

## DATA SHEET

 Version: 03  
 Revision date: 15/06/2023

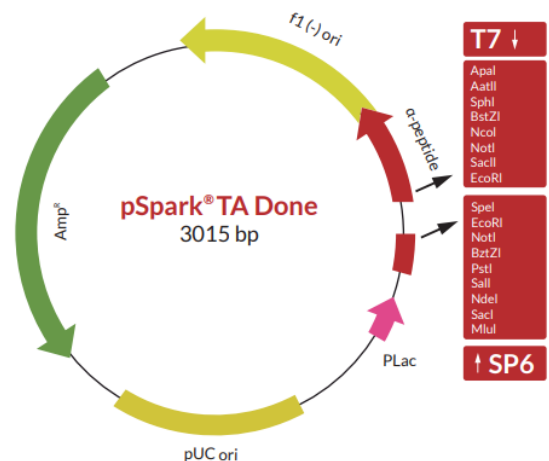
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### 1. Identification

<b>Product name</b>	<b>pSpark® TA Done</b> (50ng/uL) 20 reactions
<b>Cat No</b>	<b>C0021</b>

### 2. Description

**pSpark® TA Done** is efficient, stable and easy-to-use DNA cloning vector based on an improved TA technology that offers all of the advantages of pSpark® TA with the added convenience of recognition sites for EcoRI and NotI flanking the insertion site. Thus, several options exist to remove the desired insert DNA with a single restriction digestion



### 3. Composition

Item	Quantity
pSpark® TA Done DNA cloning vector	20 rxn (50ng/μL)
600kb control insert	5 μL
T4 DNA ligase (5 Weiss U/μL)	20 μL
10x T4 DNA ligase Buffer	100 μL

### 4. Features

- **Convenient:** recognition sites for EcoRI and NotI flanking the insertion site.
- **Flexible:** allows removing the desired insert DNA with other restriction digestion.
- **Efficient:** >600 white positive colonies expected under optimal conditions.
- **Stable:** without cloning bias due to transcription of toxic genes.
- **Easy-to-use:** eliminate screening of recombinants due to its <4% background.
- **Fast protocol:** ligation time from 60 minutes to overnight.
- **Compatible:** with direct cloning of PCR products.
- **Great versatility:** compatible with any competent cell or primer design.
- **Cost avoidance:** removes primer phosphorylation.

### 5. Storage specifications

 Store in a non-frost-free freezer at **-20°C**.

### 6. Applications

- Cloning of non-proofreading PCR fragments.



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- Production of ssDNA.
- Blue/white screening for recombinants.
- In vitro transcription from T7/SP6 dual-opposed promoters.

### 7. Recommended protocol

#### 1. Amplification

Amplify DNA with any proofreading polymerase following suggested protocol. Gel purifies the insert or check by electrophoresis the quality of amplified DNA. If a single band appears on the gel and template DNA is not a vector with Ampicillin resistance, like the pSpark® TA Done DNA cloning vector, then gel purification is not needed.

#### 2. Ligation

Suggested ratio of insert to vector is 5:1. Use the equation **ng insert = 33ng x kb** (of insert, that is, for a 1kb insert only 33 ng are needed) or use **1µL** of unpurified PCR product for directly ligation.

##### Mix:

**1 µL** of pSpark® TA Done DNA cloning vector (50ng/µL)

**1 µL** 10x T4 DNA ligase buffer

**X µL** of insert

**1 µL** of T4 DNA ligase (5 Weiss units)

H<sub>2</sub>O to **10 µL**

Incubate the reaction 60 minutes at 22°C (a ligation time from 10min to overnight could be used)

#### 3. Transformation

Transformation can be made with your regular protocol, competent cells and selective media. As pSpark® TA Done DNA cloning vector has Ampicillin resistance, this antibiotic is needed. Also, as pSpark® TA Done DNA cloning vector has blue/white screening capability, IPTG/XGal plates are recommended.

A suitable transformation protocol is:

- Thaw competent cells on ice, take **50 µL** and put into a cool 1,5 mL tube.
- Add **10 µL of the ligation mixture**, mix by gently flicking with fingertips and incubate on ice for 30 minutes.
- Heat-shock the cells at **exactly 42°C for exactly 45 seconds** (do not shake) and then transfer to ice for 2 minutes.
- Plate the cells into media with antibiotic, IPTG and XGal and incubate overnight at 37°C.

#### 4. Analysis of transformants

Either colony PCR or plasmid purification and digestion could be used for analysis of transformants. pSpark® TA Done DNA cloning vector has pUC/M13 forward and reverse sequencing primers binding site for sequencing or amplification of insert. Insert size when amplified with pUC/M13 fw and rev primers is about 200bp longer than the real size of insert.

#### 5. Expected results

Up to 2500 positive white colonies for a 60 min ligation of 33 ng of supplied control insert (600 kb) and less than 20 negative blue colonies when using cells with a transformation efficiency of  $4 \times 10^7$  cfu/ug.



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[info@canvaxbiotech.com](mailto:info@canvaxbiotech.com)[www.canvaxbiotech.com](http://www.canvaxbiotech.com)**7. Further information**

- Product Use Limitations** This product is developed, designed and sold exclusively only for research purposes use. The product was not tested for use in diagnostics or for drug development, nor is it suitable for administration to humans or animals.
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