The Importation of Hematogenous Precursors by the Thymus Is a Gated Phenomenon in Normal Adult Mice

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Abstract

Hematogenous precursors repopulate the thymus of normal adult mice, but it is not known whether this process is continuous or intermittent. Here, two approaches were used to demonstrate that the importation of prothymocytes in adult life is a gated phenomenon. In the first, age-dependent receptivity to thymic chimerism was studied in nonirradiated Ly 5 congenic mice by quantitative intrathymic and intravenous bone marrow (BM) adoptive transfer assays. In the second, the kinetics of importation of blood-borne prothymocytes was determined by timed separation of parabiotic mice. The results showed that >60% of 3–18-wk-old mice developed thymic chimerism after intrathymic injection of BM cells, and that the levels of chimerism (range, 5–90% donor-origin cells) varied cyclically (periodicity, 3 to 5 wk). In contrast, only 11–14% of intravenously injected recipients became chimeric, and chimerism occurred intermittently (receptive period ~1 wk; refractory period ~3 wk). In the intravenously injected mice, chimerism occurred simultaneously in both thymic lobes; gate opening occurred only after most intrathymic niches for prothymocytes had emptied; and the ensuing wave of thymocytopoiesis encompassed two periods of gating. These kinetics were confirmed in parabiotic mice, and in cohorts of mice in whom gating was synchronized by an initial intrathymic injection of BM cells. In addition, a protocol was developed by which sequential intravenous injections of BM cells over a 3 to 4 wk period routinely induces thymic chimerism in the apparent absence of stem cell chimerism. Hence, the results not only provide a new paradigm for the regulation of prothymocyte importation during adult life, but may also have applied implications for the selective induction of thymocytopoiesis in nonmyeloablated hosts.

Key words: thymus • bone marrow • lymphoid organization • lymphoid migration • ontogeny

Introduction

The importation of hematogenous thymocyte precursors (prothymocytes) in late fetal and early postnatal life in birds and mice (as well as in larval and postmetamorphic frogs) appears to be a gated phenomenon characterized by brief periods of receptivity interspersed by longer periods of refractivity (1–3). It has been postulated that prothymocyte gating early in ontogeny permits the sequential generation, selection, and exportation of developmentally and functionally discrete populations of thymocytes, and that differential prothymocyte generation and intrathymic processing serves to coordinate both the establishment of the immunological repertoire and the distribution of specialized populations of T cells to skin, mucosa, liver, and peripheral lymphoid tissues (for reviews, see references 4 and 5).

Unfortunately, the serial transplantation/explantation procedures used in the preceding studies cannot be conducted with adult thymic lobes, even using vascular transplantation procedures (6). Therefore, it has not been possible, until now, to experimentally assess the possibility of continued prothymocyte gating beyond the neonatal period. In addition, several cogent arguments have been advanced against this proposition. These include the: (a) possible role of intrathymic precursors in maintaining thymocytopoiesis in adult animals (for a review, see reference 7); (b) failure of large doses of bone marrow (BM) cells to establish significant thymic chimerism in nonirradiated adult recipients (8–

Abbreviations used in this paper: BM, bone marrow; FCM, flow immuno-cytometric.
stem cell engraftment.

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logical age, a protocol was devised by which repetitive in-

period (closed gate) lasts

trathymic niches for prothymocytes; and (c) the refractory

period (open gate) lasts

with the period of maximum availability of putative in-

trathymic niches. Furthermore, using

have modified our previously described intrathymic and in-

travenous BM adoptive transfer systems (17) to determine

the kinetics of prothymocyte recruitment in the thymus of

nonmyeloablated adult mice. We also conducted timed

separation experiments in parabiotic mice to detect the

progeny of recently imported donor-origin prothymocytes.

In both instances the results indicated that: (a) the importa-

tion of blood-borne prothymocytes is a gated phenomenon

during at least the first 4 mo of normal postnatal life; (b) the

receptive period (open gate) lasts ∼1 wk and coincides

with the period of maximum availability of putative in-

trathymic niches for prothymocytes; and (c) the refractory

period (closed gate) lasts ∼3 wk and includes the period of

progressive emptying of these niches. Furthermore, using

the periodicity of prothymocyte gating rather than chrono-

logical age, a protocol was devised by which repetitive in-

travenous injections of BM cells over a 3 to 4 wk period

uniformly induces a wave of thymocyte chimerism in nor-

mal adult mice, but appears not to establish hemopoietic stem cell engraftment.

Materials and Methods

Animals. Cohorts of 4 to 6-wk-old (± 3 d) male and female Ly 5 congenic C57BL/6NCR(B6) mice, obtained from the National Cancer Institute, were housed in the Center for Laboratory Animal Care (The University of Connecticut Health Center) until they reached the designated ages. Animals were maintained on commercial mouse chow and water ad libitum. In some experiments, breeding pairs were established to generate younger progeny or progeny of timed matings. Cell transfer was carried out only in sex-matched combinations and, as no gender differences were noted, the data were pooled.

Preparation of Cell Suspensions. BM cell suspensions were prepared by flushing the marrow from tibia and femur of 4 to 5-wk-old donors with cold RPMI 1640 (GIBCO BRL) supplemented with sodium bicarbonate (2 mg/ml) and 1% Hepes (1.5 M), as described (17). Repeated gentle pipetting further dispersed the cells, which were then washed in cold medium and centrifuged at 4°C for 5 min at 1,500 rpm. Thymocyte cell suspensions were prepared by gently pressing thymus lobes, stripped of attached lymph nodes, through a stainless steel screen (50 mesh), followed by washing in cold medium. Nucleated cells were counted on a Z1 Coulter Counter (Beckman Coulter).

Intrathymic Adoptive Transfer Assay for Prothymocytes. The thy-

mus was surgically exposed and one-half of the indicated number of BM cells were injected into the anterior superior portion of each lobe (10 μl/site) using a 1-ml syringe (with attached 28 gauge needle) mounted on a Tridek Stepper (Indicon Inc.), as described (17). The incision was closed with Nexaband Liquid (Veterinary Products Lab.). Control mice were injected intrathymically with RPMI alone, or one lobe was injected with BM cells and the contralateral lobe with RPMI, as indicated.

Intravenous Adoptive Transfer Assay for Prothymocytes. The in-

dicated number of BM cells suspended in 0.5 ml RPMI were in-

jected through a 28 gauge needle into the lateral tail veins of unanesthetized recipient mice. Control mice were injected intrave-

nously with RPMI alone.

Flow Immunocytometric Analysis. Thymocytes were harvested

28 d after BM cell transfer, except as indicated. The percentages

of donor and host-origin cells were determined by flow immuno-

cytometric (FCM) analysis (FACScan™; Becton Dickinson) after

Figure 1. Induction of thymic chimerism in nor-

mal adult mice by intravenous and intrathymic in-

jection of BM cells. Groups of normal adult Ly 5.1

mice, obtained from multiple cohorts, were ran-

domized for age (7–12 wk) and sex, and injected

intravenously (black bars) or intrathymically (hatched bars) with suspensions of sex-matched Ly 5.2 BM cells (20 × 10^6 intravenously or 2 × 10^6 intrathymically) from 6-wk-old donors. The fre-

quency and levels of thymic chimerism attained 28 d later were determined by FCM analysis. Data for male and female recipients are pooled, as no dif-

ferences were observed. Percentage of chimeric mice (≥5% donor-origin cells): intravenous injection = 12.7% (21 of 165); intrathymic injection = 85.9% (122 of 142).
development for immunofluorescence with anti-Ly 5.1 and anti-
Ly 5.2 monoclonal antibodies (The Jackson Laboratory), and the
expression of CD3, CD4, and/or CD8 was determined by multi-
color analysis. Dead cells and nonlymphoid cells were excluded by
gating for forward and side angle light scatter, and 10,000 via-
ble cells were collected in each file. Specificity and sensitivity of
staining were controlled by checkerboard analysis against normal
Ly 5.1 and Ly 5.2 thymocytes and purposeful mixtures thereof.
The percentage of positive cells was calculated by using the inter-
section of the fluorescence histogram with its control profile to
determine the cutoff point.

Parabiosis.  Pairs of 4 to 5-wk-old, sex- and weight-matched, 
Ly 5 congenic mice were surgically joined by cutaneous vascular
anastomosis as described previously (7). Parabiotic mice were
maintained for periods of 1–9 wk before sacrifice, or were surgically
separated at weekly intervals and killed 28 d later. Thymi
from 4–6 pairs of nonseparated or separated parabiotic partners
were harvested at the indicated time points and the respective de-
grees of chimerism were determined by FCM analysis.

Results

Frequency and Range of Thymic Chimerism Induced by In-
trathymic and Intravenous Injection of BM Cells.  If, as is gen-
erally assumed, the kinetics of importation of thymocyte
precursors in adult mice is constant under steady-state con-
ditions, <5% of sites (presumptive microenvironmental
niches) for prothymocyte engraftment should be available
in the thymus at any given time (7, 9). To test this predic-
tion, suspensions of Ly 5.2 BM cells were injected intrathy-
mically or intravenously into groups of nonirradiated Ly
5.1 mice of mixed ages (7–12 wk), using doses previously
found to be saturating in irradiated recipients (2 × 10^6) cells
intrathymically and 20 × 10^6 cells intravenously; reference
17). Thymocytes were harvested 28 d later, when peak lev-
els of chimerism occur (reference 17; also see below).

As shown in Fig. 1, 86% of recipients displayed thymic
chimerism after intrathymic injection, and the levels of chi-
erism attained were essentially random over a range of
5–90% donor origin cells. In contrast, only 13% of recipi-
ents developed significant chimerism after intravenous
injection, and, although the levels of chimerism attained
were broad (5–50% donor-origin cells), there was marked
skewing towards the lower end of the scale. In both in-
stances, >95% of the donor origin cells expressed CD3
data not shown), formally demonstrating that they were
thymocytes.

These results suggested that thymocytopoiesis in normal
adult mice was most likely to be maintained by the periodic
influx of saturating numbers of blood-borne proth-
ymocytes. This hypothesis is formally confirmed below by
age-response experiments conducted in individual cohorts
of intrathymically or intravenously injected normal mice.

Cyclical Induction of Thymic Chimerism by Intrathymically
Injected BM Cells.  In these experiments, age-matched (±
3 d) groups of 5 to 16-wk-old nonablated Ly 5.1 mice ob-
tained from a single cohort were injected intrathymically
with 2 × 10^6 Ly 5.2 BM cells. Results in Fig. 2 show that
the frequency of thymic chimerism and mean level and
number of donor origin thymocytes attained 28 d after in-
jection varied cyclically, with peaks occurring at 5, 9, and
13 wk of age. The deepest valley (week 15) occurred after
the onset of physiological thymus involution. Again, the
range of donor origin thymocytes varied from <5 to >75%
(data not shown). Furthermore, the maximal numbers of
donor-origin thymocytes generated in nonirradiated recipi-
ents (120 × 10^6) were similar to those obtained in suble-
thally irradiated recipients (17). These results suggested that

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Figure 2. The induction of thymic chimerism in adult mice is cyclical
after intrathymic (I.T.) injection of BM cells. A cohort of 5-wk-old (±
3 d) Ly 5.1 mice was divided into 12 groups (9–12 mice each) and, at
weekly intervals, a different group was injected intrathymically with a sat-
urating dose (2 × 10^6) of Ly 5.2 BM cells. The percentage of Ly 5.2 thy-
mocytes present 28 d later was determined by FCM analysis. Results for
total mice in each group (5–16 wk of age) are presented as: (A) frequency
of thymic chimerism (≥5% donor-origin cells); (B) mean percentage of
donor-origin thymocytes (± SD); and (C) mean number of donor-origin
thymocytes. Percentage of chimeric mice = 62.5%. Maximum level of
thymic chimerism = 88% (120 × 10^6 donor-origin cells). * P < 0.05 be-
tween highest and lowest values in each cycle. This experiment was re-
peated in part on two occasions using cohorts of mice varying in age from
5–8, 7–12, and 12–16 wk. Similar results were obtained, with peaks and
valleys shifting by no more than 1 wk from those illustrated.

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the age-related waves of thymic chimerism detected by intrathymic injection represented changes in the proportion of niches available for prothymocyte engraftment.

**Periodic Induction of Thymic Chimerism by Intravenously Injected BM Cells.** In these experiments, age-matched (± 3 d) groups of 3 to 18-wk-old nonablated Ly 5.1 mice were injected intravenously with $20 \times 10^6$ Ly 5.2 BM cells. Results in Fig. 3 show that the ability of intravenously injected BM cells to induce thymic chimerism was intermittent (periodicity 3–6 wk) rather than cyclical. Hence, although the spikes of receptivity in intravenously injected mice coincided roughly with the peaks of receptivity in intrathymically injected mice (Fig. 2), and the maximal percentage (78%) and number (98 $\times 10^6$) of donor-origin thymocytes generated approximated those by intrathymic injection, the differential receptivity to intravenous and intrathymic injections at most other time points indicated that the importation of blood-borne prothymocytes was gated. Furthermore, paired analysis of individual thymic lobes in intravenously injected mice (Fig. 4) suggested that gate opening was tightly coordinated between both lobes of individual thymi.

**Synchronization of Intrathymic Gating.** Despite the general correlation between maximal receptivity for the induction of thymic chimerism after intravenous and intrathymic injection of BM cells, the asynchrony within a given cohort of mice made it difficult to determine the precise relationship between these parameters. As shown in Fig. 5, this problem was not overcome by using litters of age- and sex-matched mice from timed matings. Thus, although the levels of chimerism attained after intrathymic injection were fairly uniform within individual litters, there was considerable asynchrony between litters. This was especially well illustrated in 35-d-old mice, in which the wide differences in mean levels of thymic chimerism between litters presumably reflected minor differences in the timing of gate opening for circulating host prothymocytes.

We therefore attempted to synchronize intrathymic gating in a cohort of 5-wk-old (± 3 d) Ly 5.1 mice by first filling all available niches for prothymocytes by intrathymic injection of saturating doses of Ly 5.2 BM cells. Results in Fig. 6 show that the ability of intrathymically injected BM cells to induce thymic chimerism was intermittent (periodicity 3–6 wk) rather than cyclical. Hence, although the spikes of receptivity in intrathymically injected mice coincided roughly with the peaks of receptivity in intrathymically injected mice (Fig. 2), and the maximal percentage (78%) and number (98 $\times 10^6$) of donor-origin thymocytes generated approximated those by intrathymic injection, the differential receptivity to intravenous and intrathymic injections at most other time points indicated that the importation of blood-borne prothymocytes was gated. Furthermore, paired analysis of individual thymic lobes in intravenously injected mice (Fig. 4) suggested that gate opening was tightly coordinated between both lobes of individual thymi.

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injection of host allotype BM cells. Subsequent intrathymic or intravenous injections of Ly 5.2 BM cells documented the resulting synchronization of chimerism with time. As shown in Fig. 6, A and B, the frequencies and mean levels of thymic chimerism after intrathymic injection occurred in two clearly defined cycles. Each peak was preceded by an ~2-wk period of increasing availability of putative niches for prothymocytes, followed by a 2-wk period of decreasing availability of niches (presumably due to occupation by recently imported host-origin precursors).

Similarly, results in Fig. 6, C and D, formally documented the gated entry of intravenously injected precursors at times corresponding to maximal availability of niches for intrathymically injected precursors (weeks 8 and 12 to 13). These results also suggested that gate closure (refractory period) is initiated by the occupation of these niches (descending limbs; Fig. 6, A and B), and that it persists through the subsequent period of increasing availability (emptying) of niches (ascending limbs; Fig. 6, A and B).

As anticipated, synchronization of the importation of blood-borne precursors permitted a more precise analysis of the kinetics of thymocytopoiesis during the establishment of chimerism. Thus, although a single wave of thymocytopoiesis spanning two periods of prothymocyte gating (8–10 wk) was observed after intravenous injection of BM cells into either synchronized or nonsynchronized mice (Fig. 7), peak levels of thymocytopoiesis appeared to be reached ~1 wk earlier and to persist for 1 wk less in the synchronized mice. Intermediate kinetics were observed after intrathymic injection of BM (data not shown). In addition, phenotypic analysis of the donor-origin thymocytes in intrathymically injected mice showed a progression between weeks 1 and 5 from double-negative (CD4−CD8−) to double-positive (CD4+CD8+) to single-positive (CD4+CD8− or CD4−CD8+) cells (Table I). Similar results were obtained after intravenous injection (data not shown). Hence, most of the donor-origin thymocytes in these assay systems appear to be generated by early lymphoid precursors in the BM cell inoculum.

Reproducible Induction of Thymic Chimerism by Sequential Intravenous Injections of BM Cells. Given the periodicity of prothymocyte gating (~4 wk) and the duration of the ensuing wave of thymocytopoiesis (~8 wk), it should be possible to induce thymic chimerism reliably in nonsynchr-
Table I. Phenotypic Profile of Donor-Origin Thymocytes after Intrathymic Injection

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD4⁺CD8⁻</td>
<td>59.9 ± 2.4</td>
<td>40.2 ± 6.5</td>
<td>3.4 ± 0.9</td>
<td>2.1 ± 0.4</td>
<td>2.0 ± 0.5</td>
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<tr>
<td>CD4⁺CD8⁺</td>
<td>25.7 ± 5.1</td>
<td>52.5 ± 3.6</td>
<td>88.6 ± 1.5</td>
<td>87.7 ± 2.5</td>
<td>83.0 ± 3.0</td>
</tr>
<tr>
<td>CD4⁻CD8⁻</td>
<td>2.6 ± 1.5</td>
<td>3.6 ± 1.7</td>
<td>4.2 ± 1.3</td>
<td>7.7 ± 2.4</td>
<td>10.6 ± 2.5</td>
</tr>
<tr>
<td>CD4⁻CD8⁺</td>
<td>11.7 ± 3.9*</td>
<td>3.7 ± 1.9</td>
<td>3.8 ± 1.8</td>
<td>2.5 ± 0.1</td>
<td>4.4 ± 0.8</td>
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*Mostly CD8⁺ as compared with later time points.

Figure 7. A single wave of thymocytopoiesis is induced by intravenous (I.V.) injection of BM cells into normal or synchronized adult mice. A cohort of 6-wk-old Ly 5.1 mice was injected intravenously (i.v.) with a saturating dose of Ly 5.2 BM cells. In addition, a cohort of 4-wk-old Ly 5.1 mice was synchronized by intrathymic injection of Ly 5.1 BM cells and reinjected intravenously 3 wk later with Ly 5.2 BM cells (○). Mean numbers of Ly 5.2 thymocytes present in groups of 5–10 mice were determined at weekly intervals thereafter. Results at each time point are expressed as percentage of maximal numbers of Ly 5.2 thymocytes generated to allow for differences in peak levels of chimerism.

A cohort of 5-wk-old Ly5.1 mice was injected intrathymically with 2 × 10⁶ Ly5.2 BM cells. Thymocytes were harvested from groups of five mice at the indicated times after injection and subjected to multiparameter FCM analysis. Results represent mean percentage ± SD of Ly5.2⁺ thymocytes that express the indicated CD4 and/or CD8 phenotypes.

Figure 8. Sequential intravenous (I.V.) injections of BM cells routinely induce thymic chimerism in cohorts of 7-wk-old mice. Groups of 30 7-wk-old (± 3 d) Ly 5.1 mice from a single cohort were sequentially injected intravenously with saturating doses of Ly 5.2 BM cells (A) weekly on four occasions or (B) biweekly on eight occasions. Sets of 10 mice from each group were killed 1, 2, and 3 wk after the final injection, and levels of thymic chimerism were determined by FCM analysis. Dots represent results for individual animals. Bars indicate mean levels of thymic chimerism.

Revealed that: (a) all of the multiply injected mice developed significant thymic chimerism (range of donor-origin cells, 5–32%); (b) the highest levels of chimerism in this experiment occurred at week 2 of harvest; and (c) the mean peak chimerism was significantly higher (P < 0.05) after bi-weekly (19 ± 5%; Fig. 8 B) than weekly (13 ± 5%; Fig. 8 A) injections. In contrast, only 5% of 7-wk-old control mice that had been injected for 1 wk only developed thymic chimerism 4 to 6 wk later (data not shown). These results suggested that, in most of the animals, the gate for prothymocytes opened during the second week of injection (8 to 9 wk of age), as reflected by the peak of thymocytopoiesis 4 wk later.

To determine if BM chimerism also occurred in these animals, 20 × 10⁶ BM cells were harvested from each of 20 nonablated mice 1 wk after they had been given a series of four weekly or eight biweekly intravenous injections. These cells were then transferred intravenously into sublethally irradiated (6 Gy) Ly 5.1 recipients, which were analyzed 28 d later for thymic chimerism. Only one of the secondary recipients became chimeric (data not shown). In contrast, all irradiated recipients became chimeric when they were injected with purposeful mixtures of normal BM cells at donor/host ratios as low as 1:40. Hence it appeared that stem cell chimerism capable of generating prothymocytes had not been established in the BM of the primary (nonablated) recipients.

These experiments were then expanded to encompass two gate-openings. Groups of 4 to 9-wk-old mice were injected with BM biweekly for a total of seven injections and killed 2 wk later. As shown in Fig. 9, thymic chimerism was obtained in the majority of multiply injected mice in
each age group, but not in control mice injected during week 7 only. Hence, the observed chimerism induced by intravenous injections given before or after week 7 must have resulted from separate gate openings.

**Gated Importation of Prothymocytes in Parabiotic Mice.** In the preceding experiments, mice received single or multiple intravenous injections of large numbers ($20 \times 10^6$) of BM cells. It therefore was possible that the physiological mechanisms that regulate the importation of prothymocytes were periodically overwhelmed, yielding spikes of thymic chimerism. To exclude this possibility, the kinetics of hematogenous precursors were studied in parabiotic mice, whose thymi presumably are exposed to physiological numbers of blood-borne prothymocytes at physiological intervals. 5-wk-old Ly 5 congenic mice were parabiosed for periods of 1 to 9 wk and then surgically separated to prevent further exchange of blood-borne prothymocytes, the primary mechanism for maintaining thymic chimerism in such animals (7). Under these circumstances, the occurrence of thymic chimerism 4 wk later was presumed to reflect the entry of donor-origin prothymocytes shortly before the time of separation.

Results in Fig. 10, A and B, showed two windows of receptivity for induction of thymic chimerism in the separated parabiotic partners, the first occurring at week 2 (Ly 5.2 mice) or week 4 (Ly 5.1 mice) of parabiosis, and the second 4 wk later. In addition, the mean levels of thymic chimerism attained after the first windows of receptivity closely approximated those observed 4 wk later in unseparated parabionts (data not shown). This further indicated that the initial waves of prothymocyte importation had occurred at weeks 2 and 4 of parabiosis, respectively.

**Discussion**

Two experimental approaches were used to demonstrate that thymocytopoiesis is a gated phenomenon in normal adult mice. First, age-related transfer of BM cells into non-ablated mice revealed a cyclical pattern of engraftment of intrathymically injected thymocyte precursors, and an intermittent pattern of engraftment of intravenously injected precursors. Both patterns had average periodicities of 4 wk, and receptivity for intravenously injected precursors correlated with maximum availability of putative niches for intrathymically injected precursors. Second, timed separation of parabiotic mice also revealed an intermittent pattern of importation of hematogenous precursors into thymus, again with a periodicity of ~4 wk.

Experiments in mice in which gating was synchronized by an initial intrathymic injection of host-allotype BM cells were especially useful in establishing the kinetics of prothymocyte importation. This model demonstrated that the gate for hematogenous precursors opens for ~1 wk to allow the niches to fill and then closes for 2 to 3 wk to allow them to empty. In addition, the mean duration of a wave of thymocytopoiesis in nonablated recipients of BM, as in parabiotic mice (7), was found to exceed the periodicity of gating by twofold. This is important, as proportionately shorter waves would not generate a steady-state pattern of

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**Figure 9.** Sequential intravenous (I.V.) injections of BM cells routinely induce thymic chimerism in cohorts of 4 to 9-wk-old mice. A cohort of 4-wk-old (± 3 d) Ly 5.1 mice was divided into seven groups (six mice each) and, at weekly intervals, a different group was enrolled in a course of seven biweekly intravenous injections (3 wk) of Ly 5.2 BM cells. Levels of thymic chimerism for each group were determined 2 wk after the final injection. Dots indicate results for individual animals. Bars indicate mean chimerism for each group (identified by elapsed ages during injections). The group designated “7 only” received two intravenous injections of BM cells during week 7 only and was analyzed for thymic chimerism at week 12 (i.e., at the same time as the 7–10 wk group).

**Figure 10.** The induction of thymic chimerism is periodic in parabiotic mice. Ly 5.1 and Ly 5.2 congenic mice were parabiosed at 5 wk of age, and groups of 4–6 parabiotic pairs were surgically separated at weekly intervals over a 9-wk period. The frequency of thymic chimerism (~5% donor-origin cells) in the separated (A) Ly 5.2 and (B) Ly 5.1 parabiotic partners was determined 28 d later.
thymocyte production. Furthermore, the ability of intrathymically injected BM cells both to synchronize and alter the time of gate opening in nonablated mice suggested that, under steady-state conditions, gate closing is initiated by the coordinated filling of most, if not all, niches for prothymocytes. Conversely, the temporal association of prothymocyte importation with the availability (emptying) of most, if not all, of these niches suggested that gate opening may be regulated by a threshold-dependent, downstream signal. An integrated model of thymocytopoiesis and prothymocyte gating based on these considerations is presented in Fig. 11.

Our ongoing studies suggest that the availability of saturating levels of prothymocytes to the thymus at the time of gate opening normally is assured by a feedback loop that regulates the periodic release of waves of prothymocytes from the BM (unpublished observations). Although it is tempting to speculate that this same feedback loop triggers gate opening itself, neither the nature of the gate(s) nor the signal(s) that regulates its activity is known. Thus, although gate opening occurs simultaneously in both thymic lobes (Fig. 4), the asynchronous development of thymic chimerism in parabiotic partners (Fig. 10) suggests that the signal for gate opening either does not effectively cross-circulate in the blood or is otherwise unable to stimulate the refractory partner.

It might be argued that the differential kinetics of receptivity to the establishment of thymic chimerism after intravenous and intrathymic injection of BM is not due to gating, but to occupation of different sets of binding sites by the intravenously and intrathymically injected precursors. This is highly unlikely, as differential occupation of binding sites would not explain: (a) the ability of intrathymically injected BM cells to synchronize gating for intravenously injected BM cells; (b) the origin of donor thymocytes from CD4+CD8− precursors in both assay systems; or (c) the gated entry of circulating precursors into the thymus of parabiotic mice. It is also unlikely that the periodic importation of prothymocytes in adult life is due to cyclical hormonal changes or to inapparent stress. Thus, gating is not sex-related, and dexamethasone treatment does not predispose recipients to thymocyte chimerism after intravenous or intrathymic injection (reference 9; and our unpublished observations). Rather, the fact that both the intravenous and intrathymic assays obey strict log dose saturation kinetics and generate the same maximum number of thymocytes (17) favors the existence of a finite number of specific binding sites (niches) for prothymocytes. This notion is further supported by our recent demonstration of competitive one-on-one occupancy kinetics for binding sites after combined intravenous and intrathymic injections of BM cells into radioablated mice (unpublished observation).

Pragmatically, single intravenous injections of BM cells into young adult mice of nominally receptive ages may not consistently establish thymic chimerism because of asynchronous gating, and synchronization of gating by initial intrathymic injection does not lend itself to routine use. Instead, reliance on the periodicity of gating, rather than chronological age, appears to be a more efficient approach to inducing thymic chimerism in heterogeneous groups of mice. Thus, as shown in Figs. 8 and 9, biweekly intravenous injections of BM cells over a period of 3 to 4 wk (7 to 8 injections) reproducibly generates significant thymic chimerism in parabiotic partners.

Figure 11. Integrated scheme of the kinetics of prothymocyte gating, occupation of microenvironmental niches, and generation of thymocytes in normal mice. Clusters of vertical arrows represent receptive periods (open gate) and horizontal black bars represent refractory periods (closed gate) for importation of hematogenous prothymocytes. Shaded triangles represent filling/equilibration (up slope) and emptying (down slope) phases of occupation of a finite number of intrathymic niches by prothymocytes and their immediate descendants. Dashed, dotted, and mixed symbol curves represent sequential waves of thymocytopoiesis, each generated by the gated importation of a saturating wave of prothymocytes. The lag period of thymocytopoiesis corresponds roughly to the filling/equilibration phase of occupation of the intrathymic niches. The duration of each wave of thymocytopoiesis exceeds the periodicity of gate-opening by twofold, so as to maintain relatively constant levels of total thymocytes. Gate closing appears to be initiated by occupation of intrathymic niches. Gate opening appears to be regulated in a threshold-dependent (all-or-none) manner by emptying of intrathymic niches, and to be synchronized between thymus lobes. The onset of thymic involution occurs at about week 12 and is partly related to a decrease in the number of available intrathymic niches for prothymocytes (unpublished observations). Although drawn as discrete curves, each idealized wave of thymocytopoiesis may actually consist of a series of partially overlapping waves (reference 9). Similarly, the prothymocyte “gate” may actually be a series of individual microvascular gates. The idealized time scale (weeks) approximates, but is not necessarily identical to, chronological age. The receptivity of normal neonatal mice and rats (week 0) to the induction of thymic chimerism by intravenously injected BM cells has been established in earlier studies (references 23 and 24).
ism in most recipients. Although more frequent injections or a more prolonged course of injections might further improve this protocol, at some point hemopoietic stem cell chimerism will occur in BM (14), after which the development of thymic chimerism no longer will reflect the direct importation of prothymocytes from the original inoculum.

As mentioned, gated importation of waves of prothymocytes serves to generate discrete populations of T cells in a programmed fashion in late fetal and early neonatal life (1–5). Yet, absent evidence for the continued need to generate developmentally disparate waves of thymocytes, the possible immunobiological benefits of intermittent importation of prothymocytes in adult life can only be surmised. At one extreme, the benefits of continued prothymocyte gating may relate primarily to matters of quantitative efficiency, whereby monthly restocking of the intrathymic “warehouse” with a saturating dose of prothymocytes eliminates the need for constant monitoring of inventory and available space. At the other extreme, prothymocyte gating may relate more to matters of qualitative efficiency, whereby periodic changes in the antigenic milieu can be selectively monitored (18). Such a mechanism could serve to update the immunological repertoire throughout life by optimizing the production and selection of new T cell specificities.

A particularly attractive scenario under the latter rubric would be for each wave of developing thymocytes to be selected against newly processed antigenic peptides presented by a coordinate wave of newly generated and/or imported thymid dendritic cells (19, 20). Evidence for parallel generation and interaction of waves of thymocytes and thymic dendritic cells has been provided in the fetal/neonatal period and in adult radiation BM chimeras (21). In addition, we have observed parallel waves of thymocyte and dendritic cell chimera in parabiotic mice and in non-ablated adult mice injected intravenously or intrathymically with BM cells (unpublished observations). A similar, non-exclusive scenario envisions the coordinate selection of waves of developing thymocytes by the developmentally and/or temporally regulated expression of “extrathymic” gene products by thymic epithelial cells (22).

In sum, the present results provide a potentially new paradigm for thymic function in adult life based on the regulated importation of waves of hematogenous prothymocytes in coordination with the maximal availability of intrathymic binding sites. At a basic level, these observations should expedite the fine analysis of the earliest stages of thymocytopoiesis, allow the identification of microenvironmental niches for prothymocytes, and permit the characterization of the feedback loop(s) that regulates prothymocyte gating. They should also provide a direct approach to determining if the receptor repertoire of a given wave of emerging T cells is biased towards newly introduced and/or newly expressed peptide specificities. In addition, by offering a protocol for inducing individual waves of thymic chimerism on demand in nonmyeloablated recipients, the results may have applied implications for selective prothymocyte engraftment.

We would like to thank Dr. Lynn Puddington for her expert assistance in the FCM analyses.

This study was supported in part by National Institutes of Health Grant AI33741.

Submitted: 11 May 2000
Accepted: 19 December 2000

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