

pFUSE-hIgG1e7-Fc1

Plasmid containing a human engineered IgG1 Fc region

Catalog # pfc1-hg1e7

For research use only

Version 20K05-MM

PRODUCT INFORMATION

Content:

- 20 µg of pFUSE-hIgG1e7-Fc1 plasmid provided as lyophilized DNA
- 1 ml of Zeocin™ (100 mg/ml)

Storage and Stability:

- Product is shipped at room temperature.
- Lyophilized DNA should be stored at -20°C and is stable 3 months.
- Resuspended DNA should be stored at -20°C and is stable up to 1 year.
- Store Zeocin™ at 4 °C or at -20 °C. The expiry date is specified on the product label.

Quality control:

- Plasmid construct has been confirmed by restriction analysis and sequencing.
- Plasmid DNA was purified by ion exchange chromatography and lyophilized.

GENERAL PRODUCT USE

pFUSE-Fc is a family of plasmid developed to facilitate the construction of Fc-fusion proteins by fusing the effector region of a protein to the Fc region of an immunoglobulin G (IgG).

pFUSE-Fc plasmids yield high levels of Fc-fusion proteins. The level of expression is usually in the µg/mL range. They can be transfected in a variety of mammalian cells, including myeloma cell lines, CHO cells, monkey COS cells and human embryonic kidney (HEK)293 cells, cells that are commonly used in protein purification systems.

pFUSE-Fc plasmids allow the secretion of Fc-Fusion proteins. As Fc-Fusion proteins are secreted, they can be easily detected in the supernatant of pFUSE-Fc-transfected cells by SDS-PAGE. Furthermore, functional domains can be identified by immunoblotting and ligand blotting.

Fc-Fusion proteins can be easily purified by single-step protein A or protein G affinity chromatography.

InvivoGen provides pFUSE-Fc vectors featuring Fc regions from different species and isotypes. In humans, there are four isotypes: IgG1, IgG2, IgG3 and IgG4. The Fc region mediates effector functions, such as antibody-dependent cellular cytotoxicity (ADCC) and complement-dependent cytotoxicity (CDC). IgG isoforms exert different levels of effector functions increasing in the order of IgG4<IgG2<IgG1≤IgG3. Human IgG1 displays high ADCC and CDC, and is the most suitable for therapeutic use against pathogens and cancer cells.

Under certain circumstances, for example when depletion of the target cell is undesirable, abrogating effector functions is required. On the contrary, in the case of antibodies intended for oncology use, increasing effector functions may improve their therapeutic activity¹. Modifying effector functions can be achieved by engineering the Fc regions to either improve or reduce their binding to FcγRs or the complement factors. Amino acids substitutions have been made in the human IgG1 Fc region in order to increase or reduce its ADCC and CDC.

PLASMID FEATURES

- **hIgG1e7-Fc (human IgG1 engineered Fc):** The Fc region comprises the CH2 and CH3 domains of the IgG heavy chain and the hinge region. The hinge serves as a flexible spacer between the two parts of the Fc-Fusion protein, allowing each part of the molecule to function independently. The mutations K326W/E333S has been reported to enhance CDC and binding to C1q². FUSE-hIgG1e7-Fc1 contains the K326W/E333S mutation.
- **hEF1-HTLV prom** is a composite promoter comprising the Elongation Factor-1α (EF-1α) core promoter³ and the R segment and part of the U5 sequence (R-U5') of the Human T-Cell Leukemia Virus (HTLV) Type 1 Long Terminal Repeat⁴. The EF-1α promoter exhibits a strong activity and yields long lasting expression of a transgene *in vivo*. The R-U5' has been coupled to the EF-1α core promoter to enhance stability of RNA.
- **MCS:** The multiple cloning site contains several restriction sites that are compatible with many other enzymes, thus facilitating cloning.
- **SV40 pAn:** the Simian Virus 40 late polyadenylation signal enables efficient cleavage and polyadenylation reactions resulting in high levels of steady-state mRNA⁵.
- **ori:** a minimal *E. coli* origin of replication to limit vector size, but with the same activity as the longer Ori.
- **CMV enh / hFerL prom:** This composite promoter combines the human cytomegalovirus immediate-early gene 1 enhancer and the core promoter of the human ferritin light chain gene. This ubiquitous promoter drives the expression of the Zeocin™-resistance gene in mammalian cells.
- **EM2KC** is a bacterial promoter that enables the constitutive expression of the antibiotic resistance gene in *E. coli*. EM2KC is located within an intron and is spliced out in mammalian cells.
- **Zeo:** Resistance to Zeocin™ is conferred by the *Sh ble* gene from *Streptomyces hindustanus*. The same resistance gene confers selection in both mammalian cells and *E. coli*.
- **BGlo pAn:** The human beta-globin 3'UTR and polyadenylation sequence allows efficient arrest of the transgene transcription⁶.

1. Carter PJ., 2006. Potent antibody therapeutics by design. *Nature Reviews Immunology*. Advance online publication.

2. Idusogie EE. et al., 2001. Engineered antibodies with increased activity to recruit complement. *J Immunol.* 166(4):2571-5.

3. Kim DW et al. 1990. Use of the human elongation factor 1 alpha promoter as a versatile and efficient expression system. *91(2):217-23.*

4. Takebe Y. et al. 1988. SR alpha promoter: an efficient and versatile mammalian cDNA expression system composed of the simian virus 40 early promoter and the R-U5 segment of human T-cell leukemia virus type 1 long terminal repeat. *Mol Cell Biol.* 8(1):466-72.

5. Carswell S. & Alwine JC. 1989. Efficiency of utilization of the simian virus 40 late polyadenylation site: effects of upstream sequences. *Mol Cell Biol.* 9(10):4248-58.

6. Yu J. & Russell JE. 2001. Structural and functional analysis of an mRNP complex that mediates the high stability of human beta-globin mRNA. *Mol Cell Biol.* 21(17):5879-88.

TECHNICAL SUPPORT

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METHODS

Plasmid resuspension

Quickly spin the tube containing the lyophilized plasmid to pellet the DNA. To obtain a plasmid solution at 1 µg/µl, resuspend the DNA in 20 µl of sterile H₂O. Store resuspended plasmid at -20 °C.

Plasmid amplification and cloning

Plasmid amplification and cloning can be performed in *E. coli* GT116 or in other commonly used laboratory *E. coli* strains, such as DH5α.

Zeocin™ usage

This antibiotic can be used for *E. coli* at 25 µg/ml in liquid or solid media and at 50-200 µg/ml to select Zeocin™-resistant mammalian cells.

RELATED PRODUCTS

Product	Catalog Code
Zeocin™	ant-zn-1

TECHNICAL SUPPORT

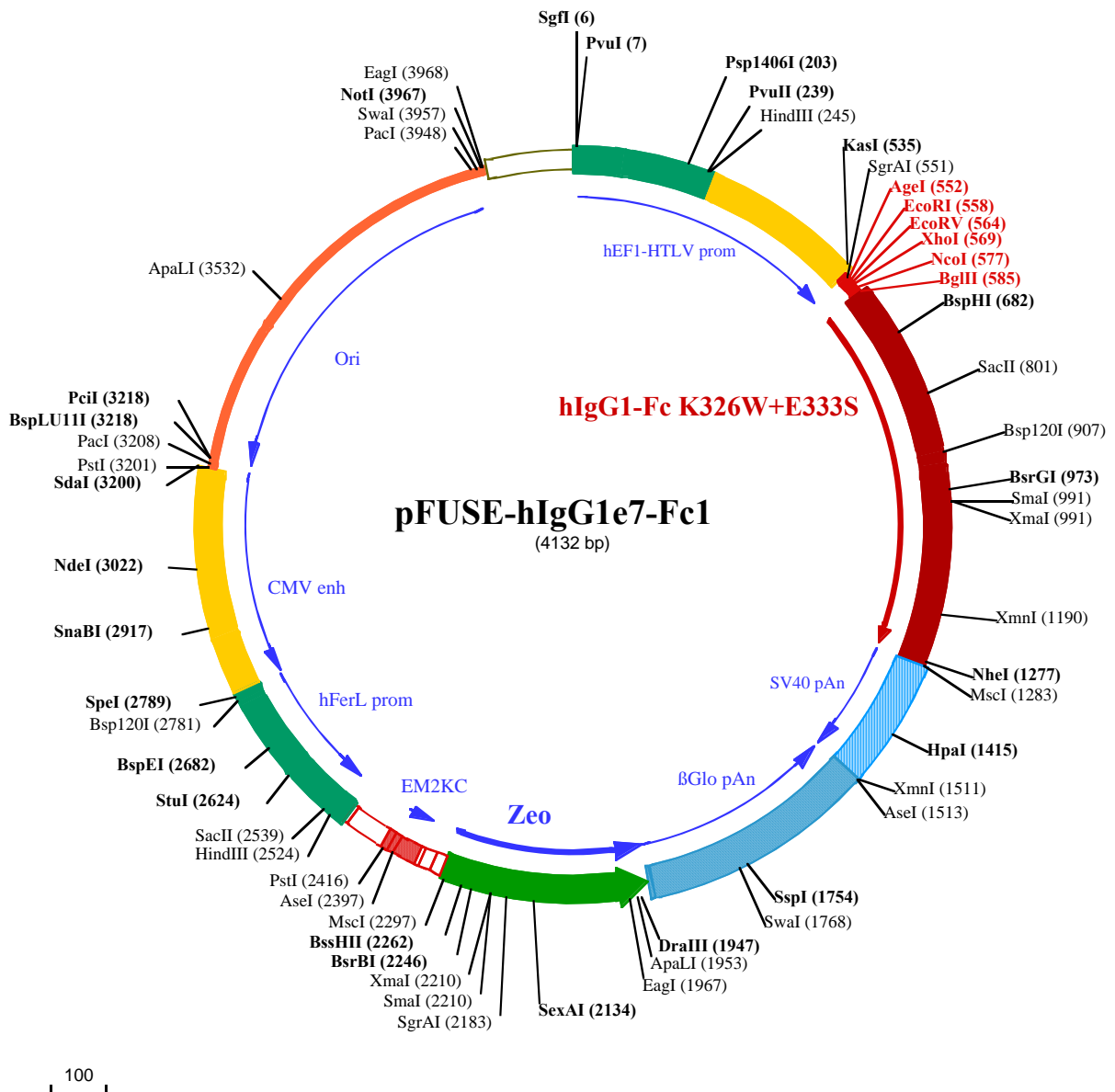
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PvuI (7)
SgfI (6)

1 GGATCTGGATCGCTCCGGTCCCCGTCAAGTGGCAGAGGCACATCCCCACAGTCCCCGAGAAGTTGGGGGAGGGTCGGCAATTGAACGGGTGCCA

101 GAGAAGGTGGCGGGGTAAACTGGAAAGTGATGTCGTACTGGCTCCGCCCTTTCCGAGGGTGGGGAGAACCGTATAAGTCAGTAGTCGCC

HindIII (245)
Psp1406I (203)
PvuII (239)

201 GTGAACGTTCTTTCCCAACGGTTGCCGCCAGAACACAGCTGAACGCTTCGAGGCGCTCCATCTCCTCACGGCCGCCCTACCTGAGGCC

301 GCCATCCACGCCGTGAGTCGCTCTGCCGCCCTGTGGTGCCTCTGAAGTGCCTCCGCTAGGTAAGTTAAAGCTCAGGTCGAGACC

401 GGCGCTTGTCCGGCTCCCTGGACCTACCTAGACTCAGCCGGCTCCACGCTTGCCTGACCTGCTGCTCAACTCTACGTCTTGTGTTCGTT

EcoRI (558)
KasI (535)
AgeI (552)
SgrAI (551)
XbaI (569)
EcoRV (564)
BglII (585)
NcoI (577)

501 TCTGTTCTGCGCCGTTACAGATCCAAGACTGTGACCGCGCCTACCTAGACTCAGCCGGCTCCACGCTTGCCTGACCTGCTGCTCAACTCTACGTCTTGTGTTCGTT

1 ▶ D K T

BspHI (682)

601 CACACATGCCACCGTGCCCAGCACCTGAACCTCTGGGGGACCGTCAGTCTCTCTTCCCCAAAACCCAAGGACACCCCATGATCTCCGGACCC

4 H T C P P C P A P E L L G G P S V F L F P P K P K D T L M I S R T

701 CTGAGGTACATGCGTGGTGGTGGACGTGAGCACGAAAGACCCCTGAGGTCAAGTCACTGGTACGTGGACGGCGTGGAGGTGCTAAATGCCAAGACAAA

37 P E V T C V V V D V S H E D P E V K F N W Y V D G V E V H N A K T K

SacII (801)

801 GCGGGGGAGGAGCAGTACAACAGCACGTACCGTGGTCAGCGTCTCACCCTGCACCAGGACTGGCTGAATGCCAAGGAGTACAAGTGCAGGTC

70 P R E E Q Y N S T Y R V V S V L T V L H Q D W L N G K E Y K C K V

XbaI (991)
BsrGI (973)
SmaI (991)

901 TCCAATGGGCCCTCCAGCCCCCATCAGAAAACCATCTCAAAGCCAAGGGCAGCCCCGAGAACCCACAGGTGTACACCTGCCCATCCGGAGG

104 S N W A L P A P I S K T I S K A K G Q P R E P Q V Y T L P P S R E

1001 AGATGACCAAGAACCAAGGTGACCGTGGCTGGTCAAAGGCTCTATCCACGGCAGATCGCCGTTGGAGTGGAGAGCAATGGCAGCCGGAAACAA

137 E M T K N Q V S L T C L V K G F Y P S D I A V E W E S N G Q P E N N

XmnI (1190)

1101 CTACAAGACCACGCCCTCCGTGCTGGACTCCGACGGCTCCCTCTCTACAGCAAGCTCACCGTGACAAGAGCAGGTGGCAGGGAAACGTCTTC

170 Y K T T P P V L D S D G S F F L Y S K L T V D K S R W Q Q G N V F

MscI (1283)
NheI (1277)

1201 TCATGCTCCGTGATGCAAGGCTCTGCACAACCAACTACACCGAGAGGCCCTCCCTGCTCCGGTAAATGAGTGTAGCTGGCCAGACATGATAAG

204 S C S V M H E A L H N Y T Q K S L S L S P G K •

1301 ATACATTGATGAGTTGGACAAACCAACTAGAATGCACTGGTGAATTTGATGCTATTGCTTATTGTAACCATTATA

HpaI (1415)

1401 AGCTGCAATAAACAGTTAACACAACAATTGCAATTCAATTATGTTAGTTCAAGGTTAGGGGAGGTGGAGGTTTAAAGCAAGTAAACCTCTACA

AseI (1513)
XmnI (1511)

1501 AATGTTGATGAAATTAACTCTAAACACAGCATAGCAAAACTTAAACCTCAAATCAAGCCTCTACTTGAATCCTTCTGAGGGATGATAAGGCATA

1601 GGCATCAGGGCTGTCGAATGTGCAATTAGCTGTTGCAGCCTCACCTCTTCAAGGTTAAGATATAGTGTATTCCCAGGTTGAACTAGCT

SspI (1754)
Swal (1768)

1701 CTTCATTTCTTATGTTAAATGCACTGACCTCCACATTCCCTTTAGTAAATATTCAAGAAATAATTAAATACATCATTGCAATGAAAATAATG

1801 TTTTTATTAGGCAGAACCTCAGATGCTCAAGGCCCTCATATAATCCCCAGTTAGTAGTTGGACTTAGGAACAAAGAACCTTAATAGAAATTGGA

ApaLI (1953)
DraIII (1947)
EagI (1967)

1901 CAGCAAGAAAGCGAGCTTAGCTTACCTCAGTCCTGCCAACAAAGTCAGCAGTTGCCGGCCGGTGCAGGGCAACTCCGCC

1901 125 ▲ • D Q E E A V F H V C N G A P D R L A F R G W

2001 ACGGCTGCTGCCGATCTCGGTCACTGGCCGGCCGGAGGGCTCCGGAGATTCTGAGCACGACCTCCGACCACTCGCGTACAGCTCGTCCAGGCC

101 P Q E G I E T M A P G S A D R F N T S V V E S W E A Y L E D L G R

SexAI (2134)
SgrAI (2183)

2101 CACCCACACCCAGGCCAGGGTGTGTCGGCACACCTGGCTGGACCGCGCTGATGAAACAGGTCACTCGTCCGGACACACCGCGAAGTCGTCC

2101 V W V W A L T N D P V V Q D Q V A S I F L T V D D R V V G A F D D

XbaI (2210)
SmaI (2210)
BsrBI (2246)
BssHII (2262)
MscI (2297)

2201 TCCACGAAGTCCGGAGAACCCGAGCCGGTGGTCAGAACCTCGACCGCTCCGGAGCGTGGACCGAACGGACTGGTCAACTGGTCAACTTGG

2201 E V F D R S F G L R D T W F E V A G A V D R A T L V P V A S T L K A

AseI (2397)

2301 CCATGATGGCTCCTCgtcaggagaggaaagagaaggtagtacaatttgCTATAGTGTAGTTGATTACTATGCAAGATATACTATGCCAATGATT

2301 M

PstI (2416)

2401 AATTGTCAAACATAGGGCTGCAgggttcatagtgccactttctgcactgccccatctctgcccaccccttccaggcatagactcagtgacttacc

HindIII (2524)
SacII (2539)

2501 AAACTCACAGGAGGGAGAAGGCAGAGCTTGAGACAGACCCGGGGCGCGAAGTGGCTAGGGCGCTCTTTATGGTGC

StuI (2624)

2601 CCTCTGGAGGCAGGGCCTCGGGAGGCCATCGCCCAATCTCGGTGGCAGGAGGGGGCCGAAGGCCGTGCCCTGACCAATCCGAGCACATAGGAGT

BspEI (2682)

2701 CTCAGCCCCCGCCCCAAAGCAAGGGGAAGTCACGCCCTGTAGCCCAGCGTGTGAAATGGGGCTTGGGGGGTTGGGCCCTGACTAGTCAAAAA

2801 CAAACTCCCCATTGACGTCAATGGGTGGAGACTTGAAATCCCCGTGAGTCACCCGCTATCCACGCCATTGATGACTGCCAAAACCGCATCATCATG

SnaBI (2917)

2901 GTAATAGCGATGACTAATACGTAGTACTGCCAAGTAGGAAAGTCCCATAAGGTATGACTGGCATAATGCCAGCGGCCATTACCGTCATTGA

NdeI (3022)

3001 CGTCATAAGGGCGTACTTGGCATATGATACACTTGTACTGCCAAGTGGCAGTTACCGTAATACTCCACCCATTGACGTCAATGGAAAGTCCC

3101 TATTGGCGTTACTATGGAACATACGTCAATTGACGTCAATGGCGGGGTCGTTGGCGGTAGCCAGGCGGCCATTACCGTAAGTTATGTAACG

PacI (3208)

PstI (3201) **PciI (3218)**

SdAI (3200) **BspLU11I (3218)**

3201 CCTGCAGGTTAATTAAGAACATGAGCAGCAAAGGCCAGCAAAGGCCAGGAACCGTAAAAGGCCGCTGCTGGCGTTTCCATAGCTCCGCC

3301 TGACGAGCATCACAAATCGACGCTCAAGTCAGAGGTGGCAAACCCGACAGGACTATAAGATAACAGCGTTCCCTGGAAAGCTCCCTGTGCGC

3401 TCTCCTGTTCCGACCCTGCCGTTACCGATAACCTGTCGCCCTTCTCCCTCGGAAGCGTGGCCTTCTCATAGCTACGCTGTAGGTATCTCAGTT

ApaLI (3532)

3501 CGGTGTAGGTCGTTCCCTCCAAGCTGGCTGTGCAAGAACCCCCCTTGTGCGCCCTTATCGGTAACACTATCGCTTGTAGTCCAACCC

3601 GGTAAGACACGACTTATGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTCTGAAGTGGTGGCCT

3701 AACTACGGCTACACTAGAAGAACAGTATTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTCGGAAAGAGTTGGTAGCTCTGATCCGCAAACAA

3801 CCACCGCTGGTAGCGTGGTTTTTGTTGCAAGCAGATTACCGCAGAAAAAAAGGATCTCAAGAAGATCCTTGATCTTCTACGGGTCTGA

EagI (3968)

PacI (3948) SwaI (3957) **NotI (3967)**

3901 CGCTCAGTGGAACGAAACTCACGTTAGGGATTTGGTCACTGGCTAGTTAATTAACATTAAACAGCGGCCGATAAAATATCTTATTTCAATTAC

4001 ATCTGTGTGTTGGTTTTGTGTGAATCGTAACAAACATACGCTCTCCATAAAACAAAACGAAACAAAACAAACTACGAAAATAGGCTGCCCCAGTGC

4101 AAGTGCAGGTGCCAGAACATTCTCTATCGAA