PCR Primer Design for DNA Assembly

Overview

The polymerase chain reaction (PCR) uses a pair of custom primers to direct DNA elongation toward each other at opposite ends of the sequence being amplified. These primers are typically between 18 and 24 bases in length and must code for only the specific upstream and downstream sites of the sequence being amplified. A primer that can bind to multiple regions along the DNA will amplify them all, eliminating the purpose of PCR.

A few criteria must be brought into consideration when designing a pair of PCR primers. Pairs of primers should have similar melting temperatures since annealing during PCR occurs for both strands simultaneously, and this shared melting temperature must not be either too much higher or lower than the reaction's annealing temperature. A primer with a T_m (melting temperature) too much higher than the reaction's annealing temperature may mishybridize and extend at an incorrect location along the DNA sequence. A T_m significantly lower than the annealing temperature may fail to anneal and extend at all.

Additionally, primer sequences need to be chosen to uniquely select for a region of DNA, avoiding the possibility of hybridization to a similar sequence nearby.

Also, there are other parameters to be taken in account - Gibbs free energy, self- and heterodimers availability, etc. All parameters are described below in details.

UGENE provides the "PCR Primer Design for DNA Assembly" feature only for nucleic sequences with the "Standard DNA" alphabet. To use it in UGENE open a DNA sequence and go to the "PCR Primer Design for DNA Assembly" tab of the Options Panel:

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2	TTIGTGGT						
3	ттатааа						
4	TTTGTGGA						
5	TTTGTGAG						
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			Find reverse	e-complem	ent		
		user primer's					
		rward 5			Form	ard 3'	
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Parame	ters of prim	ing sequence	55				
Gbbs f	ree energy	-40 kcal/mol	le		-30 kcal/m	ole	۰
Melting	point	57 °C			- 65 °C		1
Overla	p length	18 nt		¢.	- 25 nt		<u>0</u>
Parame	ters to exd	ude in whole	primers				
	ree energy		-13 kcal	/mole			1
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There are the following parameters:

- User primers the pair of forward and reverse primers user can check on the absence of unwanted connections (like hairpins, self- and heterodimers).
- Choose generated sequences as user primer's end user primers could be extended with the 8-bases sequences on 5' and 3' ends. These
 sequences do not form too simple unwanted connections (for example, "AAAATTTT", obviously, has a hairpin, so there is no such a sequence in
 the list).
- Parameters of priming sequences the range of melting temperature, Gibbs free energy and length primers SHOULD have.
- Parameters to exclude in whole primers the minimum melting temperature, the maximum Gibbs free energy and the maximum base pairs length which may be present in the self- and hetero-dimers in the considered primer.
- Select areas for priming search select areas to search forward and reverse primers in.
- Open the backbone sequence choose a sequence to be added to 5' or 3' end of the result primer.
- Other sequences in PCR reaction the list of sequence to check the unwanted connections with. The unwanted connections should fit to "Parameters to exclude in whole primers".

Designing primers

Physical quantity

The designed primers should have the set Gibbs free energy, melting temperature and length. Also, these primers could have hairpins, self- and heterodimers, but the mentioned parameters of these found dimers should be limited. Look at the pictures:

A Forward (1) A Reverse (1)	A Forward	77
24 26 28 30 32 34 36 38 40 42 44 46 48 50 52 54 CACTGGAGTTGTCCCAATTCTTGTTGAATTA	56 58 60 62 64 66 68 70 72 74 GATGGTGATGTTAATGGGCAC	(76 78 80 82 84 86 88 90 92 94 96 98 100 102 104 107]AAATTTTCTGTCAGTGGAGAGGGTGAAGGTGA

•	 Parameters of priming sequences 								
	Gibbs free energy	-40 kcal/mole		* *	-	-30 kcal/mole	* *		
	Melting point	51 ℃		* *	-	65 °C	•		
	Overlap length	18 nt		*	-	25 nt	* *		
•	 Parameters to exclude in whole primers 								
	Gibbs free energy		-7 kcal/mole				•		
	Melting point		20 °C				* *		
	Max complementary motifs length		3 nt				•		

The found primer "GATGGTGATGTTAATGGGCAC" has Gibbs free energy between -40 and -30 kcal/mol, melting temperature between 51 and 65 °C and length between 18 and 25 nucleotides. Also, there are no heirpins, self- and hetero-dimers in this primer, which has Gibbs free energy more then -7 kcal /mol, melting temperature less than 20 °C and base pairs length more than 3 nucleotides.

Regions

Forward primers are searched at the left region, reverse primers are searched at the right region. There is an amplified fragment between them:

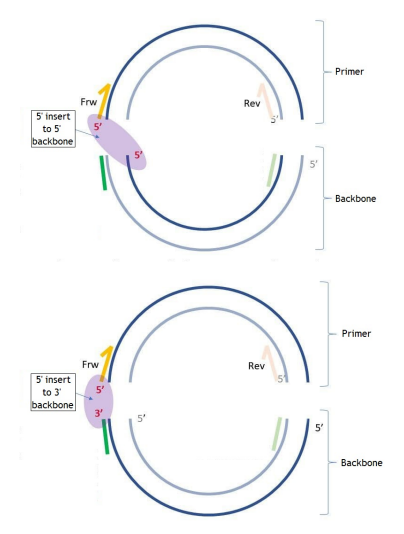
The amplified fragment The amplified fragment	
The left search area >1)	The right search area
1 20 40 60 80 100 120 140 160 180 200 220 240 260 280 300 320 34	0 360 380 400 420 440 460 480 500 520 540 560 580 600 620 640 660 680 717

Select areas for pr	iming search		
Primer search ar	eas for insert		
Left area:	1	÷ - 71	Select manually
Right area:	355	- 426	Select manually

The left search area is from 1 to 71 base, the right search area is from 355 to 426 base. The amplified fragment between them.

Backbone sequence

Backbone sequence is the sequence that is added to 5' end of the primer for the following assembly of the PCR product with the backbone molecule. This parameter is optional. You may set the backbone sequence, the sequence end you want to add backbone to and the length of the sequence from 5' and 3' ends to check if these sequences have unwanted hairpins, self- and hetero-dimers. On the pictures below you may see the differences between "5' insert to 5' backbone" ("insert" in this case is the same as "primer").



If you set "5' backbone length", then the number of bases you set from 5' end of the backbone sequence will be checked for unwanted connections. The same about "3' backbone length", but from the 3' end. If the checked end has hairpins, self- and homo-dimers you will see the following dialog:

Backbone details

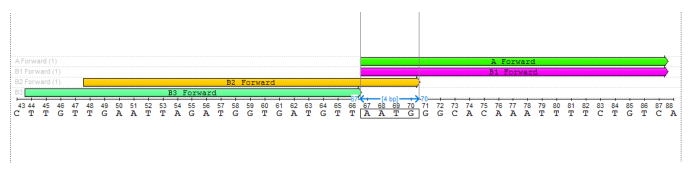
Other sequences in PCR reactions

Sequences that are supposed to be in the reaction mixture. Its important primers do not have unwanted connections to these sequences, so it will be checked that there are no hairpins, self- or hetero-dimers between the resulting primer and any sequence from the "Other sequences in PCR reaction" file.

Result

The search result is from 1 to 4 pairs of primers. The results should be as close as possible to the amplified fragment. That means, that the searching process starts from the edge of the amplified fragment to the 5' direction on the direct strand in case of the forward primer and to the 3' direction of the reverse-complementary strand in case of the reverse primer. The results are defined with special names:

- A the pair of primers closest to the amplified fragment, corresponding to the entire set of parameters.
- B1 the pair of primers closest to the amplified fragment, corresponding to the entire set of parameters if B2 and/or B3 exist.
- B2 the pair of primers corresponding to the entire set of parameters, which has a 4-nucleotide intersection with B1.
- B3 the pair of primers corresponding to the entire set of parameters, which starts immediately when B1 ends.



Also, you may see results in the result table. A single click on the result in the result table selects the corresponding annotation, double click - export the resulting primer to the separate file (with the backbone sequence, which will be added during the exporting process).

User primer

You may check the validity of the pair of primers by typing this pair into the corresponding fields. Also, you may add your primers 5' or/and 3' 8 bases endings, which do not form simple unwanted connections.

USEI	primers		
Forw	ard user primer:		
AAA		AATTTTCTGTC	
Reve	erse user primer:		
AAA		CCTTCAAACTTAAA	CACTC
Choo Filte	ose generated sequen	ces as user primer's	end
Filte			
	Sequence		
1	AAACACAG		
2	AAACACCA		
3	AAACACCC		
4	AAACACCT		
5	AAACACTC	_	-
		Find reverse	-complement
Add	sequence to a user pr	imer's end:	
Forward 5'			Forward 3'
	Reverse	3'	Reverse 5'