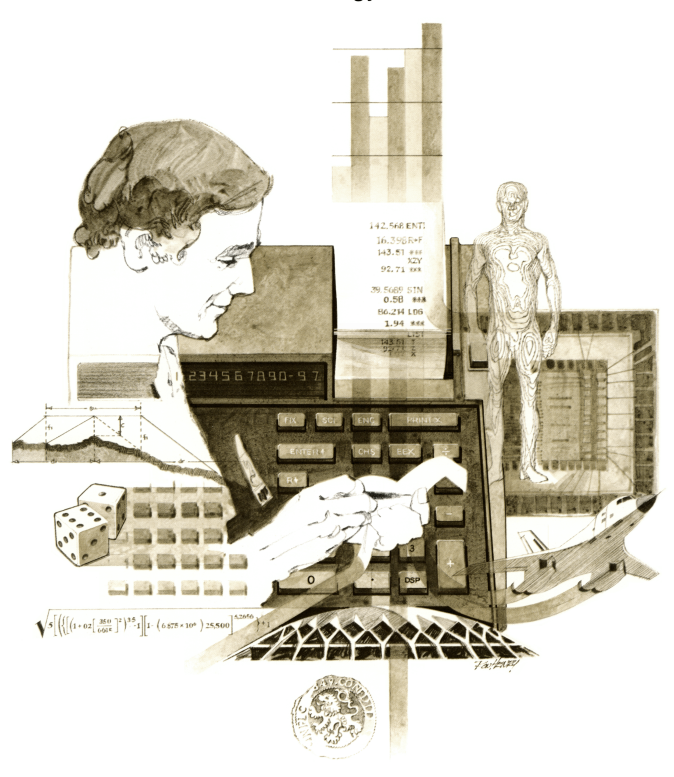
HP67 HP97

Users' Library Solutions Biology



INTRODUCTION

In an effort to provide continued value to it's customers, Hewlett-Packard is introducing a unique service for the HP fully programmable calculator user. This service is designed to save you time and programming effort. As users are aware, Programmable Calculators are capable of delivering tremendous problem solving potential in terms of power and flexibility, but the real genie in the bottle is program solutions. HP's introduction of the first handheld programmable calculator in 1974 immediately led to a request for program solutions — hence the beginning of the HP-65 Users' Library. In order to save HP calculator customers time, users wrote their own programs and sent them to the Library for the benefit of other program users. In a short period of time over 5,000 programs were accepted and made available. This overwhelming response indicated the value of the program library and a Users' Library was then established for the HP-67/97 users.

To extend the value of the Users' Library, Hewlett-Packard is introducing a unique service—a service designed to save you time and money. The Users' Library has collected the best programs in the most popular categories from the HP-67/97 and HP-65 Libraries. These programs have been packaged into a series of low-cost books, resulting in substantial savings for our valued HP-67/97 users.

We feel this new software service will extend the capabilities of our programmable calculators and provide a great benefit to our HP-67/97 users.

A WORD ABOUT PROGRAM USAGE

Each program contained herein is reproduced on the standard forms used by the Users' Library. Magnetic cards are not included. The Program Description I page gives a basic description of the program. The Program Description II page provides a sample problem and the keystrokes used to solve it. The User Instructions page contains a description of the keystrokes used to solve problems in general and the options which are available to the user. The Program Listing I and Program Listing II pages list the program steps necessary to operate the calculator. The comments, listed next to the steps, describe the reason for a step or group of steps. Other pertinent information about data register contents, uses of labels and flags and the initial calculator status mode is also found on these pages. Following the directions in your HP-67 or HP-97 **Owners' Handbook and Programming Guide**, "Loading a Program" (page 134, HP-67; page 119, HP-97), key in the program from the Program Listing I and Program Listing II pages. A number at the top of the Program Listing indicates on which calculator the program was written (HP-67 or HP-97). If the calculator indicated differs from the calculator you will be using, consult Appendix E of your **Owner's Handbook** for the corresponding keycodes and keystrokes converting HP-67 to HP-97 keycodes and vice versa. No program conversion is necessary. The HP-67 and HP-97 are totally compatible, but some differences do occur in the keycodes used to represent some of the functions.

A program loaded into the HP-67 or HP-97 is not permanent—once the calculator is turned off, the program will not be retained. You can, however, permanently save any program by recording it on a blank magnetic card, several of which were provided in the Standard Pac that was shipped with your calculator. Consult your **Owner's Handbook** for full instructions. A few points to remember:

The Set Status section indicates the status of flags, angular mode, and display setting. After keying in your program, review the status section and set the conditions as indicated before using or permanently recording the program.

REMEMBER! To save the program permanently, **clip** the corners of the magnetic card once you have recorded the program. This simple step will protect the magnetic card and keep the program from being inadvertently erased.

As a part of HP's continuing effort to provide value to our customers, we hope you will enjoy our newest concept.

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Program Title DEMOGRAPHY I: Estimates of Parameters/Rates of Increase Hewlett-Packard Company Contributor's Name Address 1000 N.E. Circle Boulevard Corvallis Oreogn State **Zip Code** 97330

Program Description, Equations, Variables Generation time is estimated using the formula: where x is age, 1, is the fractional survivorship to that age from age class zero,

m is the fecundity at age x, and R is the net reproductive rate, computed from the equation

$$R_0 = \sum_{\alpha}^{\omega} 1_x m_x$$

In both equations α is the age of first reproduction, and ω is the age of last reproduction. Any number of non-reproductive age classes is allowed, by simply keying in a constant, k, that is added to each age of reproduction internally by the program. This program is designed for use with another (next), which computes much more accurate estimates of the intrinsic rate of increase by an iterative process: This program estimates the intrinsic rate of increase, r, using the following approximate relationship:

Using the estimate of r stored in R_8 , and the products of the age-specific survivorship and fecundity schedules stored in registers 1 through 7, the program computes the sum

 $\Sigma e^{-rx} 1_{x}^{m}$

(Continued on next page)

1

Operating Limits and Warnings Only seven reproductive age classes are allowed and a constant, k, indicates the age of the oldest non-resproductive age classes.

This program has been verified only with respect to the numerical example given in Program Description II. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Program Title

Program Description I

Address	State	Zip Code
City	State	Zip Code
Program Description, Equations, Variables ($1_{\alpha}m_{\alpha}$ is stored in R_1) and ω be unity if the estimate of and if it is too large the estine the estimate of r is decrease recalculates the new sum, whi process is continued until the program terminates and display requires about 10 seconds of may take up to about 2 minute. The program also displays the If the user thinks the program with the "CLx" button and R_8 last loop.	is the age of last reproduct is good. The program compatinate of r is increased (ed). The program then stored in the sum is unity (to two decays the intrinsic rate of incalculator (computer?) times before r is displayed.	ction. This sum should pares the sum with 1.00 if the sum is too small, es this new r in R ₈ and gain as before. The imal places) and then the ncrease. Each loop e, and some parameter sets λ , which is simply e ^r .

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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SAMPLE PROBLEM: A population has the following schedules of age-specific survivorship and fecundity:

Age (x)	Survivorship (1_{χ})	Fecundity (m_{χ})
0	1.0	0.0
1	.9	0.0
2	.7	0.0
3	•5	0.0
$\frac{-4}{4}$ (α)	.4	1.0
5	.3	1.0
6	.2	1.5
7	.2	1.0
8	.1	1.0
9	.1	0.5
10 (ω)	.01	1.0

1. Compute the net reproductive rate, R, of this population and estimate its mean generation time, \hat{T} , and intrinsic rate of natural increase, \hat{r} .

Note that the data needed to compute these demographic parameters are contained entirely in the last seven (reproductive) age classes. Therefore, data are keyed in beginning with age class 4, which is the age of first reproduction. In this example k=3. Data above the dashed line may be used with the next program in this series to compute demographic parameters for non-reproductive age classes, such as expectation of life and reproductive value.

2. Calculate the intrinsic rate of increase and the finite rate of increase for the same population.

```
Solution(s) \begin{array}{ccc} R &=& 1.36 \\ T &=& 5.625 \\ \hline r &=& 0.0547 \text{ (a slight underestimate)} \end{array}
```

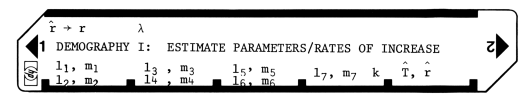
- 1. .4[ENTER ↑] 1 [ENTER ↑] .3 [ENTER ↑] 1 [A] .2[ENTER ↑] 1.5[ENTER ↑] .2 [ENTER ↑] 1 [B] .1[ENTER ↑] 1 [ENTER ↑] .1 [ENTER ↑] .5 [C] .01[ENTER ↑] 1 [ENTER ↑] 3 [D] [E] → 5.625 [R/S] → 0.0547
- [RCL] [9] \rightarrow 1.3600 2. [f] [A] \rightarrow .0552 (r)

2. [f] [A] \rightarrow .0552 (r) [f] [B] \rightarrow 1.0568 (λ)

Reference(s) Mertz, D.B. 1970. Notes on methods used in life-history studies. pp.4-17
in Connell, Mertz, and Murdoch's Readings in ecology and ecological
genetics." Harper & Row, New York, 397 p.
Pianka, E.R. 1974. Evolutionary ecology. Harper & Row. 356 pp.
Poole, R.W. 1974. An introduction to quantitative ecology. McGraw-Hill.
This program is a modification of the Users' Library Programs #03637A

and #03638A submitted by Eric R. Pianka.

User Instructions



STEP	INSTRUCTIONS	INPUT DATA/UNITS	KEYS	OUTPUT DATA/UNITS
1	Load side 1 and side 2			
2	Enter data as shown for Survivorship/Fecundity	11	ENT	
		ml	ENT	
		12	ENT	
		m ₂	A	
		13	ENT	
		m3	ENT	
		14	ENT	
		m4	В	
		1 ₅	ENT	
		m5	ENT	
		16	ENT	
		m ₆	C	
		17	ENT	
		m ₇	ENT	
	Enter age constant k	k	D	
3	Compute Î		E	Î
4	Compute r		R/S	r
5	Recall R _o from R _g		RCL 9	R
6	Iterate the solution of r		f A	r
7	Compute the finite rate of increasing λ		fB	λ

97 Program Listing I

STEP		Y ENTRY	KEY CODE	COMM	ENTS	STEP	KE	Y ENTRY	KEY CODE	COMMENTS
	001	*LBLA	21	Data anten	_		0 57	3	0 3	
	00 2	X	-35	Data entry	y		6 58	RCL8	36 0 8	
	003	STO2	35 82				0 59	+	-55	
	004	R↓	-31				060	X	-35	
	0 05	X	-35				0 61	+	-55	
	0 06	ST01	35 01				0 62	RCL4	36 04	
	0 07	RTN	24				0 63	4	04	
	00 8	*LBLB	21 12				0 64	RCL8	36 0 8	
	00 9	Х	-35				0 65	+	-55	
	010	STO4	35 04				0 66	X	-35	
	011	R↓	-31				0 67	+	-5 5	
	012	X	-35				0 68	RCL5	36 0 5	
	013	STO3	35 03				0 69	5	0 5	
	014	RTN	24				0 70	RCL8	36 0 8	
	015	*LBLC	21 13				071	+	-55	
	016	Х	-35				0 72	×	-35	
	017	ST06	35 06				0 73	+	-55	·
	0 18	R↓	-31				074	RCL6	36 06	
	019	Х	-35				075	6	06	
	0 20	STO5	35 Ø5				0 76	RCL8	36 0 8	
	0 21	RTN	24				077	+	-55	
	0 22	*LBLD	21 14				0 78	X	-35	
	0 23	ST08	35 Ø3				079	+	-55	
	024	ST00	<i>35 00</i>				080	RCL7	36 0 7	
	0 25	R↓	-31				0 81	7	0 7	
	0 26	X	-35				0 82	RCL8	36 0 8	
	027	ST07	35 07				0 83	+	-55	
•	0 28	RTN	24				084 084	×	-35	
	0 29	*LBLE	21 15				00 1 085	+	-55	
	030	DSP4	-63 04				086	RCL9	36 0 9	
	031	RCL1	36 01					RUL9 ÷	-24	
	032	RCL2	36 02				0 87			Display Î
	03 3	RCL3	36 03				889	R/S	51 36 80	Display 1
	034	RCL4	36 04				089	RCL9	36 0 9 32	
	035	+	-55				090	LN		
	036	+	-55				091	X≢Y	-41 -24	
	037	÷	-55				092	÷	-24 35.00	Store r̂ in R ₈
	038	RCL5	<i>36 05</i>				0 93	STO8	35 0 8	Display r
	039	RCL6	3 6 06				094	RTN	24	
	040	RCL7	36 07				0 95		21 16 11	
	041	+	-55				0 96	1	01	
	042	+	-55				0 97	6SB0	23 00	
	643	+	-55				098	RCL1	36 01	
	044	ST09	35 <i>09</i>				099	x	-35	
	045	RCL1	36 01				100	2	0 2	
	046	1	01				101	GSB0	23 00	
	047	RCL8	36 08				102	RCL2	36 0 2	
	048	+	-55				103	X	-35 55	
	049	X	-35				104	+_	-55	
	0 50	RCL2	36 02				105	3	0 3	
	0 51	2	02				106	GSB0	23 00	
	0 52	RCL8	36 Ø8				107	RCL3	36 03 35	
	053	÷	-55				108	×	-35 55	
	<i>0</i> 54	×	-35				109	+,	-55	
	0 55	+	-55				110	4	0 4	
	0 56	RCL3	<i>36 03</i> -				111	GSB0	23 00	
		NOLU		1-		STERS		RCL4	36 04	18 . 19
0		1 $1_{\alpha}\mathbf{m}_{\alpha}$	$\mathbf{I}_{\alpha+1^{\mathbf{m}}\alpha+1}$	$^{3}1_{\alpha+2m_{\alpha+2}}$	$ 1_{\alpha+3}\mathbf{m}_{\alpha+3} $	⁵ 1 , m		$^{\scriptscriptstyle 6 }1_{lpha+5}$ m $_{lpha}$	$+5$ $\begin{bmatrix} 7 & 1_{\omega} m_{\omega} \end{bmatrix}$	⁸ k, r̂ ⁹ R
<u></u>			S2	S3	S4	<u>*α+4</u> ** S5	α +4	<u>-α+5</u> -α	+ 5 ω ω ω	S8 S9
S0		S1	132	33	5	33			ات ا	
A		l IB		Ic Ic	L	D			E	II
 ^		٦	,	ľ					_	

97 Program Listing II

6				// Program		ing 11			
STEP	KEY	ENTRY	KEY CODE	COMMENTS	STEP	KEY ENTRY	KEY CODE	СОММІ	ENTS
	13	X	-35						
	14	+_	-55		170				
	15 12	5 cepa	85 27 aa						
	16 17	GSB0 RCL5	23 00 36 05						
	18	X	-35						
	19	+	-55						
	20	6	<i>96</i>						
	21	6SB0	23 00						
	22 23	RCL6 X	36 06 -35		ļ				
	24	, +	-55		180				
	25	7	0 7						
	26	6SB0	23 00						
	27	RCL7	36 07 						
	28 29	X +	-35 -55	Note: insertion of	_				
	29 30	ENT†	-33 -21	an "R/S" here will	-				
	31	ENT†	-21	stop each loop					
13	32	1	Ø1	and display the				l	
	33	Ū	99	sum for user's	100				
	34 35	Ø ENT#	00 -21	inspection (hit	190			ĺ	
	36	ENT† R↓	-21 -31	R/S to continue)					
	37 37	X	- 3 5						
	38	INT	16 34						
	3 9	RT	16-31						
	40 43	X=Y?	16-33						
	41 42	GT01 R∔	22 01 -31						
	43	R↓	-31						
	44	1	01		200				
	45	-	-45					ĺ	
	46	•	-62						
	47 48	2 X	02 -35					İ	
	49	RCL8	36 08						
	50	+	-55					ļ	1
	51	ST08	35 08						
	52 53	6T0a	22 16 11						
	53 54	*LBL1 RCL8	21 01 36 08		210				
	5 5	RTN	24						
15	56	*LBL@	21 00						
	5 7	RCL0	36 00	Age constant k	<u> </u>				
	58 59	+ RCL8	-55 36 08						
	55 60	KULO X	36 60 -35						
	61	CHS	-22						
16	62	ex	33						
	63	RTN	24		220				
	64 65	≭ LBLb e×	21 16 12 33	^					
	66 66	RTN	33 24	Display λ					
	67 67	R/S	5 <i>1</i>						
				LABELS	L	FLAGS	<u> </u>	SET STATUS	
A11 ↑	m ₁ †	B13	↑ m ₃ ↑ C 1 ₅	^{™5} ↑ [□] 17 ↑ m7 ↑ ^E ት	, r̂	0	FLAGS	TRIG	DISP
12	m21	$\frac{1_4}{\lambda}$	1 1114 1 16	ш6 к	, - 	1	ON OFF		
$0 \qquad r \rightarrow 1$	<u>r</u>	1	<u>c</u>	3 e		2	0 🗆 🛛 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	DEG ဩ GRAD □	FIX ☒ SCI □
5		use 6	ed 7	8 9		3	2 🗆 🛛	RAD 🗆	ENG n 2
		1					3 □ 🛛		n_ <u>∠</u>

DEMOGRAPHY II: Expectation of Life and Reproductive Value **Program Title**

Hewlett-Packard Company Contributor's Name

1000 N.E. Circle Boulevard **Address**

State Oregon **Zip Code** 97330 Corvallis City

Program Description, Equations, Variables After storing the intrinsic rate of increase r, in register 1, the user enters age (x), age-specific survivorship (1), and age-specific fecundity (m_X) sequentially, beginning with the <u>oldest</u> age class and working toward the youngest age class. User-defined key "C" is punched and working for each age class and the computed from the equation: $E_{x} = \frac{y}{1_{x}}$ for each age class and the expectation of further life, E_x , is displayed as

Fisher's reproductive value is computed from the equation:

$$v_x = \frac{e^{rx}}{1_x} \sum_{t=x}^{\infty} e^{-rt} 1_{t} m_t$$

After all age classes have been entered and E_x and v_x for each has been computed and recorded, User-defined key D computes the net reproductive rate, R_0 , and the finite rate of increase, λ , from the equations

$$R_0 = \sum_{n=0}^{\omega} 1_x m_x$$
 and $\lambda = e^r$

Generation time is computed with the approximate formula:

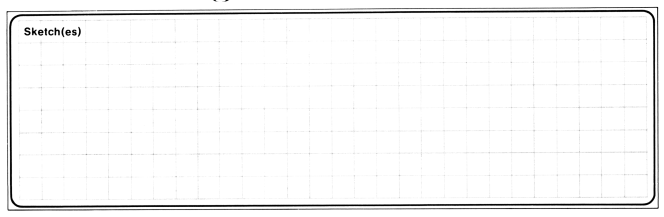
$$\hat{T} = \frac{\sum_{\alpha}^{\omega} x 1 x^{m} x}{R}$$

 $\hat{T} = \frac{\sum\limits_{\alpha}^{\omega} x^1 x^m x}{R}$ (\$\alpha\$ is age of first reproduction and \$\omega\$ is the age of last reproduction.)

Register 8 stores a sum that is used in each computation **Operating Limits and Warnings** of $v_{\mathbf{x}}$; hence an error in data entry at any age will throw off all subsequent computations for younger age groups.

This program has been verified only with respect to the numerical example given in Program Description II. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Sample Problem(s) Compute expectation of life and Fisher's reproductive value for each age group in a population with the following schedules of age-specific survivorship and fecundity:

	Age (x)	Survivorship (1 _X)	Fecundity (m_{χ})
	0	1.0	0
	1	.9	_0
	2	.7	0
the intrinsic rate \	3	.5	0
of natural increase 4	4	. 4	1
(is known to be .0552)	5	.3	1
	6	.2	1.5
	7	. 2	1
	8	.1	1
	9	.1	0.5
	10	.01	1

Also calculate net reproductive rate, the finite rate of increase and mean generation time.

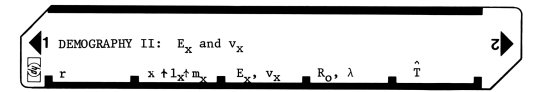
```
Reference(s) Fisher, R.A. 1930. The genetical theory of natural selection. Oxford.

Mertz, D.B. 1970. Notes on methods used in life-history studies, pp 4-17 in Connell, Mertz, and Murdoch's "Readings in ecology and ecological genetics." Harper & Row.

Pianka, E.R. 1974. Evolutionary ecology. Harper & Row.

Poole, R.W. 1974. An introduction to quantitative ecology. McGraw-Hill. Wilson, E.O. and W. Bossert. 1971. A primer of population biology. Sinauer. This program is a translation of the HP-65 Users' Library Program #03639A submitted by Eric R. Pianka.
```

User Instructions



1 Read program (in "run" mode!) 2 Enter intrinsic rate of increase in R ₁ r A	
2 Enter intrinsic rate of increase in R ₁ r	
3 Key in age (x) of oldest age group first x	
4 Key in survivorship of oldest age group, (1_x) 1_x	
first	
5 Key in fecundity of oldest age group (m _x) m _x B	(x)
first	
6 Compute and record expectation of life, E _x	E _X
7 Compute and record reproductive value, v _x	v _x
8 Repeat steps 3 through 7 for next oldest age	<u>x</u>
until all age classes have been entered and	E _X
a complete sequence of values have been	v
obtained for E _x and v _x for all ages.	
9 Calculate net reproductive rate, R _O	R _O
10 Calculate finite rate of increase, λ	λ
11 Compute estimate of mean generation time, T	Î
11 Compute estimate of mean generation time, T	1
	10 mm

97 Program Listing I

STEP	KE	Y ENTRY	KEY CODE	СОММЕ	INTS	STEP	KE	Y ENTRY	KE	Y CODE	COMM	ENTS
	001	*LBLA	21 11				057	R/S		51	Display H	30
	002	CLRG	16-53	Clear regi			05 8	RCL1	3	36 01		·
	00 3	ST01	<i>35 01</i>	Store r in	R_1		0 59	e×		33		
	004	RTN	24		_		060	RTN		24	Display 7	\
	005	*LBLB	21 12				061	*LBLE		1 15		
	006	DSP3	-63 03				062	RCL9	3	36 09		
	007	ST04	<i>3</i> 5 <i>0</i> 4	Store m _x i	n R ₄		0 63	RCL7	3	36 07		
	0 08	X≠Y	-41				<i>064</i>	÷		-24		
	009	ST03	3 5 03	Store 1 _x i	n R ₃		0 65	R∕S		51	Display 1	1
	010	X	-35				066	RCL7	3	36 07		
	011	RCL7	<i>36 07</i>				0 67	LN		32		
	012	Ť	-55				0 68	RCL1	3	36 01		
	013	ST07	35 Ø7				0 69	÷		-24	/	\
	014	R↓	-31				070	RTN		24	Disp la y T	.
	015	ST02	35 0 2	Store x in	R ₂		071	R∕S		51		
	0 16	RCL1	36 01									
	017	Х	-35									
	018	ST05	35 Ø5									
	019	RCL6	36 06									
	020	RCL3	36 03									
	021	+	-55									
(0 22	ST06	35 Ø6									
	0 23	RCL2	36 02						<u> </u>			
	0 24	RCL3	36 03			080						
	025	RCL4	36 04									
6	<i>026</i>	X	-35									
6	027	X	-35									
6	0 28	RCLS	36 39									
	0 29	+	-55									
	9 36	ST09	35 0 9									
	6 31	RCL2	36 0 2	Display age	2 X							
	032	RTN	24									
	03 3	*LBLC	21 13	1								
	934	RCL6	36 0€			090						
	035	RCL3	36 03	1								
	936	÷	-24	1								
	937	R∕S	51	Display E _x								
	938	RCL5	36 Ø5									
	0 39	CHS	-22	1					<u> </u>			
		e×	33	1			1					
	941	RCL3	36 03	1								
	042	X	-35	1								
	943	RCL4	36 04	1								
	944	Х	-35	1		100						
	945	RCL8	<i>36 0</i> 8	1								
	946	÷	-55	1								
	947	ST08	35 <i>08</i>	1					ļ			
	948	RCL5	<i>36 0</i> 5	1			4		<u> </u>			
	949	e×	33	1			1		 -			
	95 <i>0</i>	RCL3	36 Ø3	1			↓		\vdash		SET STATUS	
	05 1	÷	-24	1			+	1	\vdash	FLAGS	TRIG	DISP
	352	Х	-35	l		<u></u>	+		┵	ON OFF		
	953	RTN	24	Display v _x		110	+			$\square X$	DEG 🛛	FIX X
	3 54	*LBLD	21 14	1		110	+				GRAD 🗆	SCI □ ENG □
	355	DSP4	-63 04	1		 	+-			2 X	RAD 🗆	n_2
	356	RCL7	36 07		DEC:	CTERC				لکا ت	L	
0		t-	12	3 1	4 REGI	STERS 5		6	7		18	9
١		r	² x	$\begin{vmatrix} 3 & 1_x \end{vmatrix}$	m _X	rx		$\Sigma 1_{\mathbf{x}}$	ľ	$\Sigma 1_{\mathbf{x}} \mathbf{m}_{\mathbf{x}}$	$^{8}_{\Sigma}$ -rx $_{1_{X}}$ m $_{X}$	$\Sigma \times 1_{\mathbf{X}} m_{\mathbf{X}}$
S0		S1	S2		S4	S5		S6	S		S8	S9
1												
A		-	В	С		D			E		I	
I .												
			<u> </u>									

Program Title

DIVERSITY AND EQUITABILITY INDICES

Contributor's Name

Hewlett-Packard Company

Address

1000 N.E. Circle Boulevard

City

Corvallis

State Oregon

Zip Code 97330

Program Description, Equations, Variables

Given a data set:

$$\{n_1, n_2, n_3, \dots n_i\}$$

The program computes the following:

Diversity = 3.3219
$$(\log_{10} N - \frac{1}{N} \Sigma n_i \log_{10} n_i)$$

Equitability =
$$\frac{\text{Diversity}}{3.3219 \text{ (log}_{10}\text{N)}}$$

Where:

 n_i = number of individuals of the i th group

N = total number of individuals in the collection

 $3.3219 = conversion factor <math>log_{10}$ to log_2

Operating Limits and Warnings

n is a positive interger and n > 1.

This program has been verified only with respect to the numerical example given in Program Description II. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Sketch(es)					

Sample Proble	olem(s)	
	Compute diversity and equitability for the following data:	
The second secon	[7/0, 11 000, 000, 100, 100, 7/0, 100]	
	{ 740; 11,000; 920; 180; 180; 740; 180}	
MARKETA		
Solution(s)		
Solution(s)	[A] 740 [B] 11,000 [B] 920 [B] 180 [B] 180 [B]740 [B] 180 [B]
	[D]	L.22
	[E]	0.09
W-00.00		
Management page of Management and a second a		

```
Reference(s) Wilhm, J.L. and Dorris, T.C., 1968, Biological parameters for water quality criteria: Bioscience, v. 18, p. 477-481.

Peet, R.K., 1974, The measurement of species diversity: Annual Review of Ecology and Systematics, v. 5, p. 285-307.

This program is a modification of the Users' Library Program # 04754A submitted by W. Thomas Shoaf.
```

User Instructions

1	DIVER	SITY AND	EQUITABILITY	INDEX		2
Ini	tial	data	correct	diver	equit	/

STEP	INSTRUCTIONS	INPUT DATA/UNITS	KEYS	OUTPUT DATA/UNITS
1	Enter program			
	T		A	
2	Initialize		_A	
3	Perform step 3 for $i = 1, 2,x$	ni	В	i
	(number of individuals of the i th group)			
4	To correct entry	_	C	<i>2</i> 1
-	To correct entry	ni		i-1
5	Compute diversity		D	diversity
				1
6	Compute equitability		E	equitabiit
<u> </u>	compace equicability			
		-		
	(For a new case go to step 2)			ļ
		_		
				ļ
		-		
		_		
-				<u> </u>
		1		
				<u> </u>
				ļ
I !		1		1

97 Program Listing I

STEP KI	EY ENTRY	KEY CODE	COMMENTS	STEP	KEY ENTRY	KEY CODE	COMMENTS
001	*LBLA	21 11	Initialize, press A	A			
002	CLRG	16-53	, ,				
003	RTN	24	No individuals in	060			
004	*LBLB	21 12	No. individuals in	060			
005	ST06	35 06	ith group,				
99 6	ST+2	35-55 8 2	N = S + N				1
007 008	LOG RCL6	16 3 2 36 0 6					
009	KULU X	-35	X = S * log(s)				
010	ST+3	35-55 0 3	Z = X + Z				
011 011	1	01					
012	ST+9	35-55 09					
013	RCL9	36 0 9					
014	RTN	24		070			
015	*LBLD	21 14	Diversity				
016	RCL3	<i>36 03</i>	Z/N				
0 17	RCL2	36 02					
018	÷	-24					
019	ST04	35 04					
020	RCL2	36 0 2					
021	LOG	16 32					
022	RCL4	36 84					
0 23	-	-45	Log (N)-(Z/N)	080			
024	3	0 3		080			
025	•	-62					
026	3	03		-			
027	2	0 2		-			
028	1	0 1					
029 030	9 6701	0 9					
030 031	STO1 ×	35 Ø1 -75	3.3219*[log(N)-(Z/N	,			
0 32	RTH	-35 24	3.3219"[10g(N)=(Z/I	` 			
033	*LBLE	21 15	Equitability				
<i>034</i>	STO7	35 0 7	Iquicability	090			
035	RCL2	36 02					
0 36	LOG	16 32					
03 7	RCL1	36 01					
0 38	X	-35					
0 39	ST08	35 0 8					
949	RCL7	36 0 7					
841	RCL8	<i>36 08</i>			ļ		
04 2	÷	-24					
043	RTH	24	Commont	100			
044	*LBLC	21 13	Correct	100	 		
045 046	ST06	35 0 6		-			
046 047	ST-2	35-45 0 2		 			
94 7	LOG BCL C	16 32 76 96					
048 049	RCL6 ×	36 06 -35					
043 050	ST-3	-35 35-45 03					SET STATUS
<i>0</i> 51	1	01 01				FLAGS	TRIG DISP
0 52	ST-9	35-45 09				ON OFF	
0 53	RTN	24		110		0 □ 🔯	DEG 🔀 FIX 🖫
0 54	R/S	51		110			GRAD SCI
 		+				2 🗆 🗓	RAD ENG n 2
		1	I REGI	STERS			
0	1	2	3 4	5	6	7	8 9
	Used		ΣSlogS Used	ļ	S	Used	Used i
S0	S1	S2	S3 S4	S5	S6	S7	S8 S9
<u></u>	<u> </u>	В	C	D		_	I
Α				ا ا		_	

Program Title

NICHE BREADTH AND OVERLAP AND SHANNON'S H AND HORN'S R

Contributor's Name

Hewlett-Packard Company

Address

1000 N.E. Circle Boulevard

City

Corvallis

State Oregon

Zip Code 97330

Program Description, Equations, Variables Given resource utilization vectors for two species, this program computes niche breadths for both species using Simpson's diversity equation:

$$B = \frac{1}{\sum_{i=1}^{n} p_{i}^{2}}$$

 $B = \frac{1}{\sum_{i=1}^{n} p_{i}^{2}}$ where the p_i are the proportion of the ith resource used and n is the total number

Thus computed, B varies from 1/n to n, and can be standardized as a fraction of the maximal possible value, $B_{\mbox{max}}$, which is equal to n, by dividing by n.

Overlap is computed using Pianka's (1974) formula:

$$0 = \frac{\sum p_i q_i}{\sqrt{\sum p_i^2 \sum q_i^2}}$$

where the sums are over all i from one to n, p_i and q_i are the proportional use values of resource i by species A and B, respectively. Thus computed, overlap varies from zero to one. For some purposes, the Arc sine of this value, which varies from 0° to 90°, is useful.

Given two vectors of proportional relative importance, $\textbf{p}_{\dot{\textbf{1}}}$ and $\textbf{q}_{\dot{\textbf{1}}},$ this program computes Shannon's information theoretic index of diversity for each, according

(Continued on next page)

Operating Limits and Warnings None; zero entries are permitted.

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Program Title			
Contributor's Name			
Address			***************************************
City	State	Zip Code	

Drogram	Description.	Faustions	Variables

to the formula:

$$H_p = -\sum p_i Lnp_i \text{ or } -\sum q_i LN q_i. = H_q$$

Each H is expressed both as an absolute value and as a "J" value, which is a fraction of the maximal possible H value, H_{\max} .

The program also computes Horn's (1966) measure of overlap, $R_{\bar{0}}$, from a simplified computational formula:

Overlap =
$$R_0 = \frac{((H_p + H_q)/2) - H_r}{Ln \ 2} + 1$$

where H_r is computed from the p_i and q_i vectors, with $r_i = (p_i + q_i)/2$ according to Shannon's formula (above). H_r is stored in register 6.

Operating Limits and Warnings

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Sketch(es)

Sample Problem(s) 1. Two species (or individuals or populations or other organismic units) use four resources (microhabitats, foods, or whatever) in the following proportions:

oportions.	Resource 1	Resource 2	Resource 3	Resource 4
Species A	0.2	0.5	0.1	0.2
Species B	0.3	0.1	0.4	0.2

2. Two communities have a total of five species in the following proportional relative abundances (or importances); Community A Community A

	 Community A	Community E
Species 1	.1	0.00
Species 2	.2	. 4
Species 3	0.0	. 2
Species 4	•5	.1
Species 5	. 2	.3

What is the diversity of each and how much do they overlap?

Solution(s) 1. a. Initialize (zero all registers): [CL REG]

b. Key in p_i , q_i data as follows:

0.2 [†] 0.3 [A] 0.5 [†] 0.1 [A] 0.1 [t] 0.4 [A] $0.2 [\uparrow] 0.2 [A]$

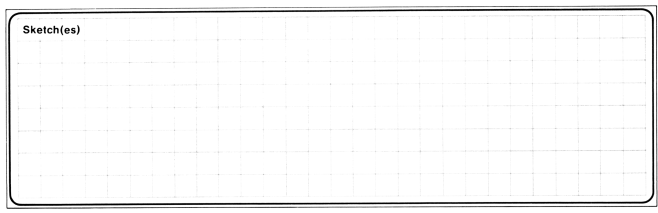
Compute niche breadths: [B] 2.94 [R/S] 0.74 (B_p/n)

[R/S] 3.33 [R/S] 0.83 (B_q/n) Compute niche overlap: [C] 0.59, Arc sine: [R/S] 36.51

Check Sums of p_i, q_i: [D] 1.00, [R/S] 1.00

(Continued on next page)

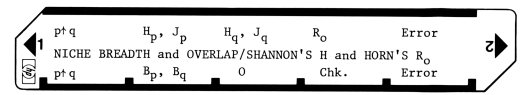
Simpson, E.H. 1949. Measurement of diversity. Nature 163: Reference(s) Pianka, E.R. 1974. Niche overlap and diffuse competition. Proc. Nat. Acad. Sci., USA 71: 2141-2145. May, R.M. 1975. Some notes on measurements of the competition matix, α . Ecology 56: in press.



Sample Problem(s)	
	Initialize (zero all registers): [CL REG] Key in the five sets of p _i and q _i data, letting i run from 1 to 5:
	0.1 [†] 0.0 [f] [A] 0.2 [†] 0.4 [f] [A]
	0.0 [↑] 0.2 [f] [A] 0.5 [↑] 0.1 [f] [A]
	0.2 [↑] 0.3 [f] [A]
c. d.	Compute diversity of community A (H_p) : [f][B] 1.28, [R/S] 0.80(H/H _{max}) Compute diversity of community B (H_q) : [f][C] 1.22, [R/S] 0.76(H/H _{max})
e.	Compute community overlap, R _o [f][D] 0.71

```
Reference(s)
Shannon, C.E. and W. Weaver, 1949. The mathematical theory of communication Univ. Illinois Press, Urbana.
Horn, H.S., 1966. Measurement of "overlap" in comparative ecological studies. Amer. Naturalist 100: 419-424.
Lloyd, M.J.H.Zar, and J. Karr. 1968. On the calculation of information theoretical measures of diversity. Amer. Midlan Naturalist 79: 257-272. This program is a modification of the Users' Library Programs # 02158A & #03150A submitted by Eric R. Pianka.
```

User Instructions



STEP	INSTRUCTIONS	INPUT DATA/UNITS	KEYS	OUTPUT DATA/UNITS
1	Enter program			
2	Initialize (zero registers)		CL REG	
3	For niche breadth and overlap, go to step 4.			
	For Shannon's H and Horn's R _O , to to step 9.			
4	Enter pairs of data, p _i , [ENTERt] \(\backslash \) repeat til	Pi	†	
	then $q_{ extbf{i}}$ finished	qi	A	
5	To remove erroneous data pairs, p; [ENTERt]	Pi	<u></u>	
	qi	qi	E	
6	After all n data pairs have been entered,	• 1		
	compute niche breadth for first species		В	Вр
	compute its standardized niche breadth		R/S	B _p /n
	compute niche breadth for second species		R/S	B _q
	compute its standardized niche breadth		R/S	B _q /n
7	Compute niche overlap		C]	0
	compute Arc sin of overlap		R/S	Arc sin(0
8	Check sum of p _i to see if it is unity		D [ΣΡί
	check sum of q _i to see if it is unity		R/S]	Σqi
9	Enter data pairs	Pi		
		q _i	f A	
	repeat 9 until all pairs of data are entered			
10	To remove erroneous data pairs, pi [ENTERt]	p _i	[ENT↑][
	q _f	qi	f [E]	
11	Compute diviersity H _B		[f B	-Σp _i LNp _i
12	Compute J (H/H _{max})		R/S	H _p /H _{max}
13	Compute diversity $H_{\mathbf{q}}$		f C	$-\Sigma q_i LNq_i$
14	Compute J (H/H _{max})		[R/S]	H _q /H _{max}
15	Calculate overlap R _o		f D	Ro

20			97	Prog	çram	Lis	ti:	ng I			
STEP K	EY ENTRY	KEY CODE		COMMEN	TS	STEP	ΚE	Y ENTRY	KEY CODE	COMI	MENTS
901	*LBLA	21 11	p ↑	<i>a</i>		0	57	RCL1	36 01		
002	ST07	35 07	P	ď		0:	58	RTN	24		
003	ST+4	35-55 04					59	*LBLD	21 14	Chk.	
004	Χ2	53				8	60	RCL2	<i>36 02</i>		
995	ST+5	35-55 <i>0</i> 5				9	61	R/S	51		
00€	X≠Y	-4 j					62	RCL4	36 04		
007	ST+2	35-55 0 2					63	RTN	24		
008	XΣ	53					64	*LBLa	21 16 11	р↑ q	
009	ST+3	<i>35-55 03</i>					65	ST07	35 0 7	1	
910	LSTX	16-63					66	X≠Y	-41		
011	RCL7	<i>36 07</i>					67	ST08	35 0 8		
012	X	-35					68	ST+4	35-55 04		
013	ST+6	35-55 <i>06</i>					69	€	00		
014	1	0 1					70	X=Y?	16-33		
015	ST+1	35-55 01					71	GT01	22 61		
816	RCL1	36 0 1					72	R↓	-31		
017	RTN	24					73	ENT†	-21		
018	*LBLB	21 12	В _р ,	B.,			74	LN	32		
019	RCL3	36 0 3	ъ,	Pq			 75	x	-35		
020	1/X	52					76	ST-5	35-45 Ø5		
021	R/S	51					77	≠LBL1	21 01		
022	RCL1	36 01					78	RCL7	36 0 7		
023	÷	-24					79 79	ST+2	35-55 0 2		
024	R/S	5 <i>i</i>					80	9	00		
025	RCL5	36 0 5					81	X=Y?	16-33		
0 26	1/X	52					82	GT02	22 0 2		
<i>027</i>	R/S	51					83	8102 R↓	-31		
028	RCL1	36 01					84 -	ENT†	-21		
8 29	÷	-24					8 5	LN	32		
030	RTN	24					86	X	-35		
031	*LBLC	21 13	0				87	ST-3	35-45 <i>0</i> 3		
0 32	RCL3	36 03	Ū				88	*LBL2	21 02		
033	RCL5	36 0 5					89 89	RCL8	36 0 8		
<i>034</i>	X	-35					90 90	RCL7	36 0 7		
035	1 X	54					91	KGL! ÷	-55		
0 36	RCL6	36 0 6					92	2	-33 0 2		İ
037	X≢Y	-41					93	÷	-24	-	
038	÷	-24									
039	R/S	51					94 95	0 X=Y?	00 16-33		
040	SIN-	16 41	Take	s Arc si	in of O		96	6T03	22 0 3		
041	RTN	24			01 0		97	6103 R↓	-31		
042	*LBLE	21 15	Erro	r			98	ENT†			
043	ST07	35 0 7	шт				20 99		-21 70		
044	ST-4	35-45 0 4						LN X	32 -35		
045	χ2	53					00 01	ST-6	-33 35-45 06		
046	ST-5	35-45 Ø5						*LBL3	21 03		
047	X≠Y	-41					02 03				
048	ST-2	35-45 02						1 ST+1	01 35-55 01		
049	X2	53					94 os				
<i>0</i> 50	ST-3	35-45 0 3					05 az	RCL1	36 01		
<i>051</i>	LSTX	16-63					06 67	RTN +I DI L	24	H _p , J _p	
052	RCL7	36 0 7					07 08	≉LBL6 RCL3	21 16 12 76 97	р, р	
0 53	X	-35						RULS R/S	36 0 3		
054	ST-6	35-45 Ø6					09 10		51 36 0 1		
0 55	1	01					16 11	RCL1 LN	36 6 1 32		
956	ST-1	35-45 Ø1				1	11 12	STO7	35 07		
					REGIS					To	
0	1 n	2 Σp_i	3 Σ 1	P_{i}^{2} 4	Σqi	Σq _i ²		⁶ Σpiqi	⁷ Used	8	9
1	n	Σqi		liLNqi	ΣP_i	$-\Sigma p_1LN$	Ip 1	-XriLN	r _i last q _i	last pi	1
SO	151	52	63	c/		C 5		C.C	C7	100	100

				nEGI.	5 . L				
0	¹ n	² Σ p _i	$^3 \Sigma p_1^2$	⁴ Σq _i	5 Σq_{i}^{2}	6 Σ Piqi	⁷ Used	8	9
	n	Σqi	$-\Sigma q_i LNq_i$	$\Sigma p_{\mathbf{i}}$	-Σp _i LNp _i	-XriLNri	last q _i	last p _i	
S0	S1	S2	S3	S4		S6	S7	S8	S9
Α	В		С		D	E		I	

97 Program Listing II

0750 45	=				COMMENTS		CTED			KEY 0005	0014	2
STEP KE 113	Y ENIRY		CODE 24		COMMENTS		STEP		Y ENTRY		COMI	MENTS
114	RTN		24					169	ENTT	-21 72		
115	*LBLc	21 16			т			178	LN	32 35	1	
116	RCL5	36		H_q ,	¹ q			171	X CT.5	-35	1	
117	R/S		51					172	ST+6	35-55 0 6	1	
118	RCL7	36						173	≠LBL 6	21 06		
119	RGL1		24					174	1	01		
120	RTN		24 24					175	ST-1	35-45 01		
121								176	RCL1	36 01		
	*LBLd	21 16		R_{o}				177	RTH	24		
122 123	RCL3 RCL5	36 76					l	178	R/S	51		
123	#CEJ	36	5 5								1	
125	2		92 9 2				180				1	
	÷			i			<u> </u>	-			1	
126			24 07								1	
127	RCL6	36										
128	-		45									
129	2		0 2							.		
130	LN		32 24								1	
131	÷		24									
132	. 1		01 EE								1	
133	† DTN		55 34				100	_			1	
134	RTN		24 15	_			190				4	
135	<i>‡LBLe</i>	21 16		Error							4	
136	STO7	35									4	
137	X‡Y czoc		41 ao					_				
138	STO8	35 ·										
139	ST-4	35-4 5										
140	<i>0</i> v_vo		00 77									
141	X=Y?	16-					ļ					
142	GT04	22						_				
143	R↓		31				200					
144	ENTT		21 20				200	-		ļ		
145	LN		32 35								4	
146	x ST+5							+				
147	*LBL4	35-55 (<u> </u>	+			4	
148		21 (1	
149	RCL7	36 (_			1	
150	ST-2	35-45						+		ļ	4	
151	0 v_vo		90 77					+-			4	
152 157		16-3						+			4	
153 154	GTO5	22 (210	+		ļ	1	
154 155	R↓ Ent†		31 21				210	+-		 	4	
155 156	LNIT		21 32					+-			4	
156 157	LN X		32 35					+-			4	
157 158	sT+3	35-55 (+		 	4	
158 159	*LBL5	21 (+-			1	
160	RCL8	36 t						+-		†	1	
161	RCL7	36 (+	•		1	
162	* *		55					+			1	
163	2		92 92					+-		†	1	
164	÷		24				220	+-		†	1	
165			e e					1		1	1	
166	X=Y?	16-									1	
167	6T06	22 (]	
168	R↓		31								L	
A	T=			LAE	BELS	T-		1	FLAGS		SET STATUS	
A Pitqi	В Вр,	$B_{\mathbf{q}}$	^C 0,A:	cc sin	D Σp _i , Σq _i	E Pi	tq _i	0		FLAGS	TRIG	DISP
2	Th	-	_		Id	T	-	11		ON OFF		1

	LABELS						SEI STATUS	
^A p _i †q _i	B $^{B}_{p}$, $^{B}_{q}$	$^{\mathrm{C}}$ 0,Arc sin	^D Σp _i , Σq _i	E p _i ∱q _i	0	FLAGS	TRIG	DISP
a Pi, qi	b H_{p} , J_{p}	c Ha, Ja	d R _O	e Error	1	ON OFF	DEG X	FIX X
0	¹ Used	² Used	³ Used	4 Used	2	1 🗆 🗓	GRAD □ RAD □	SCI □ ENG □
⁵ Used	6 Used	7	8	9	3	3 🗆 🗓	TIAD 🗆	n_2

Program Title POPULATION SIZE ESTIMATE (JOLLY'S METHOD)

Contributor's Name Hewlett-Packard Company

Address 1000 N.E. Circle Boulevard

City Corvallis State Oregon Zip Code 97330

Program Description, Equations, Variables Using Jolly's (1965) symbols, let

n = number in sample (at time t)

m = number of marked individuals in sample at time t (recaptures)

s = number of marked individuals released from sample (need not be all)

r = number of marked individuals released from sample which are subsequently
 captured again at least once (known to be alive later)

z = number of individuals marked prior to sample, but not captured in sample, and which are subsequently captured in later samples (hence the individuals known to be alive at time of sample).

The estimated size of the marked portion of the population at time t is given by

$$\hat{M} = m + z \frac{s}{r}$$

Estimated population size, N, is simply \hat{M} times the ratio of marked plus unmarked individuals in the sample (n) to marked individuals in the sample (m):

$$\hat{N} = \hat{M} \cdot \frac{n}{m}$$

Jolly's formula for the standard error of the population size is as follows:

S. E.
$$\hat{N} = \hat{N}(\hat{N} - r) \left\{ \frac{\hat{M} - m + s}{\hat{M}} \left(\frac{1}{r} - \frac{1}{s} \right) + \left(\frac{1}{m} - \frac{1}{n} \right) \right\}$$

Operating Limits and Warnings The following inequalities must be met:

 $n \geqslant m \geqslant s \geqslant r$

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Sketch(es)

Sample Problem(s) Suppose you collect a sample of 100 isopods from a marked population and recover 64 marked animals. Of these 64 you retain 16 gravid females for further observation of fecundity-size relationships (this means that 48 are released). Subsequent sampling of the population produces 28 recaptures of these 48 animals. Lastly, you knew that you had only 80 marked animals in the study plot, so that you collected all but 16 of the marked individuals. Hence Jolly's input values are

n = 100 m = 64 s = 48 r = 28 z = 16

Estimate the population size and its standard error.

Solution(s)

```
100 [ENTER \uparrow] 64 [A] 48 [ENTER \uparrow] 28 [ENTER \uparrow] 16 [B]
[C] \rightarrow 142.86 (\hat{N})
[R/S] \rightarrow 10.47 (SE)
[D] \rightarrow 91.43 (\hat{M})
```

```
Reference(s)

Jolly, G.M. 1965. Explicit estimates from capture-recapture data with both death and dilution -- stochastic model.

Biometrika 52: 225-247.

Poole, R.W. 1974. An introduction to quantitative ecology. McGraw-Hill. (pp. 307-312.)

This program is a translation of the HP-65 Users' Library

Program # 03257A submitted by Eric R. Pianka.
```

User Instructions

1 POPULATION	N SIZE ESTIMA	TE (JOLLY'	S METHOD)		2
n,m	s,r,z	N,SE	М	_ M—m+s	/

	Load side 1			1
2				
2				
	Enter n	n	ENT↑	
-				
3	Enter m	m	A	
4	Enter s	S	ENT	
5	Enter r		ENT	
	rurer +	r	ENI	
6	Enter z	z	В	
7	Compute N		C	Ñ
	_			
8	Compute SE		R/S	SE
9	Display M̂		D	м̂
10	Display \hat{M} - m + s (if desired)		E	M-m+s
\rightarrow				
1				
1				
\dashv				
\dashv				
-+				

97 Program Listing I

	KEY ENTRY	KEY CODE	COMMENTS	STEP	KEY ENTRY		25 COMMENTS
	*LBLA	21 11			057 *LBLE	21 15	
<i>002</i>		35 02			058 RCL7	36 07	
0 03		-31			059 RCL2	36 02	
004	S701	35 B1			060 -	-45	
005	RTN	24				36 83	
<i>886</i>		21 12					
0 07		<i>3</i> 5 05			062 +	-55	
008		-31			063 RTN	24	
009		35 Ø4			.064 R/S	. 51	
0 10		-31			+		4 1
011		35 03			 		- 1
011 012		23 es 24					1
		24			-		1
0 13		21 13					1
014		<i>36 02</i>		070]
015		<i>36 0</i> 5]
016		36 03]
017		36 04					1
018		-24]
019	X	-35					1
020		-55	1 .		1	1	1
0 21		35 Ø7	Store M		1		1
022		36 8 1					1
023 023		-35			1		1 1
024 024		36 0 2		080			1 1
0 25		-24	Ì		1	<u> </u>	1
025 026		35 Ø8	Chama ŵ			<u> </u>	1
0 20 027		51	Store Ñ		-	<u> </u>	1
						 	1
028		36 01 45			 	 	1
025		-45			-		1
030		36 0 8			ļ		1
031		-35			ļ		1
0 32		36 67]
0 33		36 02]
034		-45		090]
0 35	RCL3	36 <i>0</i> 3]
036	+	-55					1
0 37		<i>36 07</i>					1 1
03 8		-24					i l
039		36 04					1
040		52					1
041	RCL3	36 03					1
042		52 52	İ		†		1
042 043	1/A -	-45	1		†	†	1 1
				100	†	<u> </u>	1
044 045		-35 74 00	1		†	1	1
94 5		36 02 .	i	-	 	†	I
04 <i>6</i>	1/X	52			 	1	1
047	RCL1	36 0 1 .	1		 	†	1
0 48	1/X	52 -45			 	<u> </u>	l
04 9	-	-45		-	 	+ -	SET STATUS
0 50		-55 -35 -54 -24		-	+	+	
0 51	X	-35		-	 	FLAGS	TRIG DISP
05 2	IX	54			+	ON OFF	DE0 17 511 511
05 3	RTN	24		110	 		DEG X FIX X
054	*LBLD	21 14	n. 1 ô	110	 	1 🗆 🛚	GRAD G SCI G
05 5	RCL7	36 07 ·	Display M		+	2	RAD ENG n_2
056	RTN	24		DEGISTERS	L		
	14			REGISTERS	6	T ₇	8 9
0	\int_{0}^{1} n	2 m	$\begin{vmatrix} 3 & & 4 \\ & s & & r \end{vmatrix}$	5	0	$\hat{\mathbf{M}}$	$\begin{bmatrix} \hat{N} & & & \end{bmatrix}^g$
S0	S1	S2	S3 S4	z S5	S6	M	S8 S9
30	21	132	53 54	35	30	3'	
A			lc l	D		E	li li
A		J		ا		L	-

Program Title CELL PHASE AND CYCLE TIMES

Contributor's Name Hewlett-Packard Company

Address 1000 N.E. Circle Boulevard

City Corvallis State Oregon Zip Code 97330

Program Description, Equations, Variables

Where n_{G1} = the number of cells in G1 phase n_S = the number of cells in S phase n_{G2} = the number of cells in G2 phase n_M = the number of cells in M phase $n_{G1} + n_S + n_{G2} + n_M$ = total number of cells.

The proportion of cells in each phase $N_x = \frac{x}{n}$ is computed. The proportion of cells in each stage is related to the time required for completion of that phase:

 $N_{x} = 2 \frac{T_{x}}{T} - 1$

where $\textbf{T}_{\textbf{X}}$ is the phase time and $\,\textbf{T}$ is the cell cycle time.

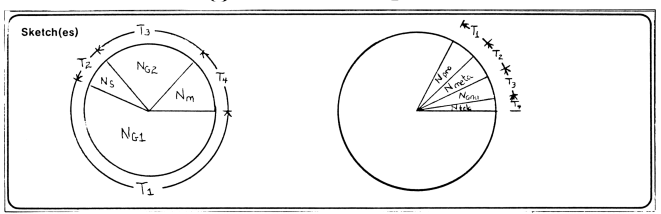
Given a starting concentration of cells, $n_{\rm O}$, and a subsequent concentration of cells, n, after a measured amount of time, t, has elapsed, the program computes the cell cycle time, T, from the equation:

$$n = n_0 2^{\frac{t}{T}}$$

Operating Limits and Warnings Data must be obtained from cells growing in the logarithmic growth phase. Viability of the cells should be confirmed as no adjustment is made for cell death. Please note that the sketches drawn employing circles to represent cell cycles are used for simplicity of expression. However, the relationship between the proportion of cells in a given phase and the amount of the cell cycle time used is not linear.

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Sample Problem(s)

- 1. In a randomly dividing cell culture with a cell cycle time of 24 hours, a sample is evaluated and 500 cells are found in M, 1000 in G2, 500 in S, and 1500 in G1 phase. Calculate the time spent in each phase: T_{G1} , T_{S} , T_{G2} , and T_{M} .
- 2. In this culture the cells in mitosis were examined, and it was found that 30 cells were in telophase, 50 in anaphase, 70 in metaphase, and 350 in prophase. As above, the cell cycle time is 24 hours and the total number of cells counted is 3500. Calculate the time spent in each phase of mitosis:

Tpro, Tmeta, Tana, and Ttelo.

(Continued on next page)

Solution(s)

1. 24 [ENTER] 3500 [f][A] 1500 [ENTER \uparrow] 500 [ENTER \uparrow] 1000 [ENTER \uparrow] 500 [A] \rightarrow 3500

 $[B] \rightarrow 8.35 \text{ hours}$

[C] \rightarrow 3.30 hours

 $[D] \rightarrow 7.73 \text{ hours}$

 $[E] \rightarrow 4.62 \text{ hours}$

2. 350 [ENTERt] 70[ENTERt] 50[ENTERt] 30[A] $\rightarrow 500$

 $[B] \rightarrow 3.17 \text{ hours}$

 $[C] \rightarrow 0.67$ hours

 $[D] \rightarrow 0.49 \text{ hours}$

 $[E] \rightarrow 0.30 \text{ hours}$

 $T_{G1} = 8.35 \text{ hours}$ $T_{S} = 3.30 \text{ hours}$

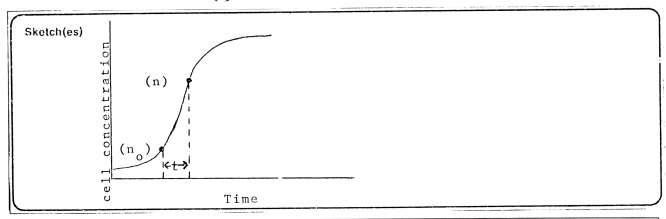
 $T_{G2} = 7.73$ hours

 $T_{\mathbf{M}}^{-}$ = 4.62 hours

 $T_{pro} = 3.17 \text{ hours}$ $T_{meta} = 0.67 \text{ hours}$ $T_{ana} = 0.49 \text{ hours}$

 $T_{telo} = 0.30 \text{ hours}$

Puck, Theodore T., and Steffen, Jan. Life cycle analysis of Reference(s) mammalian cells. I. Biophysical Journal 3: 379-397. 1963 This program is a modification of the Users' Library Programs #01522A and #01630A submitted by James M. Mason.



Sample Problem(s)

3. Find the cell cycle time, T, of cultured cells (represented above) where a starting concentration of cells, $n_{\rm O}$, a subsequent concentration of cells, $n_{\rm O}$, and the elapsed time, t, are given:

 $n_0 = 100,000 \text{ cells/ml}$

n = 500,000 cells/ml

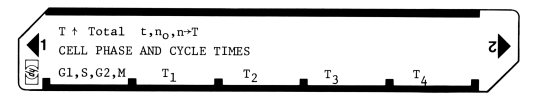
t = 48 hours

Solution(s)

3. T = 20.67 hours $48[ENTER†] 1[EEX] 5[ENTER†] 5[EEX] 5[f][B] \rightarrow 20.67 \text{ hours}$

Reference(s)

User Instructions



STEP	INSTRUCTIONS	INPUT DATA/UNITS	К	EYS	OUTPUT DATA/UNITS
1	Load side 1				
2	If cell cycle time is known go to step 3				
	If cell cycle time is to be calculate				
	go to step 5				
3	Enter cell cycle time	T	<u></u>		
	Enter total number of cells	total	f	A	T
4	Go to step 7				
5	Enter total number of cells	n total	f	A	
6	Enter elapsed time	t	[↑		
	Enter starting concentration of cells	no	[
	Enter subsequent concentration of cells	n	f	B]	T
7	Enter number of cells in four phases:				
	# cells in G1 phase or prophase	n _{G1} or n pro	[†		
	# cells in S phase or metaphase	ns or n meta		i i	
	# cells in G2 phase or anaphase	n _{G2} or n ana	1	1	
	# cells in M phase or telophase	n _M or n telo	A	i i	total 4 phase
8	Compute time for G1 or prophase	1	В	i i	T ₁
9	Compute time for S or metaphase		c		T ₂
10	Compute time for G2 or anaphase		D		T ₃
11	Compute time for M or telophase		E		T ₄
12	Find $T_1 + T_2 + T_3 + T_4$		RCL	7	ΣT ₁₋₄
	<u> </u>		Ì	i i	1-4
			i i	i i	
			j	i i	
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			i	i i	
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			[]		
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			i i		
			i		

97 Program Listing I

STEP K	EY ENTRY	KEY CODE	COMMENTS	STEP	KE	Y ENTRY	KEY CODE	COM	MENTS
001		21 11	Enter cell counts		0 57	RCL5	<i>36 0</i> 5		
002		35 01			95 8	-	-45		
003		-55			05 9	RTN	24	m	
004	STO2	35 0 2			969	*LBLD	21 14	т3	
005		-55			061	RCL1	36 0 1		
<i>006</i> 007		35 0 3 - 5 5			062	GSB0	23 00 35 05		
90 8		-55 35 0 4			063 064	STO5 RCL2	35 6 3 36 6 2		
889		24			065	GSB0	23 00		
010		21 16 11	Enter T and n total		966	RCL5	36 0 5		
011	ST08	35 0 8			067	-	-45		
012		-41			06 8	RTN	24		
013		35 0 6			069	*LBLE	21 15	^T 4	
014	RTH	24			<i>070</i>	RCL1	36 0 1	4	
015		21 16 12	Cell cycle time(T)		071	GSB0	23 0 0		
016		16 32			<i>072</i>	RTH	24		
017		-41 16 30			073	R∕S	51		
0 18 0 19		16 32 - 4 5			+-			ł	
019 020		35 Ø6			+-				
020 021	3106 R↓	-31			+				
021		02			1		1		
023 023		16 32							
024		-35		080					
<i>0</i> 25	RCL6	36 0 6							
0 26		-24			1				
0 27		<i>35 06</i>	Stores cycle time		-				
0 28		24			+				
029 070		21 00	Computational loop		╁				
030 071		36 0 8			+				
031 032	÷	-24 01			+				
03Z 033		-55			 				
<i>034</i>		16 32		090	1				
035		02							
<i>036</i>		16 32							
037	÷	-24							
0 38	RCL6	36 06			_				
0 39	X	-35			-				
040		24	m.		+				
841	*LBLB	21 12	^T 1.		+		+		
042 043	RCL3	36 03 23 00			+				
943 944	6580 ST05	23 00 35 05		100	†				
045	RCL4	36 0 4							
046	6SB0	23 00							
04 7	ST07	35 07	Total of T_1, T_2, T_3, T	4					
04 8	RCL5	36 05			1				
04 9	-	-45			+			CET CTATUS	
050	RTN	24	Tr.		+			SET STATUS	
051	*LBLC	21 13	^T 2		+		FLAGS	TRIG	DISP
952 957	RCL2	36 0 2	ł		+		ON OFF	DEG 🛛	FIX 🛛
053 054	6SB0 ST05	23 00 35 05	ļ	110			1 0 🗓	GRAD □	SCI □
054 055	RCL3	36 0 3					2 🗆 🛚	RAD 🗆	ENG □ n_2
956	GSB0	23 0 0 L					3 🗆 🛚		"
	14		REGIS	TERS		6	7	8 + 1	9
0	n_4	$ ^{2}n_{4} + n_{3}$		scra	tch	⁶ cell cycle	$^{'}_{1}$ + $_{1}$ +		ľ
			n_2 + n_2 + n_1	for ca	ions	time	T_3 + T_4	of cells	
S0	S1	S2	S3 S4	S5		S6	S7	S8	S9
Α		В	С	D			E	I	

Program Title CROSSOVER: LOCATIONS/PRODUCTS

Contributor's Name Hewlett-Packard Company

Address 1000 N. E. Circle Boulevard

City Corvallis State Oregon Zip Code 97330

Program Description, Equations, Variables This program calculates the locations of all crossovers which must occur to produce a given progeny chromatid. Two-strand crossovers between complementary parental homologues are assumed. Following a decimal point, the presence or absence of dominant alleles are represented by '1' or '0' respectively. A chromatid with alleles AbcdEFgh is expressed as .10001100. Crossover regions are numbered consecutively 1, 2, ..., n, from left to right. Crossover region 1 is the region between the two digits nearest the decimal point; region n is between the two digits farthest from the decimal point. After entering parental genotype, progeny genotype and, the total number of crossover regions, n, the program specifies each region in which a crossover occurred to produce the progeny chromatid. The presence of a crossover in region 1 is represented by 1; a crossover in region n, by n. Absence of a crossover in a region is denoted by 0. Crossover locations are calculated by an algorithm which successively compares adjacent and progeny gene pairs and determines whether progeny genes are in parental or crossover configurations.

(Continued on next page)

Operating Limits and Warnings

The maximum number of testable loci is 10. The decimal point must always be entered before the binary equivalent genotype is entered. Use only 1's and 0's. To prevent logarithmic conversion errors produced in the algorithm from appearing in the results, results must be read with calculator set to DSP 0.

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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State	Zip Code	
	State	State Zip Code

Program Description, Equations, Variables The program calculates the genotype of a chromatid produced by crossovers between specified loci of a parental chromatid and its complementary homologue. The presence or absence of a dominant parental allele is represented by '1' or '0' respectively. The parental chromatid with alleles AbcdEFgh is expressed as 10001100; its complementary homologue with alleles AbcDefGH, as 11110011. Crossover regions are numbered consecutively 1, 2, ..., n, from left to right. Crossover region 1 is the region between the leftmost two digits and crossover region n is between the rightmost two digits. After specifying crossover regions, the program provides the genotype resulting from crossovers in the designated regions. The new genotype is shown as a number consisting of 1's and 0's (a binary equivalent) with length equal to the number of parental alleles. The new genotype is calculated by means of an algorithm which successively produces binary complements of the parental strand to the right of the crossover regions specified.

Operating Limits and Warnings The number of genes, g, must be entered before calculations can be effected. Parental genes limited: $2 \leqslant g \leqslant 10$. The first digit of the parental genotype should be 1. No more than 3 crossover regions can be specified for a single calculation.

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Program Description 11

Sample Problem(s) 1. Given the paternal chromatids, Pl, observed progeny chromatids, F1, and 7 crossover regions as shown above: a) In what regions did crossovers occur producing the progeny chromatid with the genotype AbcDEfGh? b) The progeny chromatid with the genotype abCDefGH? 2. Given the parental chromatids above; a) What is the genotype of the progeny chromatid resulting from crossovers occurring in regions 2, 3 and 5? b) In regions 1, 4 and 6? 1. a) .10001100 [ENTER*] .10011010 [A] Solution(s) $7 [B] \rightarrow 34507$ Crossovers occur in regions 3, 4, 5, and 7 b) .01110011 [ENTER+] .00110011 [A] 7 [B] \rightarrow 1200000 Crossovers occur in regions 1 and 2. 2. a) 8 [E] 10001100 [ENTERt] 2 [ENTERt] 3 [ENTERt] 5 [C] \rightarrow 10101011(AbCdEfGH) 10001100 [ENTER↑] 1 [ENTER↑] 4 [ENTER↑] 6 [C] → 11111111 (ABCDEFGH)

Reference(s)

Srb, A. M., R. D. Owen and R. S. Edgar. General Genetics, 2nd Ed.,
1965. W. H. Freeman and Co., San Francisco. Chapter 6.

This program is a modification of the Users' Library Programs
#01523A and #01524A submitted by James N. Adams.

User Instructions



STEP	INSTRUCTIONS	INPUT DATA/UNITS	KEYS	OUTPUT DATA/UNITS
1	Load side 1 and side 2			
2	For crossover locations go to step 3.			
	For crossover products go to step 7.			
3	Input parental genotype, P1, following a			
	decimal point. Use 1 for a dominant and 0			
	for a recessive allele	.P1	ENT↑	
4	Input progeny genotype, F1	.F1	Α	
5	Input total crossover regions n (number of			
	genes -1 = n) and calculate regions	n	В	Xo's
	in which crossovers occur (Xo's) (0's in			
	result are ignored. E. g. Xo's=1004067 means			
	crossovers in regions 1, 4, 6 and 7.)			
6	For new case go to step 3.			
7	Input number of genes, g, in parental chromati	l g	E	g
8	Input genotype, P1, using 1 for dominant and			
	0 for recessive.	P1	↑	P1
	(For multiple crossovers go to step 12)	1		
9	Specify desired crossover region, Xo	Xo		
10	Calculate F1 genotype		D	F1
11	For an additional single Xo, go to 9			
	or			
	For a new parental type, go to 8			
	or			
	For a new case go to 7			
12	Specify first crossover region, Xol	Xo1	†	
13	Specify second crossover region, Xo2			
	or 0 for none	Xo2	↑	
14	Specify third crossover region, Xo3			
	or 0 for none	Xo3		
15	Calculate Fl genotype		С	F1
16	For an additional single Xo go to 8			
	or			
	For additional multiple Xo's go to 12			
	or			
	For a new parental type go to 8			
	or			
	For a new case go to 7			

Program Listing I

STIC STIC	STEP	KE	Y ENTRY		EY CODE		COMM	IENTS	STEP		Y ENTRY	KEY	CODE		СОМ	MENTS
602 5102 53 02 603			*LBLA		21 11	STO	F 1			0 57	RCL1			Get	new s	genes
Bell STOC	é	002	ST02			1										
805 ST01 ST0 ST0 P1	6	003	ST06		35 <i>08</i>	1				05 9					ı com	Julison
## 806 \$105 \$ 5 5 5 5 6 6 6 6 7 7 8 6 7 8 7 8 7 8 8 8 8 8 8 8	é	ØØ4	XZY		-4.	1				$\theta 6 \theta$						i
Book	6	005	5T01		35 01	STO	P1			0 61	€SB7	2	3 07			
687 KTH						510				062	ST02	3	5 02			
\$\text{Possible}{Possi						l				063	DSZI	16 2	5 46			
869										064	GT05					
10						STO	n 00	tah aaumta								
## 11			5701			310	n, es	tab.counte				_				
												2		CL.	C+ 1-6	
013						1						-		Sni	tt Tei	t one gene
## \$14 \$704 \$35 \$4 \$6 \$6 \$7 \$8 \$\$ \$7 \$5 \$6 \$6 \$6 \$7 \$8 \$\$ \$7 \$6 \$7 \$8 \$\$ \$7 \$\$ \$8 \$\$ \$9 \$\$ \$16 \$7 \$\$ \$8 \$\$ \$7 \$\$ \$7 \$\$ \$8 \$\$ \$9 \$\$ \$8 \$\$ \$7 \$\$ \$7 \$\$ \$9 \$\$ \$9 \$\$ \$7 \$\$ \$9 \$\$\$ \$9 \$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$ \$9 \$\$\$\$ \$9 \$\$\$\$ \$9 \$\$\$\$\$\$			_								_					i
			_			1										
						1						1				
## Size Size						_						1				
Second S												_				
Section Sect												- 4		Shi	ft lef	t total
615 RCL2 36 62 6256 6356 23 36 62 621 8417 16-32 622 871 16 21 61 61 61 622 624 671 16 22 61 625 625 72 36 62 625 625 72 36 62 625 625 72 36 62 625 625 72 36 62 625 625 72 36 62 625 625 72 36 62 625 625 72 36 62 625 625 72 72 625 625 72 72 625 625 72 72 625 625 72 72 625 72 72 625 72 72 625 72 72 72 72 72 72 72						Get	P1 ger	ne 1						gei	notype	1 gene
Register Register															• •	
0022 SF1 15 21 91 003						Get	F1 ger	ne 1								
023	é	021	X=Y?		16-33	Comp	oare					1				
024 CF1 16 22 61 025 RCL1 36 61 025 RCL1 36 61 025 025 RCL1 36 61 025 02	é	022	SFi	15	21 01	-										
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97 Program Listing II

36									
STEP	KE	Y ENTRY	KEY CODE	COMMENTS	STEP	KEY ENTRY	KEY CODE	COMM	MENTS
	113	10×	16 33					T	
	114	EEX	-23		170			1	
	115	5	95					1	
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	116	Х.	-35					1	
	117	ST04	<i>35 04</i>			_		4	
	118	÷	-24			ļ		4	
	119	INT	16 34					1	
	120	RCL4	36 04					_	
	121	Х	-35					1	
	122	ST05	35 Ø5	Store non-comp.reg				1	
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	124	RCL1	36 01	gene is to be	180			1	
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	129	X≠Y?	16-32						
	130	GT08	22 08						
	131	RCL3	36 03	Place uncomplement	edi			1	
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	135	EEX	-23					4	
	136	5	<i>8</i> 5					4	
	137	Х	-35]	
	138	ST+5	35-55 65						
	139	≭LBL 8	21 08	Place complemented				1	
	140	1	01	gene value in				1	
	141	ST+3	35-55 03	storage				1	
				Scorage				1	
	142	RCL2	36 02					1	
	143	RCL3	36 03		. 200			4	
	144	X=Y?	16-33	Test completion Xo	` B 200			4	
	145	GT09	22 09					1	
	146	RCL1	36 61	Get new gene				_	
	147	1	<i>01</i>					_	
	148	Ū	99		1			1	
	149	÷	-2 4					1	
		INT						†	
	150		16 34					1	
	151	ST01	35 01					1	
	152	GT02	22 02					1	
	153	*LBLE	21 15	Store number of	210			4	
	154	DSP0	-63 00	genes in P1 and	210			4	
	155	ST06	35 <i>06</i>	set display				1	
	156	RTN	24						
	157	*LBL9	21 89	Show Result					
	158	RCL5	36 Ø5	Stop]	
	159	RTN	36 63 24	1				1	
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						ľ	3 □ 🛛		n_2

Program Description I

Program Title

CHROMOSOME CLEAVAGE

Contributor's Name

Hewlett-Packard Company

Address

1000 N.E. Circle Boulevard

City

Corvallis

State Oregon

Zip Code 97330

Program Description, Equations, Variables Chromosome cleavage is a powerful experimental genetic technique which has yielded a wealth of knowledge concerning gene loci on the chromosomal DNA code. The program simulates chromosome models consisting of even hundreds or thousands of unique individually identifiable genes in linear array. By noting what happens to any two genes after chromosomal fragmentation at arbitrary break-points, entire chromosomal gene sequences may be reconstructed experimentally.

The genetic sequences are based upon the ability of a prime P, of form (8k+3) or (8k+5) for some integer k, to reproduce all integers from 1 to (P-1) in any (P-1) consecutive passes of the formula ($S_{i+1} = 2S_i$ module P), starting from any $S_1 (1 \le S_1 \le (P-1)$. Permissable chromosome lengths for $n \le 100$ genes are 10, 12, 18, 28, 36, 42, 52, 58, 60, 66, 82, 100. For example, for n=100, P=101 = (8 · (12) + 5) is a prime of form (8k + 5).

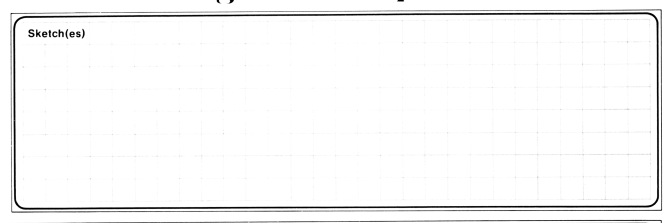
Copies of particular chromosomes are repeatedly split (each copy once) to build a statistical experimental pattern for any two genes on a chromosome. The program assumes that the 1st fragment (F_1) is identifiable – e.g. by end point radioisotope labeling – and computes how many (0, 1, 2) but not necessarily which of the two genes g_1 and g_2 are contained in F_1 for each cleavage. Powerful statistical inferences (see examples) are possible, and ultimately the whole chromosomal genes sequence may be reconstructed.

Operating Limits and Warnings

This program has been verified only with respect to the numerical example given in *Program Description II*. User accepts and uses this program material AT HIS OWN RISK, in reliance solely upon his own inspection of the program material and without reliance upon any representation or description concerning the program material.

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Program Description 11



Sample Problem(s) Do a gene-pair analysis for a chromosome of length 10 in a chromosomal DNA chain in an unknown sequence. Use .456 as the initial seed and do 10 consecutive chromosomal cleavages for each of the pairs (1,6), (1,8), and (6,9). Obtain the true gene separation for each pair and finally the true gene sequence.

```
Solution(s) .456 [f][A] 10 [A] 1 [ENTER↑] 6 [B]

Press [C] 10 times and get the sequence 2, 0, 2, 2, 0, 1, 2, 2,

[D] → 0

1 [ENTER↑] 8 [B]; Press [C] 10 times → 1, 2, 1, 2, 2, 2, 2, 2, 1, 2

[D] → -2[Disregard the sign]

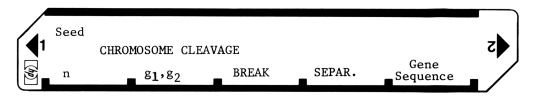
6 [ENTER↑] 9 [B]; Press [C] 10 times → 1, 0, 1, 1, 1, 1, 1, 0, 0, 0

[D] → -6 [Disregard the sign]

[E] → 7, 3, 6, 1, 2, 4, 8, 5, 10, 9 True Gene Sequence
```

Reference(s) This program is a modification of the Users' Library Program #04138A submitted by Mordecai Schwartz, M.D.

User Instructions



STEP	INSTRUCTIONS	INPUT DATA/UNITS	KEYS	OUTPUT DATA/UNITS
1	Load side 1 and side 2			
2	(Optional: enter random seed 0 < S < 1)	S	fA	
3	Select chromosome of desired length = no. of			
	genes, n, in linear sequence.	n	A	
	(Note: (n+1) must be a prime of form (8k+3)			
	or (8k+5) for some integer k. The valid			
	chromosome lengths for n<100 are n = 10, 12			
	18, 28, 36, 42, 52, 58, 60, 66, 82, and 100)			
4	Select 2 genes, g_1 and g_2 , for study $(1 \le g_i \le n)$	81	ENT [†]	
	(Note: If $g_1=7$, for example, the '7' is	g ₂	В	
	simply an identifying label (think of it as		1 []	
	the 7th gene discovered in that chromosome)		7	
	and does not necessarily indicate its			
	position in the chromosome's gene sequence.			
5	Cleave the chromosome at an arbitrary break-		7	0, 1, 2 =
	point.		C	# of genes
6	Repeat 5 as often as desired, with additional			on 1st frag
	chromosome copies, to obtain an experimental			
	statistical pattern for g_1 , g_2 separation.			
7	When ready, see actual no. of genes intervening	y		81 82
	between g ₁ and g ₂ for comparison with			separation
	deduced value.		D [(ignore sign
8	For another gene pair on same chromosome,			
	go to step 4.			
9	When experimentation is complete - ideally			
	with step 7 omitted - For all desired gene			
	pairs, compare complete chromosome with			
	your reconstruction.		[E] [
10	For a chromosome of different length go			
	to step 3.			
			SET STAT	US
			FLAGS TRIG	DISP
			0 □ 🛛 DEG	
			1 X GRAD 2 RAD	SCI 🗆 D ENG 🗆
			3 🗆 🛛	n 2

97 Program Listing I

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			I				l										i	

Program Description I

RECESSIVE GENE FREQUENCY AFTER SELECTION, MUTATION, AND INBREEDING **Program Title**

Contributor's Name Hewlett-Packard Company

1000 N.E. Circle Boulevard Address

City Corvallis State Oregon **Zip Code** 97330

Program Description, Equations, Variables The following selection and mutation model was used:

production of gametes random production of zygotes

mutation may occur

selection surviving individuals

reach maturity

From this model (see Operating Limits & Warnings) can be derived:

$$q_{n+1} = \frac{q_n^2 (ht+uht-t) + q_n (1-ht-u-uht) + u}{1 - 2ht q_n + q_n^2 (2ht-1)}$$

where:

q = recessive or mutant gene frequency u = mutation rate

u = mutation rate q = lecessive of definitions of the first of thegene frequency

 q_e is attained when $q_{n+1} = q_n$ (see pg 4, step 6 comment).

(Continued on next page)

Operating Limits and Warnings The following simplifying assumptions were made:

a) infinite population size