







**Когда известно название
гена**

Search: for

e.g. Carboxy* or chx28

Favourite genomes

 Arabidopsis thaliana TAIR10	 Oryza sativa Japonica IRGSP-1.0
 Triticum aestivum TGACv1	 Hordeum vulgare ASM32608v1
 Zea mays AGPv4	 Physcomitrella patens ASM242v1

[Edit favourites](#)

All genomes

-- Select a species --

[View full list of all Ensembl Plants species](#)

What's New in Release 35

- Updated genomes

Did you know?

New Bread Wheat Genome Assembly

A new genome assembly of *Triticum aestivum* cv. Chinese Spring is now available in Ensembl Plants. The assembly (TGACv1) and it's accompanying annotation was produced by the [Earlham Institute](#), formerly The Centre for Genome Analysis (TGAC), as part of the [Triticeae Genomics for Sustainable Agriculture](#) project.

The assembly has a scaffold N50 of 88 Kbp and a total length of 13.4 Gbp in contigs greater than 500 bp ([read more](#)). The gene model annotation consists of 217,907 loci and 273,739 transcripts. A total of 104,049 protein coding genes (154,798 transcripts) and 10,156 long ncRNAs have been annotated with high confidence ([read more](#)). Approximately 99,000 genes (99% of the total) annotated on the previous IWGSC CSS assembly (MIPS) have been mapped to the new assembly.

The Axiom 35k and 820k SNP marker sets have been provided by [CerealsDB](#) and located on the new assembly ([read more](#)).

Ensembl Plants Archive Site

Alongside release 32 we have launched a new [archive site](#), where we will keep selected previous releases of Ensembl Plants publicly available. The first release available on the archive site is release 31, and includes the previous assemblies for wheat and maize.

Ensembl Plants is developed in coordination with other plant genomics and bioinformatics groups via the EBI's role in the [transPLANT](#) consortium. The transPLANT project is funded by the [European Commission](#) within its [7th Framework Programme](#), under the thematic area "Infrastructures", contract number [283496](#).



New Search Jobs

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- New Search
- Gene (1)
- Ensembl Plants (1)

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Ensembl Plants is produced in collaboration with Gramene

Search results for 'VIT_01s0011g00960'

Showing 1 Gene found in Ensembl Plants

VIT_01s0011g00960

Description	Putative uncharacterized protein [Source:UniProtKB/TrEMBL;Acc:F6HEY8]
Gene ID	VIT_01s0011g00960
Species	Vitis vinifera
Location	1:812008-816090
Gene trees	EPIGT00820000103231 (Plant Compara) EGGT001300000085269 (Pan-taxonomic Compara)

Ensembl Plants release 35 - April 2017 © EMBL-EBI

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- Ensembl Metazoa

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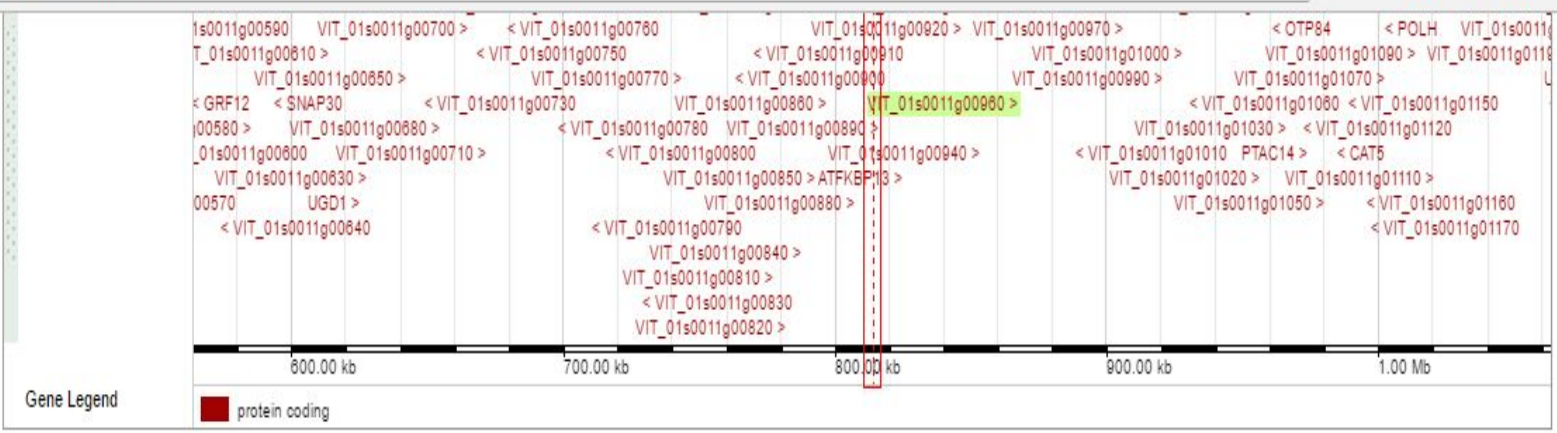
Custom tracks

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Ensembl Plants is produced in collaboration with Gramene



Location: Gene:

Navigation controls: back, forward, zoom in, zoom out, home, end

Drag/Select: [arrow icon]

812kb 813kb 814kb 815kb 816kb

Forward strand

Genes

VIT_01s0011g00960.t01 > protein coding

Contigs

< FN595752.1

Region: 1:811959-814575

Mark region (2617 bp)

Jump to region (2617 bp)

Variant - All sources

%GC

No Sequence variants (all sources)

Reverse strand

Gene Legend

■ protein coding

There are currently 46 tracks turned off.

Ensembl Plants Vitis vinifera version 88.3 (IGGP_12x) Chromosome 1: 811,192 - 816,906

Configure this page

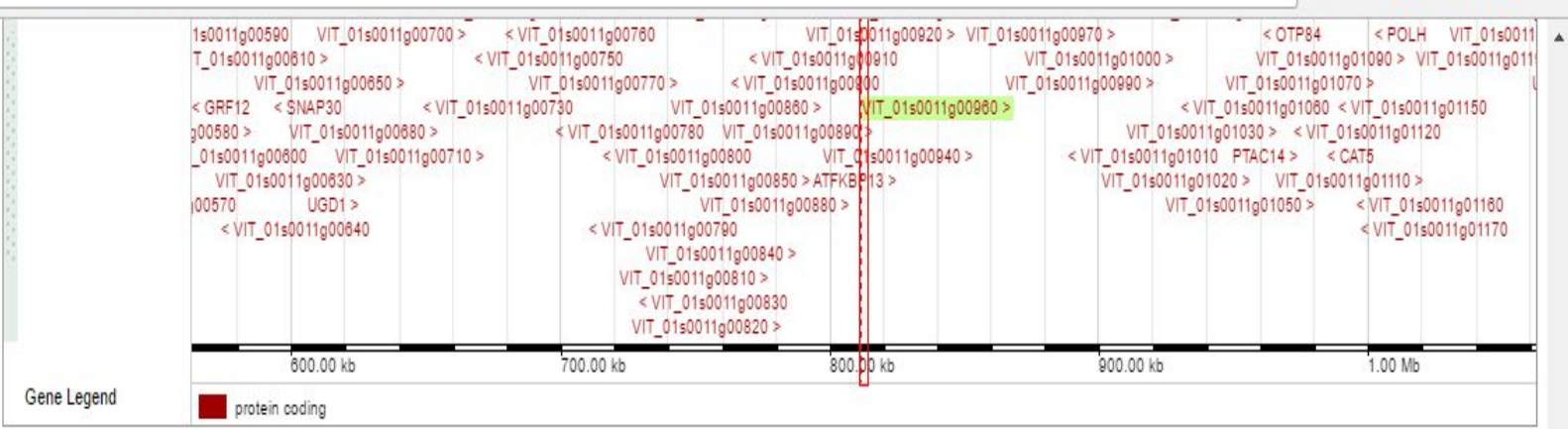
Custom tracks

Export data

Share this page

Bookmark this page

Ensembl Plants is produced in collaboration with Gramene



Location: Gene:

Navigation controls: left arrow, right arrow, zoom in (+), zoom out (-), and refresh.

Drag/Select: [icon]

Gene Legend

- Protein Coding
- protein coding

There are currently 46 tracks turned off.

Ensembl Plants Vitis vinifera version 88.3 (IGGP_12x) Chromosome 1: 811,959 - 814,575

Configure this page

Custom tracks

VIT_01s0011g00590	VIT_01s0011g00700	< VIT_01s0011g00780	VIT_01s0011g00820	VIT_01s0011g00970	< OTP84	< POLH	VIT_01s0011g01160
T_01s0011g00810	< VIT_01s0011g00750	< VIT_01s0011g00910	VIT_01s0011g01000	VIT_01s0011g01090	VIT_01s0011g01070	VIT_01s0011g01170	
VIT_01s0011g00850	VIT_01s0011g00770	< VIT_01s0011g00800	VIT_01s0011g00890				

Export data

Export Configuration - Feature List

Location to export: chromosome:IGGP_12x:1:811959:814575:1

Output: FASTA sequence *

Select location: 1 * 811959 * 814575 * 1

5' Flanking sequence (upstream): 0 * (Maximum of 1000000)

3' Flanking sequence (downstream): 0 * (Maximum of 1000000)

[Next >](#)

Fields marked * are required

Options for FASTA sequence

Genomic: Unmasked *

Fields marked * are required

TGAGGATGTTCTTGACCTGTTGAGGCTCGATCAAGAA
 GCGTCACTCCATCAACAACCTGATACAGGATATTGAC
 GACAAAGGAGAGGATCGCAGCATGGCCCTCTATTCA
 CCTTCATGTGAGAATGGCTCCTTTGTTCAATTGGTAAT
 GCCTACAATAAGCTGGTTCTTGGGCCTGGAACCG
 CGTGGTGGGGATGGCTGGACTGGGGAAGACAACCTCT
 GAAGCAGAACCTTGATTGCCATGTCTGGACCACAGCC
 TATCTTATGGACCTTGTAGTAGAGGAATTGGCTGT
 GGTGCTCTGACGCATAAGCTGCGAAAAATTTCAAC
 TGATGACTTATGGGTAAGATGTGGGAGTCCATT
 GGACAGTAGAATAATTACTACAAGGAGAGGCGAT
 TGATTCTATTGATATCCACAAGCTTCAGCCACTGTCA
 CTATAAGAAAGCTTTCTAAGAAATGGCAGATGCTCT
 ATCCATTTACAGAAATGTGATGGATTACCACTTGGG
 GTCCATTAAGGCTCAACTAAAAATGAGTGGAAAAAT
 ATTGAGAAGCAGTGGTGGCTTTCAAATATTTGAAAG
 TTTACCTTACCATCTCAAGTATTGTTCTTGTACATG
 AGTTAAGCGAAGAGACTTATTCGGCTATGGATAGCT
 AGGCAAAACGCTTGAAGAGGTTGGGGAAGAGTACCTG
 GATAAAGGCAATGAAATGGATTTGATGGAAGGCCG
 CATGTTGAAGATGATTCTCTCAGTCTCACATGAGGAG
 GGCAGTAAGAAAGTTGACTGAGAATACCCGCCCTAT
 TGTGCTCAGGACTTACCCTGTGTTGCAACCTTCTCA
 GATTGGTTCCAATTTCAAACCTATTGAAGGCTTGGAC
 ATTTCCGAGTGAATTACAGATCTTCTGCTCTGAGG
 TATAAGGAGAAATCCCTCGTCCCTCGGTGACCTTCA
 GCAGACTCTGGTAACAAAGGTACCTAAAGAAGTCTAG
 GCTGGTTTATCGCTACAATATGGAAAGTGTCTTACCT
 GGCACCAAAGAGGATGGGCGCATTGAAGAACCACAAA
 TGGGACGACAGAAATGAGTCGGCAGCACAGTATGATC
 GTTGAGGAAGCTGGGCATTGTAGAAGTGGCAAAAGAA
 AATTGTGAAGATGCGAAATCTCATTCTTGAATGTG
 TCTAGAGCTTGATGCAATGACCAATCCGCGCCCTT
 GCCTTTGAAAGGTTTCAAAGTGGGTATCTTCTAG
 AAAATGGTCTTCACTCGCAGAAGACCGATTGCAAGCT
 GGAGCTGCAGCTGCTTATGCATACACTGGAACCCAGT
 TCAGAAGCTTAAAGATATTAGACCTCAGCAATTGGAAG
 AGAAGGCACATTGCTTGCCTACAAGGCTGATCATAAGCCACTGTAGCAAGCTGGTGCA
 GGTCCCTACAGGATTGACAAGCTCATTACCTCCAAATGCTGCTGCTGCACGATATGCC
 CGAACCATTTGTGATCAGGCTGAGGAAAAATGGCGGTGATTGCGCCGTTTGGTTCACCA
 CATTCCGTGATTCAATCATATAATCAGGGCCAGTTG

PrimersDesign — Блокнот

Файл Правка Формат Вид Справка

```
>1 dna:chromosome chromosome:IGGP_12x:1:811959:814575:1
AC TTGAGGCTTTTCGCAAGTAGAGAATGGAATTTGCAGGAAAAATCAAAATGGCAGTCCA
AAATTTAGGACGCGAGCTGCGGAGCATTGAAGCCCTACTGAGAGATGCTGCTTCAAAGAA
AGAGCATGATACCAATTCACAGTCTGGATTCAAAATGTCCGTGATCAAGCTTATGCCAT
TGAGGATGTTCTTGACCTGTTGAGGCTCGATCAAGAATCAGTGTGGCGCCGCTTGAAAAAT
GCGTCACTCCATCAACAACCTGATACAGGATATTGACTGGAGTCTCAAATAATTCAGCG
GACAAAGGAGAGGATCGCAGCATGGCCCTTATTAACCAATGCTGGCAACAACACATA
CCTTCATGTGAGAATGGCTCCTTTGTTCAATTGGTAAATGTTGATACCGTGGGCATTGAGGA
GCCTACAATAAGCTGGTTTCTTGGGCCTTGAACCGAAACAGAGGCTTGAGGTGATGTT
CGTGGTGGGGATGGCTGGACTGGGGAAGACAACCTTGTCCACAGCGTGTATGAGAGGGT
GAAGCAGAACCTTGATTGCCATGTCTGGACCACAGCCTCAAAGTCCAAAACCAAACCTCGA
TATCTTATGGACCTTGTAGTAGAGGAATTGGGCTGTACAATCACACAGGGGGCCGATGT
GGTTGCTCTGACGCATAAGCTGCGAAAAATTTCTCAACAACAACGGTATGTCATAGTTCT
TGATGACTTATGGGTAAGATGTGTGGGAGTCCATTAGACTAGCCTTGCCAAATGGTAA
GGACAGTAGAATAATTACTACAAGGAGAGGCGATATAGCTAATTC TTGTAGAGATGA
TGATTCTATTGATATCCACAAGCTTCAGCCACTGTCACCGCAAAGGGCTGAGCAACTCTT
CTATAAGAAAGCTTTCTAAGAAATGGCAGATGCTCCTCAGGTTTGGAGGAAGTCTCCAA
ATCCATTTTACAGAAATGTGATGGATTACCACTTGGGATTATTGAAATCGGTAGACTTTT
GTCCATTAAGGCTCAACTAAAAATGAGTGGAAAAATATTACATGATAGCCTTGAGTCTGA
ATTGAGAAGCAGTGGTGGCTTTCAAATATTGAAAGTGTGTTGCTGCAAGTTACAATGA
TTTACCTTACCATCTCAAGTATTGTTCTTGTACATGAGCATCTTCTCAGAGCAACCC
AGTTAAGCGAAGAAGACTTATTCGGCTATGGATAGCTGAAGGTTTGTGATAGAGAAAAAG
AGGCAAAACGCTTGAAGAGGTTGGGGAAGAGTACCTGAATGAGCTGATTGACAGGAATCT
GATAAAGGCAATGAAATGGATTTTGTGGAAGGCCGACAAGTGTGGGAGTTCATAGTCT
CATGTTGAAGATGATTCTCTCAGTCTCACATGAGGAGAACTTTTGTACTGTCCGTACAGG
GGCAGTAAGAAAGTTGACTGAGAATACCCGCCCTATCTATCCAGAAGGAAGATTTTGA
TG TG TCTCAGGACTTACCCTGTGTTGCAACCTTCTCAGTTTCTGATAGGCAAAGTAAG
```

Primer3 (v. 0.4.0) Pick primers from a DNA sequence.

[Checks for mispriming in template.](#)

[disclaimer](#)

[Primer3 Home](#)

[Primer3plus interface](#)

[cautions](#)

[FAQ/WIKI](#)

There is a newer version of Primer3 available at <http://primer3.ut.ee>

Paste source sequence below (5' to 3', string of ACGTNacgtn -- other letters treated as N -- numbers and blanks ignored). FASTA format ok. Please N-out undesirable sequence (vector, ALUs, LINEs, etc.) or use a [Mispriming Library \(repeat library\)](#):

```
ACTTGAGGCTTTCGCAAGTAGAGAATGGAATTTGCAGGAAAATATCAAAATGGCAGTCCA
AAATTTAGGACGCGAGCTGCGGAGCATTGAAGCCCTACTGAGAGATGCTGCTTCAAAGAA
AGAGCATGATCACCAATTCACAGTCTGGATTCAAAATGTCCGTGATCAAGCTTATGCCAT
TGAGGATGTTCTTGACCTGTTGAGGCTCGATCAAGAATCAGTGTGGCGCCGCTTGAAAAT
GCGTCACTCCATCAACAATTGATACAGGATATTGACTGGAGTCTCCAAAATATTCAGCG
GACAAAGGAGAGGTATCGCAGCATGGCTCCTATTCAACCAATGCTGGCAACAACACATA
```

<input checked="" type="checkbox"/> Pick left primer, or use left primer below:	<input type="checkbox"/> Pick hybridization probe (internal oligo), or use oligo below:	<input checked="" type="checkbox"/> Pick right primer, or use right primer below (5' to 3' on opposite strand):
<input type="text"/>	<input type="text"/>	<input type="text"/>

[Sequence Id:](#) A string to identify your output.

[Targets:](#) E.g. 50,2 requires primers to surround the 2 bases at positions 50 and 51. Or mark the [source sequence](#) with [and]: e.g. ...ATCT[CCCC]TCAT.. means that primers must flank the central CCCC.

[Excluded Regions:](#) E.g. 401,7 68,3 forbids selection of primers in the 7 bases starting at 401 and the 3 bases at 68. Or mark the [source sequence](#) with < and >: e.g. ...ATCT<CCCC>TCAT.. forbids primers in the central CCCC.

[Product Size Ranges](#)

[Number To Return](#) [Max 3' Stability](#)

[Max Repeat Mispriming](#) [Pair Max Repeat Mispriming](#)

[Max Template Mispriming](#) [Pair Max Template Mispriming](#)

2521 CGAACCATTTGTGATCAGGCTGAGGAAAAATGGCGGTCGATTGCGCCGTTTGGTTCACCA

2581 CATTCCGTGTATTTCATCATATAATCAGGGCCAGTTG

KEYS (in order of precedence):

>>>>> left primer
<<<<< right primer

ADDITIONAL OLIGOS

	start	len	tm	gc%	any	3' seq
1 LEFT PRIMER	304	20	59.98	50.00	3.00	1.00 AAAGGAGAGGTATCGCAGCA
RIGHT PRIMER	2344	20	60.04	45.00	7.00	1.00 CTGAAATTTTCCCGACCTGA
PRODUCT SIZE: 2041, PAIR ANY COMPL: 4.00, PAIR 3' COMPL: 1.00						
2 LEFT PRIMER	304	20	59.98	50.00	3.00	1.00 AAAGGAGAGGTATCGCAGCA
RIGHT PRIMER	2345	20	60.04	45.00	7.00	1.00 TCTGAAATTTTCCCGACCTG
PRODUCT SIZE: 2042, PAIR ANY COMPL: 4.00, PAIR 3' COMPL: 2.00						
3 LEFT PRIMER	374	20	59.93	45.00	3.00	2.00 ATGGCTCCTTTGTTTCATTGG
RIGHT PRIMER	2383	20	59.99	45.00	6.00	1.00 CAGCTGTCCAATTGCTGAA
PRODUCT SIZE: 2010, PAIR ANY COMPL: 5.00, PAIR 3' COMPL: 3.00						
4 LEFT PRIMER	396	20	60.08	45.00	3.00	1.00 ATGTTGATACCGTGGGCATT
RIGHT PRIMER	2383	20	59.99	45.00	6.00	1.00 CAGCTGTCCAATTGCTGAA
PRODUCT SIZE: 1988, PAIR ANY COMPL: 3.00, PAIR 3' COMPL: 2.00						

Statistics

	con	too	in	in	no	tm	tm	high	high	high	high		
	sid	many	tar	excl	bad	GC	too	too	any	3'	poly		
	ered	Ns	get	reg	GC%	clamp	low	high	compl	compl	X		
											stab		
											ok		
Left	6394	0	0	0	0	0	1828	2695	0	1	0	105	1765
Right	6294	0	0	0	2	0	1825	2762	0	2	0	114	1589

Pair Stats:
considered 52, unacceptable product size 44, ok 8
primer3 release 1.1.4

Chromosome 1: 811,959 x plants.ensembl.org/Vitis x primer3 - Поиск в Google x Primer3 Output (primer3 x OligoCalc: Oligonucleotide x

biotools.nubic.northwestern.edu/OligoCalc.html

Oligo Calc: Oligonucleotide Properties Calculator

Enter Oligonucleotide Sequence
OD calculations are for single-stranded

Nucleotide base codes
 AAA GGA GAG GTA TCG CAG CA

Reverse Complement Strand (5' to 3') is:
 TGC TGC GAT ACC TCT CCT TT

5' modification (if any) 3' modification (if any)

50 nM Primer Measured Absorbance 1

50 mM Salt (Na⁺)

Calculate Swap Strands BLAST

Physical Constants

Length: 20 Molecular Weight: 6224.14 GC content: 50%

1 ml of a sol'n with an Absorbance of 1 at 260 nm is 4.105 microMolar and contains 25.5 micrograms.

Thermodynamic Constants Conditions: 1 M NaCl at 25°C at pH 7.

RlnK 33.404 cal/(°K*mol) deltaH 161.4 Kcal/mol
 deltaG 26.2 Kcal/mol deltaS 419.6 cal/(°K*mol)

Deprecated Hairpin/self dimerization calculations

5 (Minimum base pairs required for single primer self-dimerization)
 4 (Minimum base pairs required for a hairpin)

Check Self-Complementarity

Citation: Kibbe WA. 'OligoCalc: an online oligonucleotide properties calculator'. (2007) *Nucleic Acids Res.* 35(webserver issue): May 25. ([Abstract/Full text](#))

Oligo Self Complementarity Check - Google Chrome

about:blank

Minimum base pairs required for single primer self-dimerization: 5.
 Minimum base pairs required for a hairpin: 4.

Potential hairpin formation :

None !

3' Complementarity:
 None !

All potential self-annealing sites are marked in red (allowing 1 mis-match):
 None !

Со всеми обратным праймерами проблемы, поэтому нужно подобрать другие им на замену.

Chromosome 1: 811,959 x plants.ensembl.org/Virts x primer3 - Поиск в Google x Primer3 Input (version 0. x OligoCalc: Oligonucleotides x

bioinfo.ut.ee/primer3-0.4.0/

Primer3 (v. 0.4.0) Pick primers from a DNA sequence.

[Checks for mispriming in template.](#) [disclaimer](#) [Primer3 Home](#)
[Primer3plus interface](#) [cautions](#) [FAQ/WIKI](#)

There is a newer version of Primer3 available at <http://primer3.ut.ee>

Paste source sequence below (5'→3', string of ACGTNacgtn -- other letters treated as N -- numbers and blanks ignored). FASTA format ok. Please N-out undesirable sequence (vector, ALUs, LINEs, etc.) or use a [Mispriming Library \(repeat library\)](#):

```

ACTTGAGGCTTTCGCAAGTAGAGAATGGAATTTGCAGGAAAATATCAAAATGGCAGTCCA
AAATTTAGGACGCGAGCTGCGGAGCATTGAAGCCCTACTGAGAGATGCTGCTTCAAAGAA
AGAGCATGATCACCAATTACAGTCTGGATTCAAAATGTCCGTGATCAAGCTTATGCCAT
TGAGGATGTTCTTGACCTGTTCAAGGCTCGATCAAGAATCAGTGTGGCGCCGCTTGAAAAT
GCCTCACTCCATCAACAATTGATACAGGATATTGACTGGAGTCTCAAAAATATTCAGCG
GACAAAGGAGAGGTATCGCAGCATGGCTCCTATTCAACCAATGCTGGCAACAACACATA

```

Pick left primer, or use left primer below:
 Pick hybridization probe (internal oligo), or use oligo below:
 Pick right primer, or use right primer below (5' to 3' on opposite strand):

Sequence Id: A string to identify your output.

Targets: E.g. 50,2 requires primers to surround the 2 bases at positions 50 and 51. Or mark the [source sequence](#) with [and]: e.g. ...ATCT[CCCC]TCAT.. means that primers must flank the central CCCC.

Excluded Regions: E.g. 401,7 68,3 forbids selection of primers in the 7 bases starting at 401 and the 3 bases at 68. Or mark the [source sequence](#) with < and >: e.g. ...ATCT<CCCC>TCAT.. forbids primers in the central CCCC.

Product Size Ranges

Number To Return **Max 3' Stability**

Max Repeat Mispriming **Pair Max Repeat Mispriming**

Max Template Mispriming **Pair Max Template Mispriming**

- Ничего страшного. Просто расширяем зону поиска сайтов посадки праймеров.

2521 CGAACCATTGTGATCAGGCTGAGGAAAAATGGCGGTGATTGCGCCGTTTGGTTCACCA

2581 CATTCCGTGTATTCATTCATATAATCAGGGCCAGTTG

KEYS (in order of precedence):

>>>>> left primer
<<<<<< right primer

ADDITIONAL OLIGOS

	start	len	tm	gc%	any	3' seq
1 LEFT PRIMER	81	20	59.84	55.00	3.00	1.00 GGAGCATTGAAGCCCTACTG
RIGHT PRIMER	2414	20	60.08	45.00	4.00	1.00 GCAATGTGCCTTCTCCATT
PRODUCT SIZE: 2334, PAIR ANY COMPL: 5.00, PAIR 3' COMPL: 2.00						
2 LEFT PRIMER	146	20	59.90	40.00	4.00	2.00 TGGATTCAAATGTCCGTGA
RIGHT PRIMER	2505	20	59.70	45.00	3.00	2.00 AGCAGCATTTGGAGGTGAAT
PRODUCT SIZE: 2360, PAIR ANY COMPL: 6.00, PAIR 3' COMPL: 1.00						
3 LEFT PRIMER	81	20	59.84	55.00	3.00	1.00 GGAGCATTGAAGCCCTACTG
RIGHT PRIMER	2426	20	59.73	45.00	3.00	2.00 TTTGTAGGCAAGGCAATGTG
PRODUCT SIZE: 2346, PAIR ANY COMPL: 5.00, PAIR 3' COMPL: 1.00						
4 LEFT PRIMER	146	20	59.90	40.00	4.00	2.00 TGGATTCAAATGTCCGTGA
RIGHT PRIMER	2532	20	59.65	45.00	4.00	2.00 ACAAATGGTTCGGGCATATC
PRODUCT SIZE: 2387, PAIR ANY COMPL: 5.00, PAIR 3' COMPL: 2.00						

Statistics

	con	too	in	in	no	tm	tm	high	high	high			
	sid	many	tar	excl	bad	GC	too	too	any	3'	poly	end	
	ered	Ns	get	reg	GC%	clamp	low	high	compl	compl	X	stab	ok
Left	2886	0	0	0	0	0	821	1190	1	4	0	39	831
Right	2839	0	0	0	2	0	795	1277	0	2	0	45	718

Pair Stats:
considered 329, unacceptable product size 304, high end compl 6, ok 19
primer3 release 1.1.4

Chromosome 1: 811,959 x plants.ensembl.org/Vitis x primer3 - Поиск в Google x Primer3 Input (version 0. x OligoCalc: Oligonucleotide x

← → ↻ ⤴ ⤵ ⓘ biotools.nubic.northwestern.edu/OligoCalc.html

Oligo Calc: Oligonucleotide

Enter Oligonucleotide
OD calculations are for single

Nucleotide base codes
AGC AGC ATT TGG AGG TGA AT

Reverse Complement Strand(5' to 3') is:
ATT CAC CTC CAA ATG CTG CT

5' modification (if any) 3' modification (if any)

50 nM Primer 1 Measure
50 mM Salt (Na⁺)

Calculate **Swap Strands** **BLAS**

Physical Constants

Length: 20 Molecular Weight: 6221.14 GC con
1 ml of a sol'n with an Absorbance of 1 at 260 nm
is 4.322 microMolar and contains 26.9 micrograms.

Thermodynamic Constants Conditions: 1 M NaCl at 25°C at pH 7.

RlnK 33.404 cal/(°K*mol) deltaH 157.6 Kcal/mol
deltaG 25 Kcal/mol deltaS 411.4 cal/(°K*mol)

Depreciated Hairpin/self dimerization calculations

5 (Minimum base pairs required for single primer self-dimerization)
4 (Minimum base pairs required for a hairpin)

Check Self-Complementarity

Citation: Kibbe WA. 'OligoCalc: an online oligonucleotide properties calculator'. (2007) *Nucleic Acids Res.* 35(webserver issue): May 25. ([Abstract/Full text](#))

Oligo Self Complementarity Check - Google Chrome

about:blank

Minimum base pairs required for single primer self-dimerization: 5.
Minimum base pairs required for a hairpin: 4.

Potential hairpin formation :

None !

3' Complementarity:

None !

All potential self-annealing sites are marked in red (allowing 1 mis-match):

None !

- Остальные два праймера тоже хорошо подходят, но тут нужно выбирать в зависимости от цели, какой длины ПЦР продукт нужен.

Primer-BLAST *A tool for finding specific primers*

NCBI/Primer-BLAST: Finding primers specific to your PCR template (using Primer3 and BLAST).

PCR Template

[Reset page](#) [Save search parameters](#) [Retrieve recent results](#) [Publication](#) [Tips for finding specific primers](#)

Enter accession, gi, or FASTA sequence (A refseq record is preferred) [Clear](#)

```
ACTTGAGGCTTTGCAAGTAGAGAAATGGAATTTGCAGGAAAAATCAAAATGGCAGTCCA
AAATTTAGGACGCGAGCTGCGGAGCATTGAAGCCCTACTGAGAGATGCTGCTTCAAAGAA
AGAGCATGATACCAATTACAGTCTGGATTCAAATGTCGGTGATCAAGCTTATGCCAT
TGAGGATGTTCTTGACCTGTTCAAGGCTCGATCAAGAATCAGTGTGGCGCCGCTTGA
GCGTCACTCCATCAACAACCTTGATACAGGATATTGACTGGAGTCTCCAAAATATTCAGCG
```

Range

	From	To	
Forward primer	<input type="text"/>	<input type="text"/>	Clear
Reverse primer	<input type="text"/>	<input type="text"/>	

Or, upload FASTA file

Файл не выбран

Primer Parameters

Use my own forward primer (5'→3' on plus strand) [Clear](#)

Use my own reverse primer (5'→3' on minus strand) [Clear](#)

PCR product size

Min	Max
<input type="text" value="70"/>	<input type="text" value="1000"/>

of primers to return

Primer melting temperatures (T_m)

Min	Opt	Max	Max T _m difference
<input type="text" value="57.0"/>	<input type="text" value="60.0"/>	<input type="text" value="63.0"/>	<input type="text" value="3"/> Clear

Exon/intron selection

A refseq mRNA sequence as PCR template input is required for options in the section [Clear](#)

Exon junction span [Clear](#)

Exon junction match

Exon at 5' side	Exon at 3' side
<input type="text" value="7"/>	<input type="text" value="4"/>

Minimal number of bases that must anneal to exons at the 5' or 3' side of the junction [Clear](#)

Intron inclusion Primer pair must be separated by at least one intron on the corresponding genomic DNA [Clear](#)

Intron length range

Min	Max
<input type="text" value="1000"/>	<input type="text" value="1000000"/> Clear

Когда есть «неизвестная» последовательность ДНК

На примере гена NBS-LRR гена детекции патогенов у люцерны (давайте сделаем вид, что сиквенс действительно случайный)

BLAST/BLAT

Web Tools

- Web Tools
 - BLAST**
 - Variant Effect Predictor
 - Assembly Converter
 - ID History Converter

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BLAST search

Sequence data:


```
>
TTTCAAATGGGTGGATTGCCTTCCAACCTGGCCGACCTTTATATAGAGAATTGCCCAAAA
CTGATTCCTTCRAGAAGGAGTGGGGTTTGTTCCAACTCAATTCCTTGAAGTCATTCTTC
ATCAGTAATGAGTTTGAAAACGTGGAGTCATCCAGAGAAGAATCTGCTGCCACCACT
CTTCAAACCTCTTTGTATTAATGATTGTTCRAGCTAAGAATAATGAACAACRAGGGTTT
CTCCACCTCAATCTCTCATAGAACTACATAATTTGGAACCTGTCCTATTCTTGAGCGCTG
CCAGAGGAGGCTCTACCCAACACCCTTACTTCTATTGAAATTTCCGATTGTCCATTART
AAAGGGAAGTATGAAAAGGAGGGAGGAGAATTGGCATACAATTAGTCACATCCCTCAT
GTGACGATTGACGGAATTGACGAGAAATGAGCTAA
```

[Add more sequences](#) (1 sequence added, 29 more sequences allowed)

DNA

Protein

Search against:

 Medicago truncatula x

[Add/remove species](#)

DNA database

Protein database

Search tool:

Search Sensitivity:

- Web Tools
 - Web Tools
 - BLAST
 - Variant Effect Predictor
 - Assembly Converter
 - ID History Converter

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BLAST search

New job

Recent jobs

Refresh

Show/hide columns (1 hidden)		Filter
Analysis	Jobs	Submitted at
BLASTN	BLASTN against Medicago truncatula MedtrA17_4.0 (Genomic sequence) Done: 237 hits found View results	15/05/2017, 11:07 (BST)

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Results for BLASTN against Medicago truncatula MedtrA17_4.0 (Genomic sequence) ⓘ

Job details ⓘ

Job name: BLASTN against Medicago truncatula MedtrA17_4.0 (Genomic sequence)

Species: Medicago truncatula

Assembly: MedtrA17_4.0

Search type: BLASTN (NCBI BLAST)

Download results file

Results table ⓘ

Show All entries Show/hide columns (4 hidden) Filter

Genomic Location	Overlapping Gene(s)	Orientation	Length	Score	E-val	%ID
scaffold2053:198-652 [Sequence]	MTR_2053s0010	Forward	455 [Sequence]	455	0.0	100.0 [Alignment]
scaffold1643:3-133 [Sequence]		Reverse	131 [Sequence]	99	1.1E-48	93.9 [Alignment]
3:6733583-6733716 [Sequence]	MTR_3g022600	Reverse	134 [Sequence]	94	1.0E-45	92.5 [Alignment]
3:6941721-6941824 [Sequence]	MTR_3g023020	Reverse	104 [Sequence]	72	1.4E-32	92.3 [Alignment]
3:8571018-8571148 [Sequence]	MTR_3g027420	Forward	131 [Sequence]	67	1.3E-29	87.8 [Alignment]
3:6523624-6523683 [Sequence]	MTR_3g022230	Forward	60 [Sequence]	60	2.0E-25	100.0 [Alignment]
3:6945053-6945156 [Sequence]	MTR_3g023030	Reverse	104 [Sequence]	60	2.0E-25	89.4 [Alignment]
3:6782319-6782389 [Sequence]	MTR_3g022730	Forward	71 [Sequence]	55	1.9E-22	94.4 [Alignment]
3:8469378-8469448 [Sequence]	MTR_3g027250	Reverse	71 [Sequence]	55	1.9E-22	94.4 [Alignment]
3:6782473-6782529 [Sequence]	MTR_3g022730	Forward	57 [Sequence]	53	3.0E-21	98.2 [Alignment]
3:7585704-7585759 [Sequence]	MTR_3g024460	Reverse	56 [Sequence]	52	1.2E-20	98.2 [Alignment]
scaffold0425:9753-9823 [Sequence]	MTR_0425s0020	Forward	71 [Sequence]	51	4.7E-20	93.0 [Alignment]
3:7595470-7595524 [Sequence]	MTR_3g024460 MTR_3g024460	Reverse	55 [Sequence]	51	1.7E-20	93.0 [Alignment]

Далее возвращаемся к разделу Primer3 и делаем как делали ранее.

То есть все то же самое, только если бы у Вас не было названия гена, а был какой-то его сиквенс.

Но. Если есть название гена, который пересекается с нашим сиквенсом, то можно сделать точно также, как мы делали с самого

BLAST/BLAT

Web Tools

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- BLAST
 - Ticket
 - BLASTN against Medicago
 - Variant Effect Predictor
 - Assembly Converter
 - ID History Converter

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Results for BLASTN against Medicago truncatula Medtra17_4.0 (Genomic sequence) ?

Job details

Job name: BLASTN against Medicago truncatula Medtra17_4.0 (Genomic sequence)

Species: Medicago truncatula

Assembly: Medtra17_4.0

Search type: BLASTN (NCBI BLAST)

Download results file

Results table

Show All entries Show/hide columns (4 hidden) Filter

Genomic Location	Overlapping Gene(s)	Orientation	Length	Score	E-val	%ID
scaffold2053:198-652 [Sequence]	MTR_2053s0010	Forward	455 [Sequence]	455	0.0	100.0 [Alignment]
scaffold1643:3-133 [Sequence]		Reverse	131 [Sequence]	99	1.1E-48	93.9 [Alignment]
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3:6945053-6945156 [Sequence]	MTR_3g023030	Reverse	104 [Sequence]	60	2.0E-25	89.4 [Alignment]
3:6782319-6782389 [Sequence]	MTR_3g022730	Forward	71 [Sequence]	55	1.9E-22	94.4 [Alignment]
3:8469378-8469448 [Sequence]	MTR_3g027250	Reverse	71 [Sequence]	55	1.9E-22	94.4 [Alignment]
3:6782473-6782529 [Sequence]	MTR_3g022730	Forward	57 [Sequence]	53	3.0E-21	98.2 [Alignment]
3:7585704-7585759 [Sequence]	MTR_3g024460	Reverse	56 [Sequence]	52	1.2E-20	98.2 [Alignment]
scaffold0425:9753-9823 [Sequence]	MTR_0425s0020	Forward	71 [Sequence]	51	4.7E-20	93.0 [Alignment]

Смотрим первый слайд.

Nucleotide BLAST: Search x e! Gene: MTR_2053s0010 - x

plants.ensembl.org/Medicago_truncatula/Gene/Summary?db=core;g=MTR_2053s0010;r=scaffold2053:204-647;t=KEH15144;tl=VPv2HPtFZOqOlGFR-1785891

Medicago truncatula (MedtrA17_4.0)

Location: scaffold2053:204-647 Gene: MTR_2053s0010 Transcript: KEH15144 BLAST/BLAT results

Gene-based displays

- Summary
 - Splice variants
 - Transcript comparison
 - Gene alleles
- Sequence
 - Secondary Structure
- Gene families
- Literature
- Plant Compara
 - Genomic alignments
 - Gene tree
 - Gene gain/loss tree
 - Orthologues
 - Paralogues
- Pan-taxonomic Compara
 - Gene Tree
 - Orthologues
- Ontologies
 - GO: Molecular function
 - GO: Cellular component
 - GO: Biological process
- Phenotypes
- Genetic Variation
 - Variant table
 - Variant image
 - Structural variants
- Gene expression
- Regulation
- External references
- Supporting evidence
- ID History
 - Gene history

Gene: MTR_2053s0010

Description: NBS-LRR resistance protein

Location: [SuperContig scaffold2053: 204-647 forward strand.](#)


About this gene: This gene has 1 transcript ([splice variant](#)), [10 orthologues](#) and [7 paralogues](#).

Transcripts: [Show transcript table](#)

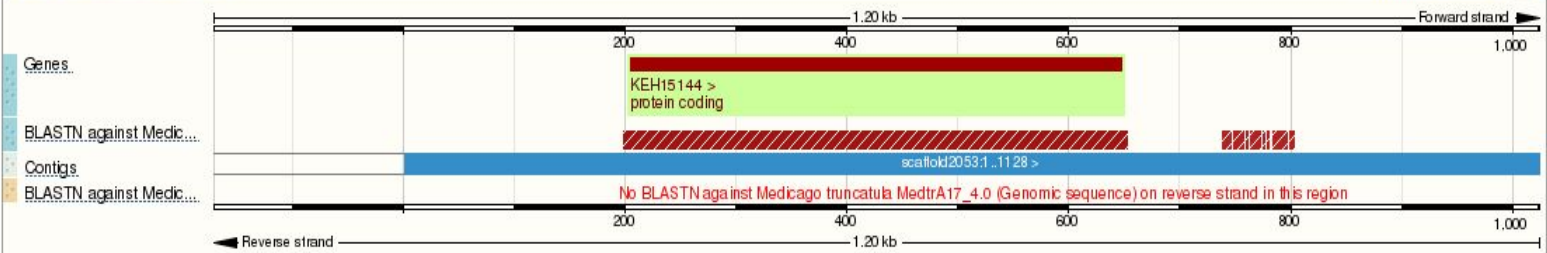
Summary

Gene type: Protein coding

Annotation method: Protein-coding genes annotation was carried out by the [International Medicago Genome Annotation Group](#) (IMGAG)

 [Go to Region in Detail](#) for more tracks and navigation options (e.g. zooming)

Drag/Select: ← →



Genes

KEH15144 > protein coding

BLASTN against Medic...

Contigs

scaffold2053:1..1128 >

BLASTN against Medic...

No BLASTN aga inst Medicago truncatula MedtrA17_4.0 (Genomic sequence) on reverse strand in this region

Gene Legend

Protein Coding

protein coding