ORIGINAL ARTICLE

Association study of the C3 gene with adult and childhood asthma

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Received: 29 November 2007/Accepted: 16 May 2008/Published online: 20 June 2008 © The Japan Society of Human Genetics and Springer 2008

Abstract Bronchial asthma (BA) is a multifactorial disorder, the development of which is affected by both environmental and genetic factors. The complement system plays an important role in immunological response against invading microorganisms. It has been shown that complement-C3-deficient mice have reduced inflammation of asthmatic airways. Previously, we reported the association of four single nuclear proteins (SNPs) in the exons of the *C3* gene with childhood and adult BA. The *C3* gene, however, is a large gene, and functional SNPs associated with susceptibility to BA have not yet been identified. We analyzed

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A. Miyatake Miyatake Asthma Clinic, Osaka, Japan 26 SNPs in the C3 gene and its promoter region to narrow down the regions showing association with childhood and adult BA. Childhood and adult atopic BA patients and healthy child and adult controls were recruited from urban cities in Japan and genotyped. In SNP analysis, an SNP (SNP24, rs11569562) located in intron 31 of the C3 gene was associated with adult BA [corrected $P(P_{cor}) = 0.030$]. In linkage disequilibrium (LD) block 4 spanning exons 24-41, the frequency of the CCC haplotype in adult BA was significantly higher than that in adult controls (P_{cor} = 0.038). Neither the SNP nor the haplotype showing association with adult BA demonstrated a significant association with serum total immunoglobulin E (IgE) level in BA patients and controls. Our results suggest that LD block 4 confers susceptibility to adult BA with mechanisms relevant to the effector phase of allergic inflammation.

Keywords Complement \cdot *C3* gene \cdot Asthma \cdot SNP \cdot Haplotype \cdot Association \cdot Total IgE

Introduction

Bronchial asthma (BA) is an inflammatory airway disease, the development of which is affected by both environmental and genetic factors (Barnes and Marsh 1998). The complement system belongs to the groups of ancient pattern-recognition systems and sensing exogenous (microorganisms) and endogenous (altered-self) danger-associated molecular patterns (Kohl and Wills-Karp 2007). The three pathways of complement activation are the classical, alternative, and lectin pathways, and all complement activation pathways converge at C3. On activation, C3 breaks into a large C3b fragment and a small C3a fragment that is called anaphylatoxin. C3b can form C5 convertase and activate another component of the complement, C5. C5 is cleaved into C5a anaphylatoxin and C5b, and both fragments exert physiological roles. Both anaphylatoxins also show proinflammatory and immunoregulatory actions. It has been shown that complement-C3-deficient mice have reduced inflammation of asthmatic airways (Barrington et al. 2001; Kohl 2001; Muller-Eberhard 1988). Mice and guinea pigs lacking the C3a receptor (C3aR) essential for C3a action were protected against bronchoconstriction, airway hyperresponsiveness (AHR), and airway inflammation developing after allergen challenge (Bautsch et al. 2000; Drouin et al. 2002; Humbles et al. 2000). Compared with wild-type mice, C3-deficient mice also exhibit diminished AHR and lung eosinophilia when challenged with an allergen (Drouin et al. 2001). They also showed decreased numbers of interleukin (IL)-4-producing lung cells and decreased serum-antigen-specific immunoglobulin E (IgE) levels. Dendritic cells (Zhou et al. 2007) and activated T cells express C3a receptor (Werfel et al. 2000), and C3 and C3a have been shown to regulate interactions between dendritic and T cells (Kawamoto et al. 2004; Zhou et al. 2007). These results suggested that the C3 pathway is involved in allergen sensitization. In BA patients, increased levels of C3a (Humbles et al. 2000; Krug et al. 2001) and C5a (Krug et al. 2001) were observed in bronchoalveolar lavage fluid after segmental allergen provocation, suggesting involvement of the complement system in allergen-induced airway inflammation in humans. Elevation of plasma C3a level was observed in BA patients on acute exacerbation (Nakano et al. 2003). These data are consistent with the notion that the C3 gene plays an important role in allergic sensitization and allergic inflammation. Furthermore, the C3 gene is located at chromosome 19p, where several studies suggested linkage to BA and related phenotypes (Blumenthal et al. 2004; Lee et al. 2000; Venanzi et al. 2001). Therefore, the C3 gene is a strong candidate gene for allergic BA.

Our previous analysis of four single nucleotide polymorphisms (SNPs) in exons of the *C3* gene suggested that variations of this gene conferred susceptibility to both childhood and adult BA (Hasegawa et al. 2004). Barnes et al. reported that a haplotype spanning introns 19–23 showed significant association with BA, log (total IgE) and log [interleukin (IL)-13]/log [interferon (IFN)- γ] (Barnes et al. 2006). The *C3* gene, however, is a large gene consisting of 41 exons and containing hundreds of SNPs; thereby, further investigation is necessary to identify functional SNPs conferring susceptibility to BA. In this study, we analyzed 26 SNPs of the *C3* gene to narrow down the regions showing association with childhood and adult BA.

Methods

Subjects

All participants were Japanese. Three hundred and fortysix childhood BA patients, 518 adult BA patients, and 550 healthy adult controls were recruited in Osaka City area. Japan. Details of these patients are described in a previous report (Nakashima et al. 2006). All participants with BA were diagnosed and selected by physicians according to the American Thoracic Society (ATS) criteria and using questionnaires based on the recommendation of the ATS, Division of Lung Disease (ATS-DLD) (Ferris 1978). In brief, patients showed repeated episodes of at least one of the following symptoms: cough, wheezing, shortness of breath, chest tightness, and sputum production. Spirometry was performed in all patients to confirm the obstructive pattern of the lung function and response to a bronchodilator. Improvement of their forced expiratory volume in 1 s (FEV₁) measurement was at least 12% after β_2 agonist inhalation. The diagnosis of atopic BA was based on one or more positive skin-scratch-test responses to a range of seven common allergens in the presence of a positive histamine control and a negative vehicle control. The seven allergens were house dust, Felis domesticus dander (Feld), Canis familiaris dander, Dactylis glomerata, Ambrosia, Cryptomeria japonica, and Alternaria alternata. The numbers of atopic childhood and adult BA were 304 and 371, respectively (Table 1). In this study, we only analyzed atopic BA patients. As child control subjects, 411 child volunteers with ages between 6 and 12 years (male:female = 1.0:1.04) were recruited in Chiba City, Japan. Total and eight specific IgE levels in serum were measured in this group (Dermatophagouides pteronyssinus, C. familiaris, F. domesticus, A. alternata, C. japonica, D. glomerata, egg white, golden/black bellied/Hungarian hamster). Questionnaires based on the International Study of Asthma and Allergies in Childhood (Asher et al. 1995) were used to exclude children with BA and/or atopic dermatitis. Three

Table 1 Clinical characteristics of patients and controls

	Childhood asthma	Adult asthma	Child control	Adult control
Number	304	371	333	550
Age mean (years)	9.69	45.13	9.22	44.57
Age range (years)	4-15	20-75	6–12	20-75
Gender (male:female)	1.54:1.0	1.11:1.0	1.0:1.04	2.81:1.0
Atopic asthma (%)	100	100		
Mean total IgE [log (IU/ml)]	2.66	2.54	1.90	ND

IgE immunoglobulin E, ND not determined

hundred and thirty-three children were used as a child control group in association studies. If atopy was defined as those who showed positive specific IgE (>0.35 IU/ml) to one or more allergens, 71% of the child controls were atopic. Details of the adult controls were described previously (Nakashima et al. 2006). Adult volunteers were interviewed by physicians, and those who were diagnosed as having BA, atopic dermatitis, and/or allergic rhinitis were excluded from the adult control group. Measurement of serum IgE levels and skin-prick tests were not performed in the adult controls. All patients and volunteers provided written informed consent to participate in the study in accordance with the rules of the process committee at the SNP Research Center (RIKEN). This study was approved by the ethics committee of Chiba University Graduate School of Medicine.

Screening and selection of SNPs

We previously investigated four SNPs in the exons of the C3 gene (Hasegawa et al. 2004): 912G/A, 1692AG, 1936GA, and 4896 CT in our previous paper, which correspond to SNP6, SNP7, SNP13, and SNP25, respectively, in this paper (Table 2). SNP25 showed association with mite-positive childhood BA and adult BA. Because SNP25 is located in exon 41, the last exon of the C3gene, SNPs located around this exon were intensively investigated (SNP17-SNP26). We searched the dbSNP database to list up SNPs with the following criteria: minor allele frequency was more than 0.3, and distance of the nearest two SNPs did not exceed 2 kbp. Considering the potential importance of the gene's promoter region for gene expression, where several regulatory elements have been reported, we searched SNPs up to 5 kbp upstream of exon 1. Four SNPs (SNP1-SNP4) were identified. In addition, we included SNP14, SNP15, and SNP16 in this study to confirm the results of Barnes et al. (2006). To estimate the linkage disequilibrium (LD) status of these SNPs, we genotyped them in 96 adult controls. Haploview 3.32 program (Barrett et al. 2005) was used to show an LD map. LD block was defined with the solid spine of the LD method implemented in the Haploview program. Of the 26 SNPs investigated, 24 composed four LD blocks (Fig. 1). We selected tag SNPs from each LD block with the aid of the Tagger routine incorporated in the Haploview program.

Genotyping

Genomic DNA was prepared from whole blood samples using a standard protocol. Whole genome amplification was performed using the illustra GenomiPhi V2

 Table 2
 Locations and allele frequencies of single nucleotide polymorphisms (SNPs) of the C3 gene

SNP	Location	Allele (1/2) ^a	Position ^b	Minor allele frequency (%) ^c	dbSNP number
SNP1	Promoter	T/C	6678365	0.330	rs171094
SNP2	Promoter	C/T	6674037	0.157	-
SNP3	Promoter	A/G	6673635	0.298	rs163913
SNP4	Promoter	A/C	6673022	0.306	rs339392
SNP5	Intron 2	A/G	6669534	0.270	rs2250656
SNP6	Exon 9	G/A	6663291	0.426	rs2230201
SNP7	Exon 14	G/A	6669848	0.468	rs2230204
SNP8	Exon 14	G/A	6663704	0.414	rs2230205
SNP9	Intron 14	G/A	6660074	0.148	rs11569429
SNP10	Intron 14	G/A	6660050	0.016	rs4807984
SNP11	Intron 17	A/G	6656246	0.403	rs11672613
SNP12	Intron 19	A/C	6648829	0.134	rs366510
SNP13	Exon 21	C/T	6648406	0.064	rs423490
SNP14	Intron 23	C/T	6647342	0.128	rs2287848
SNP15	Intron 23	C/T	6647178	0.371	rs10410674
SNP16	Intron 23	C/G	6646001	0.435	rs10402876
SNP17	Intron 33	G/A	6634846	0.436	rs344549
SNP18	Intron 33	C/G	6633953	0.441	rs344550
SNP19	Intron 33	T/-	6633534	0.446	rs11569553
SNP20	Intron 35	A/G	6631937	0.468	rs344552
SNP21	Intron 35	C/A	6631928	0.394	rs344553
SNP22	Intron 36	A/G	6630563	0.457	rs2277983
SNP23	Intron 36	G/A	6630511	0.447	rs2277984
SNP24	Intron 38	T/C	6629753	0.456	rs11569562
SNP25	Exon 41	C/T	6628989	0.447	rs4807893
SNP26	3' Downstream	T/G	6627442	0.414	rs379527

^a Base expressed in the direction the gene

^b Based on National Center for Biotechnology Information (NCBI) Build 35.1 reference group label

^c In 96 adult control subjects

amplification kit (GE Healthcare, Buckinghamshire, UK) according to the manufacturer's standard protocol. Amplified deoxyribonucleic acid (DNA) was typed by allele-specific polymerase chain reaction (AS-PCR) using either the modified TaqMan AS amplification (TaqMan-ASA) method (Fujii et al. 2000) or SYBR Green detection. The primer and TaqMan probe sequences are shown in Table 3. For the TaqMan-ASA method, $2\times$ Platinum qPCR SuperMix-UDG (Invitrogen, Carlsbad, CA, USA) was used as master mix, whereas $2\times$ Platinum SYBR Green qPCR SuperMix-UDG (Invitrogen) or $2\times$ SYBR Green Supermix (Bio-Rad, Hercules, CA, USA) (for only SNP24) was used for AS-PCR together with SYBR Green detection. For the ASA method, the PCR mixture contained 5 µl of $2\times$ PCR master mix, 0.4 µM of each PCR



Fig. 1 Structure of the *C3* gene and location of single nucleotide polymorphisms (SNPs) and their linkage equilibrium (LD) map. Exons are indicated by *black boxes*. *Numbers* starting with *E* stand for exons and those starting with *S* for SNPs. Pair-wise LD (r^2) was

primer, 0.12 µM of the TaqMan probe (when needed), and 5 ng of amplified template DNA in a final volume of $10 \ \mu$ l. The samples were analyzed using an ABI PRISM 7000 Sequence Detector System (Applied Biosystems, Foster City, CA, USA) and Chromo4 Real-Time System (Bio-Rad). The thermoprofile was 50°C for 2 min, 95°C for 2 min, then 45 cycles of 95°C for 15 s and 60°C for 30 s (for SNP4: 45 cycles of 95°C for 15 s, 55°C for 30 s, and 60°C for 30 s). For SNP2, PCR was initially performed using 0.2 µM of each first PCR primers and Taq DNA polymerase (Promega, Madison, WI, USA) according to the manufacturer's standard protocol. The thermoprofile of the first PCR was 95°C for 3 min, followed by 20 cycles at 95°C for 20 s, 60°C for 30 s, and 72°C for 7 min. In the second PCR (TaqMan-ASA method), a reaction mixture of the first PCR diluted tenfold with water was used as a template. Because an SNP was located within the sequence where the AS primer for SNP9 was designed, two primers were mixed and used in the reaction mixture for SNP9 (Table 3).

estimated from 96 control subjects. LD blocks were defined by the solid spine of LD using the Haploview program. *Number* in each cell represents r^2 (×100); *black cells* r^2 = 1; *white cells* r^2 = 0. Each cell is colored in a graduated manner according to the strength of LD

Statistical analysis

Pairwise LD in SNPs was estimated as r^2 (Hill and Robertson 1968). A 2 \times 2 contingency χ^2 test of independence was performed to evaluate the significance of an association between allele frequency and disease status. Haplotype inference and a case-control association study were performed using SNPAlyze ver. 4.1 program (DYNACOM, Mobara, Japan). The effects of genotypes on log₁₀-transformed total serum IgE levels were evaluated using analysis of variance (ANOVA). Statistical analysis was performed with SPSS software (ver. 15.0 J; SPSS Japan, Tokyo, Japan). In SNP association studies, corrected P (P_{cor}) values were calculated by multiplying the number of SNPs tested in this study (15). In haplotype association studies, P_{cor} values were calculated by multiplying the total number of haplotypes tested in the entire gene (27). Association of an SNP or haplotype with the disease was judged as significant if P_{cor} for the test of allele/haplotype frequency was <0.05.

Table 3 Primers for genotyping single nucleotide polymorphisms (SNPs) in the C3 gene

in the C3 gene		SNP genotyped	Sequence
SNP genotyped	Sequence	Sivi genotyped	Sequence
SNP1		SNP11	
F: T allele	5'-GGCATAAACAAGGGTTAAAATGT	F: A allele	5'-AGGGTCACTGGGAAAATTAGACATA
F: C allele	5'-GGCATAAACAAGGGTTAAAATGC	F: G allele	5'-AGGGTCACTGGGAAAATTAGACAGG
R: common	5'-GCTCACAAACGCCTATGA	R: common	5'-ATGGGCCAAAGGAATTACACAAT
TaqMan probe	5'-TGAAATAGATAAGTTGCTGCCACCCG	SNP12	
SNP2		F: A allele	5'-ACTECCEGACETTEACACTAC
F: C allele	5'-GCCTGGCCAACATGGCGAACC	F: C allele	5 - ACTECCUACCITUACACTAC
F: T allele	5'-GCCTGGCCAACATGGCGAAGT	K. common	5' CCCGACCACCGATCTCTCTCCCCA
R: common	5'-TGCCTCCTGGGTTCAAGTGATTCTC	SND12	5-CCC0AGCAGOGATCT010100CA
TaqMan probe	5'-TAGTGGCGCATGCCTGTGATCCCAGCTCT	SINF 15	5' GGA AGTGGA AGTCA AGGCTGGC
First PCR F:	5'-TTCCAAAGAGTGTGTCGCAA	F. C allele	5' GGAAGTGGAAGTCAAGGCTGGT
First PCR R:	5'-CCTGCTTCATAGAGTTGTCGT	P: common	5' GGGTGCCCCAAGCACTCA
SNP3		K. common	5' CONTRECECTANCE ACTE ACCEPTER ACCEPTE
F: A allele	5'-TTTGGCAATATCTAGCAAGATTACCTA	SND14	J-CEATCATTEATEATUACUUTUTEAUUAA
F: G allele	5'-TTTGGCAATATCTAGCAAGATTACCTG	D. T ollele	5' GAATGAGATGGAATTTGGCTCGA
R: common	5'-CCTTACCACCTGCTTCATAGAGTTG	R. T allele	5' GAATGAGATGGAATTTGGCTCTG
TaqMan probe	5'-TGACCCAACATATTTCCTTTCATTGCAACG	K. C allele	
SNP4		SNP15	J-CAAUTECEUUAEAEEUAUTET
F: C allele	5'-CCTGTAACCTGTAAGAATGAGAC	R: C allele	5'-CAGCGAGCTGAGGTCGGG
F: A allele	5'-CCTGTAACCTGTAAGAATGAGTA	R: T allele	5'-CAGCGAGCTGAGGTCGTA
R: common	5'-CAAAGTGCTGGTGTGAACTACTG	F: common	5'-CATCTGTGATCTGTTTTCCCTCTTTTAC
TaqMan probe	5'-TAGTATGTGCTATGTGCTGTCC	SNP16	5 chiefononeronniecerentime
SNP5		R: C allele	5'-GAGTGTCTCACTTAATAGTCAACGATG
R: A allele	5'-GCCTGCCCATTATTCTTGGTCCT	R: G allele	5'-GAGTGTCTCACTTAATAGTCAACGATC
R: G allele	5'-GCCTGCCCATTATTCTTGGTCAC	F [.] common	5'-TGGTCAGGCTGGTCTTGAACTC
F: common	5'-CCTTGTGAGCTCTTCTTTTTGAGTTC	SNP17	2 10010100010010110120010
SNP6		F: G allele	5'-CTGCCAAAGTTTTGGGATCACTG
R: G allele	5'-CACCCCGTCCAGCAGTACCTAC	F: A allele	5'-CTGCCAAAGTTTTGGGATCACTA
R: A allele	5'-CACCCCGTCCAGCAGTACCTAT	R: common	5'-CCACACCCGGCCATTTCC
F: common	5'-AACAGAGGATTTCCCTGCCTGAA	SNP18	
TaqMan probe	5'-CCCTCAAGCGCATTCCG	R: C allele	5'-AATGCCAGAAGTGAACTTCAAAGTG
SNP7		R: G allele	5'-AATGCCAGAAGTGAACTTCAAAGTC
F: A allele	5'-AACAGAGGATTTCCCTGCCTGTA	F: common	5'-CAGCAGGGTCAACATCACCATA
F: G allele	5'-AACAGAGGATTTCCCTGCCTGGG	SNP19	
R: common	5'-CACCCCGTCCAGCAGTACCTTC	F: T allele	5'-GGCTGCCTGTATTCTTGCCTAT
TaqMan probe	5'-CCCTCAAGCGCATTCCG	F: delT allele	5'-GGCTGCCTGTATTCTTGCCTCG
SNP8		R: common	5'-TGGATTCAAATTCCAGCTCTAAATAAC
F: G allele	5'-TGCTGAATAAGAAGAACAAACTGAGG	SNP20	
F: A allele	5'-TGCTGAATAAGAAGAACAAACTGAGA	F: A allele	5'-ATTCCAAGCATGAGCCACGA
R: common	5'-TGCCTCCGCCTCTTCTCA	F: G allele	5'-ATTCCAAGCATGAGCCACGG
SNP9		R: common	5'-GGAGAGGAGAAAGCCCAAATCA
F: G allele	5'-TAGGGACGAAGATGGAGATGTG	SNP21	
G allele	5'-TAGGGACGAAGATGGAGATGAG	R: A allele	5'-GATGGAGAGAAAATAACAGAAGAGTT
F: A allele	5'-TAGGGACGAAGATGGAGATGTA	R: C allele	5'-GATGGAGAGAAAATAACAGAAGAGCG
A allele	5'-TAGGGACGAAGATGGAGATG A A	F: common	5'-ATGTTGCTCAAGTTGGTCTCAAACT
R: common	5'-CITATCTCCATTTCCCCTCTGATTC	SNP22	
SNP10		R: A allele	5'-GGCCTCCCTCCAAAGACCCT
R: G allele	5'-IGATICCATCIGCATTCCCAAC	R: G allele	5'-GGCCTCCCTCCAAAGACCTC
K: A allele	5'-IGATICCATCIGCATTCCCAAT	F: common	5'-CGTGTCCCAGGAATCTATGAATTT
F: common	5-IIIICCGGAGTAGGGACGAAGA		

Table 3 continued

Table 3 continued

SNP genotyped	Sequence
SNP23	
R: G allele	5'-ACCGGGTACAGCTTTCCTCTAC
R: A allele	5'-ACCGGGTACAGCTTTCCTCTTT
F: common	5'-GGCTTCTGTGAGTTGAGAGTCTAAGAGA
SNP24	
F: T allele	5'-CATGGCCATGAGGCTACAGTATAT
F: C allele	5'-CATGGCCATGAGGCTACAGTATAC
R: common	5'-CCCATGTCACCATCCACACA
SNP25	
F: T allele	5'-ACACTTGGGTGGAGCACTGGCAT
F: C allele	5'-ACACTTGGGTGGAGCACTGGCTC
R: common	5'-GGTCCTGGCATTGTTTCTGGTTCTC
TaqMan probe	5'-AGGAGGACGAATGCCAAGACG
SNP26	
R: T allele	5'-GGTGAGAATGTGGGCAAGAAGA
R: G allele	5'-GGTGAGAATGTGGGCAAGAAGC
F: common	5'-ACCTACATCCTCTCCGGTGAGTGT

PCR polymerase chain reaction, *F* forward primer, *R* reverse primer. All TaqMan probes were labeled with 6-carboxyfluorescein (FAM, reporter dye) at the 5' end and 6-carboxy-tetramethyl-rhodamine (TAMRA, quenching dye) at 3' end

Results

Polymorphisms in the C3 gene

We selected and characterized 26 SNPs from the C3 gene to investigate mainly genetic variations of the promoter region, SNPs forming a haplotype that showed significant association with BA in African Caribbean families, and the 3' end region where a SNP showed significant association with BA in our previous study (Table 2). The location of the SNPs and LD map is shown in Fig. 1. If an LD block was defined by the solid spine of LD, out of the 26 SNPs investigated, 24 composed four LD blocks. SNP10 showed r^2 values <0.033 to any other SNPs investigated in the current study and excluded from any LD block under this definition. Although SNP16 showed moderate LD to SNP15 ($r^2 = 0.71$) and SNP11 ($r^2 = 0.60$), it did not belong to LD block 3, where SNP15 and SNP11 are located. This is because of the characteristic of the "solid spin of LD" definition. When SNP12 and SNP14, both of which showed very low r^2 (<0.01) to SNP16, were omitted from LD block 3, SNP16 was found to be included in LD block 3. As tag SNPs, four SNPs each were selected in LD blocks from 1 to 3 and three SNPs in LD block 4 (Fig. 1).

Association study of childhood and adult BA

We genotyped 15 tag SNPs in 304 childhood BA, 371 adult BA, 333 child controls, and 550 adult controls (Table 1).

All loci were in Hardy-Weinberg equilibrium in the control groups. The genotype and allele frequencies of each SNP in the patient and control groups are shown in Table 4. Results of association tests for allele frequency between the patient and control groups are also shown. Allelic frequency of SNP24 was significantly different between adult BA and adult controls (P = 0.002). The P value for the difference remained significant ($P_{cor} = 0.030$) after correction for the number of SNPs tested (15). In a recessive model, the odds ratio (OR) and its 95% confidence interval (95% CI) range for this SNP were 1.55 and 1.15-2.09, respectively. In a dominant model, the OR and 95% CI range were 1.37 and 1.01-1.88, respectively. The difference in the allele frequency of SNP24 between all BA and all control showed a similar tendency. However, the P value was not significant (P = 0.004, $P_{cor} = 0.060$). None of the other SNPs showed a significant association with either childhood or adult BA.

The frequencies of haplotypes consisting of tag SNPs in four LD blocks are shown in Table 5. A significant difference was observed in LD block 4 (Table 5). In LD block 4 containing exons 24-41, the frequency of the major haplotype CCC in adult BA (54.1%) was significantly higher than that in controls (46.4%) (P = 0.0014, $P_{cor} =$ 0.038). The frequencies of C alleles of SNP18, SNP19, and SNP24 were higher in adult BA than in adult controls. These increased frequencies were straightforwardly associated with the increased frequency of the CCC haplotype. The frequency of this haplotype in childhood BA (51.0%) was not significantly higher than that in adult controls (48.6%) ($P_{cor} > 1.0$). The OR of the CCC haplotype of LD block 4 was about 1.4 for both childhood and adult BA (Table 6). In LD block 2 containing at least exons 3–14, the frequency of the GGAG haplotype in childhood BA (5.6%) was higher than that in child controls (2.5%) (P = 0.0044). This difference, however, did not reach a genewide significance ($P_{cor} = 0.12$). In LD blocks 1 and 3, there was no haplotype showing any difference in frequency between patients and controls. LD block 1 contains the promoter region of the C3 gene; therefore, it is not likely that genetic variations of the promoter region have significant effect on susceptibility to BA in the Japanese population.

Barnes et al. reported that a 3-SNP haplotype consisting of the SNPs identical to SNP14, SNP15, and SNP16 showed significant association with BA, log (total IgE) and log (IL-13)/log (IFN- γ) in the Afro-Caribbean families (Barnes et al. 2006). To assess the reproducibility of their results, we investigated whether this 3-SNP haplotype shows association with either childhood BA, adult BA, or log (total IgE) in our samples. We found that this haplotype showed no significant association with any of these phenotypes.

	Child	nood as	thma			Adult	asthma				Child c	ontrol			-	Adult c	ontrol			Child asthm child	hood a versus	Adult asthma versus adult	All asthma versus all
	11 ^a	12 ^b	22 ^c	1^{d}	2^{d}	11^{a}	12 ^b	22 ^c	1^{d}	2^{d}	11 ^a	12 ^b	22 ^c	l ^d	2 ^d	11 ^a	12 ^b	22 ^c 1	d 2	$\frac{d}{P_{cor}}$		$P_{\rm cor}^{e}$	$P_{\rm cor}^{e}$
SNP1	0.382	0.473	0.144	0.619	0.381	0.438	0.432	0.130	0.654	0.346	0.351	0.508	0.141	0.605	0.395	0.395	0.457 (0.149 (.623 0	.377 >1		>1	>1
SNP2	0.755	0.222	0.024	0.865	0.135	0.785	0.171	0.044	0.871	0.130	0.754	0.192	0.055	0.849	0.151	0.769	0.205 (0.027 (.871 0	.129 >1		~	$\overline{}$
SNP3	0.440	0.443	0.117	0.662	0.339	0.431	0.456	0.113	0.659	0.341	0.375	0.502	0.123	0.626	0.374	0.424	0.467	0.109 (.658 0	.343 >1		~	$\overline{}$
SNP4	0.433	0.406	0.161	0.636	0.364	0.437	0.439	0.124	0.657	0.344	0.402	0.475	0.123	0.640	0.361	0.426	0.430	0.143 (.642 0	.358 >1		~	$\overline{}$
SNP5	0.542	0.405	0.054	0.744	0.256	0.616	0.323	0.061	0.778	0.223	0.562	0.369	0.069	0.747	0.254	0.628	0.310	0.062 (.783 0	.217 >1		~	$\overline{}$
SNP6	0.411	0.461	0.128	0.642	0.359	0.371	0.454	0.175	0.598	0.402	0.381	0.469	0.150	0.616	0.385	0.327	0.485 (0.188 (.570 0	.431 >1		~	$\overline{}$
SNP7	0.201	0.497	0.302	0.450	0.551	0.256	0.497	0.247	0.505	0.496	0.255	0.482	0.264	0.496	0.504	0.285	0.485 (0.230 (.528 0	.473 >1		~	$\overline{}$
SNP9	0.737	0.232	0.030	0.854	0.146	0.750	0.219	0.031	0.860	0.141	0.709	0.273	0.018	0.846	0.155	0.753	0.219 (0.028 (.863 0	.138 >1		~	$\overline{}$
SNP11	0.340	0.505	0.155	0.593	0.408	0.331	0.489	0.180	0.576	0.425	0.324	0.502	0.174	0.575	0.425	0.312	0.478 (0.210 (.551 0	.449 >1		>1	~
SNP13	0.862	0.131	0.007	0.928	0.073	0.889	0.106	0.006	0.941	0.059	0.880	0.120	0.000	0.940	0.060	0.890	0.110	0.000 (.945 0	.055 >1		>1	~
SNP14	0.769	0.228	0.003	0.883	0.117	0.779	0.202	0.019	0.880	0.120	0.778	0.213	0.009	0.885	0.116	0.786	0.206 (0.008 (.889 0	.111 >1		>1	~
SNP15	0.379	0.480	0.141	0.619	0.381	0.348	0.514	0.138	0.605	0.395	0.363	0.469	0.168	0.598	0.403	0.316	0.503 (0.181 (.568 0	.433 >1		>1	~
SNP18	0.305	0.557	0.138	0.584	0.417	0.403	0.442	0.156	0.623	0.377	0.348	0.454	0.198	0.575	0.425	0.328	0.506 (0.167 (.580 0	.420 >1		>1	~
SNP21	0.367	0.522	0.111	0.628	0.372	0.433	0.444	0.122	0.656	0.344	0.369	0.471	0.159	0.605	0.395	0.358	0.498 (0.144 (.607 0	.393 >1		0.540	0.555
SNP24	0.248	0.537	0.215	0.517	0.484	0.225	0.456	0.319	0.453	0.547	0.282	0.450	0.267	0.508	0.492	0.285	0.482	0.233 (.526 0	.474 >1		0.030	0.060
a T	,	5 L			-1-11																		

Table 4 Single nucleotide polymorphism (SNP) association study of the C3 gene

Frequency of homozygote for major allele

^b Frequency of heterozygote

^c Frequency of homozygote for minor allele

^d Allele frequency

^c Corrected P value (raw P values were multiplied by number of SNPs, 15) for allele frequency difference

Table 5 Haploty aniation study of th

study of the C3 gene	Haplotype	Frequency				P _{cor} ^a		
		Childhood asthma	Adult asthma	Child control	Adult control	Childhood asthma vs. control	Adult asthma vs. control	All-asthma vs. control
	Block 1 ^b							
	TCAA	0.441	0.483	0.436	0.467	>1	>1	>1
	CCGC	0.295	0.263	0.279	0.280	>1	>1	>1
	TTAA	0.129	0.108	0.121	0.103	>1	>1	>1
	CCAA	0.051	0.043	0.041	0.042	>1	>1	>1
	TCGC	0.032	0.051	0.040	0.021	>1	0.222	>1
	CCGA	0.009	0.014	0.032	0.020	0.192	>1	0.297
	CCAC	0.019	0.010	0.019	0.016	>1	>1	>1
	TCAC	0.018	0.009	0.004	0.017	0.324	>1	>1
	Others	0.006	0.022	0.029	0.027			
	Block 2 ^c							
	AAGG	0.322	0.361	0.333	0.403	>1	>1	>1
	AGAG	0.337	0.301	0.310	0.291	>1	>1	>1
	GGAA	0.124	0.110	0.141	0.115	>1	>1	>1
	AGGG	0.058	0.067	0.072	0.063	>1	>1	>1
	GGGG	0.049	0.055	0.053	0.051	>1	>1	>1
	GGAG	0.056	0.042	0.025	0.037	0.121	>1	0.729
	AAAG	0.016	0.018	0.015	0.015	>1	>1	>1
	AGAA	0.014	0.024	0.013	0.010	>1	0.567	>1
	GAGG	0.019	0.015	0.036	0.009	>1	>1	>1
	Others	0.006	0.006	0.002	0.005			
	Block 3 ^d							
	ACCC	0.460	0.446	0.445	0.424	>1	>1	>1
	GCCT	0.365	0.382	0.384	0.418	>1	>1	>1
^a Corrected <i>P</i> value (raw <i>P</i>	ATTC	0.066	0.058	0.055	0.054	>1	>1	>1
values were multiplied by the	ACTC	0.049	0.059	0.057	0.055	>1	>1	>1
number of haplotypes tested, 27) for haplotype frequency difference ^b This haplotype is consisted of	GCCC	0.041	0.041	0.038	0.034	>1	>1	>1
	ACCT	0.013	0.012	0.014	0.013	>1	>1	>1
	Others	0.006	0.002	0.006	0.003			
SNPs 1, 2, 3, and 4	Block 4 ^e							
^c This haplotype is consisted of	CCC	0.510	0.541	0.486	0.464	>1	0.038	0.079
SNPs 5, 6, 7, and 9	GAT	0.365	0.340	0.387	0.375	>1	>1	>1
^d This haplotype is consisted of	CCT	0.066	0.078	0.081	0.098	>1	>1	>19
SNPS 11, 13, 14, and 15	GCT	0.053	0.033	0.038	0.044	>1	>1	>1
SNPs 18, 21, and 24	Others	0.007	0.011	0.008	0.019			

Table 6 Odds ratio (OR) and its 95% confidence interval (CI) of the CCC haplotype for bronchial asthma (BA)

	Diplotype frequency	y (%)			
	Case		Control		
	CCC/CCC and CCC/other	Other/other	CCC/CCC and CCC/other	Other/other	Odds ratio (95% CI)
Childhood BA	77.8	22.2	71.2	28.8	1.42 (0.99–2.04)
Adult BA	76.9	23.1	71.0	29.0	1.36 (1.00–1.86)
All BA	77.3	22.7	71.1	28.9	1.39 (1.10–1.75)

Diplotype of each individual was inferred with SNPAlyze ver. 4.1. OR and its (95% CI) were determined with SPSS ver. 15.0

Effect of genetic variations of the *C3* gene on serum total IgE level

To examine the effect of genetic variations of the *C3* gene on serum IgE level, 15 tag SNPs and the CCC haplotype were analyzed by analysis of variance (ANOVA) for logtransformed serum total IgE values (Table 7). *P* values <0.05 were observed in SNP14 in adult BA patients (*P* = 0.042) and in SNP13 (*P* = 0.018) and SNP18 (0.041) in child controls. If we want to maintain type 1 error at 0.05 in each patient/control group, a *P* value <0.0031 (0.05/16) should be considered as significant. Thus, we deduced that these SNPs did not significantly affect serum total IgE level. The genotype of SNP24 and the CCC haplotype, both of which showed an association with adult BA, were not significantly associated with the IgE level.

Table 7	Effect	of	single	nucleotide	polymorphisms	(SNPs)	and
haplotyp	es of the	e Cŝ	gene o	n log ₁₀ -tran	sformed total imm	nunogloł	oulin
E (IgE)							

Locus	P value in AN	OVA test ^a	
	Childhood asthma	Adult asthma	Child control
Block 1			
SNP1	0.347	0.242	0.250
SNP2	0.657	0.960	0.320
SNP3	0.276	0.051	0.720
SNP4	0.329	0.058	0.182
Block 2			
SNP5	0.537	0.794	0.119
SNP6	0.860	0.533	0.805
SNP7	0.952	0.556	0.497
SNP9	0.368	0.795	0.270
Block 3			
SNP11	0.463	0.678	0.937
SNP13	0.917	0.100	0.018
SNP14	0.485	0.042	0.261
SNP15	0.486	0.816	0.688
Block 4			
SNP18	0.197	0.890	0.041
SNP21	0.065	0.280	0.097
SNP24	0.356	0.661	0.526
CCC haplotype	0.175	0.586	0.505

^a Significance of difference in the mean of \log_{10} (total IgE) among individuals with different genotypes (major allele homozygote, heterozygote, minor allele homozygote) was tested with analysis of variance (ANOVA). Three groups (childhood asthma, adult asthma, and child control) were evaluated separately. Serum IgE level was not determined in the adult control group

Discussion

In this study, we investigated the association of SNPs in the C3 gene with childhood and adult BA. We observed a significant association of SNP24 and the CCC haplotype in LD block 4 with adult BA. As the CCC haplotype was discriminated from other major haplotypes with a frequency of >1% by the SNP24 allele (C/T), this haplotype association may just be a reflection of the association of SNP24. There was no significant difference in genotype frequency of SNP24 between the adult control group and the child control group, suggesting that the significant association observed in the adult patients-control comparison was not due to skewed genotype frequency of adult controls but to changes in adult BA. The frequency of the T/T genotype in childhood BA was decreased (from 0.282 to 0.248), as in adult BA. However, the change in allele frequency in childhood BA compared with that in child controls was small (T allele: 0.517 vs. 0.508). This SNP (or that with strong LD to this SNP) may not be a risk-modifying variation for childhood BA. Although it was not significant at the gene-wide level, we observed a tendency of the frequency of the GGAG haplotype of LD block 2 to show a difference between childhood BA and child controls (P =0.0045, $P_{\rm cor} = 0.121$). To determine and definitively conclude whether this LD block confers risk for childhood BA, an association study with more childhood samples and functional analyses of this region will be necessary.

LD block 4 is about 20 kb long and contains exon 41 coding the C-terminal end of the C3 protein and 3' untranslated region as well as upstream 17 exons and introns. Because we could not find SNPs that change the amino acid sequences in exons in this LD block, functional variation(s) in this region should be those affecting either expression of the gene, ribonucleic acid (RNA) splicing, or RNA stability. To the best of our knowledge, there is as yet no study investigating elements affecting gene expression in this region of the C3 gene. Currently, we do not have sufficient data to discuss the underlying mechanisms of the association between the LD block and adult BA. Further genetic and functional analyses of the LD blocks are necessary to pinpoint genetic variation(s) responsible for differences in susceptibility to BA.

It is possible that some of the control subjects involved in this study will develop asthma. Judging from the prevalence of the asthma in Japan, the chance of developing the disease is expected to be several percent. If we exclude atopic subjects from the control group, the chance may be reduced but not completely eliminated. This issue cannot be thoroughly controlled in a case–control study. The fact that our control group contained presymptomatic BA patients reduces the statistical power to detect an association between genotypes and the disease. We must be aware of this issue when obtaining negative results. However, the main result (i.e., the significant association of the haplotype with the disease) cannot be changed by this issue. If we could eliminate presymptomatic BA patients, estimated ORs would be higher than the values we presented in this study.

We also investigated whether genetic variations of the C3 gene affect total serum IgE level using childhood and adult BA patients and child controls. The IgE level was not significantly affected by any SNPs or haplotypes, including those showing a significant association with adult BA. This was true with mite-specific IgE level (data not shown). These results suggest that genetic variations of LD block 4 showing susceptibility to BA have a slight effect on sensitization to allergens but are more relevant to the effector phase of allergic inflammation. The roles of anaphylatoxins in the pathogenesis of BA can be divided into two phases: sensitization to allergens and effector phase of allergic BA (Kohl and Wills-Karp 2007). Several roles of C5a signaling at the interface between dendritic cells and T cells are evident, but those of C3a remain unclear. In a C3aR knockout experiment, different Th2 cytokine production responses were reported in various strains of mice (C57BL/6 and BALB/c) (Drouin et al. 2002; Humbles et al. 2000), suggesting that the effect of C3a is influenced by genetic background. As observed in mice, a difference in genetic background may explain the fact that association of the haplotype consisting of SNP14, SNP15, and SNP16 with total IgE level observed in Afro-Caribbean families was not confirmed in the Japanese population. Discrepancy between our results and those of Barnes et al. may also be due to a difference in the environmental factors of the two study populations. There are a number of studies showing that the effects of genetic variation (-159C/T) of the CD14 gene on allergic sensitization and BA risk differ greatly due to environmental factors such as mite or lipopolysacharide concentration in dust (Ober et al. 2000; Simpson et al. 2006; Vercelli 2003; Zambelli-Weiner et al. 2005). It may be possible that C3 gene variations also show this type of geneenvironmental interaction and cause discrepant results in studies with different populations.

In summary, our results suggest that the LD block containing exons 24–41 of the C3 gene confer susceptibility to adult BA in the Japanese population. Because this region showed a slight effect on serum IgE level in both BA patients and normal individuals, this region may be involved in the effector phase of allergic inflammation. The effect of variations of the C3 gene on allergic sensitization and BA susceptibility may differ according to genetic background and environmental factors.

Acknowledgments We thank all patients and their families, volunteers who served as controls, and all staff members of hospitals involved in this study. We are also grateful for the excellent technical assistance of Kazuko Hatori, Mieko Yoshida, and Yoshiko Hotta. This work was supported by Grants-in-Aid for Scientific Research from the Ministry of Education, Culture, Sports, Science, and Technology, Japan, and grants from the Ministry of Health, Labor, and Welfare, Japan.

References

- Asher MI, Keil U, Anderson HR, Beasley R, Crane J, Martinez F, Mitchell EA, Pearce N, Sibbald B, Stewart AW et al (1995) International Study of Asthma and Allergies in Childhood (ISAAC): rationale and methods. Eur Respir J 8:483–491
- Barnes KC, Marsh DG (1998) The genetics and complexity of allergy and asthma. Immunol Today 19:325–332
- Barnes KC, Grant AV, Baltadzhieva D, Zhang S, Berg T, Shao L, Zambelli-Weiner A, Anderson W, Nelsen A, Pillai S, Yarnall DP, Dienger K, Ingersoll RG, Scott AF, Fallin MD, Mathias RA, Beaty TH, Garcia JG, Wills-Karp M (2006) Variants in the gene encoding C3 are associated with asthma and related phenotypes among African Caribbean families. Genes Immun 7:27–35
- Barrett JC, Fry B, Maller J, Daly MJ (2005) Haploview: analysis and visualization of LD and haplotype maps. Bioinformatics 21:263– 265
- Barrington R, Zhang M, Fischer M, Carroll MC (2001) The role of complement in inflammation and adaptive immunity. Immunol Rev 180:5–15
- Bautsch W, Hoymann HG, Zhang Q, Meier-Wiedenbach I, Raschke U, Ames RS, Sohns B, Flemme N, Meyer zu Vilsendorf A, Grove M, Klos A, Kohl J (2000) Cutting edge: guinea pigs with a natural C3a-receptor defect exhibit decreased bronchoconstriction in allergic airway disease: evidence for an involvement of the C3a anaphylatoxin in the pathogenesis of asthma. J Immunol 165:5401–5405
- Blumenthal MN, Ober C, Beaty TH, Bleecker ER, Langefeld CD, King RA, Lester L, Cox N, Barnes K, Togias A, Mathias R, Meyers DA, Oetting W, Rich SS (2004) Genome scan for loci linked to mite sensitivity: the Collaborative Study on the Genetics of Asthma (CSGA). Genes Immun 5:226–231
- Drouin SM, Corry DB, Kildsgaard J, Wetsel RA (2001) Cutting edge: the absence of C3 demonstrates a role for complement in Th2 effector functions in a murine model of pulmonary allergy. J Immunol 167:4141–4145
- Drouin SM, Corry DB, Hollman TJ, Kildsgaard J, Wetsel RA (2002) Absence of the complement anaphylatoxin C3a receptor suppresses Th2 effector functions in a murine model of pulmonary allergy. J Immunol 169:5926–5933
- Ferris BG (1978) Epidemiology Standardization Project (American Thoracic Society). Am Rev Respir Dis 118:1–120
- Fujii K, Matsubara Y, Akanuma J, Takahashi K, Kure S, Suzuki Y, Imaizumi M, Iinuma K, Sakatsume O, Rinaldo P, Narisawa K (2000) Mutation detection by TaqMan-allele specific amplification: application to molecular diagnosis of glycogen storage disease type Ia and medium-chain acyl-CoA dehydrogenase deficiency. Hum Mutat 15:189–196
- Hasegawa K, Tamari M, Shao C, Shimizu M, Takahashi N, Mao XQ, Yamasaki A, Kamada F, Doi S, Fujiwara H, Miyatake A, Fujita K, Tamura G, Matsubara Y, Shirakawa T, Suzuki Y (2004) Variations in the C3, C3a receptor, and C5 genes affect susceptibility to bronchial asthma. Hum Genet 115:295–301
- Hill WG, Robertson A (1968) Linkage disequilibrium in finite populations. Theor Appl Genet 38:226–231
- Humbles AA, Lu B, Nilsson CA, Lilly C, Israel E, Fujiwara Y, Gerard NP, Gerard C (2000) A role for the C3a anaphylatoxin receptor in the effector phase of asthma. Nature 406:998–1001

- Kawamoto S, Yalcindag A, Laouini D, Brodeur S, Bryce P, Lu B, Humbles AA, Oettgen H, Gerard C, Geha RS (2004) The anaphylatoxin C3a downregulates the Th2 response to epicutaneously introduced antigen. J Clin Invest 114:399–407
- Kohl J (2001) Anaphylatoxins and infectious and non-infectious inflammatory diseases. Mol Immunol 38:175–187
- Kohl J, Wills-Karp M (2007) A dual role for complement in allergic asthma. Curr Opin Pharmacol 7:283–289
- Krug N, Tschernig T, Erpenbeck VJ, Hohlfeld JM, Kohl J (2001) Complement factors C3a and C5a are increased in bronchoalveolar lavage fluid after segmental allergen provocation in subjects with asthma. Am J Respir Crit Care Med 164:1841–1843
- Lee YA, Wahn U, Kehrt R, Tarani L, Businco L, Gustafsson D, Andersson F, Oranje AP, Wolkertstorfer A, Berg AV, Hoffmann U, Kuster W, Wienker T, Ruschendorf F, Reis A (2000) A major susceptibility locus for atopic dermatitis maps to chromosome 3q21. Nat Genet 26:470–473
- Muller-Eberhard HJ (1988) Molecular organization and function of the complement system. Annu Rev Biochem 57:321–347
- Nakano Y, Morita S, Kawamoto A, Suda T, Chida K, Nakamura H (2003) Elevated complement C3a in plasma from patients with severe acute asthma. J Allergy Clin Immunol 112:525–530
- Nakashima K, Hirota T, Obara K, Shimizu M, Jodo A, Kameda M, Doi S, Fujita K, Shirakawa T, Enomoto T, Kishi F, Yoshihara S, Matsumoto K, Saito H, Suzuki Y, Nakamura Y, Tamari M (2006) An association study of asthma and related phenotypes with polymorphisms in negative regulator molecules of the TLR signaling pathway. J Hum Genet 51:284–291

- Ober C, Tsalenko A, Parry R, Cox NJ (2000) A second-generation genomewide screen for asthma-susceptibility alleles in a founder population. Am J Hum Genet 67:1154–1162
- Simpson A, John SL, Jury F, Niven R, Woodcock A, Ollier WE, Custovic A (2006) Endotoxin exposure, CD14, and allergic disease: an interaction between genes and the environment. Am J Respir Crit Care Med 174:386–392
- Venanzi S, Malerba G, Galavotti R, Lauciello MC, Trabetti E, Zanoni G, Pescollderungg L, Martinati LC, Boner AL, Pignatti PF (2001) Linkage to atopy on chromosome 19 in north-eastern Italian families with allergic asthma. Clin Exp Allergy 31:1220– 1224
- Vercelli D (2003) Learning from discrepancies: CD14 polymorphisms, atopy and the endotoxin switch. Clin Exp Allergy 33:153–155
- Werfel T, Kirchhoff K, Wittmann M, Begemann G, Kapp A, Heidenreich F, Gotze O, Zwirner J (2000) Activated human T lymphocytes express a functional C3a receptor. J Immunol 165:6599–6605
- Zambelli-Weiner A, Ehrlich E, Stockton ML, Grant AV, Zhang S, Levett PN, Beaty TH, Barnes KC (2005) Evaluation of the CD14/-260 polymorphism and house dust endotoxin exposure in the Barbados Asthma Genetics Study. J Allergy Clin Immunol 115:1203–1209
- Zhou W, Peng Q, Li K, Sacks SH (2007) Role of dendritic cell synthesis of complement in the allospecific T cell response. Mol Immunol 44:57–63