-02 1.440E-02 1.440E-02 1.440E-02 1.440E-02 2 1 8 7 5.000E-03 6.400E-03 5.000E-03 2.500E-03 2.500E-03 2.500E-03 2.500E-03 2.500E-03 2.500E-03 9 1 5.000E-03 5.000E-03 6.400E-03 9 37.5745 GROUPR...COMPUTE SELF-SHIELDED GROUP-AVERAGED CROSS-SECTIONS 37.6085

UNIT FOR ENDF/B TAPE UNIT FOR PENDF TAPE UNIT FOR INPUT GOUT TAPE UNIT FOR OUTPUT GOUT TAPE MAT TO BE FROCESSED NEUTRON GROUP STRUCTURE OPTION -21 -22 0 24 1395 3 0 3 PRINT OPTION (O MIN, 1 MAX) 0 1 RUN TITLE U-235 MULTIGROUP NUBAR CALCULATION TEMPERATURES (KELVIN) ZERO SIGMA ZERDES INFINITY NEUTRON GROUP STRUCTURE.....LASL 30 GROUP 1.3900E-04 - 1.5200E-01 1.5200E-01 - 4.1400E-01 1 2 4.1400E-01 - 1.1300E+00 3 . 28 1.2000E+07 - 1.3500E+07 1.3500E+07 - 1.5000E+07 1.5000E+07 - 1.7000E+07 . 29 30 WEIGHT FUNCTION..... 1/E FOR ALL L PROCESSING MAT 1395 PENDE TAPE PREPARED BY THE NJDY NUCLEAR DATA PROCESSING SYSTEM GROUP CONSTANTS AT T=ZERO DEG K 40.7915 FOR MF 3 AND MT452 TOTAL NUBAR ENRGY GROUP CONSTANTS AT GROUP INFINITE DILUTION 1 2.437E+00 2 2.437E+00 З 2.437E+00 . • 28 4.221E+00 29 4.433E+00 30 4.678E+00 41.3795 ERRORR... PRODUCE CROSS SECTION COVARIANCES 41.4265 UNIT FOR ENDF/B TAPE -21 UNIT FOR PENDE TAPE UNIT FOR INPUT GENDE TAPE 0 24 UNIT FOR DUTPUT COVARIANCE TAPE UNIT FOR INPUT COVARIANCE TAPE 25 23 MATERIAL TO BE PROCESSED 1395 NEUTRON GROUP OPTION 1 PRINT OPTION (O MIN, 1 MAX)

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READ C	OPTION (O COVARIANCI	CALC. 1 E FILE TO	READ, 2 CO D BE PROCES	MBO) SEO .	0 31			
NEUTRO	ON GROUP	STRUCTUR	EREAD	IN				
1 2 3 4 5 6 7	1.00 1.00 1.00 1.00 1.00 1.00 1.00	0000E+00 0000E+01 0000E+02 0000E+03 0000E+03 0000E+04 0000E+05 0000E+06	- 1.00000E - 1.00000E - 1.00000E - 1.00000E - 1.00000E - 1.00000E - 1.00000E	+01 +02 +03 +04 +05 +06 +07			REPACKING	
UNION STRUCTURE HAS 15 GROUPS								
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15	1.0 1.0 1.0 1.0 1.0 2.0 5.0 1.0 5.0 1.5 2.0 3.0 4.0 6.0 8.0	0000E+00 0000E+01 0000E+02 0000E+03 0000E+03 0000E+05 0000E+05 0000E+05 0000E+06 0000E+06 0000E+06 0000E+06	- 1.00000 - 1.00000 - 1.00000 - 1.00000 - 1.00000 - 2.00000 - 5.00000 - 1.50000 - 1.50000 - 3.00000 - 3.00000 - 4.00000 - 6.00000 - 8.00000 - 1.00000	+01 +02 +03 +04 +05 +05 +05 +06 +06 +06 +06 +06 +06 +06 +06 +06				
	0.0	00002.00		•			REPACKING	
COVARIANCES CALCULATED FOR 1 REACTIONS AND 15 GROUPS44.558STABLE DF MULTIGROUP CROSS SECTIONS								
GROUP ND.	LOWER ENERGY	CRDS MT45	S SECTION 2					
1 2 3 4 5 6 7	1.0000E+ 1.0000E+ 1.0000E+ 1.0000E+ 1.0000E+ 1.0000E+ 1.0000E+	00 2.43 01 2.43 02 2.43 03 2.43 04 2.43 05 2.47 06 2.91	67E+00 67E+00 67E+00 67E+00 82E+00 77E+00 77E+00 71E+00					
RELAT	IVE COVAR	NIANCE (MT452 . IG	. MT452 ,	IGP)		44.9995	
IG	IGP	+0	+1	+2				
1	1 4.8	15E-05	4.815E-05	4.536E-05	4.536E-05	4.536E-05	4.111E-05	
1 2	7 2.7	75E-05 315E-05	4.815E-05	4.536E-05	4.536E-05	4.536E-05	4.111E-05	
2 3	7 2.7	75E-05 536E-05	4.536E-05	4.298E-05	4.298E-05	4.298E-05	3.934E-05	
3	7 2.7	784E-05	4 536E-05	4.298F-05	4.298F-05	4.298E-05	3.934E-05	
4	7 2.7	784E-05	4 5265-05	4 2995-05	4 2985-05	4 2985-05	3.9345-05	
5	7 2.3	784E-05	4.5362-05	4.2982-03	4.2982 05		3.5342 05	
6	1 4.	111E-05	4.111E-05	3.934E-05	3.9341.05	J.934E-05	3.8/21-05	
6 7	7 2.0	806E-05 775E-05	2.775E-05	2.784E-05	2.784E-05	2.784E-05	2.806E-05	
7	7 3.	145E-Q5	********	********		*******	45.0915	

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REL. CDV. DPTION (O ABS, 1 REL) 1

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APPENDIX D

PREPROCESSING PROGRAM FOR IBM/CDC CODE CONVERSION

This program requires approximately 15 seconds of CDC-7600 central-processor time to convert the entire code system.

000000		<pre>PROGRAM CCC (TAPE1,TAPE2,TAPE3) ************************************</pre>
C C C 11 11		IBM DESIRED. IMACH=1
	100 1 10 120	READ AND WRITE MACHINE-INDEPENDENT LINES. READ (NIN, 10) (IA(I),I=1,23) IF (EDF(NIN)) 230,110 IF (IA(1).NE.IC) GD TD 120 IF (IA(2).EQ.IBB) GO TO 120 IF (IA(2).EQ.IIBM) GD TD 130 IF (IA(2).EQ.ICDC) GD TO 180 WRITE (NDUT,10) (IA(I),I=1,23) GD TD 100
c	130 140 150	READ AND WRITE IBM LINES. IIB=IIB+1 GD TD 170 READ (NIN,10) (IA(I),I=1,23) IF (EDF(NIN)) 230,150 IF (IA(2).EQ.IIBM) GO TO 120 IF (IMACH.NE.1) GD TD 160 IA(1)=IB GD TD 170 IA(1)=IC
i c	170	WRITE (NOUT, 10) (IA(I), I=1,23) GD TD 140
С	180 190 200	READ AND WRITE COC LINES. ICD=ICD+1 GD TD 220 READ (NIN,10) (IA(I).I=1,23) IF (EOF(NIN)) 230,200 IF (IA(2).EQ.ICDC) GO TO 120 IF (IMACH.NE.2) GD TD 210 IA(1)=IB
с	210 220	GD TD 220 IA(1)=IC WRITE (NOUT.10) (IA(I),I=1.23) GD TO 190
c c	230	CCC FINISHED. WRITE (NSYSD,20) IIB,ICD STOP
	10 20	FORMAT (A1.A2.20A4.A3) FORMAT (//I5.22H IBM BLOCKS PROCESSED 1 /I5.22H COC BLOCKS PROCESSED 2 //26H CODE CONVERSION COMPLETE.) END

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