# Instructions for Use of HEK293 Cell Residual DNA Fragment Analysis Detection Kit（qPCR） 

## The kit is intended for scientific research only and should not be used for diagnosis

Cat．No．HG－HFOO2

## Introduction

This kit is a kit specially designed for quantitative detection of the size distribution of residual HEK293 Cell DNA fragments in intermediates，bulk products and final products of various biological products．

This kit is based on the principle of PCR fluorescence probe method to quantitatively detect the size distribution of residual HEK293 Cell DNA fragments in samples．This kit designs three different amplified fragments（ $99 \mathrm{bp}, 200 \mathrm{bp}$ and 307 bp ）．The HEK293 Cell DNA quantitative reference is used to generate standard curves for different amplified fragments，and the fragment distribution of residual HEK293 Cell DNA in samples is analyzed by the ratio of different size fragments．

## Specification

$3 \times 100$ Reactions

## Main components

Table 1．Kit components

| Serial Number | Components | Fill volume／vial | Quantity | Storage conditions |
| :---: | :---: | :---: | :---: | :---: |
| 1 | HEK293 Primer\＆Probe MIX－99 | $300 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$ ，protected from light |
| 2 | HEK293 Primer\＆Probe MIX－200 | $300 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$ ，protected from light |
| 3 | HEK293 Primer\＆Probe MIX－307 | $300 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$ ，protected from light |
| 4 | HEK293 DNA quantitative reference | $60 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$ |
| 5 | $2 \times$ Probe qPCR Mix | 1.25 mL | 4 | $-18^{\circ} \mathrm{C}$ |
| 6 | DNA diluent | 1 mL | 4 | $-18^{\circ} \mathrm{C}$ |
| 7 | IPC Mix | $450 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$, protected from light |
| 8 | ROX High＊ | $50 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$, protected from light |
| 9 | ROX Low＊ | $50 \mu \mathrm{~L}$ | 1 | $-18^{\circ} \mathrm{C}$ ，protected from light |

＊Please select appropriate ROX for corresponding model．

## Storage conditions and shelf life

The shelf life is 18 months when stored at $-18^{\circ} \mathrm{C}$ and below．

## Applicable models（including but not limited to）

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－ABI PRISM 7500
－FQD－96A（Bioer）
－CFX96（Bio－Rad）
－Roche Light Cycler 480
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When using with different instrument models，please select the appropriate reference stain ROX．

| Instruments | ROX reference stain |
| :---: | :---: |
| Applied Biosystems ${ }^{\oplus} 5700,7000,7300,7700,7900 \mathrm{HT}$ ，StepOne ${ }^{\text {TM }}$ ，and StepOnePlus ${ }^{\text {TM }}$ | ROX High |
| Applied Biosystems ${ }^{\text {® }} 7500$ ，ViiA ${ }^{\text {TM }} 7$ ，QuantStudio ${ }^{\text {TM }} 12 \mathrm{~K}$ Flex，Agilent Mx3000P ${ }^{\text {TM }}$ ，Mx3005P ${ }^{\text {TM }}$ ，and Mx4000 ${ }^{\text {TM }}$ | ROX Low |
| Rotor－Gene ${ }^{T M}$ ，DNA Engine Opticon ${ }^{\top M}$ ，Opticon ${ }^{\top}{ }^{\top \mathrm{M}} 2$ ，Chromo $4^{\mathrm{TM}}$ Real－Time Detector，Mastercycler ${ }^{\circledR}$ ep realplex，Smart Cycler ${ }^{\circledR}$ ，Roche LightCycle ${ }^{\circledR} 480$ ，Roche LightCycler ${ }^{\oplus}$ Nano，Bio－Rad CFX96，and IIlumina Eco ${ }^{\text {TM }}$ | No ROX |

## Prepared by the user

－ 1.5 mL or 2 mL sterile low－attachment centrifuge tubes
－96－well qPCR plate or 8－strip PCR tubes compatible with the PCR system
－Sterile low－attachment pipette tips with cartridge， $1000 \mu \mathrm{~L}, 200 \mu \mathrm{~L}$ ，and $10 \mu \mathrm{~L}$ specifications
－Fluorescent quantitative PCR system
－Centrifuge
－Shaker
－Pipettes of various specifications（such as 1000 $\mu \mathrm{L}, 200 \mu \mathrm{~L}, 10 \mu \mathrm{~L}$ ，and $2.5 \mu \mathrm{~L}$ ）．

## Operation procedures

## I．Dilution of HEK293 DNA Quantitative Reference and Preparation of Standard Curve

HEK293 DNA quantitative reference（concentration： $3 \mathrm{ng} / \mu \mathrm{L}$ ）is subjected to gradient dilution with DNA diluent provided in the kit at concentrations of $3 \times 10^{5} \mathrm{fg} / \mu \mathrm{L}, 3 \times 10^{4} \mathrm{fg} / \mu \mathrm{L}, 3 \times 10^{3} \mathrm{fg} / \mu \mathrm{L}, 3 \times 10^{2} \mathrm{fg} / \mu \mathrm{L}, 3 \times 10^{1} \mathrm{fg} / \mu \mathrm{L}$ ．The detailed procedures are as follows：
1．Place the HEK293 DNA quantitative reference and DNA diluent in the kit on ice；after thawing completely，shake gently to mix well，and centrifuge briefly to settle the liquid to the bottom of the tube．
2．Take five 1.5 mL clean centrifuge tubes and label as ST1，ST2，ST3，ST4 and ST5，respectively．
3．Add $180 \mu \mathrm{~L}$ of DNA diluent into ST1，ST2，ST3，ST4 and ST5 tubes，respectively．
4．Perform 5 dilutions according to Table 2，ensure thorough mixing after each dilution，and then perform the next gradient dilution．For example，add $20 \mu \mathrm{~L}$ of HEK293 DNA quantitative reference to the prepared ST1 tube in Step 3，shake gently to mix well，centrifuge briefly to settle the liquid to the bottom of the tube；then add $20 \mu \mathrm{~L}$ of standard in ST1 to the prepared ST2 tube in Step 3．．．．．．dilute sequentially．

Table 2．Dilution of standards

| Standard No． | Dilution volume | Concentration |
| :---: | :---: | :---: |
| ST1 | $180 \mu \mathrm{~L}$ of DNA diluent $+20 \mu \mathrm{~L}$ of HEK293 DNA quantitative reference | $3 \times 10^{5} \mathrm{fg} / \mu \mathrm{L}$ |
| ST2 | $180 \mu \mathrm{~L}$ of DNA diluent $+20 \mu \mathrm{~L}$ of ST1 | $3 \times 10^{4} \mathrm{fg} / \mu \mathrm{L}$ |
| ST3 | $180 \mu \mathrm{~L}$ of DNA diluent $+20 \mu \mathrm{~L}$ of ST2 | $3 \times 10^{3} \mathrm{fg} / \mu \mathrm{L}$ |
| ST4 | $180 \mu \mathrm{~L}$ of DNA diluent $+20 \mu \mathrm{~L}$ of ST3 | $3 \times 10^{2} \mathrm{fg} / \mu \mathrm{L}$ |
| ST5 | $180 \mu \mathrm{~L}$ of DNA diluent $+20 \mu \mathrm{~L}$ of ST4 | $3 \times 10^{1} \mathrm{fg} / \mu \mathrm{L}$ |

## II．Preparation of Negative Control Sample（NCS）

Add $100 \mu \mathrm{~L}$ of sample matrix solution（or DNA diluent）to 1.5 mL clean centrifuge tube，and label it as negative control sample （NCS）．
The NCS can be pre－treated together with the test samples from the same batch to prepare the NCS purified solution． When adding sample for qPCR reaction，add 10uL DNA diluent which is negative control NTC．
Note：To meet the need for simultaneous analysis of fragments with three different amplification lengths，$\geq 100 \mu \mathrm{~L}$ of DNA elution volume is required for sample pre－treatment．

## III．Preparation of additive system

Standard recovery ERC：It is recommended to use 90ul sample＋10uIST3．It can also be prepared in other ways according to the actual situation．

Calculation of standard recovery rate：ERC recovery rate is between $50 \%$ and $150 \%$（standard recovery rate $=$ ERC／（0．9＊sample＋0．1＊ST3）．

## IV．Preparation of qPCR Reactions

1．Take out each reagent from the freezer and place them on ice．
2．Calculate the required number of reaction wells based on the standard curve and the number of samples to be tested． Generally， 3 replicate wells are required for each sample．That is，the number of reaction wells $=$（standard curve of 5
concentration gradients +1 NTC +1 NCS＋number of test samples）$\times 3$ ．
3．Calculate the total amount of qPCR reaction solution required for this time based on the number of reaction wells： QPCR reaction solution $=$（Number of reaction wells +2 ）$\times 20 \mu \mathrm{~L}$（containing loss in 2 wells）．
4．After each reagent is fully thawed on ice，the qPCR reaction mixture corresponding to the amplified fragment is prepared by combining the number of reaction wells as shown in Tables 3,4 ，and 5 and mixed well．

| Table 3．Preparation of qPCR Mixture for 99－bp Frag | Reaction ent | Table 4．Preparation of qPCR Reaction Mixture for 200－bp Fragment |  | Table 5．Preparation of qPCR Reaction Mixture for 307－bp Fragment |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Components | Reaction in single－well | Components | Reaction in single－well | Components | Reaction in single－well |
| 2×Probe qPCR Mix | $15 \mu \mathrm{~L}$ | $2 \times$ Probe qPCR Mix | $15 \mu \mathrm{~L}$ | $2 \times$ Probe qPCR Mix | $15 \mu \mathrm{~L}$ |
| HEK293 Primer\＆Probe MIX－99 | $3 \mu \mathrm{~L}$ | HEK293 Primer\＆Probe MIX－200 | $3 \mu \mathrm{~L}$ | HEK293 Primer\＆Probe MIX－307 | $3 \mu \mathrm{~L}$ |
| IPC Mix | $1.4 \mu \mathrm{~L}$ | IPC Mix | $1.4 \mu \mathrm{~L}$ | IPC Mix | $1.4 \mu \mathrm{~L}$ |
| ROX＊ | $0.6 \mu \mathrm{~L}$ | ROX＊ | $0.6 \mu \mathrm{~L}$ | ROX＊ | $0.6 \mu \mathrm{~L}$ |
| Total volume | $20 \mu \mathrm{~L}$ | Total volume | $20 \mu \mathrm{~L}$ | Total volume | $20 \mu \mathrm{~L}$ |

＊Select suitable ROX for corresponding model；if the model is suitable for no ROX，add an equal volume of deionized water（it is required to add deionized water without nucleic acid and nuclease contamination）．

## V．Loading

1．Mix each qPCR reaction mixture prepared by gentle shaking and centrifuge briefly to settle the liquid to the bottom of the tube．Add into the corresponding wells at $20 \mu \mathrm{~L} /$ well in typesetting design（as shown in Table 9 or adjusted by personalized design），select the corresponding amplified fragments and load as shown in Tables 6， 7 and 8，with a total volume of $30 \mu \mathrm{~L}$ per well after loading．

Table 6．Example of Loading of 99－bp Fragment in Each Reaction Well

| ST－99bp | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 99－bp fragment＋ 10 <br> $\mu \mathrm{~L}$ of ST1／ST2／ST3／ST4／ST5 |
| :---: | :---: |
| NTC | $20 ~ \mu \mathrm{~L}$ of qPCR reaction mixture for 99－bp fragment＋10 <br> $\mu \mathrm{L}$ of DNA diluent |
| NCS | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 99－bp fragment＋10 <br> $\mu \mathrm{L}$ of NCS purified solution |
| Test <br> sample | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 99－bp fragment＋10 <br> $\mu$ of test sample |

Table 8．Example of Loading of 307－bp Fragment in Each Reaction Well

| ST－307bp | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 307－bp fragment＋ <br> $10 \mu \mathrm{~L}$ of ST1／ST2／ST3／ST4／ST5 |
| :---: | :---: |
| NTC | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 307－bp fragment＋ <br> $10 \mu \mathrm{~L}$ of DNA diluent |
| NCS | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 307－bp fragment＋ <br> $10 \mu \mathrm{~L}$ of NCS purified solution |
| Test <br> sample | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 307－bp fragment＋ <br> $10 \mu \mathrm{~L}$ of test sample |

\＆This example represents a DNA standard curve of samples at five concentration gradients，a NTC，a NCS，and three test samples． prepare wells in triplicate．It can also be designed and typeset according to the user＇s own commonly used practice．

2．Seal the 96 －well plate with an optical plate sealer， gently shake to mix well，perform rapid centrifugation with a centrifuge dedicated for 96 －well plate to let all liquid gather at tube bottom，then place the plate in the qPCR system．

Table 7．Example of Loading of 200－bp Fragment in Each Reaction Well

| ST－200bp | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 200－bp fragment＋10 <br> $\mu \mathrm{L}$ of ST1／ST2／ST3／ST4／ST5 |
| :---: | :---: |
| NTC | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 200－bp fragment＋10 <br> $\mu \mathrm{L}$ of DNA diluent |
| NCS | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 200－bp fragment＋10 <br> $\mu \mathrm{L}$ of NCS purified solution |
| Test sample | $20 \mu \mathrm{~L}$ of qPCR reaction mixture for 200－bp fragment＋10 <br> $\mu \mathrm{of} \mathrm{test} \mathrm{sample}$ |

Table 9．Example of a 96－well plate layout

| $996 p$ |  |  | 200bp |  |  |  | 307 bp |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| NTC | NTC | NTC |  | NTC | NTC | NTC |  | NTC | NTC | NTC |  | A |
| NCS | NCS | NCS |  | NCS | NCS | NCS |  | NCS | NCS | NCS |  | B |
| S1 | S1 | S1 | S2 | S1 | S1 | S1 | S2 | S1 | S1 | S1 | S2 | C |
| ST1 | ST1 | ST1 | S2 | ST1 | ST1 | ST1 | S2 | ST1 | ST1 | ST1 | S2 | D |
| ST2 | ST2 | ST2 | S2 | ST2 | ST2 | ST2 | S2 | ST2 | ST2 | ST2 | S2 | E |
| ST3 | ST3 | ST3 | S3 | ST3 | ST3 | ST3 | S3 | ST3 | ST3 | ST3 | S3 | F |
| ST4 | ST4 | ST4 | S3 | ST4 | ST4 | ST4 | S3 | ST4 | ST4 | ST4 | S3 | G |
| ST5 | ST5 | ST5 | S3 | ST5 | ST5 | ST5 | S3 | ST5 | ST5 | ST5 | S3 | H |
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |  |

## VI．Run

ABI 7500 qPCR instrument（software V2．0．6）is used as an example．
1．Create a blank new procedure and select Absolute Quantitative Test Template and Taqman Probe Method．
2．Three sets of qPCR reactions are used to create new detection probes named as $99 \mathrm{bp}, 200 \mathrm{bp}, 307 \mathrm{bp}$ ，select reporter fluorophore as FAM and quench fluorophore as none；new detection probes are created and named as IPC，select reporter fluorophore as VIC，quench fluorophore as none，and detection reference fluorescence is ROX．

3．Select standard wells in the plate as Standard，assign values of $3 \times 10^{5}, 3 \times 10^{4}, 3 \times 10^{3}, 3 \times 10^{2}, 3 \times 10^{1}(\mathrm{in} \mathrm{fg} / \mu \mathrm{L})$ ，and name as ST1，ST2，ST3，ST4，ST5 in the corresponding Sample Name column；select NTC as Negative Control，NCS and test sample wells as Unknow，and name as NTC，NCS，S1，S2，S3 in the corresponding Sample Name column．

4．Set up three－step reaction program： $95{ }^{\circ} \mathrm{C}$ for $5 \mathrm{~min} ; 45$ cycles of $95{ }^{\circ} \mathrm{C}$ for $15 \mathrm{~s}, 58{ }^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 72{ }^{\circ} \mathrm{C}$ for 1 min ；and a reaction volume of $30 \mu \mathrm{~L}$ ．

5．Run the PCR program．

## VII．qPCR result analysis

1．In the＂Amplification Plot＂of Results，the user may initially check whether the amplification curve is of normal shape． Usually，the system will automatically set the threshold and baseline．Multiple thresholds may be generated in case of different target settings，leading to inaccurate result analysis．In such case，please manually set the threshold line， which must be within the exponential amplification region，for example，the threshold is usually set to 0.15 ．Select＂Auto Baseline＂，and click＂Analyze＂to view the adjusted results．

2．Data reliability evaluation：
－The difference in Ct values between 3 replicate wells shall be less than 1．0，except for wells with Ct value greater than 35 ；
－The CT values of negative controls NTC and NCS should be greater than the CT value of the lowest concentration of the standard curve，or the criteria should be set based on the laboratory＇s own validation results；
－Linear correlation coefficient $\mathrm{R}^{2}$ of the standard curve shall be equal or greater than 0.98 ，amplification efficiency shall be within $85 \%-110 \%$ ；
－The recovery of ERC shall be within $50 \%-150 \%$ ．
3．The sample value of 99bp fragment is $100 \%$ ，and the sample percentage of 200bp fragment and 307bp fragment is calculated．

## Disclaimer

Under all circumstances，the liability of our company for this product is only limited to the value of the product itself．

