IgE reactivity to α-Gal in relation to Lyme borreliosis

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Abstract

Background
An association between tick bites, the development of immunoglobulin E (IgE) antibodies to galactose-α-1, 3-galactose (α-Gal) and red meat allergy has recently been reported. Here we wanted to elucidate the relation between tick exposure, IgE antibodies to α-Gal and Lyme borreliosis (LB).

Methods
In the highly LB endemic area of Kalmar County, Sweden, serum samples and health inquiries from 518 blood donors were included. All sera were investigated for multiple IgG antibodies using a multiplex assay (recomBead, Mikrogen). In addition, three serially collected sera over a six month period from 148 patients with clinically defined erythema migrans (EM) were included. IgE antibodies against α-Gal were determined using ImmunoCAP (Thermo Fisher Scientific).

Results
In blood donors reporting previous LB (n = 124) IgE to α-Gal was found in 16%, while in donors denying previous LB but with multiple anti-Borelia antibodies (n = 94; interpreted as asymptomatic LB) 10% were IgE α-Gal-positive. Finally, in donors without Borelia antibodies denying previous LB (n = 300) 14% showed IgE to α-Gal. No significant difference in proportions among the groups were found. In EM patients, IgE to α-Gal was found in 32/148 (22%) at diagnosis, 31/148 (21%) after two-three months and 23/148 (16%) after six months. A significant reduction of proportion and level of IgE to α-Gal was found between the second and third sample (p < 0.01). A positive IgE anti α-Gal was more common among men compared with women both in blood donors and in EM patients (p ≤ 0.01).

Conclusions
IgE to α-Gal reactivity was common in a tick endemic area but showed no significant relation to previous LB. IgE anti-α-Gal reactivity in EM patients peaked within three months of
diagnosis of EM, after which it waned indicating that recent tick exposure is of importance in \( \alpha \)-Gal sensitization. Furthermore, IgE anti-\( \alpha \)-Gal was more common in men compared with women.

Introduction

In the last few years, an association between immunoglobulin E (IgE) antibodies directed towards the mammalian carbohydrate galactose-\( \alpha \)-1, 3-galactose (\( \alpha \)-Gal) and red meat allergy has been described [1–4]. The anti-\( \alpha \)-Gal response was first identified when a subset of patients treated with cetuximab (a chimeric mouse human monoclonal antibody targeting the epidermal growth factor) developed severe anaphylactic reactions at the first treatment [5]. Van Nunen and colleagues from Australia then documented an association between large local reactions to bites from the tick *Ixodes (I.) holocyclus* and adult onset red meat allergy and hypothesized a pathogenic relationship [6]. Later, Platt's Mills and co-workers from North America reported that the development of this specific antibody may be linked to the exposure of tick bites by the tick species *Amblyomma americanum* [7]. We have recently identified \( \alpha \)-Gal in the gastrointestinal tract of the principal European vector of Lyme borreliosis (LB), the tick *Ixodes (I.) ricinus* [8]. Furthermore, \( \alpha \)-Gal has also shown to be present in saliva of the ticks *Haemaphysalis longicornis* and *Amblyomma sculptum* [9, 10]. This and recent data about the prevalence of sensitization to \( \alpha \)-Gal [11, 12], strengthens the association between tick bites of various species and the development of IgE antibodies to \( \alpha \)-Gal and in turn red meat allergy [8, 13]. In addition, as \( \alpha \)-Gal is structurally related to the blood group B antigen, a protective effect of blood group B on the development of IgE anti-\( \alpha \)-Gal and red meat allergy has been reported [2, 14]. However, tick exposure is very common in many geographic areas including the south of Sweden and also in particular the Åland Islands in the Baltic Sea [15], but cases of red meat allergy are few in comparison [2]. Thus, determinants of which of the tick bitten individuals that will develop IgE anti-\( \alpha \)-Gal antibodies are still unknown. Similarly, which of the IgE \( \alpha \)-Gal-positive individuals that will develop clinical red meat allergy remains to be shown, although the level of IgE anti-\( \alpha \)-Gal seems to be of importance with cases of red meat allergy showing significantly higher IgE levels compared to \( \alpha \)-Gal positive healthy blood donors [2]. In parallel with red meat allergy, the risk of developing LB after a tick bite has been demonstrated to be small in Scandinavia, although *Borrelia* has been identified in some 26% of ticks [15, 16]. LB is the most common known tick borne infection in both Europe and North America, an infection that may present with various clinical signs and symptoms. The skin manifestation erythema migrans (EM) is the most common clinical manifestation of LB with a frequency of at least 70% of all clinical LB [17, 18]. Interestingly, an asymptomatic *Borrelia* infection, defined as a *Borrelia* IgM/IgG antibody seroconversion after tick bite in the absence of clinical symptoms, seems to be as common as developing clinical LB [15]. Our previous work revealed that 22% of patients with LB in the Stockholm area have positive IgE levels to \( \alpha \)-Gal at significantly higher frequency than healthy blood donors [2].

In this study, we investigated the IgE reactivity against \( \alpha \)-Gal and *I. ricinus* in relation to *Borrelia* IgG antibody status, blood group status, self-reported tick exposure and previous LB in Swedish blood donors collected in a LB endemic area, enabling the discrimination of previous clinical LB and asymptomatic LB. Furthermore, we wanted to elucidate the IgE anti-\( \alpha \)-Gal...
status in patients with EM in acute phase as well as in convalescent samples in order to determine the role of time in relation to IgE anti-α-Gal.

**Material and methods**

**Study subjects**

**Blood donors.** A total of 1126 healthy blood donors in Kalmar County in Sweden were consecutively included in an adjacent clinical study regarding asymptomatic LB in the spring of 2012, when scheduled for routine blood donation. Blood donor sera were collected together with health inquiries including information on previous tick exposure and previous history of LB. A complete set of health inquiry and blood sera was obtained from 1113 blood donors. Blood donors were divided in three groups according to whether they answered yes, no or undetermined to previous history of LB. To ensure a higher level of specificity, blood donors answering yes to previous history of LB were also required to have a clinical diagnosis to be included in the group.

All blood donor sera were screened for specific IgG antibodies against 13 different borrelial antigens using a commercial multiplex kit, recomBead Borrelia IgG (Mikrogen GmbH, Neuried, Germany). Analyses were performed according to the manufacturer’s instructions, whereby the results of the individual antigen reactivities were added to a final sum using a scoring system. A total sum of eight points and more were regarded as a positive test result, according to the manufacturer’s instructions.

The following selection criteria were used for selection of three subgroups of blood donors investigated in this study: All blood donors reporting previous clinical LB confirmed by medical doctor’s diagnosis, n = 124 (previous LB). All blood donors denying previous LB but with a positive test result in the multiplex Borrelia IgG antibody test, n = 94 (previous asymptomatic LB). Finally, a group of blood donors, n = 300, denying previous LB and with a negative anti-Borrelia antibody test (no known previous Borrelia exposure) was included in the following manner: The first 30 consecutively included blood donors of women and men in the following age groups were selected; 18–29, 30–39, 40–49 and 50–59 years of age. Regarding the age group of 60 years and more only 22 women were available, therefore an additional eight men were included in this age group, thereby consisting of 38 men. In total, 518 blood donors in different subgroups with serum samples and complete health inquiry results were included in this study. The blood donor serum samples had been stored at −20°C and freeze-thawed once prior to the analyses in this study.

Blood group results, AB0 and RhD, were collected from the blood donor journal system ProSang 2008 (Databyrån, Stockholm, Sweden), and the expected AB0 blood group distribution nationally was obtained from official data [19].

**Erythema migrans patients.** In a separate prospective study of patients with LB in Kalmar County in Sweden during 2003, clinical data was recorded in a study protocol and serum samples were drawn at the first consultation, after two to three months and finally after six months. EM was diagnosed by a medical doctor by typical clinical skin features and >5 cm in diameter. All EM serum samples were initially analysed for anti-Borrelia antibodies using enzyme-linked immunosorbent assay detecting C6 peptide antibodies (non-resolving assay for IgM and IgG antibodies) with analyses performed and results interpreted according to the manufacturer (optical density value >0.15) (Immunetics®, Boston, MA, USA). Serum samples from a total of 148 EM patients, each with a set of three serum samples, were available for this study [20, 21]. The serum samples from EM patients had been stored at −20°C and freeze-thawed twice before the analyses in this study.
IgE assays

IgE antibodies against α-Gal were determined by ImmunoCAP (Phadia AB/Thermo Fisher Scientific, Uppsala, Sweden) for all included blood donors as well as EM patients at all three time points. Blood donor samples and EM samples IgE positive to α-Gal (≥ 0.1 kU/L) were further investigated for IgE against I. ricinus using 5 mg of biotinylated I. ricinus antigen coupled to Streptavidin ImmunoCAP as previously described [2, 8]. Total IgE and Phadiatop®!, a mix of common inhalant allergens, (ImmunoCAP, Phadia AB/Thermo Fisher Scientific) assays were also performed in the first serum sample of all EM patients in order to assess atopy. IgE levels ≥ 0.1 kU/L against I. ricinus and ≥ 0.35 kU/L against Phadiatop®! were considered positive. All IgE analyses were performed in accordance with the instructions of the manufacturer at the Department of Clinical Immunology, Karolinska University Hospital Solna, Stockholm, Sweden.

Statistics and ethics

Statistical analyses were performed using Fisher’s exact two-tailed test for proportions and the non-parametric Mann-Whitney’s U-test for comparing age between groups. An exact test corresponding to McNemar’s paired chi-squared modification, first proposed by Liddell (1983) was used for paired measurements of proportions [22]. The Friedman ANOVA and Wilcoxon paired test were used when comparing IgE anti-α-Gal seropositivity levels in EM patients over time (Statistica 12). Comparison of total IgE and anti-α-Gal specific IgE was performed with Spearman rank correlation using Prism 7 (GraphPad Software, La Jolla, Calif). A p<0.05 was considered statistically significant. The study was approved by the regional ethical review board of Linköping University, Sweden. Written consent was obtained from the study participants.

Results

Clinical features

Descriptive data regarding sex and age distributions and self-reported tick exposure among all studied blood donors taken together (n = 518) and divided into the three identified subgroups (donors with previous LB, donors previously asymptomatically Borrelia infected and donors without known Borrelia exposure) are shown in S1 Table. In 105 of 518 (20%) blood donors the average number of tick bites annually, counting the five last years preceding the inclusion in the study in 2012, was reported to be five bites or more. Self-reported tick bites of at least five bites or more annually did not differ between women 52/234, 22% and men 53/284, 19% (data missing for two men), p = non-significant. Corresponding data regarding sex, age, duration of EM at diagnosis and self-reported previous LB for the EM patients are shown in S2 Table (n = 148). The median duration of EM at diagnosis, i.e. at the first serum sampling time, was seven days. In 46/148 (31%) of the EM patients at least one previous episode of LB was reported and they were assessed as subjects exposed to LB re-infection.

Allergen-specific IgE

Blood donors. The prevalence of IgE reactivity to α-Gal among the blood donors was 14% (71/518) with a median level of 0.30 kU/L (range 0.10–17 kU/L). Divided into subgroups according to previous Borrelia status, the rate of IgE positivity to α-Gal varied between 10–16% (Table 1). A comparison between blood donors IgE positive and negative to α-Gal revealed a male dominance among IgE positive donors, p = 0.01 (Table 2). No significant difference in tick exposure, previous LB or anti-Borrelia antibody status was shown comparing
IgE anti-α-Gal positive and negative donors. In 22 of the 71 (31%) IgE α-Gal-positive samples a positive (≥0.1 kU A/L) level of IgE to *I. ricinus* was found, with a median level of 0.54 kU A/L (range 0.10–3.8 kU A/L).

Blood group distributions among the blood donors according to IgE anti-α-Gal status are shown in S3 Table including the expected distribution of blood groups in the population in Sweden for comparison [19]. Among blood donors IgE positive to α-Gal, five (7.0%) were blood group B (including blood groups B+, AB+, B− and AB−) compared to α-Gal-negative blood donors where 60/447 (13%) were B-positive (p = 0.18). Median positive IgE reactivity to α-Gal among blood group B was 0.87 kU A/L and among B-negative 0.32 kU A/L (p = 0.50).

**Erythema migrans patients.** A total of 36/148 (24%) EM patients were IgE positive to α-Gal in at least one of the three samples. At the EM diagnosis, 32 of 148 (22%) patients were IgE positive (≥0.1 kU A/L) to α-Gal, at two to three months 31/148 (21%) and finally 23/148 (16%) were IgE positive at six months after diagnosis, Table 3. Comparing levels of IgE anti-α-Gal among those 36 individuals positive in at least one sample, a significant decrease was noted between the first and third sample as well as between the second and third samples (p < 0.01), Fig 1. The difference was also confirmed by paired measurement of proportions between the second and third sample Table 3 (p < 0.01). In Table 3 proportions of IgE to *I. ricinus*-positive samples among IgE anti-α-Gal-positive samples are also given together with levels of total IgE and the Phadiatop positive rate (26/148; 18%) for EM patients examined at diagnosis.

### Table 1. IgE anti-α-Gal positive rate in blood donors according to *Borrelia* status.

|                      | IgE α-Gal positive (≥ 0.1 kU A/L), numbers (%) |
|----------------------|-----------------------------------------------|
| All blood donors (n = 518) | 71 (14)                                      |
| Donors with previous LB (n = 124) | 20 (16)                                      |
| Donors previously asymptotically *Borrelia* infected (n = 94) | 9 (10)                                       |
| Donors without known *Borrelia* exposure* (n = 300) | 42 (14)                                      |

n = numbers
LB = Lyme borreliosis
* Donors denying previous LB also lacking serum anti-*Borrelia* antibodies

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### Table 2. Basic clinical data on blood donors according to IgE anti-α-Gal status.

| IgE α-Gal positive (≥ 0.1 kU A/L) blood donors (n = 71) | Female/Male, numbers (% females) | Median age, (range in years) | Selfreported tick exposure > 5 bites annually, numbers (%) | Previous LB, numbers (%) | Positive anti-*Borrelia* antibody test*, numbers (%) |
|-------------------------------------------------------|----------------------------------|------------------------------|---------------------------------------------------------------|--------------------------|-----------------------------------------------|
| 22/49* (23)                                           | 53 (18–69)                       | 15c (21c)                   | 20 (28)                                                      | 11 (15)                  |
| IgE α-Gal negative (<0.1 kU A/L) blood donors (n = 447) | 212/235* (47)                    | 46 (18–72)                  | 90c (20c)                                                    | 104 (23)                 | 103 (23)                                       |

LB = Lyme borreliosis
n = numbers
* recomBead Borrelia IgG
* p = 0.01
* Data missing on one blood donor

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EM patients were also divided according to IgE anti-α-Gal status, see Table 4. A positive IgE anti-α-Gal result was more common among EM men compared with women ($p < 0.001$), similar to the results found in blood donors, Table 2. No significant differences were shown comparing median age, Phadiatop status, anti-Borrelia antibody status or reported previous LB between IgE anti-α-Gal positive and negative EM patients (Table 4). In addition, the individual IgE anti-α-Gal levels in the 36 EM patients IgE positive to α-Gal at any sample time are

![IgE α-Gal positive levels in Erythema migrans patients (n=36)](https://doi.org/10.1371/journal.pone.0185723.g001)

**Fig 1.** IgE levels against α-Gal at the three different time points (at diagnosis = “0 months”; after two to three months and six months) among the 36 α-Gal IgE positive erythema migrans patients. The levels were compared between the different time points. A significant reduction in IgE anti-α-Gal levels from both the first and second sample to the third sample ($p<0.01$).

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**Table 3.** IgE anti-α-Gal positive, total IgE and Phadiatop status in erythema migrans patients.

| IgE α-Gal-positive (≥ 0.1 kU/L) | Erythema migrans patients (n = 148) | p-value* |
|---------------------------------|-----------------------------------|----------|
|                                 | At diagnosis, numbers (%)         | 2–3 months, numbers (%) | 6 months, numbers (%) |
| IgE α-Gal-positive samples, median reactivity (range) (kU/L) | 32 (22) | 31 (21) | 23 (16) | <0.01 |
| IgE to *Ixodes ricinus* (≥ 0.1 kU/L) | 0.55 (0.13–23) | 0.41 (0.10–15) | 0.54 (0.10–4.8) |
| Phadiatop positive (≥0.35 kU/L) | 10/32b (31b) | 13/31 (42) | 11/23 (48) |
| Erythema migrans patients (n = 148) | 26d (18d) |

**n = numbers**

* No significant difference at diagnosis compared with 2–3 months, but significant reduction between 2–3 months and 6 months, paired measurement

b One sample missing

c Nine samples missing

d Eight samples missing

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shown in Fig 1. Finally, a significant positive correlation between total IgE and IgE anti-α-Gal levels was noted (Fig 2).

**Discussion**

The present study of a large number of sera from blood donors sub-grouped according to previous LB status and EM patients followed over time shows that IgE anti-α-Gal is commonly

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**Table 4. Comparison of IgE α-Gal positive and negative erythema migrans patients.**

|                         | IgE α-Gal positive erythema migrans patients at diagnosis (n = 32), numbers (%) | IgE α-Gal negative erythema migrans patients at diagnosis (n = 116), numbers (%) | p-value |
|-------------------------|----------------------------------------------------------------------------------|----------------------------------------------------------------------------------|---------|
| Female/Male             | 11/21 (34/66)                                                                   | 79/37 (68/32)                                                                   | <0.001a |
| Median age, range (years)| 58 (7–78)                                                                       | 58 (16–84)                                                                      | NSb     |
| Phadiatop IgE positive  | 6c (19c)                                                                        | 20c (17c)                                                                       | NSa     |
| Anti-Borrelia antibody positive | 7 (22)                                                                           | 46 (40)                                                                          | 0.09a   |
| Previous LB             | 5d (16d)                                                                        | 41e (35e)                                                                        | 0.076a  |

n = numbers

LB = Lyme borreliosis

a Fisher’s exact two-tailed test

b Mann-Whitney U-test

c Four samples missing

d Unknown for four patients

e Unknown for one patient

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Correlation of total and α-Gal IgE titers in Erythema migrans patients (n=27)

![Correlation graph](https://doi.org/10.1371/journal.pone.0185723.g002)
found in an LB endemic area with a frequency of 10–22% with a male predominance. Furthermore, the level of IgE anti-α-Gal in α-Gal positive cases decreases over time in EM patients.

Previously, 10% of blood donors in the Stockholm area in Sweden have been shown to be IgE positive to α-Gal, thus in the same order of magnitude as found in this study with 14% [2]. For further comparison in the general population, IgE to α-Gal was found in 5.5% in Denmark and in 8.1% in Spain [12]. Considering the variation in tick abundance over different regions and geographical areas in Europe, variations in IgE anti-α-Gal reactivity rates are to be expected [23]. However, although tick exposure seems to be an important risk factor for developing IgE anti-α-Gal, sensitization may also be related to cat ownership [12]. In the present materials investigated, no data regarding pet ownership was available. In Sweden, the highest incidence of LB has been reported in Kalmar County, the county in which the samples in this study were collected [24]. As there is no systematically gathered data regarding tick exposure in the general population in Sweden, LB incidence may indicate tick prevalence to some extent. Thus, we would expect high rates of IgE α-Gal sensitization in Kalmar County in Sweden, and the 14% we found seems to be in line with this reasoning. However, LB incidence is of course also affected by a number of other variables including Borrelia abundance in ticks as well as behaviour regarding activities associated with tick exposure and tick removal routines in the population. Therefore, the interpretation of sensitization in relation to tick exposure should be made with caution. Interestingly, a considerable proportion of the blood donors in this study reported frequent tick bites, but no difference in the proportion of multiple tick bites annually was found comparing IgE α-Gal negative with positive donors. One possible explanation for this may be the sampling time; the blood donor samples were collected during the winter with an expected low tick activity in the environment. Thus, tick exposure in the blood donor group generally consisted of exposure the previous summer-autumn and historically. In contrast for the EM group, a recent tick bite must have taken place in order for the EM to occur. Therefore, our data suggests that recent tick exposure may give rise to α-Gal sensitization with decreasing titers over time approaching the general sensitization level in the population of that geographical area. In this study, 22% of the EM patients showed positive IgE levels to α-Gal at diagnosis, in fact the same level as reported from the Stockholm area [2]. Regarding sensitization depending on previous symptomatic or asymptomatic LB, no significant difference was shown. Thus, this does not seem to be a factor that determines which individuals that will develop IgE anti-α-Gal.

In addition to the association between IgE anti-α-Gal and timing of tick exposure, we found more men than women to be sensitized to α-Gal both among blood donors and EM patients. The significance of that finding remains to be elucidated, but a similar difference was also shown in Denmark but not in Spain [12]. The same sex difference has also been shown for anti-Borrelia antibodies, and may reflect a difference in tick exposure and removal strategies between men and women [25]. Data suggests that men may remove ticks later than women thus permitting greater time for exposure to tick antigens and pathogens [15].

In harmony with previous data, where a significant correlation between IgE anti-α-Gal and total IgE levels among red meat allergic patients was shown, a similar correlation was observed in this study [2]. Whereas the majority of patients with red meat allergy also showed IgE antibodies to I. ricinus, we could only demonstrate that co-sensitization in some 31–48% of the IgE anti-α-Gal-positive EM patients and blood donors. However, it is important to bear in mind that no data regarding red meat allergy was available in the present material in contrast to the study by Hamsten et al. in which red meat allergy patients were investigated.

Previously it has been shown that blood group B protects from red meat allergy, however it does not seem to inhibit from sensitization to α-Gal [2]. In the study by Hamsten et al. 13% of α-Gal-positive blood donors were blood group B and 22% of patients with a positive Borrelia
serology. In our study 7% of α-Gal-positive blood donors were blood group B, thus in line with previous findings.

The strengths of this study include the prospective design and collection of patients and blood donors together with standardised inquiry data, the sample size and clinical data regarding LB including the serially followed EM patients. The major limitation being the lack of data regarding red meat reactions and allergy.

Conclusions

In conclusion, this study confirms that IgE reactivity to α-Gal is commonly found in blood donors in an LB endemic area, but with no significant relation to previous LB, self-reported tick exposure or Borrelia antibody status. However, the IgE response to α-Gal showed a time dependent pattern in EM patients with reactivity peaking within three months of diagnosis, then declining. This indicates that recent tick exposure is of importance in sensitization to α-Gal. Furthermore, IgE anti-α-Gal was more common in men compared with women in both blood donors and EM patients, but the significance of this finding in relation to red meat allergy remains to be shown.

Supporting information

S1 Table. Basic descriptive data on blood donors and subgroups according to Borrelia status. (XLSX)

S2 Table. Basic descriptive data on erythema migrans patients. (XLSX)

S3 Table. Blood group distribution in blood donors according to IgE anti-α-Gal status. (XLSX)

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