Generating Parallel Correlated Transition Frequencies for a Markov Chain

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Abstract

This paper describes an algorithm and a FORTRAN program called MCHAIN for simulating $k$ parallel Monte Carlo replications of a Markov chain using rotation sampling. This method of sampling produces $k$ sample transition frequency vectors with a desirable structure of statistical dependence among them. In particular, these sample vectors can be used to estimate the probability that, say, $n_1, \ldots, n_r$ transitions of types $1, \ldots, r$ occur during a first passage from state $a$ to state $b$ with a variance of the estimate of $O(1/k)$ and a computation time $O(k)$ as $k \to \infty$. This compares favorably with the case of independent replications wherein the estimate would have a variance $O(1/k)$ and computation time $O(k)$ as $k \to \infty$. An example including a sample driver program are presented to illustrate how MCHAIN works in practice.

Key Words

Markov chain, Monte Carlo methods, rotation sampling, simulation, variance reduction
**Introduction**

Let $p_n = ||p_{ij}||$ denote a positive recurrent aperiodic Markov Chain with state space $S = (0,1,...,n)$. Let $N_{ij}$ denote the number of one-step transitions that occur from state $i$ to state $j$ during a first passage from state $a$ to state $b$ for $a,b \in S$ on an arbitrarily selected replication and let $N_{ij}(\ell)$ denote this quantity observed on replication $\ell$. The purpose of this paper is to describe an algorithm and a FORTRAN program called MCHAIN that generates $k$ sample data sets $\{N_{ij}(1); i,j \in S\},...,\{N_{ij}(k); i,j \in S\}$ from the chain $p_n$ with a specialized dependence among sets that allows one to estimate quantities of interest with greater statistical efficiency than independent data sets allow.

As an example of the use of these data sets, suppose one wants to compute

$$p(n_1,...,n_r) = \text{probability that during a first passage from a to b, } n_1,...,n_r \text{ one-step transitions occur from } i_1 \text{ to } j_1,...,i_r \text{ to } j_r,$$

For Markov chains with relatively general structure, convenient analytical representations are not available for (1) so that computation is not possible. In fact no convenient representations are available for chains with special structure but arbitrary $r > 1$. One way to overcome this absence of representation is to employ Monte Carlo methods. Assume there are $k$ simulated data sets $\{N_{ij}(1); i,j \in S\},...,\{N_{ij}(k); i,j \in S\}$ available to estimate (1), each simulation beginning with a departure from state $a$ and ending upon entry into state $b$. Let

$$k_{n_1,...,n_r} = \sum_{m=1}^{k} \prod_{j=1}^{r} \delta(N_{ij}(\ell) - n_{ij})$$

(2)
where
\[
\delta(x) = \begin{cases} 
1 & \text{if } x = 0 \\
0 & \text{otherwise} 
\end{cases}
\]

If \{N_{ij}(\ell); i,j \in S\} \ell = 1,\ldots,k \text{ are independent, then}

\[
\hat{p}(n_1,\ldots,n_r) = \frac{K_{n_1,\ldots,n_r}}{k}
\]

is an unbiased estimator of \( p(n_1,\ldots,n_r) \) with variance proportional to \( 1/k \). Using a specialized sampling procedure called rotation sampling, MCHAIN induces statistical dependence among the data sets \{N_{ij}(\ell); i,j \in S\} \ell = 1,\ldots,k \text{ that preserves the unbiasedness of (3) but changes the convergence rate of its variance to } O(1/k^2) \). Fishman (1982b) illustrates the significance of this improved convergence for the estimation of the probability density function of first passage time in a semi-Markov process. More generally, the improved convergence is clearly beneficial when estimating functions of Markov chains that are linear combinations of \( p(n_1,\ldots,n_r) \) for varying \( n_1,\ldots,n_r \). Complete details about rotation sampling can be found in Fishman (1982a).

Section 1 describes how one constructs a simulation model to induce the desired dependence and shows its consequences in terms of the improved efficiency. For comparison a procedure is first described for simulating independent replications of a Markov chain. Section 2 describes MCHAIN and, in particular, its input requirements. Section 3 presents an example of how MCHAIN performs.

1. Serial and Parallel Simulation

To put the proposed sampling scheme into perspective we first describe the more conventional simulation of a Markov chain in which data on \( k \) independent replications are collected in serial order.
Serial Simulation

1. Start in state \( i = a \).

2. For \( m = 1, \ldots, k \) and \( i, j = 0, 1, \ldots, n \): \( N_{ij}^{(m)} = 0 \).

3. \( m + 1 \).

4. Sample \( U \) from \( U(0,1) \).

   Use \( U \), \( \{p_{ij}; j=0,1,\ldots,n\} \) and the alias sampling method
   (Walker 1977 and Kronmal and Peterson 1979) to determine
   entered state \( j \).

5. \( N_{ij}^{(m)} = N_{ij}^{(m-1)} + 1 \).

6. If \( j \neq b \), \( i + j \) and go to 4.

7. If \( m = k \), deliver \( \{N_{ij}^{(m)}; i,j=0,1,\ldots,n\} \) \( m=1,\ldots,k \).

8. \( m = m + 1 \).

9. \( i = a \).

10. Go to 4.

Here \( U(0,1) \) in step 4 denotes the uniform distribution on the half-open interval \([0,1)\). The alias sampling method is a procedure for sampling from a tabled discrete distribution at a cost independent of the length of the table. For example, if the entered state is \( i \neq b \), then sampling occurs from the distribution \( p_{i0}, p_{i1}, \ldots, p_{in} \) with constant execution time regardless of the value of \( n \).

In the case of parallel simulation based on rotation sampling all \( k \) replications begin at the same moment in time. The steps are:
Parallel Simulation

Definitions: $i_m$ = state occupied by replication $m$

$K_j$ = number of replications in state $j$ at the beginning of a step.

$K^*_j$ = number of replications in state $j$ at the end of a step.

1. For $m=1,\ldots,k$:
   1a. $i_m = a$.
   1b. For $i=0,1,\ldots,n; j=0,1,\ldots,n$, $N_{ij}^{(m)} = 0$.

2. For $i=0,1,\ldots,n$ : $K_i = 0$.

3. For $i=0,1,\ldots,n$ : $K^*_i = 0$.

4. Sample $U_0, U_1,\ldots,U_n$ independently from $U(0,1)$.

5. For $m=1,\ldots,k$:
   5a. If $i_m = b$, go to 5h.
   5b. Determine $j$ using $U_i$ and $\{p_{ij}; j=0,1,\ldots,n\}$
   5c. $K^*_j = K^*_j + 1$.
   5d. $N_{ij}^{(m)} + N_{ij}^{(m)} + 1$.
   5e. $U_{i_m} = U_{i_m} + 1/K_{i_m}$.
   5f. If $U_{i_m} \geq 1$, $U_{i_m} = U_{i_m} - 1$.
   5g. $i_m = j$.
   5h. Continue.


7. If $K_b = k$, deliver $\{N_{ij}^{(m)}; i,j=0,1,\ldots,n\}$.

8. For $j=1,\ldots,k$ and $j \neq b$ : $K_j = K^*_j$.

9. Go to 3.
Here rotation sampling is implemented in step 5e so that for \( K_{i_m} \) replications in state \( i_m \) the \( K_{i_m} \) uniform deviates used to determine the newly entered states have the uniform distribution on \([0,1)\) but are not independent. Fishman (1981, 1982b) contain more detailed descriptions of rotation sampling. Step 5b uses an alternative method to the alias to determine the entered state. In particular, this cutpoint method uses the inverse transform method to select the entered state for replication \( m \) as

\[
j = (r: \sum_{s=0}^{r-1} p_{i_m} s \leq U_i < \sum_{s=0}^{r} p_{i_m} s, \quad p_{i_m,-1} = 0; \quad r = 0, 1, \ldots, n) \tag{4}
\]

at a cost independent of \( n \). This procedure for sampling from the Markov chain together with the rotation sampling induces the desired dependence. Since the alias sampling method, which is slightly less costly, does not use the inverse transform method, it cannot be used to induce the desired correlation.

To appreciate the significance of the dependence that rotation sampling and the cutpoint method together induce, we first focus on (2). Let \( w = n_1 + \ldots + n_r \). Then

\[
K_w = \sum_{n_1=0}^{w} \ldots \sum_{n_r=0}^{w-n_1-\ldots-n_{r-1}} K_{n_1, \ldots, n_r} \tag{5}
\]

is the number of replications absorbed on the \( w \)th transition. Making use of the fact that \( \text{var} K_w = O(1) \), Proposition 1 in Fishman (1982b) shows that \( \text{var} K_{n_1, \ldots, n_r} = O(1) \). Therefore, \( \text{var} \hat{p}(n_1, \ldots, n_r) = O(1/k^2) \).
Computation Time Complexity

Although the accelerated convergence of the variances of estimators is important, one also needs to consider how the computation time complexity of the parallel simulation (PS) compares with that of the serial simulation (SS). Both are $O(K)$. Although for a given $k$ parallel simulation takes more time than serial simulation does, the ratio of these times converges to a constant as $k \to \infty$, thereby demonstrating the superiority of parallel simulation as measured by the common variance reduction measure (see Hammersley and Handscomb 1964)

$$VR = \frac{\text{variance using SS}}{\text{variance using PS}} \cdot \frac{\text{computation time using SS}}{\text{computation time using PS}} = O(k) \text{ as } k \to \infty. \quad (6)$$

2. The MCHAIN Program

Figure 1 lists a FORTRAN subroutine called MCHAIN that generates the transition frequency data for a finite-state Markov chain using rotation sampling. The program uses pointers (M(*)) to indicate the S(*) states that can be entered from *, thus eliminating the nonzero entries in the transition matrix and reducing storage space.

Of special interest is the method of determining the state to be entered by a replication at each transition, as in (4). MCHAIN uses the cutpoint method developed in Fishman and Moore (1981) whose execution time is independent of $S(J)$, the number of states that can be entered from $J$. Therefore, using this method makes the cost of state transition determination independent of the number of nonzero entries in each row of $p_n$. MCHAIN uses the random number generator GGUBS in the IMSL(1982) library, but this can be changed at the user's discretion.
MCHAIN allows for three different levels of output through the variable DETAIL. Setting DETAIL=2 results in a printout of transition frequencies for each replication and summations of these frequencies across replications. Setting DETAIL=1 leads to a printout only of the summations across replications and DETAIL=0 suppresses all printed output. In practice, we envision that MCHAIN may be used to generate input for other simulation, as exemplified in Fishman (1982b). In this case a user needs to alter MCHAIN to store or pass the frequency data and may have limited or no interest in the printout of the frequency data themselves. DETAIL's options enable him to specify his level of interest.

MCHAIN allows for two types of simulation runs, macroreplications and microreplications. MCHAIN runs $I$ macroreplications each consisting of $K$ microreplications based on parallel simulation using rotation sampling. The need for these macroreplications arises when estimating the variances of estimators. See Fishman (1982b). In general, it is advisable to make $K$ large relative to $I$ to gain the benefit of rotation sampling. Space requirements for the arrays in MCHAIN are roughly $4 \times (\text{SIZE} + 8 \times \text{NP} + 2 \times K + 6 \times \text{ALL})$.

3. **Example**

Consider a single server queueing model for which the time between successive arrivals are independent and exponentially distributed with rate $\lambda$, service times are independent and exponentially distributed with rate $\omega$, the queue discipline is first-come-first-served and there is a finite capacity $n$. Corresponding to this formulation is an embedded nearest neighbor Markov chain. These states are the number of
jobs in the system and whose nonzero transition probabilities are

\[ p_{01} = 1 \]
\[ p_{i,i-1} = \frac{\omega}{\lambda + \omega} \quad i=2, \ldots, n \]
\[ p_{i,i+1} = \frac{\lambda}{\lambda + \omega} \quad i=1, \ldots, n-1 \]
\[ p_{n,n} = \frac{1}{\lambda + \omega} . \]

We show a run of \textsc{MChain} for a first passage from state \( a=4 \) to state \( b=17 \) for a single macroreplication containing 15 microreplications based on rotation sampling with \( \lambda = .9 \), \( \omega = 1 \) and \( n=19 \). The input for \textsc{MChain} is \( \text{NP}=20, \text{INITIAL}=4, \text{ABSORB}=17, \text{I}=1, \text{K}=15, \text{SEED}=1556203872, \text{SIZE}=1000 \) and \( \text{DETAIL}=2 \). The quantities ALL, KK and N and the \( M, P, S \) and SUMS arrays are computed in the sample driver program in Fig. 2. Figure 3 shows the output for \textsc{MChain}. The entries in the table are the frequencies of transitions for all possible transition type by microreplication.
References


**SUBROUTINE MCHAIN**

This subroutine simulates \(K\) replications of an \((N+1)\)-state Markov chain in parallel using rotation sampling.

For reference see:

- **AND "Generating parallel correlated transition frequencies for a Markov chain,"** Report UNC/ORS/A/TR 82/8, Curriculum in Operations Research and Systems Analysis, University of North Carolina at Chapel Hill.

**SUBROUTINE MCHAIN** (NP, INITAL, ABSORB, P, Q, N, S, SUMS, ALL, KK, L, K,
1 SEED, SIZE, U, V, DETAIL, CUTP, ROTATE, CN, KSTAR, KPRIME, STATE)

**DESCRIPTION OF VARIABLES:**

- **AORB** = AORBSING STATE
- **ALL** = SUM OF S(J) FOR ALL J
- **CN(*)** = HOLDS TRANSITION COUNTS FOR EACH TRANSITION TYPE AND EACH MICROREPLICATION; LAST COLUMN CONTAINS TOTALS BY TRANSITION TYPE FOR ALL MICROREPLICATIONS
- **COLMAX** = MAXIMUM NUMBER OF COLUMNS TO PRINT PER OUTPUT PAGE
- **CCUNT** = UNIFORM DEVIATE COUNTER
- **CUTP(*)** = POINTERS FOR CUTOPT SAMPLING METHOD
- **SEE FISHMAN, G. S. AND L. R. MOORE, III (1981).** "SAMPLING FROM A DISCRETE DISTRIBUTION WHILE PRESERVING MONOTONICITY", TECHNICAL REPORT 81/7,
- **OPERATIONS RESEARCH AND SYSTEMS ANALYSIS, UNIVERSITY OF NORTH CAROLINA AT CHAPEL HILL.**
- **DETAIL** = DETAIL FLAG, VALUES:
  - 0 FOR NO PRINTING, 1 FOR SUMMARY TOTALS ONLY,
  - 2 FOR FULL DETAIL PRINTING
- **EPSEED** = RANDOM NUMBER GENERATOR SEED (DOUBLE PRECISION)
- **EP** = STEP VARIABLE FOR SAMPLING TRANSITION TYPE
- **UIMS** = IMSL SUBROUTINE FOR RANDOM NUMBER GENERATION
- **I** = NUMBER OF MACROREPLICATIONS
- **I1** = INDEX USED BY OUTPUT SECTION
- **IP** = INDEX USED BY OUTPUT SECTION
- **I** = NUMBER OF FIRST COLUMN TO PRINT ON CURRENT OUTPUT PAGE
- **II** = INDEX USED BY OUTPUT SECTION
- **IIS** = S(J+1)
- **LAST** = NUMBER OF LAST COLUMN TO PRINT ON CURRENT OUTPUT PAGE
- **IND** = FLAG, INITIALLY SET TO 0 TO ALLOW FIRST TRANSITION IF STARTING IN ABSORBING STATE
- **INITIAL** = INITIAL STATE
- **IP** = INDEX USED BY OUTPUT SECTION

Fig. 1 MCHAIN FORTRAN Program
C *** ISEED = SAVED INITIAL SEED
C *** IX = INDEX
C *** J = INDEX FOR STATES
C *** J1 = INDEX FOR MICROREPLICATIONS
C *** JA = INDEX FOR ALL
C *** JJ = INDEX USED FOR INDIRECT SUBSCRIPTING
C *** K = NUMBER OF PARALLEL MICROREPLICATIONS
C *** K1 = K+1
C *** KA = NUMBER OF MICROREPLICATIONS THAT HAVE RETURNED TO ABSORBING STATE
C *** KPRIME(*) = NUMBER IN STATE AT END OF TRANSITION
C *** KSTAR(*) = NEXT KPRIME(*)
C *** KTEST = TEMPORARY VARIABLE USED IN OUTPUT SECTION
C *** KTOTAL = TOTAL NUMBER OF TRANSITIONS
C *** LCOUNT = INDEX USED BY OUTPUT SECTION TO COUNT OUTPUT LINES
C *** LI = NUMBER OF CURRENT MACROREPLICATION
C *** LMAX = MAXIMUM NUMBER OF LINES TO PRINT PER OUTPUT PAGE
C *** LX = CUTOPOINT TO ACCELERATE SAMPLING PROCEDURE
C *** M(*) = STATES THAT CAN BE REACHED FROM J-1
C *** N = NUMBER OF HIGHEST STATE (NP-1)
C *** NP = NUMBER OF STATES
C *** NSKIP = NUMBER OF OUTPUT PAGES TO BE PRINTED
C *** CH = USED IN SETTING UP STATE TRANSITION PROBABILITIES
C *** P(*) = TRANSITION MATRIX
C *** Q(*) = VECTOR OF CUMULATIVE PROBABILITIES FOR EACH TRANSITION TYPE
C *** K = INDEX FOR STATES
C *** ROTATE(*) = ROTATION INDEX FOR SAMPLING FROM EACH STATE
C *** S(J) = NUMBER OF STATES THAT CAN BE ENTERED FROM J-1
C *** SEED = RANDOM NUMBER GENERATOR SEED
C *** SIZE = BLOCK SIZE FOR RANDOM NUMBER GENERATION
C *** SJ = S(J)
C *** SK = SUMS(J)
C *** STATE(*) = CURRENT STATE OCCUPIED BY MICROREPLICATION J1
C *** SUMS(J) = SUM OF S(1) THRU S(J-1), AND SUMS(1)=0
C *** U = CURRENT UNIFORM DEVIATE
C *** UU(*) = CURRENT UNIFORM DEVIATE FOR TRANSITION TYPE *
C *** V(*) = UNIFORM DEVIATE ARRAY
C ***
INTEGER CUTP(ALL),ABSORB,ALL,CH(KK),COUNT,DETAIL,KPRIME(NP),I,
1   I1,IND,INITIAL,ISEED,IX,J,J1,JA,JK,K,KK1,K,KKA,LI,LX,LCOU,
2   LMAX,M(ALL),KSTAR(NF),NP,R,S(NP),SEED,SIZE,SJ,SK,
3   STATE(K),SUMS(NP)
INTEGER CLMAX,IFIRST,II,ILAST,IP,ITOTAL,KTEST,KTOTAL,N,NSKIP
DOUBLE PRECISION RCTATE(NF),EP,OM,UU(NP),U,DSEED,P(ALL),Q(ALL)
C ***
C *** INITIALIZE DOUBLE PRECISION SEED
C ***
ELSEED=SEED
K1=K+1

Fig. 1 (Continued)
**Fig. 1 (Continued)**

```plaintext
C *** DETERMINE TRANSITION PROBABILITIES
C ***
    DO 100 J=1,NP
        SK = SUMS(J)
        Q(SK+1) = P(SK+1)
        IF(S(J).LT.2) GO TO 100
        SJ = S(J)
        DO 90 R=2,SJ
            Q(SK+R) = Q(SK+R-1) + P(SK+R)
        END DO
        CONTINUE
        Q(SK+SJ) = 1.0 DO
        CONTINUE
C ***
C *** SET UP INDIRECT INDEXING FOR TRANSITION ARRAYS
C ***
    DO 200 J=1,NP
        SJ=S(J)
        EP=(1.0/SJ)
        OM=0
        LX=SUMS(J)+1
        DO 200 R=1,SJ
            CUTP(SUMS(J)+R)=LX
            IF(Q(LX).GT.OM) GO TO 195
            LX=LX+1
            GO TO 150
        195 OM=OM+EP
        CONTINUE
        DO 950 LI=1,I
C ***
C *** SET STARTING SEED
C ***
    ISEED=DSEED
C ***
C *** SET FLAG TO ALLOW INITIAL TRANSITION FROM ABSORBING STATE
C ***
    IND=0
C ***
C *** GET ARRAY OF RANDOM DEVIATES
C ***
    CALL GGOES(ISERED,SIZE,V)
C ***
C *** GET INITIAL RANDOM DEVIATE
C ***
    U(INITIAL+1)=V(1)
    U=V(1)
    COUNT=2
C ***
C *** START MICROREPLICATIONS IN INITIAL STATE
C *** AND INITIALIZE TRANSITION COUNTERS
```
C ***
KPRIME(INITAL+1)=K
DO 210 J1=1,K
  STATE(J1)=INITAL
DO 210 JA=1,ALL
  CN(ALL*(J1-1)+JA)=0
210 CONTINUE
DO 215 J=1,NP
  ROTATE(J)=0.0D0
215 KSTAR(J)=0
KA=0
C ***
C *** BEGIN SIMULATION OF STATE TRANSITIONS
C ***
220 DO 300 J1=1,K
C ***
C *** DO NOT MOVE FROM ABSORBING STATE EXCEPT DURING INITIAL TRANSITION
C ***
  IF(STATE(J1).EQ.ABSORB.AND.IND.EQ.1) GO TO 300
  IX=STATE(J1)
C ***
C *** TEST TO PREVENT DIVISION BY ZERO
C ***
  IF(KPRIME(IX).EQ.0) GO TO 300
C ***
C *** GET RAW DCMM DEVIATE FOR THIS MICROREPLICATION
C ***
  U=UU(IX)+(ROTATE(IX)/KPRIME(IX))
  IF (U.GE.1.0) U=U-1.
C ***
C *** FIND CUTPCINT FOR SEARCH OF SAMPLING DISTRIBUTION
C ***
  LX=(S(IX)*U)
C ***
C *** DETERMINE THE DIRECTION OF TRANSITION
C ***
  JJ=CUTP(SUMS(IX)+LX+1)
  IF(U.LE.Q(JJ)) GO TO 250
  JJ=JJ+1
  GO TO 225
C ***
C *** INCREMENT THE APPROPRIATE TRANSITION COUNTER
C ***
  CN(ALL*(J1-1)+JJ)=CN(ALL*(J1-1)+JJ)+1
C ***
C *** SET THE STATE FOR MICROREPLICATION J1
C ***
  STATE(J1)=M(JJ)
C ***
C *** INCREMENT THE TEMPORARY COUNTER AND

Fig. 1 (Continued)
THE ACTIVE MICROREPLICATION COUNTER

KSTAR(M(JJ)+1) =KSTAR(M(JJ)+1)+1
ROTATE(IX) =ROTATE(IX)+1.0

SET FLAG TO PREVENT TRANSITIONS FROM ABSORBING STATE

IND=1

ADD NEWLY ABSORBED MICROREPLICATIONS TO COUNTER

KA=KA+KSTAR(ABSORB+1)

IF ALL K MICROREPLICATIONS HAVE BEEN ABSORBED, EXIT

IF(KA.EQ.K) GC TO 500
KSTAR(J) =KA

REinitialize COUNTERS FOR NEXT SET OF TRANSITIONS

DO 400 J=1,NP
    ROTATE(J) =0.0D0
    KPRIME(J) =KSTAR(J)
    KSTAR(J) =0
    IF(KPRIME(J)*(J-ABSORB-1).EQ.0) GO TO 400

GET NEXT RANDOM DEVIATE

UU(J) =V(COUNT)
CCOUNT=COUNT+1
IF(COUNT.LE.SIZE) GO TO 400

IF NECESSARY, GET NEW ARRAY OF RANDOM DEVIATES

CALL GGBUBS(DSEED,SIZE,V)
COUNT=1
CONTINUE
GO TO 220
500 SEED=DSEED
IF(DETAIL.EQ.0) GC TO 990

PRINT RESULTS OF SIMULATION

WRITE(3,9000) LI
9000 FORMAT(1//' RESULTS FROM MCHAIN FOR MACROREPLICATION NO.' ,I5//)
550 WRITE(3,9010) NP,INITIAL,ABSORB,ALL,K,I,ISEED,SEED,SIZE
9010 FORMAT(///,
1 ' NO. OF STATES ' =',I10//,
2 ' INITIAL STATE ' =',I10//.

Fig. 1 (Continued)
ABSORBING STATE
TOTAL NO. OF (I,J) PAIRS
NO. OF CORRELATED MICROREPLICATIONS
NO. OF INDEPENDENT MACROREPLICATIONS
INITIAL SEED
FINAL SEED
BLOCKING FACTOR

**CALCULATE TOTALS ACROSS ALL CHAINS AND STORE IN CN(*,K1)**

**DETERMINE OUTPUT FORMAT PARAMETERS (PAGE WIDTH AND LENGTH)**

**PRINT SUMMARY TOTALS ONLY**

**DETERMINE NUMBER OF OUTPUT PAGES**

Fig. 1 (Continued)
C *** IF(KI.GT.COLMAX) GO TO 700
   COLMAX=K1
   NSKIP=1
   GO TO 750
700   NSKIP=K1/CCLMAX
   KTEST(NSKIP*COLMAX
   IF((K1-KTEST).GT.0) NSKIP=NSKIP+1
C ***
C *** PRINT TRANSITION COUNTS IN TABLE FORMAT
C ***
75C   DO 900 I=1,NSKIP
       IFIRST=ILAST+1
       ILAST=I*COLMAX
       IF(I1.EQ.NSKIP) ILAST=K1
C ***
C *** PRINT PAGE HEADING
C ***
   WRITE(3,2100)
   2100   FORMAT(//'TRANSITION COUNTS IN TABLE FORMAT'
               /'TRANSITION COUNTS IN TABLE FORMAT')
   WRITE(3,2110) (IX,IX=IFIRST,ILAST)
   2110   FORMAT(' FROM TO',16(1X,I5))
   WRITE(3,2120)
   2120   FORMAT('-------------------------------------------------------'
               /'
               /')
   IP=0
   N=NF-1
   LCOUNT=0
C ***
C *** PRINT TRANSITION COUNTS
C ***
   DC 850 J=0,N
         I1S=5(J+1)
   DC 800 II=1,IIS
   LCOUNT=LCOUNT+1
   IP=IP+1
   IA=ALL*(IFIRST-1)+IP
   IB=ALL*(ILAST-1)+IP
   WRITE(3,2200) J,M(IP), (CN(IX),IX=IA,IB,ALL)
   2200   FORMAT(//'TRANSITION COUNTS IN TABLE FORMAT'
               /'TRANSITION COUNTS IN TABLE FORMAT')
   CONTINUE
   IF(LCOUNT.LE.LMAX) GO TO 850
   WRITE(3,2100)
   WRITE(3,2110) (IX,IX=IFIRST,ILAST)
   WRITE(3,2120)
   LCOUNT=0
850   CONTINUE
900   CONTINUE
   WRITE(3,2300) K1
   2300   FORMAT('0 ***COLUMN I5, CONTAINS THE RCW TOTALS')

Fig. 1 (Continued)
950 WRITE(3,2400) KTOTAL
2400 FORMAT(10 ***TOTAL FOR ALL ROWS IS: 'I6)
990 RETURN
END

Fig. 1 (Continued)
C *** THIS IS A SAMPLE DRIVER PROGRAM FOR SUBROUTINE MCHAIN
C ***
C *** THIS DRIVER PROGRAM PROVIDES THE SETUP FOR USE OF MCHAIN WITH
C *** THE M/M/1/N QUEUEING MODEL WITH ARRIVAL RATE LAM, SERVICE
C *** RATE W, ONE SERVER AND CAPACITY N. WITH THE EXCEPTION OF
C *** M, P, S AND SU MS, ALL OTHER ARRAYS ARE USED IN MCHAIN.
C ***
C *** DESCRIPTION OF VARIABLES -
C ***
C *** ABSORB = ABSORBING STATE
C *** ALL = SUM OF S(J) FOR ALL J
C *** CN(*) = HOLDS TRANSITION COUNTS FOR EACH TRANSITION TYPE AND
C *** EACH MICROREPLICATION; LAST COLUMN CONTAINS TOTALS
C *** CUTP(*) = POINTERS FOR CUTFPOINT SAMPLING METHOD
C *** DETAIL = DETAIL FLAG, VALUES:
C *** 0 FOR NO PRINTING, 1 FOR SUMMARY TOTALS ONLY,
C *** 2 FOR FULL DETAIL PRINTING
C *** I = DESIRED NUMBER OF INDEPENDENT MACROREPLICATIONS
C *** INITIAL = INITIAL STATE
C *** J = INDEX FOR STATES
C *** K = NUMBER OF PARALLEL MICROREPLICATIONS/MACROREPLICATION
C *** KPRIME(*) = NUMBER IN STATE AT END OF TRANSITION
C *** KSTAR(*) = NEXT KPRIME(*)
C *** LAM = ARRIVAL RATE (<W)
C *** M(*) = STATES THAT CAN BE REACHED FROM J-1
C *** N = NUMBER OF HIGHEST STATE (<50)
C *** NP = NUMBER OF STATES (N+1)
C *** P(*) = TRANSITION MATRIX
C *** Q(*) = VECTOR OF CUMULATIVE PROBABILITIES FOR EACH TRANSITION TYPE
C *** ROTATE(*) = ROTATION INDEX FOR SAMPLING FROM EACH STATE
C *** SEED = RANDOM NUMBER GENERATOR SEED
C *** SIZE = BLOCK SIZE FOR RANDOM NUMBER GENERATOR (<1001)
C *** S(J) = NUMBER OF STATES THAT CAN BE ENTERED FROM J-1 (<NP)
C *** SUMS(J) = SUM OF S(1) THRU S(J-1), AND SUMS(1)=0 (<100)
C *** UU(*) = CURRENT UNIFORM DEVIATE FOR TRANSITION TYPE *
C *** V(*) = UNIFORM DEVIATE ARRAY
C *** W = SERVICE RATE
C ***
INTEGER CUTP(100), ABSORB, ALL, CN(2500), DETAIL, INITIAL, I, J, 1, JA,
1   K, KK, KPRIME(50), KSTAR(50), M(100), N, NP,
2   S(50), SEED, SIZE, STATE(50), SUMS(50)
REAL V(1000)
DOUBLE PRECISION RCTATE(50), LAM, P(100), Q(100), UU(50), W
C ***
C *** INITIALIZE VALUES
C ***
READ (1,1000) NP, INITIAL, ABSORB, I, K, SEED, SIZE
1000 FORMAT (15/15/15/15/15/15/15)
WRITE (3,7060) INITIAL, ABSORB

Fig. 2  Sample Driver Program
**Fig. 2 (Continued)**

7080 FORMAT(' INITIAL STATE: ','I5', ' AESORBING STATE: ','I5')
WRITE(3,7081) K,NP
7081 FORMAT(' NUMBER OF REPLICATES: ','I5', ' NUMBER OF STATES: ','I5')
WRITE(3,7082) SEED,SIZE
7082 FORMAT(' INITIAL SEED: ','I12', ' BLOCKING FACTOR: ','I6')
READ(1,1002) DETAIL
1002 FORMAT(I5)
READ(1,1001) LAM,W
1001 FORMAT(2F5.2)
WRITE(3,7085) LAM,W,DETAIL
7085 FORMAT(' LAM: ','F5.2',' W: ','F5.2',' DETAIL: ','I3')
C ***
C *** FOR EACH STATE, DETERMINE THE NUMBER OF STATES THAT CAN BE ENTERED
C ***
S(1)=1
DO 80 J=2,NP
   S(J)=2
80 CONTINUE
WRITE(3,2015) (S(J),J=1,NP)
2015 FORMAT(' S(J) =1,2015)
SUMS(1)=0
ALL=S(1)
DO 90 J=2,NP
   SUMS(J)=ALL
   ALL=ALL+S(J)
90 CONTINUE
KK=ALL*(K+1)
C *** RESTRICTIONS ON PARAMETER SIZES BELOW ARE FOR THIS PROGRAM.
C ***
C *** COMPUTE TRANSITION PROBABILITIES
C ***
DC 100 J=2,NP
   P(SUMS(J)+1) = W/(LAM+W)
   P(SUMS(J)+2) = LAM/(LAM+W)
100 CONTINUE
P(1)=1.000
WRITE(3,2016) (P(JA),JA=1,ALL)
2016 FORMAT(' P(*) =1,20F5.2)
C ***
C *** FOR EACH STATE J DETERMINE STATES THAT CAN BE ENTERED
C ***
M=NP-1
DC 400 J=2,N
   M(SUMS(J)+1) = J-2
   M(SUMS(J)+2) = J
400 CONTINUE
M(1)=1
M(SUMS(NP)+1) = M-1
M(SUMS(NP)+2) = M
WRITE(3,2017) (M(JA),JA=1,ALL)
2017 FORMAT(' M(JA) = 1,20I5.1')
2017 FORMAT(* M(*) =', 20I5)
C ***
C *** CALL SUBROUTINE MCHAIN
C ***
   CALL MCHAIN(WP,INITAL,ABSORB, P, Q, M, S, SUMS, ALL, KK, I, K,
      1 SEED, SIZE, UU, V, DETAIL, CUTO, ROTATE, CN, KSTAR, KPRIME, STATE)
C ***
C *** END OF PROGRAM
C ***
   STOP
   END

Fig. 2  (Continued)
INITIAL STATE:  4  ABSORBING STATE:  17
NUMBER OF REPLICATES:  15  NUMBER OF STATES:  20
INITIAL SEED:  1556203872  BLOCKING FACTOR:  1000
LEN:  0.95  N:  1.00  OMNITOLE:  2
S(i) =  1  2  2  2  2  2  2  2  2  2  2  2  2  2  2  2  2  2  2
F(*) =  1.00  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51  0.49  0.51
H(*) =  1  0  2  1  3  2  4  3  5  4  6  5  7  6  8  7  9  8  10  9
H(*) =  11  10  12  11  13  12  14  13  15  14  16  15  17  16  18  17  19  18  19

RESULTS FROM BCHAIN FOR MACROREPLICATION NO.  1

NO. OF STATES =  20
INITIAL STATE =  4
ABSORBING STATE =  17
TOTAL NO. OF (I,J) PAIRS =  39
NO. OF CORRELATED MACROREPLICATIONS =  15
NO. OF INDEPENDENT MACROREPLICATIONS =  1
INITIAL SEED =  1556203872
FINAL SEED =  1950653926
BLOCKING FACTOR =  1000

Fig. 3  MCHAIN and DRIVER Program Output
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***COLUMN 16 CONTAINS THE ROW TOTALS

***TOTAL FOR ALL ROWS IS:  3897

Fig. 3 (Continued)
This paper describes an algorithm and a FORTRAN program called MCHAIN for simulating \( k \) parallel Monte Carlo replications of a Markov chain using rotation sampling. This method of sampling produces \( k \) sample transition frequency vectors with a desirable structure of statistical dependence among.
them. In particular, these sample vectors can be used to estimate the probability that, say, \( n_1, \ldots, n_r \) transitions of types \( 1, \ldots, r \) occur during a first passage from state \( a \) to state \( b \) with a variance of the estimate of \( O(1/k^2) \) and a computation time \( O(k) \) as \( k \to \infty \). This compares favorably with the case of independent replications wherein the estimate would have a variance \( O(1/k) \) and computation time \( O(k) \) as \( k \to \infty \). An example including a sample driver program are presented to illustrate how MCHAIN works in practice.
END