Altered ratio of circulating follicular regulatory T cells and follicular helper T cells during primary EBV infection

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Abstract
Follicular help T cells (Tfh) play an important role in the activation and differentiation of B cells, while follicular regulatory T cells (Tfr) control Tfh and resulting humoral immune responses. Accumulating evidence has demonstrated that the dysregulation of Tfr contributed to the pathogenesis of infectious diseases. However, the role of Tfr in Epstein–Barr virus (EBV) infection remains lacking. Fifty-five EBV-infected infectious mononucleosis (IM) patients and 21 healthy individuals (HIs) were recruited in the study. We investigated the number of Tfr (FoxP3+CXCR5+PD-1+CD4+) and Tfh (FoxP3−CXCR5+PD-1+CD4+) of peripheral blood in IM patients at diagnosis (D0) and day 15 after diagnosis (D15) via multicolor flow cytometry. Results revealed that circulating Tfh (cTfh) and Tfr (cTfr) of IM at D0 were both increased compared to HIs, and cTfr began to decline and return to normal at D15, while cTfh was still higher than those of HIs. More interestingly, the cTfr/cTfh ratio of IM at D0 and D15 was lower than that of HIs, suggesting that the balance between cTfh and cTfr was disturbed during primary EBV infection. Correlation analysis showed a positive correlation between cTfr with CD19+IgD+CD27− naive B cells, CD19+IgD−CD27hi plasmablasts or CD19+CD24hiCD27hi B cells. Moreover, both cTfr and the cTfr/cTfh ratio of IM at D0 were negatively correlated with EBV DNA virus load. These results indicate that an imbalance of cTfr and cTfh cells may be involved in the immunopathogenesis of EBV-infected IM patients and may provide novel strategies for controlling EBV-related disease.

Keywords Infectious mononucleosis · Epstein–Barr virus · Follicular regulatory T cells · Follicular helper T cells · Flow cytometry

Introduction
Epstein–Barr virus (EBV) preferentially infects naïve B cells at oropharyngeal lymphoid tissues and subsequently establishes a persistent infection in the circulating memory B cells [1–5]. EBV is strongly associated with nasopharyngeal carcinoma, lymphoma and autoimmune diseases [6–8]. Many T cell subsets and NK are suggested to be involved in immune response to EBV infection. Nevertheless, the mechanisms the human immune system in the control of EBV infection is not completely understood. EBV-infected infectious mononucleosis (IM) patients provide an excellent model for the study of immune responses against EBV.
Follicular helper T cells (Tfh), a novel subset of CD4+ T helper cells, are essential for the formation of germinal centers (GCs) and humoral response [9]. Tfh bind to B cells via CXCR5 on the surface of Tfh and its ligand CXCL13 in the follicular GCs of B cells, then migrate to B cell follicles/GCs in the lymph nodes and support B cell activation,
expansion and differentiation into plasma cells and generate B-cell memory in GCs [9]. In contrast, follicular regulatory T cells (Tfr) are an effector subset of regulatory T cells (Treg), which control GCs cell number and Tfh function but have varied effects on the quality and antigen specificity of the response [10–14]. Studies suggest that partial or temporary disruption of Tfr function leads to increased availability of costimulatory molecules and increase in the antigen-specific immune response [15–18]. Accumulating evidence has demonstrated that circulating Tfr (cTfr) and Tfh (cTfh) have been found in blood and display phenotypic and functional feature of classical Tfr and Tfh [19–21]. Recent studies revealed that cTfr expand during chronic viral and parasitic infections such as human immunodeficiency virus (HIV), hepatitis B virus (HBV), hepatitis C virus (HCV), and Schistosoma japonica [22–24]. Increased cTfr frequency in patients with chronic HBV or HCV was strongly associated with serum viral load of both infections. Miles et al. [24] found Tfr contribute to inefficient GCs responses and inhibit HIV and SIV clearance. Botta et al. [25] found that Tfr were elevated in the late immune response to influenza infection and inhibited the production of autoimmune antibodies. Moreover, Dhaeeze et al. [26] showed that cTfr increased after influenza vaccination and were correlated with anti-flu Ab responses. An altered balance of cTfh and cTfr has been associated with autoimmune diseases such as systemic lupus erythematosus [27], rheumatoid arthritis [28].

To date, few studies have focused on the role of cTfr during EBV infection. Our previous study found that cTfh cells were high in IM patients, which was negatively correlated with naive B cells and positively correlated with memory B cells and plasmablasts [29]. How cTfr arise and evolve over time during EBV infection and the role of cTfr and the effect of cTfr and cTfh balance on B cell differentiation during primary EBV infection remain largely unknown. Here, we observed the balance of cTfr and cTfh and the association cTfr or the cTfr/cTfh ratio with different B subsets in primary EBV-infected IM patients.

Materials and methods

Subjects

Fifty-five IM patients and 21 healthy individuals (HIs) from Zhejiang Provincial People’s Hospital of Hangzhou Medical College were recruited in this study. Serological screening was performed to confirm their EBV infection and to exclude other virus or bacteria infection, including herpes simplex virus 2, rubella virus, cytomegalovirus, toxoplasma, rotavirus, coxsackie virus, mycoplasma, chlamydia, and hepatitis A, B, C and D. Patients with irrelevant chronic diseases or other autoimmune diseases were excluded. Clinical characteristics of the IM patients and HIs are shown in Table 1. EDTA-K₂ anticoagulated peripheral blood samples were collected at diagnosis (D0) and day 15 after diagnosis (D15) and detected within 24 h. Informed parental written consent was obtained in accordance with the Ethics Committee of Zhejiang Provincial People’s Hospital and the Declaration of Helsinki.

Flow cytometry

Peripheral blood mononuclear cells (PBMCs) were acquired by Ficoll-Hypaque density separation (Dakewe, China) and then were stained for the following surface markers of cTfr and cTfh for 20 min: CD4-PC7 (Clone 13B8.2, Beckman Coulter), CXCR5-Alexa Fluor®488 (Clone RF8B2, BD Biosciences) and PD-1-PerCP-Cy5.5 (Clone EH12.1, BD Biosciences). After surface marker staining, cells were fixed and permeabilized using a Cytofix/Cytoperm kit (BD Biosciences, San Diego, CA, USA) and then incubated intracellularly with FoxP3-PE (Clone 259D/ C7, BD Biosciences) for 30 min. Cells were washed and were acquired on FACScano™II flow cytometer, and data were analyzed by FACSDiva software (BD Biosciences, USA). Isotype-matched antibodies were used in all procedures. The strategy is shown in Supplementary Figure 1. FoxP3⁺CXCR5⁺PD-1⁺CD4⁺ cells were defined as Tfr, whereas FoxP3⁺CXCR5⁺PD-1⁺CD4⁺ were as Tfh. The CD4⁺ T cell absolute number of each patient in peripheral blood was obtained by peripheral blood routine lymphocyte subsets analysis and the use of fluorescent bead system. The absolute number of Tfr and Tfh was multiplied by the absolute number of CD4⁺ T cells and the corresponding percent of Tfr and Tfh. PBMCs were stained for the following surface markers to characterize B cell subsets as following:

Table 1 Clinical characteristics of IM patients and healthy individuals

|                | HIs | IM  | P value |
|----------------|-----|-----|---------|
| Sex            |     |     |         |
| Male           | 12  | 34  |         |
| Female         | 9   | 21  |         |
| Age (years)    | 6.33±4.26 | 4.72±2.87 | 0.066   |
| WBC (×10⁹/L)   | 8.31±2.42 | 13.13±5.32 | 0.0002  |
| LYM (×10⁹/L)   | 3.07±1.11 | 8.15±4.48 | <0.0010 |
| EBV DNA (copies/mL) | – | 256740.5±853540.2 | 0.000 |
| EB-VCA-IgM (IU/L) | – | 11.99±5.86 | 0.000 |

WBC white blood cell, LYM lymphocyte, EBV DNA Epstein–Barr virus deoxyribonucleic acid, EB-VCA-IgM Epstein–Barr viral capsid antigens immunoglobulin M

Analysis of T test, Chi-squared test; P < 0.05 was considered statistically significant

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IgD-APC (Clone 11-26c, BD Biosciences), CD24-PerCP-Cy5.5 (Clone ML5, BD Biosciences), CD27-PE (Clone L128, BD Biosciences) and CD19-FITC (Clone J4.119, Beckman Coulter). The strategy of different B cells is shown in Supplementary Figure 2. CD19+IgD+CD27− cells were defined as naïve B cells, CD19+IgD+CD27+ cells were as memory B cells, while CD19+IgD−CD27hi cells were as plasmablasts.

EBV serology and EBV DNA quantification

Serum samples were measured immunoglobulin (Ig) M (IgM) viral capsid antigen (VCA) using enzyme immunoassays (Liason VCA IgM, DiaSorin S.p.A.) according to the manufacturer’s instructions. EBV DNA was measured using a commercial real-time PCR kit, amplifying a 191 bp region of the EBNA-1 gene (BioQuant EBV, Biodiversity), according to the manufacturer’s protocol, and detected by the ABI PRISM 7500 Sequencer Detection System (Applied Biosystems, USA).

Statistical analysis

Statistical analysis was performed with GraphPad Prism 5.01 software. The quantitative data were presented as the mean values ± standard deviations. Statistical tests for data analysis included one-way ANOVA test and Spearman r correlation. P value < 0.05 were considered to be statistically significant.

Results

cTfr and cTfh were significantly increased, while the cTfr/cTfh ratio was significantly decreased in IM patients

The number of CD4+ T cells of IM patients was no different from that of HIs, though the frequency of CD4+ T cells was significantly decreased in IM patients compared to HIs (Fig. 1a). Besides increased levels of cTfh cells, we also observed a significant increase in the cTfr cells during acute EBV infection (D0) in IM compared to HIs (Fig. 1b, c). Then, we observed that both cTfr and cTfh began to decline and were significantly lower at D15 than those at D0, but Tfh at D15 were still higher than HIs (Fig. 1b, d). More importantly, when we further analyzed the cTfr/cTfh ratio, the results showed that cTfr/cTfh ratio at D15 is significantly lower than HIs, and cTfr/cTfh ratio at D0 is low compared with HIs but had no statistical significance (Fig. 1f). These data suggested the imbalance between cTfr and cTfh during acute EBV infection.

Positive correlation between cTfr and CD19+IgD+CD27− naïve B cells, CD19+IgD−CD27hi plasmablasts and CD19+CD24hiCD27hi B cells

B cells play a central role in the pathogenesis of infectious disease. CD19+CD24hiCD27hi B cells have been demonstrated to downregulate inflammatory reactions and induce tolerance by production of IL-10 and/or TGF-β [30, 31]. Our previous study found cTfh were negatively correlated with naïve B cells and positively correlated with memory B cells and plasmablasts. Whether increased cTfr or low cTfr/cTfh ratio will affect the B cell differentiation during primary EBV infection is unknown. Herein, we analyzed the correlation between cTfr or the cTfr/cTfh ratio and different B cell subsets, which included CD19+ B cells, CD19+IgD+CD27− naïve B cells, CD19+IgD+CD27+ memory B cells, CD19+IgD−CD27hi plasmablasts, CD19+CD24hiCD27hi B cells. Our data showed a positive correlation between cTfr and CD19+IgD+CD27− naïve cells, CD19+IgD−CD27hi plasmablasts or CD19+CD24hiCD27hi B cells (Fig. 2d, h, j). However, no correlation between the cTfr/cTfh ratio and these B cell subsets was found (Fig. 2k–t).

Upregulation of cTfr was negatively correlated with EBV DNA viral load in IM patients

cTfr were increased, but the cTfr/cTfh ratio was decreased during primary EBV infection, which prompts us to wonder whether cTfr or the cTfr/cTfh ratio is related to EBV DNA viral load during primary EBV infection. Correlation analysis showed a significant negative correlation between cTfr or the cTfr/cTfh ratio and EBV DNA viral load (Fig. 3a–c). However, we found no correlation between cTfr or the cTfr/cTfh ratio and EB-VCA-IgM in IM patients (Fig. 3d–f).

Discussion

Tfh are important for helping B cell activation and differentiation, while Tfr suppress B cell responses through modulation of Tfh and GCs development [6]. However, Laidlaw et al. [32] observed high IL-10-producing Tfr to support GCs responses in mouse models of acute lymphocytic choriomeningitis virus infection. Our findings extend the previous observations of a high Tfh in IM patients [29]. In this study, we also found a significant increase in cTfr in IM patients during primary EBV infection, indicating that an increased cTfr may be a marker of ongoing humoral activity, similar to other study [27, 28, 32]. Most importantly, we found that cTfr were positively correlated with CD19+IgD+CD27− naïve B cell and CD19+IgD−CD27hi plasmablasts. Therefore, in this observation study, these
Fig. 1 Elevated levels of cTfr and cTfh but low cTfr/cTfh ratio in IM patients. 

a Frequency in lymphocytes (left) and the number (right) of CD4+ T cells among HIs (n=21) and IM patients (n=55) at D0 and D15, 
b frequency in CD4+ T cells (top) and the number of cTfr cells (bottom) from the peripheral blood of HIs and IM patients at D0 and D15, 
c representative flow cytometric dot plots of cTfr on CD4+FoxP3+ T cells, 
d frequency in CD4+ T cells (left) and the number (right) of cTfh from the peripheral blood of HIs and IM patients at D0 and D15, 
e representative flow cytometric dot plots of cTfr on CD4+FoxP3+ T cells, 
f the cTfr/cTfh ratio in HIs and IM patients at D0 and D15. 

D0: at diagnosis; D15: day 15 after diagnosis.
Fig. 2 Positive correlation between cTfr and CD19+IgD+CD27− naive B cells, CD19+IgD+CD27hi plasmablasts or CD19+CD24hiCD27hi B cells in IM patients at D0. Correlation between the frequency and number of cTfr and CD19+ B cells (a, b), CD19+IgD+CD27− naive B cells (c, d), CD19+IgD+CD27+ memory B cells (e, f), CD19+IgD−CD27hi plasmablasts (g, h), CD19+CD24hiCD27hi B cells (i, j); correlation between the cTfr/cTfh ratio and frequency and number of CD19+ B cells (k, l), CD19+IgD+CD27− naive B cells (m, n), CD19+IgD+CD27+ memory B cells (o, p), CD19+IgD−CD27hi plasmablasts (q, r), CD19+CD24hiCD27hi B cells (s, t).
finding suggest high cTfr may inhibit B cell differentiation in the immune response to primary EBV infection. Further study is needed to investigate the function of this elevated Tfr.

CD19⁺CD24hiCD27hi B cells, a distinct subset of regulatory B cells (Bregs), could be potent suppressors of immunity which can suppress proinflammatory Th1/Th17 responses and induce Treg by production of IL-10 and/or TGF-β [33]. Li et al. [34] found that IL-10⁺ regulatory B cells were significantly higher following incubation with Tfr than with non-Tfr Treg, which suggested that Tfr were more potent at inducing IL-10⁺ regulatory B cells. Siwe et al. [35] found that high in IL-10⁺ regulatory B cells were shown to be positively correlated with HIV viral load in HIV-infected patients. Our data showed a positive correlation between cTfr and CD19⁺CD24hiCD27hi B cells. Most interesting, we found a negative correlation between cTfr or the cTfr/cTfh ratio and EBV DNA viral load. These results suggest that cTfr may induce the production of CD19⁺CD24hiCD27hi B cells, further lead to immune dysfunction during primary EBV infection and hinder EBV clearance.

Given that Tfh and Tfr are reciprocal and antagonistic regulators of GCs responses, a balance of Tfh and Tfr is critical for immune homeostasis. A disordered cTfr/cTfh ratio is associated with the development of infectious disease [12]. In this study, we found significantly low cTfr/cTfh ratio during primary EBV infection, which indicates an imbalance between cTfr and cTfh during primary EBV infection. During the course of an immune response to primary EBV infection, both cTfh and cTfr begin to proliferate; however, cTfh proliferate faster and skew the proportion in favor of helper capacity. By the D15, cTfr have return to normal, while cTfh proliferation continues, which cause a shift from immune tolerance state to immune responsive.

Our previous study has demonstrated that Tfh cells were positively correlated with EBV DNA load during primary EBV infection [29]. Most interestingly, we here also observed a negative correlation between cTfr or the cTfr/ cTfh ratio and EBV DNA viral load, which indicated that the viral control was probably hindered by the development of immunosuppressive Tfr cells. Therefore, we think the balance of cTfr and cTfh cells may play a role in controlling EBV and cTfr may play a role in controlling the Tfh response and limiting the spontaneous immune damage induced by excessive immunity during primary EBV infection. Our data showed a positive correlation between cTfr and CD19⁺IgD⁺CD27hi naïve B cell, CD19⁺IgD⁺CD27hi plasmablasts. These results suggest that Tfr contribute to
inefficient GCs responses and hinder EBV clearance, similar to the observation of Miles et al. [24].

Taken together, our study indicates that low cTfr/cTfh ratio may contribute to immunopathogenesis in the immune response to EBV infection and hinder EBV clearance. These results provide unique insight into the underlying immune response of primary EBV infection and suggest potential strategies for controlling EBV-related disease through suppressing cTfr.

Authors contribution QLN designed the experiments, and JQ wrote the manuscript. YQH and WMX conducted flow cytometry. ZYY identified the patients and obtained the samples. ZZ conducted RT-PCR and IgM test. All authors read and approved the final manuscript.

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Compliance with ethical standards

Conflict of interest The authors declare no conflicts of interest.

Informed consent Written informed consent was obtained from all enrolled patients and healthy individuals.

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References

1. Thorley-Lawson DA, Hawkins JB, Tracy SI, et al. The pathogenesis of Epstein–Barr virus persistent infection. Curr Opin Virol. 2013;3(3):227–232.
2. Babcock GJ, Decker LL, Volk M, et al. EBV persistence in memory b cells in vivo. Immunity. 1998;9(3):395–404.
3. Rickinson AB, Long HM, Palendiru U, et al. Cellular immune controls over Epstein–Barr virus infection: new lessons from the clinic and the laboratory. Trends Immunol. 2014;35(5):159–169.
4. Li ZX, Zeng S, Wu HX, Zhou Y. The risk of systemic lupus erythematosus associated with Epstein–Barr virus infection: a systematic review and meta-analysis. Clin Exp Med. 2019;19(1):23–36.
5. Coskun O, Sener K, Kilic S, Erdem H, Besirbellioglu AB, Gull HC, Egyiun CP. Stress-related Epstein–Barr virus reactivation. Clin Exp Med. 2010;10(1):15–20.
6. Longnecker R, Neipel F. Introduction to the human γ-herpesviruses. In: Human herpesviruses: biology, therapy, and immunoprophylaxis. 2007;Chapter 22.
7. Posnett DN. Herpesviruses and autoimmunity. Curr Opin Investig Drugs. 2008;9(5):505–514.
8. Szkaradkiewicz A, Kruk-Zagajewska A, Wal M, Jopek A, Wierzbiacka M, Kuch A. Epstein–Barr virus and human papillomavirus infections and oropharyngeal squamous cell carcinomas. Clin Exp Med. 2002;2(3):137–241.
9. Craft JE. Follicular helper t cells in immunity and systemic autoimmunity. Nat Rev Rheumatol. 2012;8(6):337–347.
10. Wing JB, Tekguc M, Sakaguchi S. Control of germinal center responses by t-follicular regulatory cells. Front Immunol. 2018;9:1910.
11. Miles B, Connick E. Control of the germinal center by follicular regulatory t cells during infection. Front Immunol. 2018;9:2704.
12. Fazilleanu N, Alouolu M. Several follicular regulatory t cell subsets with distinct phenotype and function emerge during germinal center reactions. Front Immunol. 2018;9:1792.
13. Sage PT, Sharpe AH. T follicular regulatory cells in the regulation of b cell responses. Trends Immunol. 2015;36(7):410–418.
14. Xie MM, Dent AL. Unexpected help: follicular regulatory t cells in the germinal center. Front Immunol. 2018;9:1536.
15. Niu Q, Huang ZC, Wu XJ, et al. Enhanced in-6/phosphorylated STAT3 signaling is related to the imbalance of circulating T follicular helper/T follicular regulatory cells in patients with rheumatoid arthritis. Arthritis Res Ther. 2018;20(1):200–208.
16. Chen M, Lin X, Cheukfai L, et al. Advances in t follicular helper and t follicular regulatory cells in transplantation immunity. Transplant Rev (Orlando). 2018;32(4):187–193.
17. Fonseca VR, Aguaz-Doce A, Maceiras AR, et al. Human blood Tfr cells are indicators of ongoing humoral activity not fully licensed with suppressive function. Sci Immunol. 2017;2(14):eaan1487.
18. Baptista D, Mach F, Brandt KJ. Follicular regulatory t cell in atherosclerosis. J Leukoc Biol. 2018;104(5):925–930.
19. He J, Tsai LM, Leong YA, et al. Circulating precursor CCR7(lo) Pd-1(hi) CXCR5(+) CD4(+) T cells indicate Thf cell activity and promote antibody responses upon antigen reexposure. Immunity. 2013;39(4):770–781.
20. Vinuesa CG, Cook MC. Blood relatives of follicular helper t cells. Immunity. 2011;34(1):10–12.
21. Sage PT, Alvarez D, Godec J, et al. Circulating t follicular regulatory and helper cells have memory-like properties. J Clin Investig. 2014;124(12):5191–5204.
22. Wu X, Su Z, Cai B, et al. Increased circulating follicular regulatory t-like cells may play a critical role in chronic hepatitis b virus infection and disease progression. Viral Immunol. 2018;31(5):379–388.
23. Wang L, Qiu J, Yu L, et al. Increased numbers of CD5+ CD19+ CD1d high IL-10+ Bregs, CD4+ Foxp3+ Tregs, CD4+ CXCR5+ Foxp3+ follicular regulatory T (Tfr) cells in CHB or CHC patients. J Transl Med. 2014;12:251.
24. Miles B, Miller SM, Folkvord JM, et al. Follicular regulatory T cells impair follicular T helper cells in HIV and SIV infection. Nat Commun. 2015;6:8608.
25. Botta D, Fuller MJ, Marquez-Lago TT, et al. Dynamic regulation of t follicular regulatory cell responses by interleukin 2 during influenza infection. Nat Immunol. 2017;18(11):1249–1260.
26. Dhaeze T, Peelen E, Hombrouck A, et al. Circulating follicular regulatory t cells are defective in multiple sclerosis. J Immunol. 2015;195(3):832–840.
27. Xu B, Wang S, Zhou M, et al. The ratio of circulating follicular t helper cell to follicular t regulatory cell is correlated with disease activity in systemic lupus erythematosus. Clin Immunol. 2017;183:46–53.
28. Liu C, Wang D, Lu S, et al. Increased circulating follicular treg cells are associated with lower levels of autoantibodies in patients
with rheumatoid arthritis in stable remission. Arthritis Rheumatol. 2018;70(5):711–721.
29. Liu J, Zhou Y, Yu Q, et al. Higher frequency of CD4+ CXCR5+ ICOS+ PD1+ T follicular helper cells in patients with infectious mononucleosis. Med (Baltim). 2015;94(45):e2061.
30. Iwata Y, Matsushita T, Horikawa M, et al. Characterization of a rare IL-10-competent B-cell subset in humans that parallels mouse regulatory B10 cells. Blood. 2011;117(2):530–541.
31. Blair PA, Norena LY, Flores-Borja F, et al. CD19(+) CD24(hi) CD38(hi) B cells exhibit regulatory capacity in healthy individuals but are functionally impaired in systemic lupus erythematosus patients. Immunity. 2010;32(1):129–140.
32. Laidlaw BJ, Lu Y, Amezquita RA, et al. Interleukin-10 from CD4+ follicular regulatory T cells promotes the germinal center response. Sci Immunol. 2017;2(16):eaan4767.
33. Carter NA, Vasconcellos R, Rosser EC, et al. Mice lacking endogenous IL-10-producing regulatory B cells develop exacerbated disease and present with an increased frequency of Th1/Th17 but a decrease in regulatory T cells. J Immunol. 2011;186(10):5569–5579.
34. Li H, Zhou R, Wang C, et al. T follicular regulatory cells infiltrate the human airways during the onset of acute respiratory distress syndrome and regulate the development of B regulatory cells. Immunol Res. 2018;66(4):548–554.
35. Siewe B, Stapleton JT, Martinson J, et al. Regulatory B cell frequency correlates with markers of HIV disease progression and attenuates anti-HIV CD8(+) T cell function in vitro. J Leukoc Biol. 2013;93(5):811–818.

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