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A kinetic study of the murine mixed lymphocyte reaction by 5,6-carboxyfluorescein diacetate succinimidyl ester labeling

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11 Abstract

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12Alternatives to the use of radioisotopes to measure cell proliferation in mixed lymphocyte reactions (MLR) are desirable to 13 avoid the hazards and costs associated with radioisotope use. The versatile fluorochrome 5,6-carboxyfluorescein diacetate 14 succinimidyl ester (CFSE) has been used to measure MLR and provides the opportunity to measure several different growth 15parameters. This study was aimed at determining which growth parameter is most practical and suitable for measuring murine 16MLR. The parameters measured were: the relative number of daughter T-cells, the relative number and frequency of reactive Tcell precursors and the relative number of mitotic events. Responder cells were CFSE-labeled unfractionated splenocytes from 17C57BL/6 mice. Stimulator cells included irradiated splenocytes from C57BL/6 (control), B6D2F1 (haplo-allogeneic) or FVB/N 18(allogeneic) mice. Cultures were harvested daily for 1 week. Stimulator T-cells rapidly declined to less than 0.2–0.3% of the 1920mixed population by day 2 of culture. Experimental groups had a significantly higher number of daughter T-cells and mitotic 21events after 2 days of culture with the number of daughter T-cells climbing exponentially after 5 days of culture. The number 22and frequency of reactive T-cell precursors were significantly higher in experimental groups on days 2-3, but this difference 23became insignificant by day 4. Among all the parameters, the relative number of daughter T-cells was the most practical for 24measuring MLR, after 5 days of culture, based upon the growth kinetics of responder T-cells and the survival of the stimulator 25cells.

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28Keywords: 5,6-carboxyfluorescein diacetate succinimidyl ester; Mixed lymphocyte reaction; Mouse

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Abbreviations: MLR, mixed lymphocyte reactions; CFSE, 5,6-Carboxyfluorescein diacetate succinimidyl ester; 3H-TdR, 3Hthymidine; PBS, phosphate buffer saline; BSA, bovine serum albumin; FBS, fetal bovine serum; PI, propidium iodide.

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1. Introduction

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The mixed lymphocyte reaction (MLR) measures 32the proliferative response of responder T cells against 33 antigens present on allogeneic stimulator cells. It is 34presumed that the MLR is an in vitro analog of in vivo 35alloreactivity, and is widely used in transplantation 36

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immunology to measure recipient T cell responses 37 against donor tissues due to the mismatch of MHC 38 antigens. Cell proliferation is an essential part of an 39immune response which can be measured in an 40 41 allogeneic mixed lymphocyte culture as an indicator 42of immune reaction. There are numerous methods to measure changes in cell number. The most widely 43used method to measure proliferation is ³H-thymidine 44 (³H-TdR) incorporation. However, due to the hazards 45and cost associated with the use and disposal of 46tritium, nonradioactive alternatives have been pur-47 sued. 48

Nonradioactive methods include the use of a range 49of fluorochromes, which have been used for measur-50ing the proliferation of lymphocytes as well as for 51tracking migration and positioning in vivo (Parish, 521999). Among them, 5,6-carboxyfluorescein diacetate 53succinimidyl ester (CFSE) stands out as being the 54most versatile in terms of long-term cell tracking and 55quantifying proliferation either in vivo or in vitro 5657(Lyons, 1999; Parish, 1999). CFSE can spontaneously and irreversibly couple to intracellular proteins and is 58equally distributed between two daughter cells when 59cells divide. Thus, proliferating cells can be tracked 6061 by flow cytometry based upon the sequential loss of fluorescence intensity (Lyons, 2000; Lyons and Par-62 ish, 1994). Furthermore, multiparameter flow cytom-63 etry allows for the examination of cell subsets within 64 the dividing cell population as well as monitoring 65 phenotypic changes associated with activation and 66 cell division. 67

68 CFSE staining has been used as a replacement for ³H-TdR incorporation to measure the MLR of cells of 69 human and nonhuman-primate origins (Matthews et 70al., 2000; Popma et al., 2000; Nitta et al., 2001). In 7172most cases, the outcome of the MLR has been 73determined based on the total numbers of daughter T-cells generated. However, the ability of CFSE to 74track cell division allows for other parameters of cell 75proliferation to be measured. These include estimates 7677 of the number and frequency of T-cell precursors that have responded in the MLR as well as the number of 7879total mitotic events that have occurred. The utility of 80 measuring these parameters in analyzing murine MLR has not been closely examined. Therefore, we studied 81 the kinetics of murine T-cell responses in allogeneic 82 83 MLR using CFSE labeling and multiparameter flow 84 cytometry.

2. Materials and methods 85

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2.1. Responder and stimulator 87

Two inbred strains of mice, C57BL/6 (H-2^b) and 88 FVB/N (H- 2^{q}), and one hybrid strain of mice B6D2F₁ 89 $(C57BL/6 \times DBA/2, H-2^{b/d})$ were purchased from 90 Charles River (Wilmington, MA) or Simonsen (Gilroy, 91CA). They were 8-30 weeks of age when sacrificed 92 and their spleens were harvested. Responder T-cells 93were from the C57BL/6 strain, whereas stimulators 94 included C57BL/6 (syngeneic), B6D2F1 (haplo-allo-95geneic) and FVB/N (allogeneic) strains. Splenocytes 96 were harvested under sterile conditions and passaged 97through 70 µm cell strainers (Becton Dickinson, Frank-98 lin Lakes, NJ). Red cells were depleted by chemical 99lysis using ACK buffer, pH 7.2-7.4, consisting of 0.15 100 M NH₄CL, 1.0 mM KHCO₃ and 0.1 mM Na₂EDTA 101(Sigma, St. Louis, MO). The cell suspension was then 102washed twice with phosphate buffer saline (PBS) 103containing 0.3% bovine serum albumin (BSA) (PBS/ 104BSA) (Roche Molecular Biochemicals, Indianapolis, 105IN). Responder cells were suspended at 2×10^7 cells/ 106ml in PBS/BSA for CFSE labeling, whereas stimulator 107cells received 3000 cGy irradiation and were then 108resuspended at 6×10^6 cells/ml in culture medium. 109

2.2. CFSE labeling of responder cells

CFSE labeling of responder cells was undertaken as 112previously described (Lyons, 2000; Lyons and Parish, 113 1994). Immediately before labeling, 5 mM CFSE stock 114(Molecular Probes, Eugene, OR) in DMSO (Fisher 115Scientific, Fair Lawn, NJ) was thawed and diluted to 116 $10 \,\mu\text{M}$ in a volume of PBS/BSA equal to that in which 117the responder cells were suspended. The two equal vo-118 lumes were mixed to initiate labeling and periodically 119agitated at room temperature for 10 min. The labeling 120process was quenched by adding an equal volume of 121heat-inactivated fetal bovine serum (FBS, Hyclone, 122Logan, UT) to the sample. After 1 min, the CFSE-la-123beled cells were washed twice, recounted and adjusted 124to a concentration of 2×10^6 cells/ml in culture media. 125126

The MLR was performed in 96-well U-bottom 128 microtiter plates (Costar, Cambridge, MA). Culture 129

medium consisted of RPMI 1640 supplemented with 13010% heat-inactivated FBS, sodium pyruvate (110 mg/ 1311), nonessential amino acids, L-glutamate (1 mM), 2-132mercaptoethanol (5 \times 10⁻⁵ M), penicillin (50 U/ml) 133and streptomycin (50 µg/ml) and N-acetyl cysteine 134135(Sigma; 10 mmol/l, pH adjusted to 7.2). CFSE-labeled C57BL/6 responder cells were plated at 1×10^6 cells/ 136ml in a volume of 250 µl per well and cocultured at a 137ratio of 1:3 with 3000 cGy irradiated C57BL/6, 138139B6D2F₁ or FVB/N stimulator cells. The plates were then placed in a humidified 37 °C, 5% CO₂ incubator. 140 The control experiment for accessing background fluo-141 rescence was set up by coculture of unlabeled C57BL/ 1426 responder cells and irradiated stimulator cells under 143the same conditions. In order to assess the kinetic 144 division of the responder cells and the viability of sti-145mulator cells, cells were harvested from quadruplet 146wells on a daily basis. 147

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149 2.4. Flow cytometric analysis

Cells harvested from each well were suspended in 150culture supernatant from the clone 2.4G2 hybridoma 151cell line (ATCC, Manassas, VA) that produces mAb 152153against FcyII/FcyIII receptors to block nonantigen-154specific binding of immunoglobulins. CFSE-labeled cells were incubated on ice for 30 min with PE 155conjugated anti-CD3 (Caltag Laboratories, Burlin-156game, CA). Cells were then washed three times in 157158PBS/BSA with 0.01% NaN₃ (Sigma). Cells from each well were suspended in 150 µl PBS/BSA with 0.01% 159NaN₃ and 2 µg/ml propidium iodide (PI, Molecular 160 161Probes). Three-color flow cytometry was performed on a FACScan (BD Biosciences, San Jose, CA). 162Events for each well were acquired at a fixed speed 163164(high setting) for 1 min to measure an equal volume 165from each sample, thereby providing a basis for the 166relative comparison of data collected for each sample. Dead cells were excluded from analysis based on 167staining with PI. 168

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170 2.5. Data and statistical analyses

Data were analyzed using CellQuest acquisition
and analysis software (BD Biosciences). The total
numbers of events (cells) were determined by analyzing the data using dot plots and rectangular regions to
define the cell populations. Histograms were used to

track the divisions of CFSE-labeled cells where each 176 peak of CFSE-fluorescence defined a single round of 177 cell division. All data presented are from analyses of 178 live (PI⁻) cells only. 179

The relative numbers of T-cell precursors required 180for generating these daughter cells under each division 181peak was calculated by dividing the number of daugh-182 ter-cell events by 2 raised to the power of the given 183round of division (2^n) . The sum of all the calculated 184 numbers of precursors from each division peak was 185used to represent the number of reactive T-cell pre-186cursors. Furthermore, the frequency of reactive T-cell 187 precursors was estimated by dividing the number of 188reactive T-cell precursors by the number of total T-189cells, which equals the number of undivided T-cells 190 plus the number of reactive T-cell precursors. Alterna-191tively, the number of total live T-cells measured on the 192first day of culture was also used to estimate the 193frequency of reactive T-cell precursors. The frequency 194values were multiplied by 100 to be represented as a 195percentage of all T cells. In addition, the mitotic events 196 under each peak can also be determined by subtracting 197the number of reactive T-cell precursors from the 198number of daughter T-cells under each peak (Wells 199et al., 1997). Thus, the relative number of total mitotic 200events can be calculated as the sum of individual 201mitotic events under each peak, which equals the 202number of daughter T-cells minus the number of 203reactive T-cell precursors. 204

Independent sample *t*-test was used for comparing 205means of events and ratios between control and ex-206perimental groups. Paired-samples t-test was used to 207make a comparison between the two precursor fre-208quencies generated on the basis of different denomi-209 nators, one derived from the total T-cell precursors on 210 the indicated day and the other from the mean undi-211vided T-cells on day 1. A P-value < 0.05 was consid-212ered statistically significant. 213

3. Results

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3.1. Proliferation of CD3⁺ responder cells 216

Mixed lymphocyte cultures were harvested after 1–217 7 days and stained with CD3-PE to identify the T-cell 218 component of the cultures. CD3 staining and CFSE 219 fluorescence were detected by flow cytometry (Fig. 1).220





Fig. 1. Flow cytometric analysis of mixed lymphocyte cultures. Progressive loss of CFSE fluorescence by C57BL/6 responder CD3⁺ T-cells, indicated by the regions R3 and R4, is shown for days 1, 3 and 6 of culture. Viable FVB/N stimulator cells are indicated by regions R5 and R6. The stimulator cells were mostly located at the area of low fluorescent intensity ($<10^{1}$).

Daughter T-cells, derived from the responder spleno-221cytes, could be differentiated from undivided T-cells 222223by the intensity of CFSE staining. The relative numb-224ers of daughter T-cells could be easily determined by electronic gating, as shown in Fig. 1. In the case of 225226alloreactive MLR, the numbers of daughter T-cells showed a slow increase until day 4 and then an expo-227nential expansion after day 5 (Fig. 2A). The syngeneic 228control group (C57BL/6) generated significantly lower 229numbers of daughter T-cells than the haploidentical 230



Fig. 2. Kinetics of expansion and survival of responder T-cells in mixed lymphocyte cultures. Accumulations of daughter T-cells from the responder population are shown in (A) and survival of undivided T-cells are shown in (B). Data are shown as the mean \pm 2.0 standard errors of the mean (SE) of measurements made on replicate cultures.

 $B6D2F_1$ and completely MHC-mismatched FVB/N231groups beginning on the second day of culture on-
wards. There were no significant differences in the
numbers of daughter T-cells between the $B6D2F_1$ and
FVB/N groups at any time point. Thus, a T-cell pro-
liferative response to MHC antigens could be readily
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detected by this method.231

Survival of the undivided CD3⁺ cells in the mixed 238cultures was also analyzed (Fig. 2B). A linear decrease 239 in the number of unresponsive T-cells was observed in 240the C57BL/6 group over the 7 days of culture ana-241lyzed. However, the B6D2F1 and FVB/N groups had 242 comparatively steady numbers of undivided CD3⁺ 243cells until day 3. A slight drop followed in these 244groups on day 4, and then a linear decrease in the 245number of unresponsive T-cells was seen as in the 246C57BL/6 control group. 247

The kinetics of CD3⁺ cell proliferation was ana-248lyzed in greater detail using histograms to view the 249levels of CFSE fluorescence (Fig. 3). On day 1 (24 h of 250culture), CD3⁺ responder cells were uniformly stained 251with CFSE, indicating that these cells had not yet 252begun to divide. By the second day of culture, CD3⁺ 253responders, stimulated by B6D2F1 or FVB/N spleno-254cytes, had begun to divide for up to three rounds of 255mitosis (Fig. 3 and data not shown). In contrast, 256responder T-cells stimulated with syngeneic C57BL/ 2576 cells did not show any evidence of cell division for 258the first 2 days (Fig. 3). On days 3 and 4, four to six 259rounds of divisions could be detected in all the three 260 groups. On day 5, CD3⁺ cells of all three groups had 261divided at least for seven to eight times, resulting in a 262loss of fluorescent intensity on some responder cells to 263background levels. Thus, at this time point further 264resolution of subsequent cell division could not be 265accurately measured. On days 6-7, almost all the 266daughter T-cells, stimulated by B6D2F1 and FVB/N 267splenocytes, had reached the area of background 268fluorescence due to many rounds of divisions. 269

Measured estimates of the relative number and 270frequency of reactive T-cell precursors are listed for 271a representative experiment in Table 1. These calcu-272lated measures are based on measurements of viable 273(PI⁻) CD3⁺ cells. A significantly higher number and 274frequency of reactive T-cell precursors were observed 275in the B6D2F1 and FVB/N groups compared to the 276C57BL/6 group on days 2-3. However, the frequency 277of reactive T-cell precursors did not differ between 278

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t1.1 Table 1

Relative numbers and frequencies of reactive T-cell precursors as well as the number of mitotic events from MLR analyzed at 1-4 days of t1.2 culture

Day	Stimulator	Relative Number of T-cell Precursors	Estimated T-cell Precursor Frequency (%) ^a	Relative Number of Mitotic Events
B6D2F1	2.38 ± 0.32	$0.03 \pm 0.01/0.03 \pm 0.00$	2.38 ± 0.31	
FVB/N	3.38 ± 0.55	$0.05 \pm 0.01/0.04 \pm 0.01$	3.38 ± 0.55	
2	C57BL/6	5.00 ± 0.79	$0.07 \pm 0.01/0.06 \pm 0.01^{\rm b}$	5.00 ± 0.79
	B6D2F1	$54.50 \pm 1.85^{\circ}$	$0.68 \pm 0.02^{\circ}/0.65 \pm 0.02^{\circ}$	$80.75 \pm 2.39^{\circ}$
	FVB/N	$47.25 \pm 2.46^{\circ}$	$0.64 \pm 0.02^{\circ}/0.60 \pm 0.03^{\circ}$	$67.25 \pm 2.25^{\circ}$
3	C57BL/6	40.50 ± 2.06	$0.63 \pm 0.04 / 0.47 \pm 0.02^{\rm b}$	140.00 ± 10.79
	B6D2F1	$152.75 \pm 11.18^{\circ}$	$1.87 \pm 0.10^{\circ}/1.82 \pm 0.13^{\circ}$	$759.75 \pm 75.93^{\circ}$
	FVB/N	$129.00 \pm 3.49^{\circ}$	$1.69 \pm 0.04^{\circ}/1.65 \pm 0.04^{\circ}$	$573.50 \pm 32.17^{\circ}$
4	C57BL/6	141.00 ± 15.15	$2.37 \pm 0.18 / 1.63 \pm 0.18^{\rm b}$	591.00 ± 76.73
	B6D2F1	157.75 ± 3.59	$2.08 \pm 0.04/1.88 \pm 0.04^{\rm b}$	$1411.75 \pm 59.31^{\circ}$
	FVB/N	160.25 ± 6.20	$2.24 \pm 0.10/2.05 \pm 0.08^{b}$	$1720.75 \pm 101.25^{\circ}$

t1.16 Data are shown as the mean \pm standard error of the mean for quadruplicate analyses performed on live cells.

^a T-cell precursor frequencies were calculated by two methods. The first number represents the frequency calculated using the sum of reactive T-cell precursors and non-responsive T-cells measured on the day when the culture was harvested as the denominator. The second value t1.17 was calculated using the mean number of undivided T-cells on day 1, when responder T-cells had not yet divided, as the denominator.

t1.18 ^b P < 0.05 comparison between first and second numbers, paired-samples t-test.

t1.19 $^{c}P < 0.05$ compared with the C57BL/6 control group, independent-samples t-test.

control and experimental groups by the fourth day of 279culture. The frequencies of T-cell precursors were 280estimated by two methods, as described in Materials 281and methods. The linear decrease in undivided CD3^{*} 282283cells in the control group resulted in an overestimation of T-cell precursor frequency on days 2-4; whereas 284the relatively slow decrease of undivided CD3⁺ cells in 285286 the experimental groups did not notably influence the frequency calculations until day 4. 287

The relative numbers of mitotic events that occurred 288in the cultures could also be calculated (Table 1). The 289B6D2F₁ and FVB/N groups exhibited higher division 290of T-cells on days 2-4 compared with the control 291292group. This pattern was consistent with the measures of the relative number of daughter T-cells (Fig. 2A), 293which measures the outcome of these early mitotic 294events as a latter accumulation of daughter T-cells. By 295the 5th day of culture, it was difficult to accurately 296297estimate the number and frequency of reactive T-cell precursors as well as the number of mitotic events. This298is because T cells under peak 6 might contain daughter299cells that had undergone more than six rounds of300division and can no longer be discriminated from301background fluorescence as indicated using an un-302stained control (Fig. 3).303

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3.2. Proliferation of responder CD3⁻ cells

Responder CD3⁻ cells also started to proliferate in 306 the mixed culture beginning on day 2, reached a peak 307 around days 3-4, and subsequently declined to a 308 plateau (Fig. 4A). This proliferation pattern did not 309 parallel that of responder $CD3^+$ cells. In B6D2F₁ and 310 FVB/N groups, the ratio of CD3⁺ to CD3⁻ responder 311cells was usually less than 1 over the first 3 days, 312gradually increased to five on day 5, and abruptly 313amplified to over 10 on day 7. In contrast, the ratios 314from C57BL/6 control group sustained below 1 until 315

Fig. 3. Histogram plots gated on $CD3^+$ cells show the kinetics of cell division associated with an MLR. Histograms of CFSE fluorescence for Tcells stimulated by haploallogeneic B6D2F₁ splenocytes are shown in the left column and histograms for the syngeneic control group are shown for comparison in the right column. A peak of low fluorescence (arrow) represents viable stimulator T-cells observed during the first 2 days of culture. On days 5 and 7, cells with ≥ 6 rounds of division had reached a fluorescent intensity similar to the background autofluorescence of unstained responder T-cells (thin line). This overlap limited the resolution of further cycles of cell division and marked the maximal cycle that could be distinguished.

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Fig. 4. Proliferation and contribution of CD3⁻ responder cells to mixed lymphocyte cultures. Accumulations of daughter CD3⁻ cells (non-T-cells) from the responder population are shown in (A). The ratio of daughter CD3⁺ to CD3⁻ cells is shown in (B). Data are shown as the mean \pm 2.0 SE.

- day 4, slightly increased to the peak of 3 on day 6, andthen went down on day 7 (Fig. 4B).
- 318
- 319 3.3. Viability of stimulator cells

Viable stimulator cells in the mixed cultures were
observed in the early days of culture and could be identified by their lack of CFSE staining. They remained
after 1 day of culture, comprising both CD3⁻ cells and
CD3⁺ cells (Fig. 1). CD3⁻ stimulator cells exhibited a

much slower decline and represented 2-3% of the total 325cell population on day 3 of culture (Fig. 5A). However, 326 a rapid decline of viable $CD3^+$ cells to 0.2-0.3% of all 327 viable cells was observed by the second day of culture, 328 and less than 0.1% stimulator CD3⁺ cells were present 329on days 3-4 (Fig. 5B). An MLR, in which the 330 responders were not stained with CFSE, was also 331 undertaken to trace the viability of stimulator cells by 332 flow cytometry using differences in MHC antigens (H-333 2K^{b, d, and q}). Total viable stimulator cells in the mixed 334culture accounted for 11.5-23.6% on day 1, 2.9-7.6% 335 on day 2, 0.5-0.7% on day 3, 0.5-1% on day 4, 0.2-336 0.5% on day 5 and 0.1–0.3% on days 6-7 (data not 337 shown). 338



Fig. 5. Survival of stimulator cells in mixed lymphocyte cultures. The percentages of viable $CD3^-$ stimulator cells are shown in (A) and the percentages of viable stimulator $CD3^+T$ – cells are shown in (B). Data are shown as the mean ± 2.0 SE.

339 4. Discussion

340 Cell proliferation encompasses DNA synthesis, mitosis and an increase in cell number. ³H-TdR incor-341poration measures cell proliferation by the incorpora-342 343 tion of ³H-TdR into DNA while cells are dividing, and has been a standard method for measuring cell prolif-344 eration in mixed lymphocyte cultures for decades. In 345view of the increasing costs of waste disposal and 346 protective precautions associated with radioisotope 347 348 use, developing an alternative for quantify cell prolif-349eration, which can be as effective as, but less hazardous than ³H-TdR, is welcomed. CFSE labeling stands 350out in many aspects as a versatile alternative to 351conventional ³H-TdR incorporation. Most importantly, 352this method was shown to correlate well with ³H-TdR 353354incorporation (Fulcher and Wong, 1999; Popma et al., 2000). To extend the adoption of CFSE for use in 355measuring murine MLR, we closely examined a num-356 ber of parameters associated with cell division to 357 358determine the optimal timing and methods to measure murine MLR. 359

³H-TdR incorporation only measures cell division 360 during a brief period of culture and provides no 361 362 information on the types of cells growing or their degree of response. In contrast, CFSE labeling can 363 provide more information such as the relative number 364 of reactive T-cell precursors, the estimated frequency 365 of reactive T-cell precursors, the relative number of 366 mitotic events and relative number of daughter T-cells 367 associated with an MLR. The proliferation of CFSE-368 369 labeled responder cells could be traced for at least 1 week, providing a detailed picture of the different 370 stages of an MLR. Moreover, immunophenotypic 371 analyses of the dividing cells insured that the mea-372373 surement was specific to CD3⁺ T-cells. This might be 374crucial in some cases because unfractionated tissues such as spleen, lymph node or fetal liver (Harris et al., 375376 1994) have been used as the source of responder T cells and substantial non-T-cell proliferation might 377 occur from such tissues. Under theses circumstances, 378 data obtained by ³H-TdR incorporation might vari-379 380 ously reflect proliferation of both T and non-T cells. According to our results, non-T-cells displayed a 381different pattern of proliferation from T-cells. Clearly, 382non-T-cells are most apt to contribute to overall 383 proliferation in the early days of a splenic MLR. 384 385Furthermore, cell proliferation in the control group

tended to be more the result of non-T-cells than in the 386 experimental groups, which could be readily distinguished and disregarded by the flow cytometric 388 method. 389

The simplest method of quantifying an MLR using 390 CFSE was found to be measurement of the accumula-391 tion of daughter CD3⁺ cells. The magnitude of the T-392 cell proliferative response elicited by various stimula-393 tor conditions mainly depended upon the rounds of 394division the T-cell precursors had undergone rather 395 than upon the frequency of reactive T-cell precursors. 396 This argument was supported by the presence of two 397 distinct peaks on days 6 and 7 in the histogram plots 398 shown in Fig. 3. One peak represented quiescent T-399 cells and the other was daughter T-cells that had 400 undergone at least seven to eight rounds of division. 401There was a relative paucity of daughter cells in 402between. T-cell proliferation was asynchronous, to a 403 degree, likely due to variation in the time of entry into 404 the first division cycle (Hasbold et al., 1999). None-405theless, new T-cell precursors did not appear to be 406 recruited after day 5 of culture. Accordingly, the 407 alloresponse to MHC-mismatched stimulators led to 408 an exponential expansion of daughter T-cells by elicit-409 ing more rounds of division than the nonspecific 410 response in syngeneic group. Modest T-cell prolifera-411 tion in the control group appeared later than in the 412experimental groups, becoming evident on day 3. 413

The number and frequency of reactive T cell pre-414cursors in an MLR could also be estimated using flow 415cytometry. Our estimated frequencies of reactive T-cell 416 precursors was about 2%, which was within some 417 published estimates of the percentage of allorespon-418 sive precursors (Ford and Atkins, 1973; Ford et al., 4191975; Matzinger and Bevan, 1977; Sherman and 420Chattopadhyay, 1993). These measurements were best 421 performed on the third day of culture rather than at 422 later time points that are better suited for measuring the 423accumulation of daughter T cells. This is because 424 differences between the control and experimental 425 groups were evident on days 2 and 3, but by day 4, 426 these differences disappeared. The weak background 427 proliferation observed in the controls by day 4 of 428culture might be due to nonspecific activation of the 429 T cells by cytokines produced in the cultures or FBS 430used to supplement cultures. Another reason for early 431analysis of the cultures is that the death of some of the 432 daughter cells and nonresponsive T-cells affects the 433

estimates of the size and frequency of the precursor 434pool. Dead (PI⁺) daughter and nonresponder cells were 435observed, but it was necessary to disregard these cells 436for analyses as the PI staining and high nonspecific 437 binding of antibody by dead cells interferes with the 438analysis of CD3 expression. Since the estimation of the 439frequency of T-cell precursors relies on an accurate 440 measurement of the total number of live T cells used to 441 442initiate the cultures, we tested two methods by which 443to calculate precursor frequencies. One method relied on a single flow cytometric analysis, which was used 444 to estimate the total pool of T cells based on the 445calculated number of reactive T-cell precursors plus 446 the number of undivided T-cells. A second method 447 448 used an additional analysis after 1 day of culture, when the T-cell precursor pool had not yet begun to divide, 449to measure the total number of live T-cells. We chose 450to make this analysis after 1 day rather than on day 0 of 451culture as the CFSE staining procedure may result in 452the death of some T cells that is not immediately 453evident. Both methods of calculating the size of the 454total T-cell pool resulted in similar estimates of reac-455tive T-cell precursor frequencies for the experimental 456groups, but the control group had a faster loss of 457nonresponsive T cells leading to higher estimates of 458459precursor frequencies beginning on the second day of culture. Thus, choosing which method to use to 460calculate the size of the total T-cell pool depends on 461 the types of MLR being compared. 462

463 Being able to estimate the number or frequency of reactive T-cell precursors may have specific applica-464 tion in studies aimed at measuring these values. 465466 However, this study clearly indicates that measuring the relative number of daughter T-cells is the best 467 parameter for analyzing the magnitude of an MLR 468 (Matthews et al., 2000; Popma et al., 2000). It is simple 469470as well as straightforward without the requirement of extensive calculations. The number of mitotic events 471can also be used for comparing in vitro MLR because 472its pattern generally paralleled that of the accumulation 473of daughter T-cells at least during the first 4 days of 474 475cultures. However, it offered no clear advantages over the measuring the number of daughter T-cells and 476 required labor-intensive calculations. Using the num-477 ber of daughter T-cells, the differences between the 478 control and experimental groups could be distin-479480guished beginning on the second day of culture on-481 wards, and were more pronounced after 5 days of culture. Furthermore, irradiated stimulators were near-
ly absent from culture after 3 days. Thus, quantifying482MLR by the number of daughter T-cells after 5 days of
culture could be easily performed with minimal inter-
ference from stimulator cells. These findings support486the use of a flow cytometry as a reliable method to
analyze murine MLR based on CFSE staining.487

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