Effects of Breeding Environments on Generation and Activation of Autoreactive B-1 Cells in Anti-red Blood Cell Autoantibody Transgenic Mice

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Summary

In anti-red blood cell autoantibody transgenic (autoAb Tg) mice almost all B cells are deleted except for B-1 cells in the peritoneal cavity and the gut. About one-half of the auto Ab Tg mice suffer from autoimmune hemolytic anemia (AIHA) in the conventional condition. Oral administration of lipopolysaccharides activates B-1 cells and induces autoimmune symptoms in the Tg mice, suggesting that the autoimmune disease in anti-RBC autoAb Tg mice is triggered by infections. To examine the association of bacterial infections with the generation of B-1 cells and the occurrence of the autoimmune disease, we analyzed anti-RBC autoAb Tg mice bred in germ-free and specific pathogen-free conditions. In germ-free conditions, few peritoneal B-1 cells were detected, while a significant number of peritoneal B-1 cells existed in specific pathogen-free conditions. In both conditions, no mice suffered from AIHA. However, when these Tg mice were transferred to the conventional condition or injected with lipopolysaccharide, peritoneal B-1 cells expanded and some of these mice suffered from AIHA. These results clearly showed that bacterial infections are responsible for both the expansion of B-1 cells and the onset of the autoimmune disease in these Tg mice.

Several lines of evidence indicate that bacterial infection may activate autoreactive lymphocytes in vivo and induce autoimmune diseases. LPS derived from gram-negative bacteria is known to activate B cells polyclonally and the administration of LPS induces the production of autoantibodies (autoAbs) even in normal mice (1). Enterotoxins derived from bacteria serve as superantigens that can bind T cell receptors, resulting in stimulation and expansion of a wide population of T cells (2). In some cases, autoreactive lymphocytes may be activated after the infection of bacteria and viruses, whose antigens have certain structural similarities with self-antigens (molecular mimicry) (3). Recently, the induction of autoimmune disease by infection was suggested in experimental allergic encephalomyelitis (EAE) transgenic (Tg) model mice, in which all T cells express myelin basic protein–specific T cell receptors. EAE did not occur in the Tg mice bred in specific pathogen-free (SPF) conditions, but some of the mice housed under conventional conditions showed the autoimmune disease (4). On the other hand, there exist several arguments against the involvement of infection in autoimmune diseases. The frequency of autoimmune gastritis in day 3 thymectomized mice remained the same under germ-free (GF) as well as under conventional conditions (5) and the frequency of autoimmune insulitis is known to increase in NOD mice bred under SPF conditions (6). Thus, it remains elusive whether or how infection is involved in autoimmunity.

B-1 cells constitute a distinct B cell population that differs in several of its functional properties from conventional B cells (7–9), and has a strong capacity for self-renewal (10). They are found predominantly in peripheral tissues such as the peritoneal and pleural cavities (11). B-1 cells preferentially use VH segments proximal to the DJ segments and produce IgM against self- and bacterial antigens (7–9). They are characterized by expression of surface antigens such as IgM hi, IgD lo, B220 lo, M ac-1+, and CD23 lo (7–9). Although B-1 cells are suggested to play an important role in autoimmune diseases (7, 8), the physiological and pathological roles of B-1 cells still remain unclear.

A key to solve these questions about B-1 cells and autoimmunity lies in creating suitable animal models to simplify genetic and pathogenic factors of autoimmunity. Previously, we established and characterized a Tg mouse line carrying the Ig genes derived from the anti-red blood cell (RBC) autoAb 4C8 (12–15). In these mice, almost all B cells were deleted in the periphery but a normal number of B-1 cells survived in the peritoneal cavity, which is sequestered from RBCs (12, 13). Under conventional breeding condi-
tions, about one-half of the animals of this Tg line suffer from autoimmune hemolytic anemia (AIHA), even though they have the same genetic background and the same number of B-1 cells (12, 15). The production of the autoAb by peritoneal B-1 cells was shown to be responsible for AIHA in the Tg mice because apoptotic death of B-1 cells by repeated exposure to RBCs completely cured AIHA (13) and because AIHA did not occur in the anti-R BC autoAb Tg mice bearing the X-linked immunodeficiency (xid) mutation, which deletes B-1 cells (14). On the other hand, oral administration of LPS to the nonsymptomatic Tg mice caused activation of peritoneal B-1 cells and induction of AIHA (14). These lines of circumstantial evidence suggest that enteric bacteria or some infectious agents activate peritoneal B-1 cells and induce the autoimmune disease in the Tg mice (15).

In the present study, we examined whether bacterial infections have influences on the generation and activation of B-1 cells and on the occurrence of autoimmunity in the anti-R BC autoAb Tg mice that are bred under different conditions, i.e., the GF, SPF, and conventional conditions. We report that bacterial infections most likely via the gut are necessary for the generation of B-1 cells and the occurrence of the autoimmune disease in this Tg model.

Materials and Methods

Mice. Homozygous heavy chain and light chain Tg mice of the anti-R BC autoAb were established and maintained under conventional breeding conditions in the Center for Molecular Biology and Genetics, Kyoto University. By mating homozygous heavy and light chain Tg mice, we obtained conventionally bred anti-R BC autoAb Tg mice that carry both heavy and light chain transgenes of autoAb (12). To make GF animals from anti-R BC autoAb Tg mice, two conventional pregnant mice (a heavy chain Tg mouse and a light chain Tg mouse) were aseptically hysterecto-

| Table 1. Occurrence of AIHA in Anti-RBC AutoAb Tg Mice in Different Breeding Environments |
|---------------------------------------------|-----------------|-----------------|-----------------|-----------------|
| Frequency of autoimmune hemolytic anemia | ConV | SPF | GF | ConV* (LPS injected) | GF/ConV | SPF/ConV | GF* (LPS injected) |
| Number of tested mice | 63 | 26 | 30 | 13 | 6 | 12 | 8 |
| Number of mice with autoimmune hemolytic anemia | 29 | 0 | 0 | 13 | 2 | 4 | 2 |
| Number of mice with autoimmune hemolytic anemia % | 46 | 0 | 0 | 100 | 33 | 33 | 25 |

We considered Tg mice with hematocrit values below 40% as AIHA mice and examined the occurrence of AIHA in 6- to 8-wk-old anti-RBC autoAb Tg mice in different breeding conditions: GF, SPF, and conventional conditions (ConV).

*Anti-R BC autoAb Tg mice bred in the conventional or GF condition were examined 7 d after oral administration of 100 μg of LPS. GF Tg mice or SPF Tg mice were transferred to the conventional condition, bred there for 4-6 wk, and then examined.

Flora of stool from GF mice were examined under standard methods (16). The stool was cultured for aerobic or anaerobic bacteria and fungi as described before (12–14). Single cell suspensions were made from the samples of stool, and the bacterial and fungal contamination were checked. Bacterial and mycological contaminations were absent in GF mice, feces derived from GF mice were aerobically cultured for 2 wk in thiglycollate broth (Eiken, Tokyo, Japan) at 37°C, and a small amount of the cultured fluid was inoculated on the surfaces of two brain–heart infusion agars (Nissui, Tokyo, Japan) and a Sabouraud's agar (Eiken). Each agar plate of the former was incubated for 1 wk at 37°C under aerobic or anaerobic conditions, and the latter was aerobically incubated for 1 wk at room temperature. All GF mice were negative in all these tests. To check the SPF conditions, materials were collected from blood, rhinal swab, cecum contents, duodenum contents, and anus, and were examined by ELISA, culture, or microscopic test for the following pathogenic organisms: Sendai virus, mouse hepatitis virus, Mycoplasma pulmonis, Tyzzer's organism, Corynebacterium Kutscheri, Pasteurella pneumotropica, Salmonella species, Giardia musis, Trichomonas species, Syphacia species, and Spironucleus muris.

Flow Cytometry. Cells were isolated from spleen or peritoneal cavities as described before (12–14). Single cell suspension of 10⁶ mononuclear cells was pelleted and resuspended in 10 μl of normal mouse serum for blocking the nonspecific binding of antibodies. After 10-min incubation, one of the first reagent antibodies, i.e., rat anti-mouse B220 (RA3-6B2) mAb or rat anti-Mac-1 Ab (M1/70.15.115.5H1), was added directly at the appropriate dilutions in PBS. After incubation, cells were washed twice with 1 ml of PBS containing 5% FCS and 0.05% sodium azide. The PE-conjugated anti-mouse IgM Ab (Southern Biotechnology Associates, Birmingham, AL) was added as the second reagent Ab in 10 μl of PBS at appropriate concentrations. After excluding dead cells by propidium–iodine staining and gating the lymphoid cells in forward and side scatter analysis, 10⁶ viable lymphoid cells were analyzed on FACScan® and were plotted on quadruple logarithmic scales.
Results and Discussion

Breeding Conditions Determine the Occurrence of AIHA in Anti-RBC AutoAb Tg Mice. We previously reported that only one-half of anti-RBC autoAb Tg mice suffered from AIHA when housed under conventional breeding conditions (12). AIHA was induced in nonsymptomatic Tg mice by the administration of LPS (14), suggesting that some environmental factors such as infections are involved in the occurrence of AIHA. To demonstrate the involvement of infection in the onset of autoimmunity, we first examined the occurrence of AIHA in Tg mice bred in the GF and SPF conditions. GF Tg mice contained no or few B cells, whereas SPF Tg mice had a lesser but significant number of B-1 cells (B220<sup>+</sup>, IgM<sup>+</sup>, Mac-1<sup>-</sup>) as compared with Tg mice bred in the conventional condition (Fig. 1). Moreover, when GF Tg mice were transferred to the conventional condition, a small but significant number of B-1 cells appeared in the peritoneal cavity (Fig. 1).

To demonstrate more clearly the relationship between the occurrence of AIHA and infection, we transferred GF and SPF Tg mice into the conventional condition. Interestingly, one-third of the transferred Tg mice suffered from AIHA when transferred to the conventional facility (Table 1). Oral administration of LPS also induced AIHA in GF Tg mice. These results imply that autoAb-producing cells or their progenitors exist in GF and SPF Tg mice, even though their number is very small. A negligible number of peritoneal B-1 cells in GF Tg mice contained no or few B cells, whereas SPF Tg mice hardly detected them by flow cytometry after staining with fluorescence-conjugated mAbs. In spleens and bone marrows, B cells, including B-1 cells and conventional B cells, were hardly detected in Tg mice bred in the SPF and GF conditions as well as in the conventional condition (data not shown). By contrast, the numbers of B-1 cells in the peritoneal cavity of Tg mice are different depending on environmental conditions. The peritoneal cavity of GF Tg mice contained no or few B cells, whereas SPF Tg mice had a lesser but significant number of B-1 cells (B220<sup>+</sup>, IgM<sup>+</sup>, Mac-1<sup>-</sup>) as compared with Tg mice bred in the conventional condition (Fig. 1). Moreover, when GF Tg mice were transferred to the conventional condition, a small but significant number of B-1 cells appeared in the peritoneal cavity (Fig. 1).

Table 2. Examination of Microorganisms in Conventionally Bred Anti-RBC AutoAb Tg Mice

| Organism                  | Method     | Material       | R results |
|---------------------------|------------|----------------|-----------|
| Sendai virus              | ELISA      | Serum          | +         |
| Mouse hepatitis virus     | ELISA      | Serum          | +         |
| M ycoplasma pulmonis      | ELISA      | Serum          | –         |
| Tyzzer’s organism         | ELISA      | Serum          | +         |
| Corynebacterium kutscheri | Culture    | R hinal swab   | –         |
| Pasteurella pneumonia      | Culture    | R hinal swab   | –         |
| Salmonella species        | Culture    | Cecum contents | –         |
| Giardia muris             | Microscopic test | Cecum contents | –         |
| Trichomonas species       | Microscopic test | Cecum contents | –         |
| Syphacia species          | Microscopic test | Anus          | +         |
| Spiroplasma muris         | Microscopic test | Duodenum contents | –         |

Figure 1. The number of peritoneal B-1 cells in anti-RBC autoAb Tg mice depends on breeding conditions. Peritoneal cells of 8- to 16-wk-old anti-RBC autoAb Tg mice bred in different environments were examined by flow cytometric analysis with anti-IgM and anti-B220 or anti-Mac-1 Abs. GF/ConV: anti-RBC autoAb Tg mice bred in the GF condition were transferred to the conventional condition (ConV) and kept there for 4-6 wk before examination. Percentages of cell numbers in each quadrant are indicated.
duce the autoimmune disease. In other words, in this Tg model, the expansion and activation of peritoneal B-1 cells may be induced by two steps: enteric bacteria increases the number of peritoneal B-1 cells and pathogenic infection induces peritoneal B-1 cells to produce autoAbs.

B-1 Cells, Infection, and Autoimmunity. The anti-R BC Tg line clearly provided clear evidence for the involvement of B-1 cells and intestinal bacterial infection in autoimmunity. It remains to be seen whether this conclusion can be more generalized in other types of autoimmune diseases, although similar observations were reported for the EAE Tg model (4). It is also important to indicate that the generation of peritoneal B-1 cells is also dependent on bacterial infection in this Tg line. However, this finding contradicts a previous report that the number of peritoneal B-1 cells in GF BALB/c mice was the same as that of conventionally bred BALB/c mice (16). In fact, the H chain Tg mouse bred in the GF condition contained the same level of B-1 cells as anti-R BC Tg bred in the conventional condition (Fig. 1). There are two distinct properties of GF anti-R BC Tg mice as compared with GF BALB/c mice: autoAb-producing B-1 cells and clonal deletion of conventional B cells. The above discrepancy is likely due to the absence of conventional B cells in anti-R BC Tg, although it is totally unknown whether the presence of conventional B cells and their interaction with other immunocytes like B and T cells provide growth stimulation factors of self-reactive B-1 cells under the GF condition. It is intriguing that enteric bacteria can stimulate and activate the peritoneal B-1 cells. We previously showed that there is cell traffic between B-1 cells in the peritoneal cavity and the lamina propria of the gut (14). Bacterial polyclonal stimulants may directly stimulate B-1 cells in the lamina propria. Alternatively, bacterial infection activates other immune cells like T cells and dendritic cells, resulting in secretion of lymphokines such as IL-5 and IL-10 (17).

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