

Nomenclature for factors of the Dog Major Histocompatibility System (DLA), 1998: first report of the ISAG DLA Nomenclature Committee

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Summary

A Nomenclature committee for Factors of the Dog Major Histocompatibility System or Dog Leukocyte Antigen (DLA) has been convened under the auspices of the International Society for Animal Genetics (ISAG) to define a sequence based nomenclature for the genes of the DLA system.

The remit of this committee includes:

- assignment of gene names
- rules for naming alleles
- assignment of names to published alleles
- assignment of names to new alleles
- rules for acceptance of new alleles

DLA Nomenclature Committee, rules for acceptance, DLA genes and alleles, sequence based nomenclature

Introduction

This first ISAG DLA Nomenclature report considers the rules for acceptance of DLA genes and alleles, together with an appropriate sequence based nomenclature. Names have been assigned to existing sequences, where appropriate. The report also includes sequence alignments (both nucleotide and amino acid) for DLA class II alleles, including data which has not been previously published.

There is a table for each class II locus, which lists all published and unpublished sequence data, with accession numbers and references, plus equivalents for new and previously used allele names.

The Committee would like to acknowledge the contribution of the IUIS Committee on Nomenclature of DLA Determinants (Vriesendorp *et al.* 1977) (a subcommittee of the Nomenclature Committee of the International

Union of Immunological Societies), which started this work over 20 years ago.

Klein (Klein *et al.* 1990) proposed the use of the term *Cafa* (from *Canis familiaris*) to describe the class I and II genes in the dog MHC region. However, many publications before and since have continued to use the term DLA. The Nomenclature Committee has affirmed the use of the term DLA. This proposal has the approval of Dr Ronald Bontrop, the editor of the journal *Immunogenetics*, who maintains the register of MHC symbols for all species.

Assignment of DLA gene names

Class I

The assignment of official names for the class I genes has been considered by the committee. Studies suggest that four class I genes can be transcribed, and Table 1 lists those that have been identified to date.

Evidence from other species; cattle, horse, mouse, rat, suggests that there are different numbers of class I genes on different haplotypes. In cattle there are no clear locus specific characteristics so that it is not possible to assign alleles to particular loci just from the sequence data (S. Ellis, personal communication).

Although there appear to be locus specific characteristics which distinguish the different dog class I genes, the Committee considers it premature to assign official DLA names to these genes and needs to examine more data before naming the genes. Evidence from family studies may be needed in the dog to confirm that particular alleles belong to the same allelic series, and more data on the mapping of genes in the dog MHC is necessary.

The original serological data in the dog may have to be ignored, as it was assumed that there were three expressed class I alleles, whereas later evidence (Deeg *et al.* 1982; Doxiadis *et al.* 1989) showed that the originally defined DLA-B locus codes for a class II molecule, thus making the serology difficult to interpret. An attempt

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Accepted 26 April 1999

Table 1. Genes in the DLA complex

Official name	Locally assigned name	Molecular characteristics	References
–	DLA-A	?	(Sarmiento & Storb 1990a)
–	DLA-79	Non-classical class I gene associated with 7.9 kb Hind III fragment	(Burnett & Geraghty 1995; Burnett <i>et al.</i> 1997; Graumann <i>et al.</i> 1998)
–	DLA-88	Class I gene associated with 8.8 kb Hind III fragment	(Burnett <i>et al.</i> 1997; Graumann <i>et al.</i> 1998)
–	DLA-12	Non-classical class I gene associated with 12 kb Hind III fragment	(Burnett <i>et al.</i> 1997; Graumann <i>et al.</i> 1998)
–	DLA-12a	Class I pseudogene associated with 12 kb Hind III fragment	(Burnett <i>et al.</i> 1997)
–	C1 pg-26	Not in DLA region. Class I processed gene associated with 2.6 kb Hind III fragment	(Burnett <i>et al.</i> 1997)
–	DLA-53	Class I pseudogene associated with 5.3 kb Hind III fragment	(Burnett <i>et al.</i> 1997)
–	DLA-64	Non-classical class I gene associated with 6.4 kb Hind III fragment	(Burnett <i>et al.</i> 1997; Graumann <i>et al.</i> 1998)
DLA-DRA1	DRA	DR alpha chain	(Sarmiento & Storb 1988; Wagner <i>et al.</i> 1995)
DLA-DRB1	DRB1	DR beta chain	(Sarmiento & Storb 1988, 1990a)
DLA-DRB1	DRBB1		(Wagner <i>et al.</i> 1996b,c)
DLA-DRB2	DRB2	DRB pseudogene	(Sarmiento & Storb 1988, 1990b)
DLA-DRB2	DRBB2		(Wagner <i>et al.</i> 1996b,c)
DLA-DQA1	DQA1	DQ alpha chain	(Sarmiento & Storb 1988; Wagner <i>et al.</i> 1996b)
DLA-DQB1	DQB1	DQ beta chain	(Sarmiento & Storb 1988; Wagner <i>et al.</i> 1998a)
–	DQB2	?pseudogene	(Sarmiento & Storb 1988; Wagner <i>et al.</i> 1998a)
–	DPA	DP alpha chain	(Sarmiento & Storb 1988)
–	DPB1	DP beta chain	(Sarmiento & Storb 1988)
–	DPB2		(Sarmiento & Storb 1988)
–	DOB	DO beta chain	(Sarmiento & Storb 1988)
LMP2	LMP2		J.A. Gerlach, personal communication

will be made to correlate the original serological data with the current molecular data, by DNA sequencing some of the class II alleles from dogs used in the serology studies of the First, Second and Third Canine Immunology workshops (Vriesendorp *et al.* 1973, 1976; Deeg *et al.* 1986; Bull *et al.* 1987).

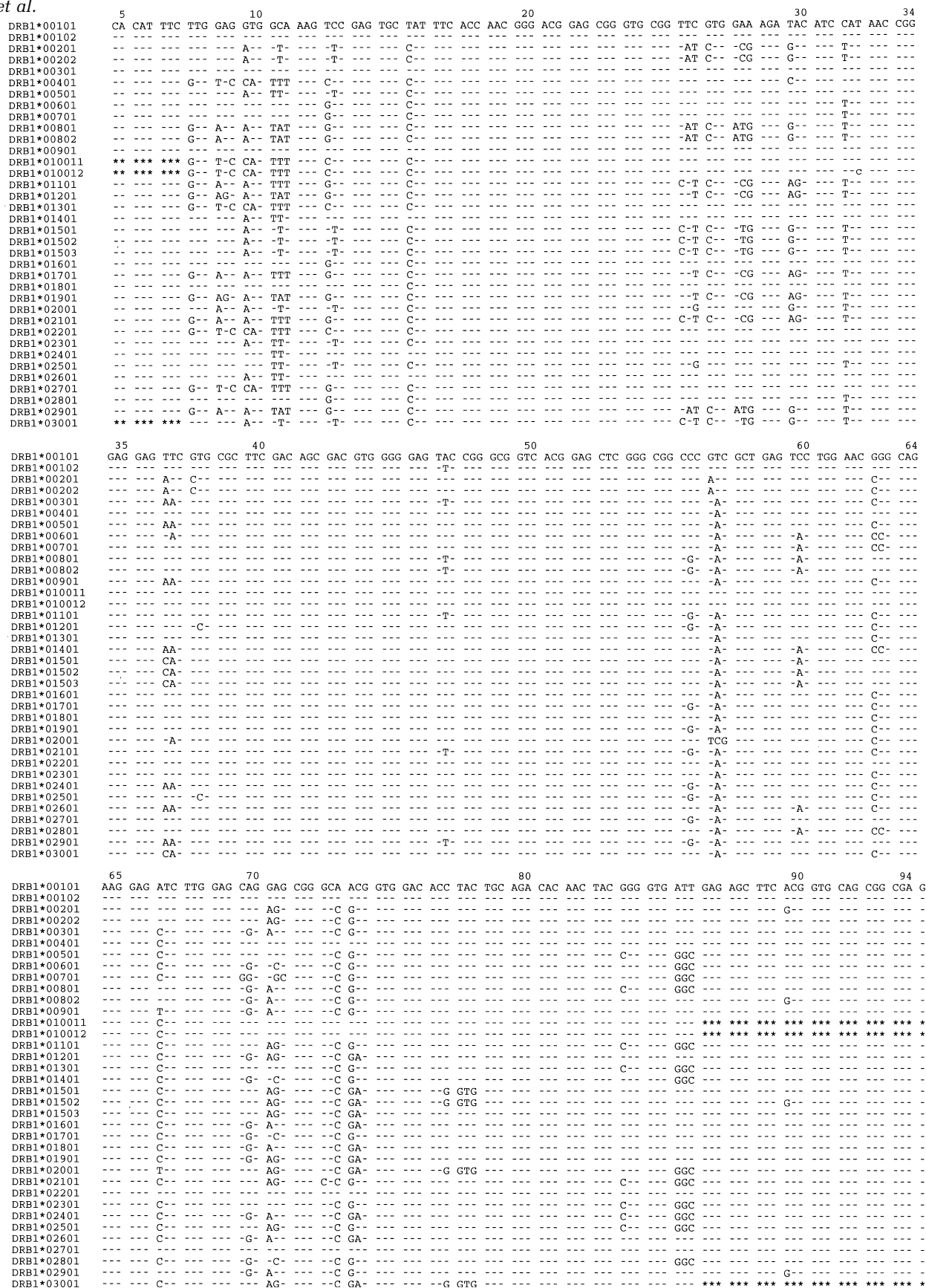
Class II

Within the dog class II region, clear homologues have been identified for human HLA-DRA, HLA-DRB, HLA-DQA and HLA-DQB genes. These will be named similarly, but the numbering of genes will be sequential, based on those numbers already in common use, and with no attempt to co-ordinate numbers between dog and human homologues. Thus DLA-DRB1 may or may not be a homologue of HLA-DRB1, as the identification of exact gene homologues may require considerably more sequence data than is currently available.

Table 1 also lists the DLA class II genes that have been identified to date. Where sufficient cloning and sequencing data have confirmed homology between the human and dog genes, official names have been assigned. Where such data are not yet available, such as for the putative DPA and DPB genes, no official names have been assigned.

Table 2. Definition of the hypervariable region boundaries for DLA class I and class II loci

DLA locus	Inclusive codon boundaries for the hypervariable regions		
	HVR 1	HVR 2	HVR 3
Class I: DLA-88	62–77	91–116	152–158
DRB1	8–16	26–38	56–74
DQA1	25	55	68–82
DQB1	9–13	26–37	57–75



	10	20	30	40	50	60	70	80	90	
DRB1*00101	HPLEV	AKSECYTFNG	TERVRVERVY	IHNREEFVRF	DSDVGEYRAV	TELGRPVAES	WNGQKEILEQ	ERATVDTYCR	HNVGVIESFT	VQRR
DRB1*00102					F-					
DRB1*00201	--M	V-F-H--	--YLA-D	-Y--IL-		I-	R--R-	R-A-		-A-
DRB1*00202	--M	V-F-H--	--YLA-D	-Y--IL-		I-	R--R-	R-A-		
DRB1*00301				N-	F-	D-	R-L-R	K-A-		
DRB1*00401	--VQV	F-P-H--	H			D-	L-			
DRB1*00501	--M	L-F-H--		N-		D-	R-L-	A-		R-G-
DRB1*00601	--M	-A-H--		-Y-Y-		D-Y	P-L-R	A-A-		-G-
DRB1*00701		-A-H--		-Y-		D-Y	P-L-G	A-		-G-
DRB1*00801	--VKM	Y-A-H--	--YLM-D	-Y-	F-	RD-Y		R-K-A-		-R-G-
DRB1*00802	--VKM	Y-A-H--	--YLM-D	-Y-	F-	RD-Y		R-K-A-		-A-
DRB1*00901				N-		D-	R-F-R	K-A-		
DRB1*010011	**VQV	F-P-H--					L-			****
DRB1*010012	**VQV	F-P-H--					L-			****
DRB1*01101	--VKM	F-A-H--	--LLA-S	-Y-	F-	RD-	R-L-L	R-A-		-R-G-
DRB1*01201	--VRM	Y-A-H--	--LA-S	-Y-	A-	RD-	R-L-R	R-E-		
DRB1*01301	--VQV	F-P-H--				D-	R-L-	A-		-R-G-
DRB1*01401	--M	L-		N-		D-Y	P-L-R	A-A-		-G-
DRB1*01501	--M	V-F-H--	--LLV-D	-Y-H-		D-Y	L-	R-E-V-		
DRB1*01502	--M	V-F-H--	--LLV-D	-Y-H-		D-Y	L-	R-E-V-		-A-
DRB1*01503	--M	V-F-H--	--LLV-D	-Y-H-		D-Y	L-	R-E-		
DRB1*01601		-A-H--				D-	R-L-R	K-E-		
DRB1*01701	--VKM	F-A-H--	--LA-S	-Y-		RD-	R-L-R	A-A-		
DRB1*01801		-A-H--				D-	R-L-R	K-E-		
DRB1*01901	--VRM	Y-A-H--	--LA-S	-Y-		RD-	R-L-R	R-E-		
DRB1*02001	--KM	V-F-H--	--L-Y-	-Y-Y-		S-	R-L-	R-E-V-		-G-
DRB1*02101	--VKM	F-A-H--	--LLA-S	-Y-	F-	RD-	R-L-	R-PA-		-R-G-
DRB1*02201	--VQV	F-P-H--				D-				
DRB1*02301	--M	L-F-H--				D-	R-L-	A-		-R-G-
DRB1*02401				N-		RD-	R-L-R	K-E-		-R-G-
DRB1*02501		L-F-H--	--L-	-Y-A-		RD-	R-L-	R-A-		-R-G-
DRB1*02601	--M	L-		N-		D-Y	R-L-R	K-E-		
DRB1*02701	--VQV	F-A-H--				RD-				
DRB1*02801		-A-H--		-Y-N-		D-Y	P-L-R	A-A-		-G-
DRB1*02901	--VKM	Y-A-H--	--YLM-D	-Y-N-	F-	RD-		R-K-A-		-A-
DRB1*03001	**M	V-F-H--	--LLV-D	-Y-H-		D-	R-L-	R-E-V-		****
	--		--			--				
	HVR 1		HVR 2			HVR 3				

Rules for naming alleles

done for the dog MHC at the current time. We have considered the use of dendrograms to aid such a naming system for major types, but found that this tended to base the assignment of major types on the first HVR only. Since we wanted a system based on all three HVR, this was not acceptable to the committee. Fifthly, a precedent for such a system for naming alleles already exists in the human HLA nomenclature, for HLA-DPB, where a system based on six variable regions is used.

The extent of the HVR is well known for HLA-DRB1, but has not been defined for other HLA loci. We defined the HVR in DLA-DRB1 to be the same as those in HLA-DRB1. The DLA-DQB1 gene is very like DLA-DRB1, and thus we defined very similar regions as the HVR. For DLA-DQA1, which is much less variable, we selected the most polymorphic areas as the HVR. The extent of these regions are given in Table 2.

This system for naming alleles will result in more major types than in the HLA or BoLA systems. Since the current HLA system may soon run out of new allele numbers for some loci (e.g. HLA-DPB), we have introduced the use of an extra digit in the allele names, as compared to HLA and BoLA.

The rules used for the establishment of allele names are:

1. Names will be based on the amino acid sequences.
2. Allele names will consist of five or six digits; the first three digits indicating the major type, the fourth and fifth digits indicating the subtype, and the sixth digit (if

present) indicating non-expressed variation (silent substitutions). To aid recognition of DLA alleles, we recommend that alleles should be verbalized as in the following example: DRB1*01501 = “DRB1* zero-15, zero-one”.

3. Class I alleles within a single major type will be identical for the three hypervariable regions (HVR) in exons 2 and 3. Differences outside the HVR will be indicated as subtypes of the major type.

4. Class II alleles within a single major type will be identical for the three HVR in exon 2. Differences outside the HVR will be indicated as subtypes of the major type.

5. If a name is given on the basis of a partial sequence the first full length sequence that includes the original partial sequence will assume the allele name.

6. If minor sequence errors are identified the sequence will be corrected.

Assignment of names to be published and new alleles

Using the above guidelines, and the HVR definitions, each locus was considered in turn, and allele names were assigned.

Class I
No official names for the class I alleles will be given until it has been clarified that they all belong to the allelic series as published (Burnett & Geraghty 1995; Burnett et al. 1997; Graumann et al. 1998).

Class II
Nucleotide and amino acid alignments are given for DLA-DRB1, DQA1 and DQB1, in Figs 1–6.

References and accession numbers for each sequence for DRB1, DQA1 and DQB1, are given in Tables 3–5. The equivalents for all previous names are also indicated. Due to some partial sequences, some of the equivalents for DQB are probable rather than definite.

DRB1 references: (Sarmiento et al. 1990; Wagner et al. 1996c; Francino et al. 1997; Kennedy et al. 1998), J. M. Angles confirmatory sequence for DRB1*02401 (personal communication).

DQA1 references: (Sarmiento et al. 1992; Wagner et al. 1996a; Polvi et al. 1997).

DQB1 references: (Sarmiento et al. 1993; Polvi et al. 1997; Wagner et al. 1998a).

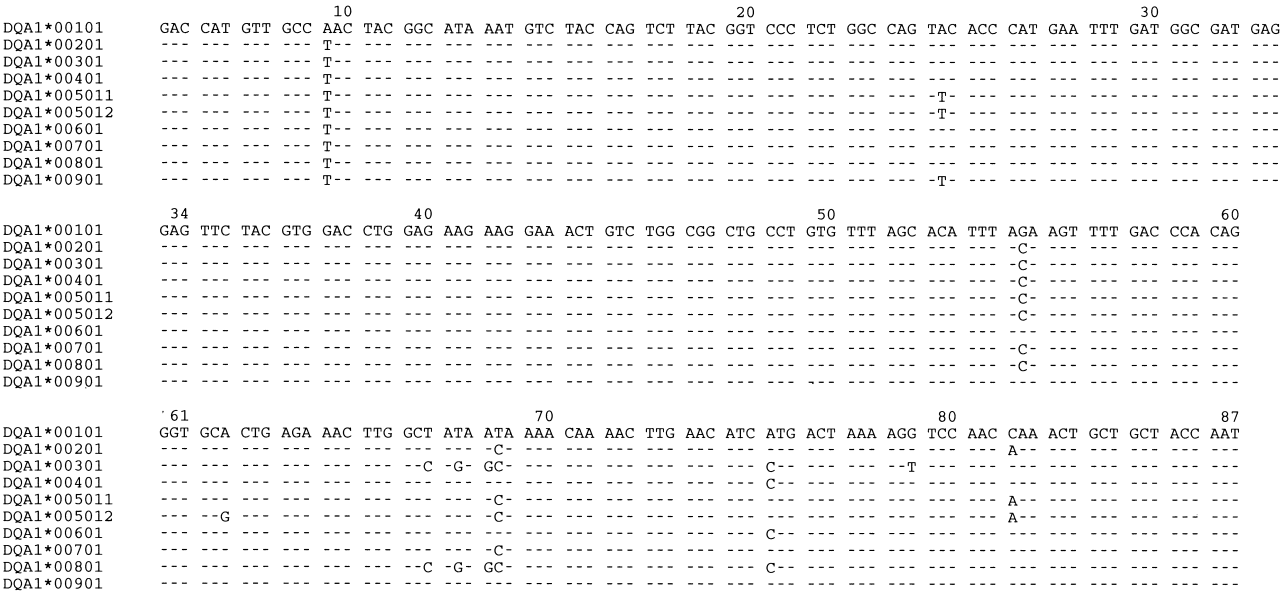


Fig. 3. DLA-DQA1 nucleotide sequence alignment

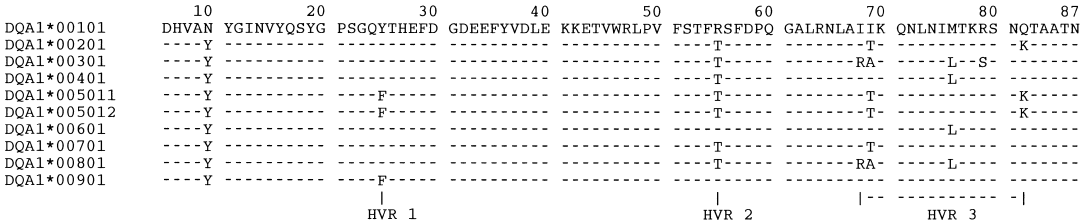


Fig. 4. DLA-DQA1 amino acid sequence alignment.

Conditions for acceptance of new sequences

1. For class I genes full length exon 2 and exon 3 sequences are required.
2. For class II genes the sequence for the first domain (exon 2) must be included.

3. Sequencing should be performed on both strands of the template DNA.

4. Where a sequence is obtained from cDNA or where PCR products are subcloned prior to sequencing, a minimum of three clones must be sequenced. Alternatively, identical sequences from two different dogs are acceptable.

Table 3. Accession numbers and references for DRB1 sequences

Official allele name	Previous allele names					Accession number	Reference
	Sarmiento ^a	Wagner ^b	Francino ^c	Kennedy ^d	Other		
DRB1*00101	Dw4	D1	—	—	—	M57529	a
DRB1*00102	Dw3	D3	—	—	—	M57528	a
DRB1*00102	—	—	—	—	Dw3	S76138	h
DRB1*00201	Dw1	D2	—	—	—	M57537	a
DRB1*00202	—	D2a	—	—	—	U44777	b
DRB1*00301	—	—	—	0902	—	AJ003012	d
DRB1*00401	D4	D4 m*	—	—	—	M57532	a, b
DRB1*00501	—	—	—	2302	—	AJ003017	d
DRB1*00501	—	D24†	—	—	—	AF098496	e
DRB1*00601	D6	D6 m*	—	—	—	M57534	a, b
DRB1*00701	D7	—	—	—	—	M57533	a
DRB1*00801	D8	D8 m*	—	—	—	M57535	a, b
DRB1*00802	—	—	—	08var2	—	AJ012456	f
DRB1*00901	—	D9	—	—	—	M57531	a
DRB1*010011	—	—	D25†	—	—	AF016910	g
DRB1*010012	—	—	Cafa-10	—	1102†	X93572	c
DRB1*01101	—	—	Cafa-11	—	1112†	X93573	c
DRB1*01201	—	—	—	1902	—	AJ003015	d
DRB1*01301	—	D13	—	—	—	U44778	b
DRB1*01401	—	D14	—	—	—	U44779	b
DRB1*01501	D15/Dw8	D15 m*	—	—	—	M57536	a, b
DRB1*01502	—	—	—	1502	—	AJ003013	d
DRB1*01503	—	—	—	1503	—	AJ003014	d
DRB1*01601	—	—	—	18var1	—	AJ012454	f
DRB1*01701	—	D17	—	—	—	U44780	b
DRB1*01801	—	D18	—	—	—	U44781	b
DRB1*01901	—	D19	—	—	—	U44782	b
DRB1*02001	—	D20	—	—	—	U58684	b
DRB1*02101	—	D21	—	—	—	U44783	b
DRB1*02201	—	D22	—	—	—	U58685	b
DRB1*02301	—	—	—	2301	—	AJ003016	d
DRB1*02401	—	—	—	2401	—	AJ003018	d
DRB1*02501	—	—	—	2501	—	AJ003019	d
DRB1*02601	—	—	—	2601	—	AJ003020	d
DRB1*02701	—	drb 26	—	—	—	AF061039	i
DRB1*02801	—	drb 25	—	—	—	AF061038	i
DRB1*02901	—	—	—	08var1	—	AJ012455	f
DRB1*03001	—	—	D23†	—	—	AF016911	g
Partial sequence	—	—	D24†	—	—	AF016912	g
Partial sequence	—	—	—	—	1-Dob-A†	M30129	j
Partial sequence	—	—	—	—	1-Dob-B†	M30130	j
Partial sequence	—	—	—	—	2-Dob†	M30131	j
Partial sequence	—	—	—	—	3-Lab†	M30132	j
Partial sequence	—	—	—	—	4-Pood†	M30133	j

†Unpublished alleles.

*Modification of original Sarmiento sequence, corrected by Wagner, same accession number.

References: a = (Sarmiento *et al.* 1990), b = (Wagner *et al.* 1996c), c = (Francino *et al.* 1997), d = (Kennedy *et al.* 1998), e = Wagner, GenBank 1998, unpublished, f = Kennedy *et al.*, *European Journal of Immunogenetics*, in press, g = Francino, GenBank 1997, unpublished, h = (He *et al.* 1994), i = (Wagner *et al.* 1998b), j = Motoyama, GenBank 1996, unpublished.

5. If direct sequencing of PCR amplified material is performed from a homozygous animal, products from at least two separate PCR reactions should be sequenced.

6. If direct sequencing of PCR amplified material is performed from a heterozygous animal, products from at least two separate PCR reactions should be sequenced, and the sequence should also be confirmed by cloning and sequencing.

7. Where possible sequences should be confirmed by another laboratory.

8. An accession number in a nucleotide sequence database should be obtained.

9. Submission of a sequence to the Nomenclature Committee should include a computer-readable copy of the sequence.

10. DNA (if possible from an animal homozygous for the allele), should be made available for a central repository maintained by the

	6	10										20										30										35									
DQB1*00101	GAT	TTC	GTG	TAC	CAG	TTT	AAG	GGC	GAG	TGC	TAT	TTC	ACC	AAC	GGG	ACG	GAG	CGG	GTG	CGG	CTT	CTG	ACG	AGA	GAC	ATC	TAT	AAC	CGG	GAG											
DQB1*00201	---	---	---	-T-	---	---	-A-	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*00301	---	---	---	---	---	---	---	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*00401	---	---	---	-T-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---										
DQB1*00501	---	---	---	-T-	---	-A-	---	-C-	---	-T-	-A-	T--	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*00701	---	---	---	-T-	---	---	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	G-	---	---	---	---	---	---	---	---										
DQB1*00801	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*01101	---	---	---	---	---	---	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*01301	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*01302	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*01303	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T-	-A-	T--	---	---	---	---	---	---										
DQB1*01501	---	---	---	---	-G-	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	T-	-	G-T	-A-	T--	---	---	---	---	---										
DQB1*01601	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	AG-	---	---	---	---	---	---										
DQB1*01701	---	---	---	-T-	-G-	---	-C-	---	-	---	---	---	---	---	---	---	---	---	---	---	---	T-	-	G-T	-A-	T--	---	---	---	---	---										
DQB1*01801	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	T-	-	-T-	-A-	T--	---	---	---	---	---										
DQB1*01901	---	---	---	-T-	-G-	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	T-	-	G-T	-A-	T--	---	---	---	---	---										
DQB1*02001	---	---	---	---	---	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	AG-	---	---	---	---	---	---										
DQB1*02002	---	---	---	---	---	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	AG-	---	---	---	---	---	---										
DQB1*02101	---	---	---	---	---	---	-C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	AG-	---	---	---	---	---	---										
DQB1*02201	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T	-A-	T--	---	---	---	---	---										
DQB1*02301	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-T	-A-	T--	---	---	---	---	---										

	36	40										50										60										65									
DQB1*00101	GAG	CAC	GTG	CGC	TTC	GAC	AGC	GAC	GTG	GGG	GAG	TAC	CGG	GCG	GTC	ACG	GAG	CTC	GGG	CGG	CCG	GAC	GCT	GAG	TAC	TGG	AAC	GGG	CAG	AAG											
DQB1*00201	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	---	C-A	---											
DQB1*00301	---	T-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	TCG	---	---	---	---	---	CC	---											
DQB1*00401	---	T-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	TGG	---	---	---	---	---	CC	---											
DQB1*00501	---	T-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	TGG	---	---	---	---	---	CC	---											
DQB1*00701	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---										
DQB1*00801	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---										
DQB1*01101	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	C-A	---	---										
DQB1*01301	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	TCG	---	---	---	---	---	CC	---	---										
DQB1*01302	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	CC	---	---										
DQB1*01303	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	CC	---	---										
DQB1*01501	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	CC	---	---										
DQB1*01601	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	TCG	---	---	---	---	---	---	---	---										
DQB1*01701	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	-T-	---	-C-	---	---	---	---	---	---										
DQB1*01801	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	---	---	---	---	---	CC	---	---											
DQB1*01901	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	TCG	---	---	---	---	---	---	---	---										
DQB1*02001	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	-T-	---	---	---	---	---	---	---	---										
DQB1*02002	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	-T-	---	---	---	---	---	---	---	---										
DQB1*02101	---	TT-	---	---	---	---	---	---	---	---	---	-T-	---	---	---	---	---	---	---	---	---	-C	-T-	---	---	---	---	---	---	---	---										
DQB1*02201	---	TT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	TCG	---	---	---	---	CC	---	---											
DQB1*02301	---	T-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	-C	TCG	---	---	---	CC	---	---	---											

	66	70										80										90										94									
DQB1*00101	-GAG	CTC	TTG	GAG	CGG	AGT	CGG	GCC	GAG	GTG	GAC	ACG	GTG	TGC	AGA	CAC	AAC	TAC	GGG	AGG	GAA	GAG	CTC	ACC	ACG	TTG	CAG	CGG	CGA												
DQB1*00201	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*00301	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*00401	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											
DQB1*00501	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											
DQB1*00701	-	---	A--	-	---	A-	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											
DQB1*00801	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*01101	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											
DQB1*01301	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*01302	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*01303	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	GT-	---	---	---	---	---	---	---	---	---											
DQB1*01501	-	A-	---	-	A-	---	-A	AC-	---	---	---	---	---	---	---	---	---	---	---	---	GT-	---	---	TA-	---	---	---	---	---	---											
DQB1*01601	-	A-	---	-	A-	---	-A	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*01701	-	T-	---	-	A-	---	-A	AC-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*01801	-C	GAG	A--	-C	---	GT-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	GT-	---	---	TA-	---	---	---	---	---	---											
DQB1*01901	-	---	---	-	AA	---	-A	---	-A	AC-	---	---	---	---	---	---	---	---	---	---	GT-	---	---	T-	---	---	---	---	---	---											
DQB1*02001	-	A-	---	-	A-	---	-A	---	-C	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*02002	-	A-	---	-	A-	---	-A	---	-C	---	---	-G-	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*02101	-	A-	---	-	A-	---	-A	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---											
DQB1*02201	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											
DQB1*02301	-C	GAG	A--	-C	---	GTA	---	---	---	C-	---	---	---	---	---	---	---	---	---	---	TT-	---	---	---	---	---	---	---	---	---											

Fig. 5. DLA-DQB1 nucleotide sequence alignment.

	6	10	20	30	40	50	60	70	80	90	94
DQB1*00101	DFVYQ	FKGECYFTNG	TERVRLTDR	IYNREEHVRF	DSDVGEYRAV	TELGRPDAEY	WNGQKELLER	RRAEVDTVCR	HNHYGREELTT	LQRR	
DQB1*00201	---	-F-	---	---	---	---	---	---	---	---	
DQB1*00301	---	-A-	---	---	---	---	---	---	---	---	
DQB1*00401	---	-F-	---	---	---	---	---	---	---	---	
DQB1*00501	---	-F-	---	---	---	---	---	---	---	---	
DQB1*00701	---	-F-	---	---	---	---	---	---	---	---	
DQB1*00801	---	-F-	---	---	---	---	---	---	---	---	
DQB1*01101	---	---	---	---	---	---	---	---	---	---	
DQB1*01301	---	---	---	---	---	---	---	---	---	---	
DQB1*01302	---	---	---	---	---	---	---	---	---	---	
DQB1*01303	---	---	---	---	---	---	---	---	---	---	
DQB1*01501	---	---	---	---	---	---	---	---	---	---	
DQB1*01601	---	---	---	---	---	---	---	---	---	---	
DQB1*01701	---	---	---	---	---	---	---	---	---	---	
DQB1*01801	---	---	---	---	---	---	---	---	---	---	
DQB1*01901	---	---	---	---	---	---	---	---	---	---	
DQB1*02001	---	---	---	---	---	---	---	---	---	---	
DQB1*02002	---	---	---	---	---	---	---	---	---	---	
DQB1*02101	---	---	---	---	---	---	---	---	---	---	
DQB1*02201	---	---	---	---	---	---	---	---	---	---	
DQB1*02301	---	---	---	---	---	---	---	---	---	---	

Fig. 6. DLA-DQB1 amino acid sequence alignment.

Table 4. Accession numbers and references for DQA1 sequences

Official allele name	Previous allele names			Accession number	Reference
	Sarmiento ^a (partial sequences)	Polvi ^b (partial sequences)	Wagner ^c		
DQA1*00101	0101		–	M74907	a
DQA1*00101	–	–	Dqa2	U44786	c
DQA1*00201	0201	–	–	M74909	a
DQA1*00201	–	–	Dqa9†	U75455	d
DQA1*00301	–	0301	–	Y07944	b
DQA1*00401	–	0203	–	Y07943	b
DQA1*00401	–	–	Dqa4	U44788	c
DQA1*005011	0202	–	–	M74910	a
DQA1*005011	–	–	Dqa3	U44787	c
DQA1*005012	–	–	Dqa5	U44789	c
DQA1*00601	–	0103	–	Y07942	b
DQA1*00601	–	–	Dqa6	U44790	c
DQA1*00701	–	–	Dqa7	U44842	c
DQA1*00801	–	–	Dqa8†	U61400	d
DQA1*00901	0102	–	–	M74908	a
DQA1*00901	–	–	Dqa1	U44785	c

†Unpublished sequences.

References: a = (Sarmiento *et al.* 1992), b = (Polvi *et al.* 1997), c = (Wagner *et al.* 1996a), d = Wagner, GenBank 1996, unpublished.

Nomenclature Committee. This DNA will be amplified and made available as reference material for other researchers.

Submission of new sequences

Sequences of new DLA genes or alleles should be submitted to the chairman of the DLA Nomenclature Committee, Lorna Kennedy, to receive official names. The sequence data or accession number(s) should be sent by e-mail: to the address given. Electronic submissions of sequence data are preferred. All sequence information will remain confidential until published or available on sequence databases. The Committee encourages the use of DLA as a keyword to ensure sequences may be found in database searches.

The use of numbers or names for alleles, genes or specificities which pre-empt formal designations such as: “DLA-E”, “DQA1*00401” or “DLA-DM” before consideration by the Nomenclature Committee is strongly discouraged.

Sequence database

The committee plans to establish a database of DLA allele sequences which would also include other data, such as how the sequence was

obtained, and how many clones were sequenced, etc. We will also record the breed of dog in which the sequence was found. This data will not be made public at this time, however, as such information may cause assumptions to be made about the restriction of particular alleles to certain breeds. (This is in keeping with the policy of the human HLA nomenclature committee, which also records the ethnic origin of all sequences submitted, but does not release that information.) Most of the DLA alleles listed in this report have been found in more than one breed, and many have been found in most breeds tested ($n = 60$). To date, only one allele (DLA-DRB1*02401) has been found to be limited to a single breed (Japanese Akita). However, as there are over 250 known breeds, there are still many breeds that have not been sampled, so DRB1*02401 may yet be found in another breed.

Acknowledgments

The ISAG DLA Nomenclature Committee met in UC Davis, California on 2nd August 1998. The meeting was jointly supported by the Center for Companion Animal Health, School of Veterinary Medicine, University of California, and the Canine Health Foundation (American Kennel Club).

Table 5. Accession numbers and references for DQB1 sequences

Official allele name	Previous allele names		Wagner ^c #	Francino ^d	Accession number	Reference
	Sarmiento ^a (partial sequences)	Polvi ^b (partial sequences)				
DQB1*00101	0101#	—	—	—	M90802	a
DQB1*00101	—	—	dqb2#	—	AF043147	c
DQB1*00101	—	—	—	dqb0102†	AF016905	d
DQB1*00201	0201#	—	—	—	M90803	a
DQB1*00201	—	—	dqb3#	—	AF043148	c
DQB1*00201	—	—	—	dqb0203†	AF016908	d
DQB1*00301	0301#	—	—	—	M90804	a
DQB1*00301	—	—	dqb6#	—	AF043151	c
DQB1*00401	0401#	—	—	—	M90805	a
DQB1*00401	—	—	dqb5#	—	AF043150	c
DQB1*00501	—	0501#	—	—	Y07947	b
DQB1*00501	—	—	dqb12#	—	AF043157	c
DQB1*00701	—	0701#	—	—	Y07949	b
DQB1*00701	—	—	dqb4#	—	AF043149	c
DQB1*00701	—	—	—	dqb1001†	AF016907	d
DQB1*00801	—	0801†#	—	—	AF043492	e
DQB1*00801	—	—	dqb1#	—	AF043167	c
DQB1*01101	—	—	—	dqb1101†	AF016904	d
DQB1*01301	—	—	dqb13	—	AF043158	c
DQB1*01302	—	—	dqb14	—	AF043159	c
DQB1*01303	—	—	dqb7	—	AF043152	c
DQB1*01303	—	—	—	dqb0901†	AF016906	d
DQB1*01501	—	—	dqb15	—	AF043160	c
DQB1*01601	—	—	dqb16	—	AF043161	c
DQB1*01701	—	—	dqb17	—	AF043162	c
DQB1*01801	—	—	dqb18	—	AF043163	c
DQB1*01901	—	—	dqb9	—	AF043154	c
DQB1*02001	—	—	dqb20	—	AF043165	c
DQB1*02002	—	—	dqb19	—	AF043164	c
DQB1*02101	—	—	dqb11	—	AF043156	c
DQB1*02201	—	—	dqb10	—	AF043155	c
DQB1*02301	—	—	dqb8	—	AF043153	c
DQB1*02301	—	—	—	dqb0303†	AF016909	d
Partial sequence	—	0302	—	—	Y07946	b
Partial sequence	—	0601	—	—	Y07948	b
Partial sequence	—	0202	—	—	Y07945	b

†Unpublished sequences.

#Sequence identities confirmed by J.L. Wagner, October 1998.

References: a = (Sarmiento *et al.* 1993), b = (Polvi *et al.* 1997), c = (Wagner *et al.* 1998a), d = Francino, GenBank 1997, unpublished, e = Polvi, GenBank 1998, unpublished.

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