

Names for DO (ISBT 014) Blood Group Alleles

Intro

General description: The Dombrock blood group system consists of 10 antigens carried on a GPI-linked glycoprotein (DO, ART4, CD297) that consists of 314 amino acids. It has a leader sequence and a GPI motif, both of which are cleaved from the membrane bound protein. The DO gene consists of 3 exons distributed over 18 kb of gDNA.

Gene name: *ART4 (DO)*

Number of exons: 3

Initiation codon: Within exon 1

Stop codon: Within exon 3

Entrez Gene ID: 420

LRG: LRG_807

LRG sequence: NG_007477.2 (genomic)

NM_021071.4, ENST00000228936.6 (transcript)

Reference allele: *DO*01* (shaded)

Acceptable: *DO*A*, or *Do^a* if inferred by haemagglutination

Reference allele

*DO*01* encodes: DO1, DO3, DO4, DO5, DO6, DO7, DO8, DO9, DO10

Antithetical antigens: [DO1 DO2]

(ISBT 014) DO blood group alleles v5.0 30-JUN-2021

Phenotype	Allele name	Nucleotide change	Exon Intron	Predicted amino acid change	(Reference No.) PMID	Accession number	rs number
DO:1+ or Do(a+)	<i>DO*01</i> or <i>DO*A</i>				PMID: 11552072	NM021071 AF290204	n.a.
DO:2 or Do(b+)	<i>DO*02</i> or <i>DO*B</i>	c.793A>G	2	p.Asn265Asp	PMID: 11552072	NM021071 AF290204	rs11276
DO:-4 or Hy-	<i>DO*02.-04</i>	c.323G>T c.793A>G	2	p.Gly108Val p.Asn265Asp	PMID: 11896313	AH011615 AH011616	rs28362797 rs11276
DO:-5 or Jo(a-)	<i>DO*01.-05</i>	c.350C>T	2	p.Thr117Ile	PMID: 11896313	AH011617	rs28362798
DO:-6 or DOYA-	<i>DO*01.-06</i>	c.547T>G	2	p.Tyr183Asp	PMID: 20088839	n.a.	n.a.
DO:-7 or DOMR-	<i>DO*02.-07</i>	c.431C>A c.432C>A c.793A>G	2	p.Ala144Glu p.Ala144Glu p.Asn265Asp†	PMID: 20412531	GU724770	rs1355202105 rs1210078970 rs11276
DO:-8 or DOLG-	<i>DO*01.-08</i>	c.674T>A	2	p.Leu225Gln	(1), Abstract	n.a.	rs532592412
DO:-9 or DOLC-	<i>DO*01.-09</i>	c.566C>T	2	p.Thr189Met	(2), Abstract	n.a.	rs28362800
DO:-10 or DODE-	<i>DO*01.-10</i>	c.405C>A	2	p.Asp135Glu	(3), Abstract	n.a.	rs28362799
Null phenotypes							
DO:-3 or Gy(a-)	<i>DO*01N.01</i>	c.442C>T	2	p.Gln148Ter‡	PMID: 11552072	AH011373	rs56340844
DO:-3 or Gy(a-)	<i>DO*01N.02</i>	c.343_350del	2	p.Met115Hisfs*18	PMID: 11552072	AH011373	rs587777832
DO:-3 or Gy(a-)	<i>DO*01N.03</i>	c.219delT	2	p.Val73Valfs*5	(4), Abstract	n.a.	n.a.
DO:-4 or Gy(a-)	<i>DO*01N.04</i>	c.730dupG (published as c.728_729insG)	2	p.Glu244Glyfs*8	PMID: 33190238	MN082686	rs769684528
DO:-5 or Gy(a-)	<i>DO*01N.05</i>	c.93G>A c.370delT	2	p.Leu31Leu p.Leu124Cysfs*5	PMID: 33206405	MT747635	rs4106889023 rs2137544113
DO:-6 or Gy(a-)	<i>DO*01N.06</i>	c.201delA	2	p.Gly68Alafs*10	(5), Abstract	n.a.	n.a.
DO:-3 or Gy(a-)	<i>DO*02N.01</i>	c.145-2A>G c.793A>G	2	Aberrant splicing	PMID: 11724986	AY029516	rs587777831 rs11276
DO:-3 or Gy(a-)	<i>DO*02N.02</i>	c.144+2T>C c.793A>G	2	Aberrant splicing	PMID: 12028057	AH011372	rs587777833 rs11276
DO:-3 or Gy(a-)	<i>DO*02N.03</i>	c.185T>C c.793A>G	2	p.Phe62Ser p.Asn265Asp	PMID: 17655578	EF178609	rs150640567 rs11276
DO:-3 or Gy(a-)	<i>DO*02N.04</i>	c.268C>T c.793A>G	2	p.Gln90Ter p.Asn265Asp	PMID: 25865759	LC011479	rs759901596 rs11276

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- Abstract (2) Karamatic Crew V, Thornton N, Bullock T, et al. Serological and molecular characterization of DOLC, a novel high incidence antigen in the Dombrock blood group system. *Vox Sang* 2013; 105 (Suppl.1), 30
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- PMID 11552072 Rios M, Storry JR, Hue-Roye K, et al. Two molecular bases for the Dombrock null phenotype. *Br J Haematol* 2002; 117:765-767
- Abstract (4) Vrignaud C, Ramelet S, Laiguillon G, et al. Characterization of a novel DO*01 silent allele caused by a nucleotide deletion mechanism and responsible for a Gy(a-) phenotype in a patient of French European ancestry with anti-Gy^a. *Transfusion* 2019, 59 (Suppl), 18A
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- PMID 33206405 Morin,P.-A., Ethier,C., Lavoie,J., Robitaille,N. and Baillargeon,N. A novel variant DO*A allele with a c.370delT mutation leading to a DO null phenotype in a Syrian family. *Transfusion* 2020, online, DOI 10.1111/trf.16193

- Abstract (5) Lubenow N, Petersen B, Sandberg M, Claesson-Linder Y, Jöud M, Storry J. Novel single nucleotide deletion in ART4 accounts for the Gy(a-) phenotype in a woman of Lebanese origin. *Vox Sang* 2020; 115(Suppl)
- PMID 11724986 Lucien N, Celton J-L, Le Pennec P-Y, et al. Short deletion within the blood group Dombrock locus causing a Do_{null} phenotype. *Blood* 2002; 100:1063-1064
- PMID 12028057 Rios M, Hue-Roye K, Storry JR, et al. Molecular basis of the Dombrock null phenotype. *Transfusion* 2001; 41:1405-1407
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- PMID 25865759 Onodera T, Tsuneyama H, Ogasawara K, et al. A novel DO null allele with a c.268C>T (p.Gln90Stop) mutation in Japanese. *Vox Sang* 2015, 109:191-3

Track of changes		from version	to version
1	Version	v4.1	v5.0 30-JUN-2021
2	Author	created: Lilian Castilho, v4.1	Lilian Castilho, June 2021
3	Review	reviewed: n.a.	Barbera Veldhuisen, June 2021
4	General	Last word version publised on ISBT website	First Excel map version. Spread-sheets "Intro", "Allele Table", "References", and "Versioning" created.
5	Intro	Text changed	The Dombrock blood group system consists of 10 antigens carried on a GPI-linked glycoprotein (DO, ART4, CD297) that consists of 314 amino acids. It has a leader sequence and a GPI motif, both of which are cleaved from the membrane bound protein.
6	Intro	LRG ID line added: n.a.	LRG_807
7	Intro	Reference allele line moved from Allele Table to Intro: n.a.	Reference allele <i>DO*01</i> encodes DO1, DO3, DO4, DO5, DO6, DO7, DO8, DO9, DO10
8	Intro	Antithetical Antigens line created in Intro: n.a.	Antithetical antigens: [DO1 DO2]
9	Allele Table		Table columns "(Reference No.) PMID", "Accession number" and "rs-number" added, content added.
10	Allele Table	Text change: Reference allele <i>DO*01</i> encodes DO1, DO3, DO4, DO5, DO6, DO7, DO8, DO9, DO10 Line moved to Intro:	see above
11	Allele Table	Allele added: n.a.	<i>DO*01N.03</i>
12	References	Abstract added n.a.	Abstract. Vrignaud C, Ramelet S, Laiguillon G, et al. Characterization of a novel <i>DO*01</i> silent allele caused by a nucleotide deletion mechanism and responsible for a Gy(a-) phenotype in a patient of French European ancestry with anti-Gya. Transfusion 2019, 59 (Suppl), 18A

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1	Version	v4.1	v5.0 30-JUN-2021
13	Allele Table Allele added:	n.a.	<i>DO*01N.04</i>
14	References PMID added	n.a.	PMID: 33190238. Bub CB, Aravechia MG, Santos L, et al. A novel DO*01 silent allele associated with a nucleotide insertion in a Brazilian patient with anti-Gya. Transfusion 2020, online , DOI: 10.1111/trf.16190
15	Allele Table Allele added:	n.a.	<i>DO*01N.05</i>
16	References PMID added	n.a.	PMID: 33206405. Morin,P.-A., Ethier,C., Lavoie,J., Robitaille,N. and Baillargeon,N. A novel variant DO*A allele with a c.370delT mutation leading to a DO null phenotype in a Syrian family. Transfusion 2020, online, DOI 10.1111/trf.16193
17	Allele Table Allele added:	n.a.	<i>DO*01N.06</i>
18	References Abstract added	n.a.	Abstract. Lubenow N, Petersen B, Sandberg M, Claesson-Linder Y, Jöud M, Storry J. Novel single nucleotide deletion in ART4 accounts for the Gy(a-) phenotype in a woman pf Lebanese origin. Vox Sang 2020; 115(Suppl)
19	References References new:	n.a.	All references from abstract (4) to PMID 25865759 added for the first time.
20	End Version	v4.1	v5.0 30-JUN-2021