



Jul-28-2021 **WFSR** 

Report: test

## <u>Allergen Results</u>

#### Allergen database.

The Allermatch database (AllergenDB) was constructed combining three different external sources. UniProt (SwissProt) allergens (https://www.uniprot.org/docs/allergen), COMPARE (Comprehensive Protein Allergen Resource) (http://db.comparedatabase.org/) and the WHO/IUIS Allergen database (http://www.allergen.org/). Overlap between the three database sources were removed using the order: UniProt > WHO/IUIS > COMPARE. If possible, the GenBank accessions were converted to UniProt accessions. This resulted in the AllergenDB with the size of 2277.

Last database update: 02/14/2021

#### FASTA.

1

For homology search Allermatch uses FASTA(v36.3.8h) with default the parameters(matrix = BLOSUM50(15:-5), open = -10, extend = -2, ktup = 2)

#### Identity search to known allergens.

Each of the ORF sequences has been compared with the AllergenDB using three different in-sillico similarity search methods:

- A full FASTA alignment, where each ORF is compared to the AllergenDB, and the top hits are retained. All the hits are sorted on E-value and the top 100 with the lowest E-value are shown. For alignments shorter than 80 amino acids, a recalculated identity(Recalc) has been calculated to a hypothetical 80-amino-acid window.
- An 80-mer sliding window search, where each ORF is chopped up in 80-amino-acid windows with steps of a single residue. Each 80-amino-acid window is compared with all sequences in the AllergenDB. Each 80-amino-acid window with a identity of >35% is shown. For alignments shorter than 80 amino acids, a recalculated identity score over a hypothetical 80 amino acid window has been calculated. For example, 40% identity on a stretch of 40 aa converts to 20% identity on an 80 aa window.
- An 6-mer search, will look for an exact hit of >6 or more contiguous amino acids to a sequence in the AllergenDB.

#### Top 100 full-FASTA search results

Based on E-Value

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	LYSC_CHICK	P00698	129	2.3e-61	129	100.0	na

<u>kvfgr</u>celaaamkrhgldnyrgyslgnwvcaakfesnfntqatnrntdgstdygilqinsrwwcndgrtp

KVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTP

GSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGCR 147 GSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGCR 129





2

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	XP_014705584	XP_01470558	<b>4</b> 148	2.3e-32	150	48.0	na

VQAWIRGCR 147 ....: LSEYLASCN 148

3

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	LYSC1_HORSE	P11376	129	1.6e-30	130	49.2	na

4

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	LALBA_BOVIN	P00711	123	1.4e-18	120	40.0	na

 25
 RCELA
 RAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTPGSRN

 5
 KCEVFRELK--DLKGYGGVSLPEWVCTTFHTSGYDTQAIVQNND-STEYGLFQINNKIWCKDDQNPHSSN

5

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	TLP_ACTCC	P83958	201	1.8	60	31.7	23.77





6

7

8

9

10

11

12

13

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	4X9U_A	4X9U_A	189	2.3	53	26.4	17.49

94 CNIPOSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

Hit ID Identity Length Accession Length E-value Overlap Recalc >LYSC\_CHI 147 TLP\_ACTDE P81370 201 2.4 32.3 25.03

FNTQATNRNTDGSTDYGI---LQINSRWWC-NDGRTPGSRNLCNIPCSALLSSDITA-SVNC 112

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	NLTP_MAIZE	P19656	93	2.5	41	34.1	17.48

85 GRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMN 125

Hit ID Accession Length E-value Overlap Identity Recalc >LYSC\_CHI 147 L7TY87\_ACTDE L7TY87 189 3.1 53 26.4 17.49

<u>94 CNIPO</u>SALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	KIWEL_ACTDE	P84527	189	3.1	53	26.4	17.49

<u>% CNIP</u>SALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

4 CNGPCRDL--NDCDGQLICIKGKCNDDPQVGTHI----CRGTTPSPQPGGC 49

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	L7TUI7_ACTDE	L7TUI7	189	3.7	53	26.4	17.49

<u>% CNIP</u>SALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	E7CLQ1_PRUAV	E7CLQ1	91	3.9	52	30.8	20.02

86 GSRNICNIPCSALLSSDITASVNCAKKIVSDGNGMNA--WVAWRNRCKGTD 138

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	NLTP_PRUAV	Q9M5X8	91	3.9	52	30.8	20.02

86 GSRNICNIPCSALLSSDITASVNCAKKIVSDGNGMNA--WVAWRNRCKGTD 138

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1	4

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	E7CLQ2_PRUAV	E7CLQ2	91	3.9	52	30.8	20.02

89 GSRNICNIPCSALLSSDITASVNCAKKIVSDGNGMNA--WVAWRNRCKGTD 138

	ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
15	>LYSC_CHI	147	Q1M0Y6_PERAM	Q1M0Y6	331	4.5	29	41.4	15.01
				-	-	-	-	•	

9% ILQINSRWWCNDGRTPGSRNLCNIPCSAL 101

16

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	E7BQV5_PERAM	E7BQV5	331	4.5	29	41.4	15.01

55 ILQINSRWWCNDGRTPGSRNLCNIPCSAL 101

17

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	L7TUJ3_ACTER	L7TUJ3	190	5.1	52	26.9	17.48

<u>%4 CNIPd</u>SALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRG 145

18

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	L7TRW9_ACTER	L7TRW9	189	6	52	26.9	17.48

<u>% CNIP</u>SALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRG 145

19

ID Le	ength	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI 14	47	L7TY92_ACTER	L7TY92	190	6.1	52	26.9	17.48

64 CNIPOSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRG 145

20

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	L7TV12_ACTDE	L7TV12	189	8.3	53	24.5	16.23

<u>%4 CNIP</u>dSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

21

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	KIWEL_ACTCC	P85261	189	8.3	53	24.5	16.23

<u>%4 CNIP</u>dSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146





22

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	AMP1_FAGES	PODKH7	40	8.6	24	37.5	11.25

23

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	AMP2_FAGES	PODKH8	40	8.6	24	37.5	11.25

24

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity	Recalc
>LYSC_CHI	147	VA5_VESMG	P86870	202	9.1	36	33.3	14.98





#### Exact word and sliding window results

Shown per ORF

#### Exact word results:

Name	Accession ID	Description	Species	No exact words	Hit %
LYSC_CHICK	P00698	Lysozyme C	Gallus gallus	124	87.32

18

KVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDI TASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGCRL 147

Name	Accession ID	Description	Species	No exact words	Hit %
	k	l	DGSTDY&LQINSRWWCNDGR	IPGSRNLCNIPCSA:	LLSSDI
TASVNCAKKIVSDO	SNGMNAWVAWRNI	RCKGTDVQAWIRGCRL 129	•		

52 ESNFNT 58 ::::::

34 ESNFNT 40

Name	Accession ID	Description	Species	No exact words	Hit %
XP_014705584	XP 014705584	PREDICTED: lysozyme C, mi	Equus asinus	1	0.7
AP_014705564	XP_014703364	lk isozyme	Equus asinus	1	0.7

52 ESNFNT 58 53 ESNFNT 59

#### Sliding window results:

Name	Accession ID	Description	Species	No windows	Hit %
LYSC_CHICK	P00698	Lysozyme C	Gallus gallus	68	100.0

Alignments containing 68 window(s): (1:81-1:62) 77.5% identity (2:82-1:63) 78.8% identity (3:83-1:64) 80.0% identity (4:84-1:65) 81.2% identity (5:85-1:66) 82.5% identity (6:86-1:67) 83.8% identity (7:87-1:68) 85.0% identity (5:85-1:66) 82.5% identity (6:86-1:67) 83.8% identity (7:87-1:68) 85.0% identity (8:88-1:69) 86.2% identity (9:89-1:70) 87.5% identity (10:90-1:71) 88.8% identity (11:91-1:72) 90.0% identity (12:92-1:73) 91.2% identity (13:93-1:74) 92.5% identity (14:94-1:75) 93.8% identity (15:95-1:76) 95.0% identity (15:95-1:76) 95.0% identity (16:96-1:77) 96.2% identity (17:97-1:78) 97.5% identity (19:99-1:80) 100.0% identity (20:100-2:81) 100.0% identity (21:101-3:82) 100.0% identity (22:102-4:83) 100.0% identity (23:103-5:84) 100.0% identity (24:104-6:85) 100.0% identity (25:105-7:86) 100.0% identity (25:105-7:86) 100.0% identity (26:106-8:87) 100.0% identity (27:107-9:88) 100.0% identity (28:108-10:89) 100.0% identity (30:110-12:91) 100.0% identity (31:111-13:92) 100.0% identity (31:111-13:92) 100.0% identity (31:113-15:94) 100.0% identity (33:113-15:94) 100.0% identity (33:113-15:94) 100.0% identity (33:115-17:96) 100.0% identity





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(36:116-18:97) 100.0% identity (37:117-19:98) 100.0% identity (38:118-20:99) 100.0% identity (39:119-21:100) 100.0% identity (40:120-22:101) 100.0% identity (41:121-23:102) 100.0% identity (42:122-24:103) 100.0% identity (42:122-24:103) 100.0% identity (44:124-26:105) 100.0% identity (44:124-26:105) 100.0% identity (45:125-27:106) 100.0% identity (46:126-28:107) 100.0% identity (47:127-29:108) 100.0% identity (49:129-31:110) 100.0% identity (50:130-32:111) 100.0% identity (51:131-33:112) 100.0% identity (52:132-34:113) 100.0% identity (53:133-35:114) 100.0% identity (55:135-37:116) 100.0% identity (56:136-38:117) 100.0% identity (56:136-38:117) 100.0% identity (56:136-38:117) 100.0% identity (59:139-41:120) 100.0% identity (61:141-43:122) 100.0% identity (62:142-44:123) 100.0% identity (62:142-44:123) 100.0% identity (63:143-45:124) 100.0% identity (63:143-45:124) 100.0% identity (64:144-46:125) 100.0% identity (66:146-48:127) 100.0% identity (67:147-49:128) 100.0% identity (67:147-49:128) 100.0% identity (67:147-49:128) 100.0% identity (68:148-50:129) 100.0% identity
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YGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQA

WIRGCRL 148 :::::: WIRGCRL 129





Name	Accession ID	Description		Species	No windows	Hit %
XP_014705584	XP_014705584	PREDICTED: lysozyme milk is	С,	Equus asinus	68	100.0
		ozyme				

# Alignments containing 2 window(s): (1:81-1:82) 51.8% identity (2:82-2:83) 51.8% identity

STDYGILQINSRWW 82 SSDYGLFQLNNKWW 83

## Alignments containing 1 window(s): (3:83-4:84) 51.2% identity

- 3 SLLILVL-CFLPLAALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTD

YGILQINSRWWC 83 ::..:.YGLFQLNNKWWC 84

Alignments containing 8 window(s): (4:84-6:85) 49.4% identity (5:85-7:86) 50.6% identity (6:86-8:87) 50.6% identity (7:87-9:88) 50.6% identity (8:88-10:89) 49.4% identity (9:89-11:90) 49.4% identity (10:90-12:91) 49.4% identity (11:91-13:92) 49.4% identity

ILQINSRWWCNDGRTPGS 91 LFQLNNKWWCKDNKRSSS 92





Alignments of	contain	ing 3 window(s): identity identity
(12:92-13:93)	48.1%	identity `´
113-93-14-94	48 1%	identitý

(14:94-15:95) 48.1% identity

WCNDGRTPGSRNL 94 WCKDNKRSSSNAC 95

# Alignments containing 1 window(s): (15:95-16:95) 50.6% identity

- 15 AALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCN

DGRTPGSRNLC 95 :.. ::: DNKR-SSSNAC 95

#### Alignments containing 1 window(s): (16:96-17:96) 51.9% identity

- 16 ALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCND
- 17 YEAKVFSKCELAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKD

GRTPGSRNLCN 96 NKRSSS-NACN 96

Alignments containing 4 window(s): (17:97-18:97) 53.1% identity (18:98-19:98) 53.1% identity (19:99-20:99) 54.3% identity (20:100-21:100) 54.3% identity

- 17 LGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDG
  18 EAKVFSKCELAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDN

RTPGSRNLCNIPCS 100 . .: : ::: :: KR-SSSNACNIMCS 100





## Alignments containing 1 window(s): (21:101-22:101) 53.1% identity

SRNLCNIPCSA 101 : : ::: :: S-NACNIMCSK 101

Alignments containing 4 window(s): (22:102-23:102) 53.1% identity (23:103-24:103) 54.3% identity (24:104-25:104) 54.3% identity (25:105-26:105) 53.1% identity

22 GRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGS

23 SKCELAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SS

RNLCNIPCSALLSS 105 : ::: :: ::. SNACNIMCSKLLDD 105

## Alignments containing 1 window(s): (26:106-27:106) 51.9% identity

27 LAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKRSSS-NAC

NIPCSALLSSD 106 1: :: ::... NIMCSKLLDDN 106

Alignments containing 4 window(s): (27:107-28:107) 51.9% identity (28:108-29:108) 50.6% identity (29:109-30:109) 50.6% identity (30:110-31:110) 50.6% identity

IPCSALLSSDITAS 110 : :: ::...: IMCSKLLDDNIDDD 110





## Alignments containing 1 window(s): (31:111-32:111) 50.6% identity

ALLSSDITASV 111 KLLDDNIDDDI 111

Alignments containing 4 window(s): (32:112-33:112) 49.4% identity (33:113-34:113) 50.6% identity (34:114-35:114) 51.9% identity (35:115-36:115) 53.1% identity

LLSSDITASVNCAK 115 ::...: LLDDNIDDDISCAK 115

## Alignments containing 1 window(s): (36:116-37:116) 53.1% identity

DITASVNCAKK 116 .: ...::. NIDDDISCAKR 116

Alignments containing 4 window(s): (37:117-38:117) 51.9% identity (38:118-39:118) 53.1% identity (39:119-40:119) 53.1% identity (40:120-41:120) 54.3% identity

ITASVNCAKKIVSD 120 : ...:: : : : : : IDDDISCAKRVVRD 120





## Alignments containing 1 window(s): (41:121-42:121) 53.1% identity

- 42 YSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFOLNNKWWCKDNKRSSS-NACNIMCSKLLDDNIDDD

VNCAKKIVSDG 121 .:::.:: ISCAKRVVRDP 121

Alignments containing 4 window(s): (42:122-43:122) 51.9% identity (43:123-44:123) 51.9% identity (44:124-45:124) 51.9% identity (45:125-46:125) 51.9% identity

42 SLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASV

NCAKKIVSDGNGMN 125 .:::..: : .::.
SCAKRVVRDPKGMS 125

## Alignments containing 1 window(s): (46:126-47:126) 51.9% identity

KIVSDGNGMNA 126 . . : : . : : . : RVVRDPKGMSA 126

Alignments containing 4 window(s): (47:127-48:127) 51.9% identity (48:128-49:128) 50.6% identity (49:129-50:129) 50.6% identity (50:130-51:130) 51.9% identity

47 VCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKK

48 VCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDDNIDDDISCAKR

IVSDGNGMNAWVAW 130 .: : .::.:: :: VVRDPKGMSAWKAW 130





## Alignments containing 1 window(s): (51:131-52:131) 50.6% identity

GNGMNAWVAWR 131 .::.:: PKGMSAWKAWV 131

Alignments containing 4 window(s): (52:132-53:132) 50.6% identity (53:133-54:133) 50.6% identity (54:134-55:134) 50.6% identity (55:135-56:135) 50.6% identity

- 52 FESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDG
- 53 YESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDDNIDDDISCAKRVVRDP

NGMNAWVAWRNRCK 135 .:..: :: ..:: KGMSAWKAWVKHCK 135

## Alignments containing 1 window(s): (56:136-57:136) 49.4% identity

AWVAWRNRCKG 136 :: :: .:: AWKAWVKHCKD 136

Alignments containing 3 window(s): (57:137-58:137) 48.1% identity (58:138-59:138) 48.1% identity (59:139-60:139) 46.9% identity

WVAWRNRCKGTDV 139 : :: ..:: :.
WKAWVKHCKDKDL 139





Alignments containing 1 window(s): (60:140-62:140) 45.0% identity

- Alignments containing 1 window(s): (61:141-63:141) 45.0% identity

Alignments containing 4 window(s): (62:142-64:142) 45.0% identity (63:143-65:143) 45.0% identity (64:144-66:144) 45.0% identity (65:145-67:145) 43.8% identity

- 62 NRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRN
- 64 GKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDDNIDDDISCAKRVVRDPKGMSAWKAWVK

RCKGTDVQAWIRG 145 HCKDKDLSEYLAS 145

#### Alignments containing 1 window(s): (66:146-68:146) 45.0% identity

- 66 DGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146 68 NGSSDYGLFQLNNKWWCKDNKRSSS-NACNIMCSKLLDDNIDDDISCAKRVVRDPKGMSAWKAWVKHCKDKDLSEYLASC 146
- Alignments containing 2 window(s): (67:147-69:147) 45.0% identity (68:148-70:148) 45.0% identity

DVQAWIRGCRL 148 :...:: DLSEYLASCNL 148





Name	Accession ID	Description	Species	No windows	Hit %
LYSC1_HORSE	P11376	Lysozyme C, milk isozyme	Equus caballus	68	100.0

# Alignments containing 14 window(s): (1:81-1:63) 43.2% identity (2:82-1:64) 44.4% identity (3:83-1:65) 45.7% identity (4:84-1:66) 45.7% identity (5:85-1:67) 46.9% identity (6:86-1:68) 46.9% identity (7:87-1:69) 46.9% identity (8:88-1:70) 46.9% identity (9:89-1:71) 46.9% identity (10:90-1:72) 46.9% identity (11:91-1:73) 48.1% identity (11:91-1:73) 48.1% identity (12:92-1:74) 48.1% identity (13:93-1:75) 48.1% identity (14:94-1:76) 48.1% identity 1 MRSLLILVLCFLPLAALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGST SINCE STORM STORM

## Alignments containing 1 window(s): (15:95-1:76) 50.6% identity

## Alignments containing 1 window(s): (16:96-1:77) 51.9% identity





Alignments containing 4 window(s): (17:97-1:78) 53.1% identity (18:98-1:79) 53.1% identity (19:99-1:80) 54.3% identity (20:100-2:81) 54.3% identity

RTPGSRNLCNIPCS 100 KR-SSSNACNIMCS 81

## Alignments containing 1 window(s): (21:101-3:82) 53.1% identity

21 FGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPG

FSKCELAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKRSS

SRNLCNIPCSA 101 : : ::: :: S-NACNIMCSK 82

Alignments containing 4 window(s): (22:102-4:83) 53.1% identity (23:103-5:84) 54.3% identity (24:104-6:85) 54.3% identity (25:105-7:86) 53.1% identity

SKCELAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SS

RNLCNIPCSALLSS 105 SNACNIMCSKLLDE 86

## Alignments containing 1 window(s): (26:106-8:87) 51.9% identity

26 LAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLC

LAHKLKAQEMDGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKRSSS-NAC

NIPCSALLSSD 106 NIMCSKLLDEN 87





Alignments containing 4 window(s): (27:107-9:88) 51.9% identity (28:108-10:89) 50.6% identity (29:109-11:90) 50.6% identity (30:110-12:91) 50.6% identity

IPCSALLSSDITAS 110 : :: ::..: IMCSKLLDENIDDD 91

# Alignments containing 1 window(s): (31:111-13:92) 50.6% identity

31 KRHGLDNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCS

ALLSSDITASV 111 ::..: .. KLLDENIDDDI 92

Alignments containing 4 window(s): (32:112-14:93) 49.4% identity (33:113-15:94) 50.6% identity (34:114-16:95) 51.9% identity (35:115-17:96) 53.1% identity

LLSSDITASVNCAK 115 LLDENIDDDISCAK 96

## Alignments containing 1 window(s): (36:116-18:97) 53.1% identity

36 DNYRGYSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSS

18 DGFGGYSLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKRSSS-NACNIMCSKLLDE

DITASVNCAKK 116 .: ...::. NIDDDISCAKR 97





Alignments containing 4 window(s): (37:117-19:98) 51.9% identity (38:118-20:99) 53.1% identity (39:119-21:100) 53.1% identity (40:120-22:101) 54.3% identity

ITASVNCAKKIVSD 120 

# Alignments containing 1 window(s): (41:121-23:102) 53.1% identity

41 YSLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITAS

VNCAKKIVSDG 121 .:::..:: ISCAKRVVRDP 102

Alignments containing 4 window(s): (42:122-24:103) 51.9% identity (43:123-25:104) 51.9% identity (44:124-26:105) 51.9% identity (45:125-27:106) 51.9% identity

42 SLGNWVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASV

24 SLANWVCMAEYESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDENIDDDI

NCAKKIVSDGNGMN 125 SCAKRVVRDPKGMS 106

## Alignments containing 1 window(s): (46:126-28:107) 51.9% identity

46 WVCAAKFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAK

KIVSDGNGMNA 126 : : .::.: RVVRDPKGMSA 107





Alignments containing 4 window(s): (47:127-29:108) 51.9% identity (48:128-30:109) 50.6% identity (49:129-31:110) 50.6% identity (50:130-32:111) 51.9% identity

IVSDGNGMNAWVAW 130 .: : .::.:: ::
VVRDPKGMSAWKAW 111

# Alignments containing 1 window(s): (51:131-33:112) 50.6% identity

51 KFESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSD 

GNGMNAWVAWR 131 PKGMSAWKAWV 112

Alignments containing 4 window(s): (52:132-34:113) 50.6% identity (53:133-35:114) 50.6% identity (54:134-36:115) 50.6% identity (55:135-37:116) 50.6% identity

52 FESNFNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDG 34 YESNFNTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDENIDDDISCAKRVVRDP

NGMNAWVAWRNRCK 135 KGMSAWKAWVKHCK 116

## Alignments containing 1 window(s): (56:136-38:117) 49.4% identity

56 FNTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMN

AWVAWRNRCKG 136 :: :: ..:: AWKAWVKHCKD 117





Alignments co	ntaining 3 window(s):
(57:137-39:118)	) 48.1% identity ` ` (
(58:138-40:119)	48.1% identity
750-130-41-120	

57 NTQATN-RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNA 39 NTRAFNGKNANGSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDENIDDDISCAKRVVRDPKGMSA

WVAWRNRCKGTDV 139 WKAWVKHCKDKDL 120

# Alignments containing 1 window(s): (60:140-43:121) 45.0% identity

 $60\ \mathtt{ATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQ}\ 140$ 

## Alignments containing 1 window(s): (61:141-44:122) 45.0% identity

Alignments containing 4 window(s): (62:142-45:123) 45.0% identity (63:143-46:124) 45.0% identity (64:144-47:125) 45.0% identity (65:145-48:126) 43.8% identity

62 NRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRN

45 GKNANGSSDYGLFOLNNKWWCKDNKR-SSSNACNIMCSKLLDENIDDDISCAKRVVRDPKGMSAWKAWVK

RCKGTDVOAWIRG 145 HCKDKDLSEYLAS 126

## Alignments containing 1 window(s): (66:146-49:127) 45.0% identity

66 DGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDVQAWIRGC 146

# Alignments containing 2 window(s): (67:147-50:128) 45.0% identity (68:148-51:129) 45.0% identity

67 GSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGT 50 GSSDYGLFQLNNKWWCKDNKR-SSSNACNIMCSKLLDENIDDDISCAKRVVRDPKGMSAWKAWVKHCKDK

DVQAWIRGCRL 148 DLSEYLASCNL 129





Name	Accession ID	Description	Species	No windows	Hit %
LALBA_BOVIN	P00711	Alpha-lactalbumin	Bos taurus	59	86.76

# Alignments containing 1 window(s): (9:89-3:67) 36.2% identity 3 LT------KCEVFRELK-DLKGYGGVSLPEWVCTTFHTSGYDTQAIVQNND-STEYGLFQINNKIWCKDDQNP 67 Alignments containing 1 window(s): (11:91-2:69) 37.5% identity Alignments containing 3 window(s): (12:92-1:70) 37.5% identity (13:93-2:71) 38.8% identity (14:94-3:72) 38.8% identity 12 LPLAALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWW EQLT-----KCEVFRELK--DLKGYGGVSLPEWVCTTFHTSGYDTQAIVQNND-STEYGLFQINNKIW CNDGRTPGSRNL 94

Alignments containing 3 window(s): (15:95-1:73) 41.2% identity (16:96-2:74) 42.5% identity (17:97-3:75) 43.8% identity

CKDDONPHSSNI 72

15 AALGKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCND GRTPGSRNLCNI 97 DONPHSSNICNI 75

# Alignments containing 1 window(s): (18:98-1:76) 41.2% identity

18 GKVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIP 98 





```
Alignments containing 7 window(s): (19:99-1:77) 42.5% identity (20:100-2:78) 42.5% identity (21:101-3:79) 42.5% identity (22:102-4:80) 42.5% identity (23:103-5:81) 43.8% identity (24:104-6:82) 43.8% identity (25:105-7:83) 42.5% identity
 GSRNLCNIPCSALLSS 105
            : ::::::::
        HSSNICNISCDKFLDD 83
Alignments containing 17 window(s): (26:106-6:84) 41.2% identity (27:107-7:85) 41.2% identity (28:108-8:86) 42.5% identity (29:109-9:87) 42.5% identity (30:110-10:88) 42.5% identity (31:111-11:89) 42.5% identity (32:112-12:90) 42.5% identity (33:113-13:91) 43.8% identity (34:114-14:92) 43.8% identity (35:115-15:93) 45.0% identity (36:116-16:94) 45.0% identity (37:117-17:95) 46.2% identity (38:118-18:96) 46.2% identity (39:119-19:97) 45.0% identity (40:120-20:98) 45.0% identity (41:121-21:99) 43.8% identity (42:122-22:100) 43.8% identity
 26 LAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCN
       CEVFRELKDLKGYGGVSLPEWVCTTFHTSGYDTOAIVONND-STEYGLFOINNKIWCKDDONPHSSNICN
         IPCSALLSSDITASVNCAKKIVSDGN 122
         ISCDKFLDDDLTDDIMCVKKILDKVG 100
Alignments containing 16 window(s): (43:123-23:100) 45.0% identity (44:124-24:101) 43.8% identity (45:125-25:102) 45.0% identity (46:126-26:103) 45.0% identity (47:127-27:104) 45.0% identity (48:128-28:105) 43.8% identity (49:129-29:106) 43.8% identity (50:130-30:107) 43.8% identity (51:131-31:108) 43.8% identity (52:132-32:109) 43.8% identity (53:133-33:110) 43.8% identity (54:134-34:111) 45.0% identity (55:135-35:112) 43.8% identity (56:136-36:113) 43.8% identity (57:137-37:114) 43.8% identity (58:138-38:115) 43.8% identity
 43 LGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNC
                                                                  . : . : ` : : . : : . . . . . . .
 23 LPEWVCTTFHTSGYDTQAIVQNND-STEYGLFQINNKIWCKDDQNPHSSNICNISCDKFLDDDLTDDIMC
        AKKIVSDGNGMNAWVAWRNRCKGTD 138
         VKKIL-DKVGINYWLAHKALCSEKL 115
```





## Alignments containing 1 window(s): (59:139-39:117) 43.2% identity

39 QAIVONND-STEYGLFQINNKIWCKDDQNPHSSNICNISCDKFLDDDLTDDIMCVKKIL-DKVGINYWLA

WRNRC-KGTDV 139 HKALCSEKLDQ 117

Alignments containing 2 window(s): (60:140-40:117) 42.5% identity (61:141-41:118) 41.2% identity

60 ATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAW

RNRCKGTDVQA 141 KALCSEKLDQW 118

Alignments containing 3 window(s): (62:142-42:118) 41.2% identity (63:143-43:119) 41.2% identity (64:144-44:120) 41.2% identity

42 VONND-STEYGLFOINNKIWCKDDONPHSSNICNISCDKFLDDDLTDDIMCVKKIL-DKVGINYWLAHKA

RCKGTDVQAWIR 144 LCS-EKLDOWLC 120

# Alignments containing 1 window(s): (65:145-44:121) 38.8% identity

# Alignments containing 3 window(s): (66:146-45:120) 40.0% identity (67:147-46:121) 40.0% identity (68:148-47:122) 40.0% identity

66 DGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKG ::.::..

TDVQAWIRGCRL 148 EKLDOWL--CEK 122





Jul-28-2021 **WFSR** 

Report: test

#### **Celiac Results**

#### Celiac database.

The epitope database is constructed by the combination of three different sources. Epitopes from Sollid and coauthors (https://pubmed.ncbi.nlm.nih.gov/31735991/), ProPepper (https://www.propepper.net/epitope) and AllergenOnline (http://www.allergenonline.org/celiacbrowse.shtml) were combined and duplicates were removed.

Last database update: 02/14/2021

#### FASTA.

For homology search Allermatch uses FASTA(v36.3.8h) with default the parameters(matrix = BLOSUM50(15:-5), open = -10, extend = -2, ktup = 2)

#### Identity search to known celiac epitopes.

Each of the ORF sequences has been compared with the epitope database using three different in-sillico similarity search methods:

- A partial epitope match, where each epitope from the epitope database is compared to each ORF and the top hits are retained. All the hits are sorted on E-value and the top 100 with the lowest E-value are shown.
- An identical epitope match, where each epitope from the epitope database is compared to each ORF and only 100% hits are retained.
- A Q/E-X1-P-X2 motif search, where each of the possible 50 motif combinations is matched to each ORF. Additional a motif occurence is calculated, which indicates how many times a certain motif is present in the epitope database.

#### Top 100 epitope match results

Based on E-Value

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Glt-156 minimal epit	15	0.013	7	57.1	71.4	AllergenOnline
		-	ubmed.n	 cbi.nlm.nih	l n.gov/15714	<u> </u>  306		

53 ESNFNTQ 59 ESPFSQQ 15

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glt-156 minimal epit						
K	147	ope in considered De	15	0.013	7	57.1	71.4	AllergenOnline
		amidated form						
	https://pubmod.nghi.nlm.nih.gov/15714206							

53 ESNFNTQ 59 ESPFSQQ 13

1





3

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glt-156 minimal epit						
K	147	ope in considered De	15	0.016	7	57.1	71.4	AllergenOnline
		amidated form						
	200 // 200 200 // 200 200 // 20							

https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNFNTQ 59 ::::: 8 ESPFSQQ 14

4

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		LMW glutenin-glt-156						
K	147	(p40-p59; E48 and E	20	0.024	7	57.1	71.4	AllergenOnline
		51)						
	https://pubmed.nchi.nlm.nih.gov/12055577							

53 ESNFNTQ 59

:: :. : 12 ESPFSQQ 18

5

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	LMW glutenin-glt-156 (p40-p59; E51)	20	0.025	7	57.1	71.4	AllergenOnline
	https://pubmod.nghi.nlm.nih.gov/12055577							

53 ESNFNTQ 59 ::::: 12 ESPFSQQ 18

6

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	alpha gliadin 123-13 2	10	0.03	3	100.0	100.0	ProPepper
https://www.iedb.org/epitope/51376								

96 IPC 98 ::: 5

7

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	alpha-gliadin(pl23-p 132)	10	0.031	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/12594302								
https://pubmed.nchi.nlm.nih.gov/18395083								

96 IPC 98

::: 3 IPC 5





8

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	gamma5-gliadin (p227 -237; E232)	11	0.031	4	75.0	100.0	AllergenOnline
	https://pubmed_nchi_nlm_nih_gov/15972656							

72 GILQ 75 ::.: 1 GIIQ 4

9

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	Wheat peptide W09	14	0.034	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

11 FLP 13 :::

5 FLP 7

10

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		gamma5-gliadin (p227						
К	147	-p237) ; gamma-II ep	11	0.035	4	75.0	100.0	AllergenOnline
		itope						

https://pubmed.ncbi.nlm.nih.gov/15972656

72 GILQ 75 ::.: 1 GIIQ 4

11

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	DQ2-gamma-II peptide	11	0.037	4	75.0	100.0	ProPepper
https://www.iedb.org/epitope/20305								

72 GILQ 75 ::.: 1 GIIQ 4

12

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	alpha-Glia AG11 (p78	17	0.039	2	100.0	100.0	AllergenOnline
K	14/	-p95)	' '	0.039	3	100.0	100.0	Affergenomine
	https://pubmed.nghi.nlm.nih.gov/10200712							

11 FLP 13 ::: 6 FLP 8





13

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Wheat peptide W09	12	0.039	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

11 FLP 13 ::: 4 FLP 6

14

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Wheat peptide W09	14	0.04	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

11 FLP 13 ::: 5 FLP 7

15

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	alpha-Glia AG11 (p78 -p95; E86)	17	0.042	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.goy/19299713								

11 FLP 13 :::

6 FLP 8

16

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	alpha gliadin 78-95	17	0.046	2	100.0	100.0	DroDonnor	
K		AG11	1 /	0.046	3	100.0	100.0	ProPepper	
	https://www.iedb.org/epitope/109723								

11 FLP 13 ::: 6 FLP 8

**17** 

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		alpha gliadin P11	15	0.046	3	100.0	100.0	ProPepper
К	147	aipha giladin Pil	15	0.046	3	100.0	100.0	Propepper
https://www.iedb.org/epitope/139589								

11 FLP 13 ::: 7





18

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	1 47	gamma gliadin 1		0.049	7	42.9	71.4	7.1.1	
К	147	gamma giladin i	9	0.049	′	44.7	/1.4	AllergenOnline	
	https://pubmed.ncbi.nlm.nih.gov/32728397								
	https://pubmed.ncbi.nlm.nih.gov/22726570								

53 ESNFNTQ 59 :..: : 2 EQSFPQQ 8

19

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC K	147	gamma gliadin	9	0.051	7	42.9	71.4	AllergenOnline		
	https://pubmed.ncbi.nlm.nih.gov/32728397									
https://pubmed.ncbi.nlm.nih.gov/22726570										

53 ESNFNTQ 59

20

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Rye peptide R12	12	0.057	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

12 LPL 14

::: 10 LPL 12

21

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	Wheat peptide W36	15	0.063	9	44.4	88.9	AllergenOnline	
https://pubmed.ncbi.nlm.nih.gov/20650871									

1 MRSLLILVL 9

.:::...: 5 IRSLVLRTL 13

22

ID	Length	Hit ID		Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Wheat peptide	W36	15	0.064	9	44.4	88.9	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871									

1 MRSLLILVL 9

.:::...: 5 IRSLVLRTL 13





23

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Wheat peptide W09	20	0.067	2	100.0	100.0	AllergenOnline
K	147	wheat peptide wos	20	0.007	3	100.0	100.0	Affergenonfine
https://pubmed.nchi.nlm.nih.gov/20650871								

11 FLP 13

::: 10 FLP 12

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glia-gamma30-gliadin						
K	147	(p222-236; E225 and	15	0.067	4	75.0	100.0	AllergenOnline
		E231)						

https://pubmed.ncbi.nlm.nih.gov/12055577

72 GILQ 75 ::.: 5 GIIQ 8

25

24

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	Wheat peptide W36	16	0.07	9	44.4	88.9	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

1 MRSLLILVL 9

:::...: 5 IRSLVLRTL 13

26

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC K	147	Rye peptide R09	12	0.071	3	100.0	100.0	AllergenOnline	
https://pubmed.ncbi.nlm.nih.gov/20650871									

12 LPL 14 ::: 2 LPL 4

27

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	R12E	12	0.072	3	100.0	100.0	ProPepper
https://www.iodh.org/opitops/22002								

12 LPL 14

::: 10 LPL 12





28

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Rye peptide R12	12	0.073	3	100.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

12 LPL 14 ::: 10 LPL 12

29

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	w09	20	0.075	3	100.0	100.0	ProPepper	
https://www.iedb.org/epitope/238729									

11 FLP 13 ::: 10 FLP 12

30

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC K	147	W36s	16	0.076	9	44.4	88.9	ProPepper	
https://www.iedb.org/epitope/238963									

1 MRSLLILVL 9 :::...: 5 IRSLVLRTL 13

31

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Wheat peptide W37	12	0.076	Ω	37.5	62.5	AllergenOnline		
K	111	wheat peptide W3/	12	0.070	0	37.3	02.5	ATTELGENOMITME		
	https://pubmed.ncbi.nlm.nih.gov/20650871									

119 DGNGMNAW 126 : .:. : DPSGQVQW 10

32

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Glia-gamma30-gliadin	1.5	0.076	4	75.0	100.0	AllergenOnline		
К	14/	(p222-236; E225)	15	0.076	4	75.0	100.0	Allergenonline		
	https://pubmed.ncbi.nlm.nih.gov/12055577									

72 GILQ 75 ::.: 5 GIIQ 8





33

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	DQ2-y -II y-Glia (p	15	0.077	A	75.0	100.0	AllergenOnline		
K		222-p236; E229)	13	0.077	7	75.0	100.0	Allergenonline		
	https://www.doc.doc.doc.doc.doc.doc.doc.doc.doc.doc									

https://pubmed.ncbi.nlm.nih.gov/19299713

72 GILQ 75 ::.: 3 GIIQ 6

3	4
J	4

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glt-156 minimal epit						
K	147	ope in considered na	15	0.077	7	42.9	71.4	AllergenOnline
		tive form						

https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNFNTQ 59 .: .: .: 7 QSPFSQQ 13

35

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC K	147	Glia-gamma30-gliadin (p222-236; E231)	15	0.081	4	75.0	100.0	AllergenOnline		
	https://pubmed.nchi.nlm.nih.gov/12055577									

72 GILQ 75 ::.: 5 GIIQ 8

36

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		Glt-156 minimal epit							
K	147	ope in considered na	15	0.081	7	42.9	71.4	AllergenOnline	
		tive form							
https://pubmed.nebi.nlm.nih.gov/15714306									

53 ESNFNTQ 59

.: :. ? QSPFSQQ 14

37

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	DQ2-y -II y-Glia (p	1 5	0.085	4	75.0	100.0	AllergenOnline	
К	14/	222-p236)	13	0.065	4	75.0	100.0	Affergenomine	
	https://pubmed.nghi.nlm.nih.gov/19290712								

72 GILQ 75 ::.: 3 GIIQ 6





38

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	HMW glutenin-glt04 (	36	0.085	29	13.8	62.1	AllergenOnline		
K		p707-p742)	30	0.085	29	13.0	02.1	Allergenonline		
	https://pubmed.ncbi.nlm.nih.gov/10540324									

39 RGYSLGNWVCAAKFESNFNTQATNRNTDGS 67

3 QGQRPGQWLQPGQGQQGYYPTSPQQSGQGQ 31

39

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database			
>LYSC_CHIC	147	Wheat peptide W36	20	0.086	9	44.4	88.9	AllergenOnline			
	https://pubmed.ncbi.nlm.nih.gov/20650871										

1 MRSLLILVL 9

.:::...: 9 IRSLVLRTL 17

40

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		LMW glutenin-glt-156						
K	147	(p46-p54; E48 and E	9	0.086	4	75.0	75.0	AllergenOnline
		51)						
https://www.durabi.nlm.nib.mov/16001025								

https://pubmed.ncbi.nlm.nih.gov/16091925 https://pubmed.ncbi.nlm.nih.gov/12055577 https://pubmed.ncbi.nlm.nih.gov/32728397

53 ESNF 56

ESPF 9

41

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Rye peptide R01, R09	20	0.087	3	100.0	100.0	AllergenOnline
K		kye peptide kui, ku9	20					
https://pubmed.ncbi.nlm.nih.gov/20650871								

12 LPL 14 :::

3 LPL 5

42

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	alpha2-gliadin 1448 (p231-p245)	15	0.093	14	28.6	71.4	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/16878175								

55 NFNTQATNRNTDGS 68

2 SFQPSQQNPQAQGS 15





43

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	R01	20	0.094	3	100.0	100.0	ProPepper
https://www.iedb.org/epitope/238942								

12 LPL 14 :::: 3 LPL 5

44

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC K		alpha-gliadin 4037	17	0.094	9	44.4	66.7	AllergenOnline	
https://pubmed.ncbi.nlm.nih.gov/21091908									

71 YGILQINSR 79 .::.:: 9 FGIFGTNYR 17

45

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Glia-gamma30-gliadin (p222-p236)	15	0.094	4	75.0	100.0	AllergenOnline
https://pubmed.ncbi.nlm.nih.qov/12055577								

72 GILQ 75 ::.: 5 GIIQ 8

46

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		LMW glutenin-glt-156						
К	147	(p45-p54; E48, E49 and E51) minimal epi	10	0.095	4	75.0	75.0	AllergenOnline
		tope						
https://pubmed.ncbi.nlm.nih.gov/15714306								

53 ESNF 56 :::: 7 ESPF 10

47

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		alpha gliadin 251-27	20	0.097	0	44.4	66.7	DwoDonnow
K	147	0	20	0.097	9	44.4	00.7	ProPepper
https://www.iedb.org/epitope/72058								

71 YGILQINSR 79

12 FGIFGTNYR 20





48

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Sec-gammal in Deamid	11	0.099	7	42.9	71.4	AllergenOnline
K	14/	ated form	14	0.099	,	42.9	71.4	Allergenonline
https://pubmed.ncbi.nlm.nih.gov/14517794								

53 ESNFNTQ 59

:..: <del>:</del> 5 EQSFPEQ 11

49

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	LMW glutenin-glt-156 (p40-p59)	20	0.1	7	42.9	71.4	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/12055577								

53 ESNFNTQ 59 .: : .: 12 QSPFSQQ 18

50

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	LMW glutenin-glt-156 (p40-p59; E48)	20	0.1	7	42.9	71.4	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/12055577								

53 ESNFNTQ 59 .:..: 12 QSPFSQQ 18

51

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Glt-156 minimal epit	15	0.1	5	60.0	80.0	AllergenOnline
K	14/	ope (p41-p55)						
https://pubmed.ncbi.nlm.nih.gov/15714306								

53 ESNFN 57

:: :. 11 ESPFS 15

52

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Sec-gammal in Deamid	1.4	0 1	7	42.9	71.4	AllergenOnline		
K	14/	ated form	14	0.1	/	42.9	/1.4	Affergenomine		
	https://pubmed.ncbi.nlm.nih.gov/14517794									

53 ESNFNTQ 59





53

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	Sec-gammal in Deamid ated form	14	0.1	7	42.9	71.4	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/14517794								

53 ESNFNTQ 59

:..: <del>:</del> 5 EQSFPEQ 11

54

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	gamma gliadin 222-23	15	0.11	1	75.0	100.0	ProPepper	
K		6 DQ2-gamma-II	13	0.11	4	75.0	100.0	Propepper	
	https://www.iedb.org/epitope/109223								

72 GILQ 75 ::.: 3 GIIQ 6

55

	ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
ł	>LYSC_CHIC K	147	1448	15	0.11	14	28.6	71.4	ProPepper	
	https://www.iedb.org/epitope/22327									

55 NFNTQATNRNTDGS 68 .:..: ...:: 2 SFQPSQQNPQAQGS 15

56

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	W36	20	0.11	٥	44.4	88.9	ProPepper	
K	14/	WJU	20	0.11	)	11.1	00.9	Liorephei	
https://www.iedb.org/epitope/238943									

1 MRSLLILVL 9

.:::...: 9 IRSLVLRTL 17

57

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	DQ2.2-glut-L-1 conta	20	0.11	7	42.9	71.4	DwoDonnow		
К		ining peptide	20	0.11	<b>'</b>	42.9	71.4	ProPepper		
	https://www.iedb.org/epitope/167143									

53 ESNFNTQ 59 .: : .: 12 QSPFSQQ 18





58

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	DQ2.2-glut-L-1 conta	20	0.11	7	42.9	71.4	ProPepper	
К		ining peptide	20	0.11	,	42.9	71.4	Propepper	
	https://www.jedb.org/epitope/52130								

53 ESNFNTQ 59 .: : .: 12 QSPFSQQ 18

59

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC K	147	gliadin-gamma30 (222 -236)	15	0.11	4	75.0	100.0	ProPepper	
	https://www.iedb.org/epitope/70564								

72 GILQ 75 ::.: 5 GIIQ 8

60

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	alpha-gliadin CAB769	20	0.11	a	44.4	66.7	AllergenOnline	
K	147	61 (p251-p270)	20	0.11	9	11.1	00.7	Allergenonline	
	https://pubmed.ncbi.nlm.nih.gov/17629515								

71 YGILQINSR 79 .::. :: 12 FGIFGTNYR 20

61

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		CAUTION 100% match t						
K		o Archaea protein lo						
	147	wer to others and to	9	0.11	7	42.9	57.1	AllergenOnline
		gamma5 (p63-p71; E6						
		3, E68 and E71)						

https://pubmed.ncbi.nlm.nih.gov/15972656

53 ESNFNTQ 59 

62

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Rye peptide R12	20	0.11	2	100.0	100.0	AllergenOnline		
K	14/	kye peptide kiz	20	0.11	3	100.0	100.0	Allergenonline		
	https://pubmed.ncbi.nlm.nih.gov/20650871									

12 LPL 14 ::: 16 LPL 18





63

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Rye peptide R12	16	0.11	3	100.0	100.0	AllergenOnline		
	https://pubmed.nebi.nlm.nib.goz/20650971									

12 LPL 14 ::: 12 LPL 14

64

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC K	147	Sec-gammal in Deamid ated form	14	0.11	7	42.9	71.4	AllergenOnline	
https://pubmed.ncbi.nlm.nih.gov/14517794									

53 ESNFNTQ 59

65

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC K	147	DQ2.5-glut-L2	9	0.11	4	75.0	75.0	Sollid		
	https://pubmed.ncbi.nlm.nih.gov/15714306									
	https://pubmed.ncbi.nlm.nih.gov/12055577									

53 ESNF 56 :::: 9

66

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	Wheat peptide W37	20	0.12	8	37.5	62.5	AllergenOnline		
	https://pubmed.ncbi.nlm.nih.gov/20650871									

119 DGNGMNAW 126

: .:.

Г	ח		Length	ні+
	8	DPS	SGQVQW	15

67

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glt-156 minimal epit						
K	147	ope in considered De	15	0.12	5	60.0	80.0	AllergenOnline
		amidated form						
https://pubmed.nchi.nlm.nih.gov/15714306								

53 ESNFN 57

:: :. 10 ESPFS 14





68

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		LMW glutenin-glt-156						
К	147	(p45-p54; E48 and E	10	0.12	4	75.0	75.0	AllergenOnline
		51) minimal epitope						

https://pubmed.ncbi.nlm.nih.gov/12055577

https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNF 56 :::: 10

69

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		LMW glutenin-glt-156							
K	147	(p45-p54; E49 and E	10	0.12	4	75.0	75.0	AllergenOnline	
		49) minimal epitope							
	heter at 1/2 about 1 about 1 about 1573 4206								

https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNF 56

:::: 7 ESPF 10

70

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	LMW glutenin-glt-156	9	0.12	4	75.0	75.0	AllergenOnline		
K (p46-p54; E51) https://pubmed.ncbi.nlm.nih.gov/16091925										
	https://pubmed.ncbi.nlm.nih.gov/12055577									
	https://pubmed.ncbi.nlm.nih.gov/32728397									

53 ESNF 56 :::: : 6 ESPF 9

71

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	alpha-gliadin p19 (p	20	0.13	12	25.0	66.7	AllergenOnline	
K	14/	21-p40)	20	0.13	12	25.0	00.7	Affergenonfine	
	https://pubmed.ncbi.nlm.nih.gov/8315377								

12 LPLAALGKVFGR 23

.::. .:. 2 VPLVQQQQFLGQ 13

**72** 

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	gamma5 (p62-p72; E68 , E63 and E71)	11	0.13	7	42.9	57.1	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/15972656								

53 ESNFNTQ 59 : . : :

EQPFPEQ 8





73

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	gamma-gliadin 1383 (p141-p160)	20	0.13	6	50.0	83.3	AllergenOnline	
	https://pubmed.ncbi.nlm.nih.gov/16878175								

97 PCSALL 102

::..: 10 PCKNFL 15

74

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		LMW glutenin-glt-156						
K	147	(p45-p54; E51) mini	10	0.13	4	75.0	75.0	AllergenOnline
		mal epitope						

https://pubmed.ncbi.nlm.nih.gov/12055577 https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNF 56

:: : ESPF 10

75

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	1 4 7	glt04 (p723-p735; E7	1.2	0.13	7	42.9	57.1	AllergenOnline		
K	14/	24)	13	0.13	′	42.9	57.1	Allergenonline		
	https://pubmed.ncbi.nlm.nih.gov/22342873									
https://pubmed.ncbi.nlm.nih.gov/10540324										

84 DGRTPGS 90

EGYYPTS 8

76

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	R12	20	0.14	3	100.0	100.0	ProPepper	
https://www.iedb.org/epitope/238384									

12 LPL 14 ::: 16 LPL 18

77

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		gamma-gliadin M23 M3	20	0.14	10	40.0	70.0	AllergenOnline
K	147	6999 (221-240) homol						
		ogous to DQ2-gamma-I						
		I						
https://pubmod.nabi.nlm.nib.goz/12109706								

https://pubmed.ncbi.nlm.nih.gov/12198706 https://pubmed.ncbi.nlm.nih.gov/16878175

13 PLAALGKVFG 22 :: :...: 3 PLFQLAQGLG 12





78

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		Wheat peptide W05	20	0.14	7	42.9	71.4	AllergenOnline	
https://pubmed.ncbi.nlm.nih.gov/20650871									

53 ESNFNTQ 59

:. :. : 11 EQPFSQQ 17

79

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	Rye peptide R21	12	0.14	8	37.5	62.5	AllergenOnline
https://pubmed.ncbi.nlm.nih.gov/20650871								

122 GMNAWVAW 129 : .. : :

3 GPSGQVEW 10

80

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	W37s	16	0.15	8	37.5	62.5	ProPepper
https://www.iedb.org/epitope/238519								

119 DGNGMNAW 126

: .:. : 5 DPSGQVQW 12

81

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC K	147	gamma gliadin peptid 1383	20	0.15	6	50.0	83.3	ProPepper
https://www.iedb.org/epitope/52318								

97 PCSALL 102

::..: 10 PCKNFL 15

82

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	R21E	12	0.15	o	37.5	62.5	ProPepper
К	14/	RZIE	12	0.15	0	37.5	62.5	Propepper
https://www.iedb.org/epitope/238962								

122 GMNAWVAW 129

3 GPSGQVEW 10





83

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	alpha-gliadin (proli	16	0.15	7	42.9	42.9	AllergenOnline
K		ne-rich domain)	10	0.13	,	12.9	12.9	Allergenonline
https://pubmod.nghi.nlm.nih.gov/1720424								

48 CAAKFES 54

: :: 1 CPQPFPS 7

84

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	gamma5 (p62-p74; E63	13	0.15	7	42.9	57.1	AllergenOnline
K , E68 and E71) https://pubmed.ncbi.nlm.nih.gov/15972656								

53 ESNFNTQ 59

EQPFPEQ 8

85

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	DQ8-glut-H1, DQ8.5-g	0	0.16	6	50.0	50.0	DweDenner
K	14/	lut-H1	9	0.16	6	30.0	50.0	ProPepper
https://www.iedb.org/epitope/161144								

85 GRTPGS 90 : ::

2 GYYPTS 7

86

87

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	alpha-gliadin CAB769	20	0.16	4	50.0	100.0	AllergenOnline	
K	14/	60 (p253-p272)	40	0.10	7	30.0	100.0	ATTELGENORITINE	
	https://pubmed.ncbi.nlm.nih.qov/17629515								

93 LCNI 96

2 MCNV 5

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Glutenin-Glt-17 (p50						
K	147	-p58; E52, E53 and E	9	0.16	2	100.0	100.0	AllergenOnline
		55)						

https://pubmed.ncbi.nlm.nih.gov/16091925

13 PL 14

:: 8 PL 9





88

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		Deamidated Glt-156 m						
К	147	inimal epitope (p40-	15	0.16	4	75.0	75.0	AllergenOnline
		p59)						

https://pubmed.ncbi.nlm.nih.gov/15714306

53 ESNF 56

12 ESPF 15

89

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		Naturally occurring							
K	147	glutenins (p722-p736	15	0.16	7	57.1	57.1	711 augum 0m1 i m a	
	14/	) (homolog of glt04)	12	0.16	/	57.1	57.1	AllergenOnline	
	https://pubmed.nchi.nlm.nih.gov/10540324								

97 PCSALLS 103 : : :: 7 PTSPQLS 13

90

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	147	DQ8-glut-H1	9	0.16	6	50.0	50.0	Sollid		
	https://pubmed.ncbi.nlm.nih.gov/10540324									

85 GRTPGS 90 : ::

2 GYYPTS 7

91

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		DO8.5-glut-H1	۵	0.16	6	50.0	50.0	Sollid	
K	14/	DQ0.5-GIUU-HI	٦	0.10	0	30.0	30.0	201110	
https://pubmed.ncbi.nlm.nih.gov/22013116									

85 GRTPGS 90 : ::

2 GYYPTS 7

۵	2

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC		gamma-gliadin 1391 (						
K		p231-p250) ; gamma-g						
	147	liadin M24 M36999 (2	20	0.17	6	50.0	100.0	AllergenOnline
		31-250) identical to						
		DQ2-gamma-II						

https://pubmed.ncbi.nlm.nih.gov/12198706 https://pubmed.ncbi.nlm.nih.gov/16878175

1 MRSLLI 6

::::.. 15 IRSLVL 20





93

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC		Glutenin-Glt-17 (p50		0.17	2	100.0	100.0	AllergenOnline	
K	147	-p58; E52 and E55)	9	0.17	2	100.0	100.0	Allergenonline	
	https://wwh.al.a.hi.al.a.hi.a.nib.a.n/16001035								

13 PL 14

:: 8 PL 9

94

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	Glutenin-Glt-17 (p50 -p58; E53 and E55)	9	0.17	2	100.0	100.0	AllergenOnline	
	https://pubmed.ncbi.nlm.nih.gov/16091925								

13 PL 14 8 PL 9

95

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC K	147	Barley peptide B03	16	0.17	13	30.8	46.2	AllergenOnline		
	https://pubmed.ncbi.nlm.nih.gov/20650871									

85 GRTPGSRNLCNIP 97 : : : : : : : 1 GQQPFPQPEQPIP 13

96

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database
>LYSC_CHIC	147	DO2.2-glia-al	a	0.17	1	50.0	100.0	Sollid
K	147	DQ2.2-911a-a1	9	0.17	7	30.0	100.0	301114
https://pubmed.ncbi.nlm.nih.gov/25261484								

56 FNTQ 59

:..: 2 FSVQ 5

97

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	LMW glutenin 17 (53-	0	0.18	4	50.0	75.0	DwoDonnow	
К	14/	60)	8	0.18	4	50.0	75.0	ProPepper	
	https://www.iedb.org/epitope/47569								

56 FNTQ 59

:.: 2 FSQQ 5





98

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	G4-9A gliadin (p62-p	11	0.18	E	60.0	60.0	AllergenOnline	
K	14/	75; E65 and A70)	14	0.10	5	00.0	80.0	Arrergenomine	
	https://pubmod.ngbi.nlm.nih.gov/12524402								

https://pubmed.ncbi.nlm.nih.gov/12524403

12 LPLAA 16 :::: 5 LPYPA 9

99

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database		
>LYSC_CHIC	-   147   glt04 (p723-p734)   12   0.18   6   50.0   50.0   AllergenOnl.									
	https://pubmed.ncbi.nlm.nih.gov/10917890									

85 GRTPGS 90 : ::

3 GYYPTS 8

100

ID	Length	Hit ID	Length	E-value	Overlap	Identity	Similarity	Database	
>LYSC_CHIC	147	HMW glutenin (p724-p	11	0.18	6	50.0	50.0	AllergenOnline	
K	14/	734)							
	https://pubmed.ncbi.nlm.nih.gov/16091925								
	https://pubmed.ncbi.nlm.nih.gov/10540324								
	https://pubmed.ncbi.nlm.nih.gov/22013116								

85 GRTPGS 90 : : : 2 GYYPTS 7









Jul-28-2021 **WFSR** 

Report: test

# **Toxin Results**

#### Toxin database.

For the toxin database sequences have been retrieved from Swiss-Prot's animal toxin annotation project(https://www.uniprot.org/program/Toxins). If annotated, signal- and propeptides are removed. This results in a database with a size of 7452.

Last database update: 01/28/2021

#### FASTA.

For homology search Allermatch uses FASTA(v36.3.8h) with default the parameters(matrix = BLOSUM50(15:-5), open = -10, extend = -2, ktup = 2)

### <u>Identity search to known toxins.</u>

A full FASTA alignment search has been performed, where each ORF is compared to the sequences in the toxin database. For each ORF the top hits are retained. All the hits are sorted on E-value and the top 100 with the lowest E-value are shown.

## Top 100 full-FASTA search results

Based on E-Value

1

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	TX219_APOSC	P49272	76	0.036	82	26.8

8 RNTDGSTDYGILQINSRWWCNDGRTPG---SRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVA

ONLGSDIPHDIIKLPNGOWC---KTPGALCSSRSEC---CKAKHSDSVTYSSGCSROW-SDOOGL----

WRNRCKGTDVQA 140 FINOCRTCNVES 73

2

>LYSC_CHI 147	ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
	>LYSC_CHI	147	SCX12_CENNO	P63019	67	0.7	46	34.8

DGRTEGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAW 129

3

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	SX15F_RHOJU	E7CLP4	65	0.97	59	33.9

PGSRNLCNIPCSALLSSDITASVNC-AKKIVSDGNGMNAWVAWRNRCKGT----DVQAW 141 





4

5

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	PA2A2_HELSU	P80003	142	1	54	29.6

<u>& laaam</u>krhgldnyrgyslgnwvcaakfesnfntqatnrntd--gstdygilqin 77

43 MAALEYKHGMRNYRPHTVSHCDCDNQFRSCL-MNVKDRTADLVGMTYFTVLKIS 95

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	CRVP2_DISTY	Q2XXQ5	219	3.3	103	25.2

YRGYSLGNWVCAAKF--ESNFNTQATNRNTDGSTDYGILQI--NSRWWCNDGRTPGSRNLCNIPCSALLS 

SDITASVNCAKKIVSDGNGMNAWVAWRNRCKGT 136 EDVF--MNC-KSLVAOSNCODDYI--RKNCPAT 213

6

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	CRVP1_DISTY	Q2XXQ6	219	3.3	103	25.2

YRGYSLGNWVCAAKF--ESNFNTQATNRNTDGSTDYGILQI--NSRWWCNDGRTPGSRNLCNIPCSALLS 

SDITASVNCAKKIVSDGNGMNAWVAWRNRCKGT 136 EDVF--MNC-KSLVAQSNCQDDYI--RKNCPAT 213

7

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	SX17F_RHOJU	E7CLP5	65	4.2	55	29.1

88 PGSRNLCNIPCSALLSSDITASVNC-AKKIVSDGNGMNAWVAWRNRCKGTDVQAW 141

8

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	TX216_APOSC	P49270	76	4.4	81	22.2

8 RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIP---CSALLSSDITASVNCAKKIVSDGNGMNAWVAW

RNRCKGTDVQA 140 :.:. ::.. -NQCRTCNVES 73





9

10

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	TOPC5_PHONI	P84093	80	4.7	77	27.3

55 NTQAINRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRC 133 2 NCIELNNDCDGSKD-DCQCCRDNAYCSCYNFFGIKSGCK--CS-VGNSGTGYSV-CLKKL--ECPNRRAWTSWKKEC 71

ID :	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	PA2A3_TROCA	Q45Z28	124	6.2	54	31.5

8½ CNDGRTPGSRNLCNIPCSALLSS-DITASVNCAKKIVSDGNGMNAW-VAWRNRC 133

77 CNDGELTCKDN--NDECKAFICNCDRTAAICFARTPYNDAN----WNINTKTRC 124

Length Hit ID E-value Accession Length Overlap Identity 11 >LYSC\_CHI 147 TX214\_APOSC P49269 76

68 RNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIP---CSALLSSDITASVNCAKKIVSDGNGMNAWVAW

4 QNLGSGIPHDRIKLPNGQWC---KTPG--DLCSSSSECCKAKHSNSVTYASFCSREW-SGQQGL----F

INOCRTCHVES 73

12

13

14

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	PA2A1_TROCA	Q45Z30	124	7.5	54	31.5

82 CNDGRTPGSRNLCNIPCSALLSS-DITASVNCAKKIVSDGNGMNAW-VAWRNRC 133

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	PA2A2_TROCA	Q45Z29	124	7.5	54	31.5

82 CNDGRTPGSRNLCNIPCSALLSS-DITASVNCAKKIVSDGNGMNAW-VAWRNRC 133

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	CRVP_PHIOL	Q09GJ9	221	8.3	103	23.3

<u>YRGY</u>SLG--NWVCAAKFESNFNT--QATNRNTDGSTDYGILQINSRWWCNDGRTPGSRNLCNIPCSALLS 

SDITASVNCAKKIVSDGNGMNAWVAWRNRCKGT 136

...: :: ...:...: :. :: :: :: NEFT---NC-NELVQQSSCQDDWI--KSNCAAT 213

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15

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	CRVP_OPHHA	Q7ZT98	221	8.3	103	24.3

SDITASVNCAKKIVSDGNGMNAWVAWRNRCKGT 136 ..: :: ......: ...: ... NEYT---NC-DSLVKQSSCQDEWI--KSKCPAS 213

16

ID	Length	Hit ID	Accession	Length	E-value	Overlap	Identity
>LYSC_CHI	147	VM3B1_BOTJA	Q0NZY0	166	9.5	57	29.8

8 CNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSD----GNGMNAWVAWRNRC 133