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(54) **Titre :** PREPARATION ET UTILISATION DE BANQUES GENOMIQUES D'ANTICORPS SYNTHETIQUES  
("BIBLIOTHEQUES D'ANTICORPS SYNTHETIQUES")

(54) **Title:** PREPARATION AND USE OF GENE BANKS OF SYNTHETIC HUMAN ANTIBODIES ("SYNTHETIC HUMAN-  
ANTIBODY LIBRARIES")

(57) **Abrégé/Abstract:**

The invention relates to the preparation and use of gene banks of synthetic human antibodies (huAb) or parts of antibodies which contain the antigen-binding domain. Starting from a huAb framework in a suitable vector, the hypervariable regions of the antibody cDNA are formed by almost "randomly" combined oligonucleotides. Relatively conserved amino acids in the hypervariable regions have here been taken account of in the choice of appropriate nucleotides during the oligonucleotide synthesis and the ratio of the nucleotides used is likewise chosen such that a nonsense codon is to be expected at most in every 89th position. Expression of this synthetic huAb cDNA in microbial expression systems, e.g. in E. coli in the vector pFMT which is described below, thus makes a synthetic huAb library with a comprehensive repertoire for screening using selected antigens available in vitro.



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Abstract of the disclosure

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          antibodies ("synthetic human-antibody libraries")

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20       vector pFMT which is described below, thus makes a  
synthetic huAb library with a comprehensive repertoire  
for screening using selected antigens available in vitro.

25       The human or mammalian immune system comprises an esti-  
mated number of between  $10^6$  and  $10^8$  different antibodies.  
This number of antibodies seems to be sufficient to cause  
an immune reaction of the body both against all naturally  
occurring antigens and against artificial antigens. If it  
is furthermore taken into account that often different  
antibodies react with the same antigen, the repertoire of  
30       antibodies that are really different would rather be in  
the region from  $10^6$  to  $10^7$ .

Up to now specific antibodies have always been obtained  
starting from an immunization with the particular anti-  
gen, for example injection of the antigen into the body  
or in vitro incubation of spleen cells with this antigen.  
35       In the case of polyclonal antibodies, the immunoglobulins



can then be isolated from the serum and the specific antibodies can be isolated therefrom, e.g. by absorption methods. Monoclonal antibodies are isolated from the cell supernatants or from the cell lysate of spleen tumor cells (hybridoma cells) which have been fused with individual B lymphocytes and cloned. The abovementioned methods are unsuitable in particular for the preparation of specific human antibodies or human monoclonal antibodies.

10 The present invention therefore has the object of developing a generally usable method for generating specific human monoclonal antibodies (huMAbs) or parts of antibodies, which contain synthetic hypervariable domains.

15 It has now been found that by using almost randomly synthesized oligonucleotides coding for the three hypervariable regions of each variable part of heavy or light chains (called CDR1, 2 and 3, CDR meaning complementary determining region) synthetic human gene banks can be generated. The synthesized antibody DNA was then preferably ligated into an antibody expression vector especially constructed for this purpose, namely the vector pFMT, preferably after amplification using the polymerase chain reaction (PCR).

25 The oligonucleotides which are used for the synthesis of the variable domains of heavy and light chains are compiled in Tab. 1. Set A here contains fewer limitations than set B. The limitations below the synthesis of the hypervariable regions (see CDR regions in Tab. 4) being random were made in H3, H4, H6, L2, L3 and L5 in set A in order firstly to allow for positions in the sequence for certain conserved amino acids, secondly to reduce the number of stop codons, and thirdly to incorporate a new restriction site.

35 (a) In order to reduce the probability of the stop codon

occurring, only half the amount of the three other nucleotides was allowed for T at the first position of each codon and A was omitted at the third position of each codon, in each case. As a statistical average, only every 89th codon will thus be a stop codon.

(b) For the 2nd codon in the CDR1 region of the light chain, only those nucleotides were allowed which code for the amino acids V, A or G.

(c) Likewise, only those combinations coding for V, I or M were allowed for codon No. 10 in the CDR1 region of the light chain and for codon No. 4 in the CDR1 region of the heavy chains.

(d) In the CDR3 region of the light chain, only those nucleotides coding for the amino acid glutamine were allowed for codon No. 1.

(e) In the CDR2 region of the heavy chain, only those nucleotides coding for the amino acid tyrosine were allowed for codon No. 11.

(f) An A was advantageously incorporated at the third position of the last codon in the CDR2 region of the heavy chain in order to introduce a restriction site for MluI.

The random nature of these oligonucleotides was preferably limited even further in those positions where predominantly one or few amino acids occur (set B in Tab. 1, the limitations here are based on the tables by Kabat et al. (1987), Sequences of Proteins of Immunological Interest-U.S. Dept. of Health and Human Services, U.S. Government Printing Offices). A list of the corresponding nucleotides and brief notes on the codon combinations are compiled in Tab. 1 and in the notes for Tab. 1.



After ligation of equimolar amounts of the oligonucleotides H1 to H7 and L1 to L5, these are ligated into the pretreated expression vector pFMT. Preferably, a PCR step using the primers H1 and H8, or L1 and L6 should be carried out beforehand in order to amplify the amount of DNA. After producing suitable restriction sites at the ends of the antibody DNA using appropriate restriction enzymes, the DNA is ligated into the antibody expression vector pFMT in the same manner as above (see examples).

10 The expression pFMT makes possible the expression of antibody cDNA and the subsequent secretion of the expression products in bacteria (E.coli). The antibody operon of the plasmid contains the sequences of the variable parts of both the heavy and light chain of an antibody.

15 Suitable leader sequences from the amino terminal part of a bacterial protein makes secretion of the antibody parts possible. The leader sequences are cleaved off by a bacterial enzyme during the secretion. During the secretion of the antibody cDNA products, the light and heavy

20 chains of the antibody (with or without an adjacent constant domain) become associated. This results in the formation of an antibody or antibody fragment which, in either case, contains a functional antigen binding site. Similar constructs for individual antibodies have also

25 been described by other authors (Better et al. (1988), Science 240, 1041, and Skerras & Plückthun (1988), Science 240, 1038).

In the synthetic human-antibody library formed by the expression in, for example, E. coli, the desired human

30 antibodies or antibody parts are found by screening bacterial clones using the selected antigen. In a preferred embodiment, an additional sequence which codes for a marker peptide, for example a TAG sequence, is incorporated so that the expression products can be detected

35 in a simple way using established monoclonal antibodies against the marker peptide (Wehland et al. (1984), EMBO J. 3, 1295).

The abovementioned exemplary formulations and the examples below shall be understood as illustrating but not restricting the invention.

5 The invention therefore relates to gene banks of synthetic huAb or antigen-binding parts thereof, obtained by means of (1) cDNA for the hypervariable regions generated on a random basis, where the random sequences are limited by clauses (a) to (e) set A or in accordance with Tab. 1, set B, (2) preferably a subsequent  
10 amplification step of these random sequences and (3) ligation of the said cDNA into a suitable expression vector, preferably pFMT, an additional coding sequence for a marker peptide being incorporated in a preferred embodiment.

15 The invention also relates to a process for the separation of the abovementioned gene banks, and the process and the use thereof for the isolation of clones which secrete specific antibodies or antigen-binding parts thereof.

20 Finally, the invention is explained in detail in the examples and contained in the patent claims.

**Examples:**

**Example 1: Preparation of an antibody expression vector**

25 The plasmid pKK233-2 (Amann and Brosius, (1985) Gene 40, and Straus and Gilbert (1985) Proc. Natl. Acad. Sci. 82, 2014) was chosen as base vector for the construction of the antibody expression vector (Fig. 1).

30 Before the incorporation of the antibody operon, the plasmid was cut with SalI and BamHI, the ends were filled in with Klenow polymerase and ligated. By doing so, the two restriction sites and the DNA between them were



deleted.

5 Additionally, the plasmid was cleaved with HindIII, the  
ends were filled in with Klenow polymerase and ligated  
using BamHI linkers. By this procedure, the HindIII  
restriction site was removed and a BamHI site inserted.  
The antibody DNA was inserted into this modified plasma.  
A simplified structure of the antibody operon coding for  
a dicistronic antibody mRNA is shown in Tab. 2. In order  
to make possible the secretion of the antibody, the  
10 leader sequence of the bacterial enzyme pectate lyase was  
used. The leader sequence of this enzyme has already been  
used for the expression and secretion of a chimeric  
murine/human antibody (Fab fragment, Better et al., loc.  
cit.), and of the variable region of a "humanized"  
15 antibody (Ward et al., loc. cit.; Huse et al., loc.  
cit.). DNA for the first leader sequence ( $P_1$  upstream of  
the heavy chain), and the sequence for a second ribosome  
binding site (RBS) and a second leader sequence ( $P_2$   
upstream of the light chain) were synthesized from  
20 several oligonucleotides (Tab. 3).

Antibody cDNAs which code for the variable regions of the  
heavy and light chains of a human antibody (HuVhlys or  
HuVllys; Riechmann et al., (1988) J. Mol. Biol. 203, 825)  
were obtained from Dr. G. Winter (Cambridge, UK). The  
25 restriction sites HindIII (HuVhlys) and EcoRV (HuVllys)  
were introduced to make possible the insertion of the  
antibody cDNA into the expression vector. Further  
restriction sites for BanII (HuVhlys) and BstEII or KpnI  
(HuVllys) were introduced to exchange hypervariable  
30 regions en bloc. At the end of the HuVhlys cDNA sequence  
a stop signal was incorporated. A BanII site in the light  
chain was removed. These alterations were carried out by  
means of site directed mutagenesis in the bacteriophage  
M13mp18 (Zoller and Smith, Meth. Enzymol. 100, 468-500).  
35 The sequence of the completed antibody DNA is shown in  
Tab. 4.



For the insertion of the leader sequence  $P_1$  (Tab. 3) the modified plasmid pKK233-2 was digested using the restriction enzymes NcoI and PstI, and  $P_1$  was inserted in between these sites (pKK233-2- $P_1$ ). Further cloning steps,  
5 apart from the last step, were carried out using the plasmid pUC18. The reason is that the presence of individual parts of the antibody operon in the expression vector adversely influences the growth of the bacterial host.

10 Before the cloning in pUC18, its BamHI restriction site had to be removed. After digesting with BamHI, the single-stranded ends were filled in using the Klenow fragment and were religated. This modified plasmid was then digested using PstI and HindIII, and  $P_2$  plus RBS was  
15 ligated in between the restriction sites (pUC18- $P_2$ ). During this process, the original HindIII restriction site of the plasmid disappears and a new HindIII restriction site is incorporated. pUC18- $P_2$  was then digested using PstI and HindIII, and the DNA of the heavy chain  
20 (PstI-HindIII insert from M13) was ligated into these two sites (pUC18-HP<sub>2</sub>). This plasmid was then digested using EcoRV and BamHI, and the DNA of the light chain (EcoRV-BamHI insert from M13) was ligated in (pUC18-HP<sub>2</sub>L).

The PstI-BamHI insert was then recloned in pUC18 after  
25 the restriction sites for HindIII, BanII and KpnI therein had previously been removed. The HindIII restriction site was removed as above for pKK233-2, the religation taking place without an insertion of BamHI linkers, however. Subsequently, the resulting plasmid was digested using  
30 SmaI and BanII, and, after filling in the protruding ends by means of T4 DNA polymerase, religated. The insertion of the PstI-BamHI restriction fragment results in pUC-HP<sub>2</sub>L. In a preferred embodiment, a Tag sequence was additionally inserted in the BanII and HindIII restriction  
35 sites (Tab. 3). The Tag sequence encodes the recognition sequence Glu-Gly-Glu-Glu-Phe of the monoclonal antibody Y1 1/2 (Wehland et al., (1984), EMBO J. 3, 1295). Because

of this peptide marker the expression product of the resulting plasmid pUC-HTP<sub>2</sub>L is readily detectable.

5 For the insertion of HP<sub>2</sub>L or HTP<sub>2</sub>L in the expression vector, the two plasmids were cut using PstI and BamHI, and the PstI-BamHI HP<sub>2</sub>L insert from pUC-HP<sub>2</sub>L or the HTP<sub>2</sub>L insert from pUC-HTP<sub>2</sub>L was ligated into the modified plasmid pKK233-2-P<sub>1</sub> into these two restriction sites. A diagrammatic representation of the completed expression vector pFMT is shown in Tab. 5.

10 **Example 2: Synthesis of antibody DNA containing random sequences in hypervariable regions**

15 The synthesized oligonucleotides for the synthesis of the variable parts of antibody DNA are compiled in Tab. 1. For the synthesis of the hypervariable regions almost random nucleotide sequences were used. Limitations on the random nature are illustrated in Tab. 1. Two different sets of oligonucleotides were synthesized. In set A the hypervariable regions are predominantly random apart from those few positions where almost exclusively certain amino acids occur. In set B, the random nature of the nucleotide sequences in those positions where pre-  
20 dominantly one or few amino acids occur was additionally limited.

25 The oligonucleotides were purified by HPLC chromatography or polyacrylamide gel electrophoresis, and then 5'-phosphorylated.

**Example 3: Ligation of the synthetic oligonucleotides**

30 The oligonucleotides in Tab. 1 were ligated together stepwise on an antibody DNA template. For this purpose, large amounts (about 1 mg) of single-stranded M13mp=18 DNA containing the antibody DNA inserts were isolated. In order to separate the antibody DNA from the vector, the inserts were made double-stranded on the two ends using



two appropriate oligonucleotides and were digested using the enzymes PstI and HindIII (heavy chain) or using EcoRV and BamHI (light chain). The antibody DNA was then purified using agar gel electrophoresis.

5 On these DNA templates, first only three oligonucleotides were ligated: H1, pH2 and pH3 (heavy chain), and L1, pL2 and pL3 (light chain), H1 and L1 having been marked first with <sup>32</sup>P at their 5' end ("p" designates 5'-phosphorylated). Amounts of 100pmol of each oligonucleotide were  
10 used. The hybridized oligonucleotides were purified on 2% agarose gels and analyzed on a sequencing gel. The amount was determined by a radioactivity measurement. Equimolar amounts of pH4 and pH5 (heavy chain), and pL4 and pL5 (light chain) were then ligated onto the already ligated  
15 oligonucleotides on each particular template. These DNAs were then purified as in the preceding step and the procedure was repeated up to the purification step, using equimolar amounts of pH6 and pH7. Finally, the ligated oligonucleotides were purified by means of a denaturing  
20 polyacrylamide gel and preferably amplified using the polymerase chain reaction (PCR). Alternatively or in order to avoid losses caused by the last purification step, the oligonucleotides were amplified using PCR directly after the last ligation step. The primers H1 and  
25 H8 (heavy chain), and L1 and L6 (light chain) were used under standard conditions for the PCR. Amplified template DNA was digested selectively using KpnI (light chain) or using AluI (heavy chain). Where appropriate, a second amplification step using the PCR was subsequently carried  
30 out.

**Example 4: Insertion of the antibody DNA into the expression plasmid**

The synthesized antibody DNA was cut using the restriction enzymes PstI and BanIII (heavy chain), and BstEII  
35 and KpnI (light chain). The bands having the expected molecular weight were purified by agar gel



electrophoresis, precipitated using ethanol and then, in two steps (first the DNA of the light chain and then the DNA of the heavy chain), ligated into the pUC-HP<sub>2</sub>L (see above) which had been cut and purified in the same way.

5 The HP<sub>2</sub>L insert was then ligated into the restriction sites PstI and BamHI of the plasmids pKK233-2-P<sub>1</sub> (see Example 1). An analogous way was used for the HTP<sub>2</sub>L fragment. The antibody library is therefore established in the antibody expression plasmid (Tab. 6). The reason  
10 for intermediate cloning in pUC is that the presence of individual parts of the antibody operon in the expression vector has an adverse influence on the growth of the bacterial host (see above also).

15 **Example 5: Expression and screening of antibodies in E. coli**

Competent E. coli are transfected with pFMT plasmids containing the inserted antibody-DNA library, grown on agarose plates and then incubated using nitrocellulose filters coated with the desired antigen. After removing  
20 non-specifically bound antibodies, the active clones are identified with a labeled antibody against the human immunoglobulins secreted from E. coli. In the preferred embodiment, the antibody YL 1/2 which is directed against the Tag sequence is used for this purpose.

Legend for Fig. 1:

Restriction map of the expression vector pKK233-2 (Amann and Brosius, loc. cit.).

P<sub>trc</sub> denotes hybrid tryptophan lac promoter

5 RBS denotes ribosome binding site

rrnB denotes ribosomal RNA B operon

5S denotes gene for 5S RNA

Before cloning antibody DNA in the expression vector, the following alterations were carried out:

- 10
- 1) The SalI and EcoRI restriction sites were removed together with the DNA between them.
  - 2) The HindIII restriction site was converted to a BamHI restriction site.

TAB. 1

Oligonucleotide for the synthesis of a library of anti-body DNA (variable parts)

## Set A

H1	5'CCAGGTCCAACCTGCAGGAGAGCGGTCCAGGTCTTGTGAGACCTAG3
H2	5'CCAGACCCTGAGCCTGACCTGCACCCGTG3'
H3	5'TGTCTGGCTTCACCTTCAGC T1/2 TT CTTT1/2TTGGGTGCGCCAGCCACCTGGAC3' C C   C CC  A CC CC   A AG  G AA AG   G G   3 GG G
H4	5'GAGGTCTTGAGTGGATTGGT T1/2TT TAT T1/2TT T1/2TACGCGTGACAATGCTGGTAGAC3'   C CC    C CC  C C   A AG    A AG  A A   G G   10   G G   5 G G
H5	5'ACCAGCAAGAACCAGTTCAGCCTGCGTCTCAGCAGCGTGACAGC3'
H6	5'CGCCGACACCGCGGTCTACTACTGTGCGCGC T1/2 TT TGGGGTCAGGGCT3'   C CC    A AG    G G   10
H7	5'CCCTCGTCACAGTCTCCTCA3'
H8	5'CTGTGACGAGGCTGCCCTGACCCCA3'
L1	5'GCGCCAGCGTGGGTGACAGG3'
L2	5'GTGACCATCACCTGTT1/2TTGTT T1/2TT CTTT1/2TTTGGTAACAGCAGAAGCCAGGT3' C CC CC  C CC  A CC CC A AG GA  A AG  G AA AG G G G  G G   7 GG G
L3	5'AAGGCTCCAAAGCTGCTGATCTAC T1/2TT GGTGTGCCAAGCCGTTTCAGCGGTAGCGGT3'   C CC    A AG    G G   7
L4	5'AGCGGTACGGACTTCACCTTCACCATCAGCAGCCTCCAGCCAGAGGAC3'
L5	5'ATCGCCACCTACTACTGCCAG T1/2TT TTCGGCCAAAGGTAC3'   C CC    A AG    G G   8
L6	5'CCACCTTGGTACCTTGGCCGAA3'



- 13 -

## Set B

H1, H2, H5, H7, H8 and L1, L4 and L6 are identical to those in set A.

H3 5'TGTCTGGCTTCACCTTCAGC AC10%T95% TT20%C TC20%G T5% TA20% T2% T5% T70%  
C C GA45%A5% C10% GG80% A85% G80% C28%A75%G30%  
G45% A70% G10% A50%G20%  
G20%

TGGGTGCGCCAGCCACCTGGAC3'

H4 5'GAGGTCTTGAGTGGATTGGT T14.5%TT AT90%C T14.5%TT T5% T10%T70% T14.5%TT  
C28.5%CC G10% C28.5%CC C70%C80%G30% C28.5%CC  
A28.5%AG A28.5%AG A20%A10% A28.5%AG  
G28.5%G G28.5%G G5% G28.5%G

AAT A15%A20%C ACT AT16%A A70%C10%C TAT A80%C20%C CCC AC10%C T40%TC C5% T5% G  
GG G85%G80% GA C80% G30%A70%A G20%A60% GAA A40%A C20% A90%A90%  
G A4% G20% G20% G50% G40% G5% G5%

A20%A10%T CGCGTGACAATGCTGGTAGAC3'  
G80%G90%

H6 5'CGCCGACACCGCGGTCTACTACTGTGCGCGC T1/2 TT GC25%C TATTGGGGTCAGGGCT3'  
C CC A75%  
A AG  
G G8

L2 5'GTGACCATCACCTGT CAA T30%CG AGT C75%AA T30%T10%C C10%T40%T90%  
AG G70% A10% C10%C30% A50%C10%A10%  
G15% A30%A60% G40%A10%  
G30% G40%

A70%T20%T70% C5% T5% T90% T90%T10%C C70%TA20% AC40%T TGGTAACAGCAGAAGCCAGGT3'  
G30%A40%A30% A90%C20%A10% C2% C5% A20% G80% GA40%  
G40% G5%.A40% A6% A85% G10% G20%  
G35% G2%

L3 5'AAGGCTCCAAAGCTGCTGATCTAC T14.5%TT A40%T20%T AC5% C AC20%T70% CTA  
C28.5%CC G60%C70% A10% A60%A30% G  
A28.5%AG A7% G85% G20%  
G28.5%G G3%

C20%C70%C20% T70%CT GGTGTGCCAAGCCGTTTCAGCGGTAGCGGT3'  
G80%A30%A80% C15%  
A15%

L5 5'ATCGCCACCTACTACTGC CT10%A CT20%T10% T60%C30%G T35%T5% T T15%T5% C  
A90% A80%G90% A10%G70% C5% C20% C10%C20%  
G30% A40%A50% A60%A75%  
G20%G25% G15%

T14.5%TT CT15%C90% T14.5%TT ACGTTCCGCCAAGGTAC  
C28.5%CC C70%A10% C28.5%CC  
A28.5%AG A15% A28.5%AG  
G28.5%G G28.5%G

**Notes for Tab. 1**

The random nature of the oligonucleotides of set B was limited in a manner which generates approximately the relevant amount of frequent amino acids in each position of the hypervariable regions (in accordance with the tables of Kabat et al, loc. cit.). In this strategy the number of expected stop codons was also reduced even further. In contrast with the oligonucleotides in set A, a restriction site for MluI was not introduced.

TAB. 2

CONSTRUCTION OF THE VECTOR pFMT FOR THE EXPRESSION AND  
SECRETION OF ANTIBODIES IN BACTERIA

DNA OF THE VARIABLE DOMAIN OF A HUMAN LYSOZYME ANTIBODY

⇓

INTRODUCTION OF RESTRICTION SITES BY SITE DIRECTED MUTAGENESIS

⇓

SYNTHESIS OF THE LEADER SEQUENCE OF PECTATE LYASE AND OF THE RIBOSOME BINDING SITE

⇓

LIGATION INTO BACTERIAL EXPRESSION PLASMIDS

⇓



P/O: promoter/operator, RBS: ribosome binding site, P2: leader sequence of pectate lyase, VH: variable domain of the heavy chain, VL: variable domain of the light chain



TAB. 3

Sequences of the leader sequences P1 and P2 in the antibody operon, and of the Tag sequences

P1

Leader sequence of pectate lyase (P1)

M K Y L L P T A A A G L L L L A A Q P A M A Q V Q L Q  
CATGAAATACCTCTTGCCTACGGCAGCCGCTGGCTTGCTGCTGCTGGCAGCTCAGCCGGCGATGGCGCAAGTTCAGCTGCA(G)  
PstI

P2

RBS

Leader sequence of pectate lyase (P2)

M K Y L L P T A A A  
(C)TGCAGCCAAGCTTGAATTCATTAAAGAGGAGAAATTAAGTCCATGAAGTACTTACTGCCGACCGCTGCGGCG  
PstI HindIII

G L L L L A A Q P A M A D I  
GGTCTCCTGCTGTTGGCGGCTCAGCCGGCTATGGCTGATATCGGATCCAGCT  
EcoRV BamHI

The nucleotides in parentheses are the adjacent nucleotides of the plasmid

The leader sequences were synthesized by hybridization of the following oligonucleotides.

P1

a. 5'CATGAAATACCTCTTGCCTACGGCAGCCGCTGGCTTG3'

b. 5'TTAAGTCCATGAAGTACTTACTGCCGACCGCTGCG3'

c. 3'ACGTCGGTTCGAACCTAAGTTTAACTCCTCTTTAATTGAGGTACTTCATGAATGACGGCTGGCGACGCCGCCAGAGGA

CGACAACCGCCGAGTCGGCCGATACCGACTATAGCCTAGGTCGA5'

d. 5'GCTCAGCCGGCTATGGCTGATATCGGATCC3'

e. 5'GCGGGTCTCCTGCTGTTGGCG3'

The Tag sequences were synthesized by hybridization of the following sequences:

a. 5'CCTTAGTCACAGTATCCTCAGAAGGTGAAGAATTCTA3'

b. 5'AGCTTAGAATTCTTCACCTTCTGAGGATACTGTGACTAAGGAGCC3'

TAB. 4

Nucleotide sequences of antibody DNA

a) Heavy chain (variable domain), HuVhlys HindIII.....

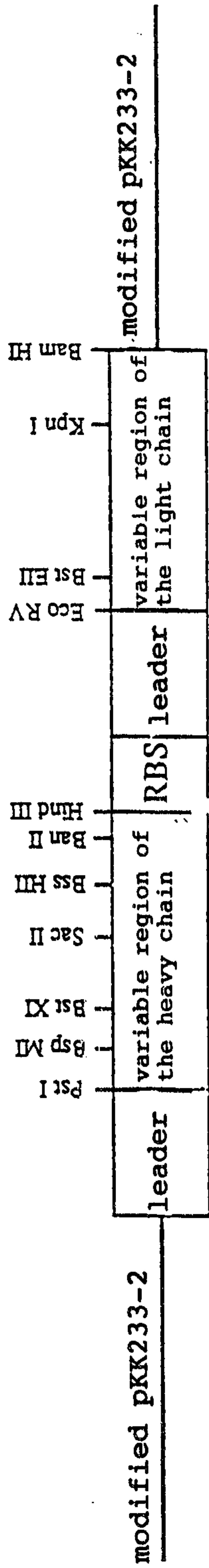
.....G V H S Q V Q L Q E S G P G L V R  
CTCTCCACAGGTGTCCACTCCCAGGTCCAACTGCAGGAGAGCGGTCCAGGTCTTGTGAGA  
PstI  
P S Q T L S L T C T V S G F T F S /G//Y//G/  
CCTAGCCAGACCCTGAGCCTGACCTGCACCGTGTCTGGCTTCACCTTCAGCGGTATGGT  
BspMI  
/V /N /W V R Q P P G R G L E W I G /M/ I/ W/ G/  
GTAAACTGGGTGAGACAGCCACCTGGACGAGGTCTTGAGTGGATTGGAATGATTGGGGT  
CDR2 60 70  
/D /G /N /T /D /Y /N /S /A /L /K /S R V T M L V D T  
GATGGAACACAGACTATAATTCAGCTCTCAAATCCAGAGTGACAATGCTGGTAGACACC  
80 90  
S K N Q F S L R L S S V T A A D T A V Y  
AGCAAGAACCAGTTCAGCCTGAGACTCAGCAGCGTGACAGCCGCCGACACCGCGGTCTAT  
SacII  
Y C A R E /R /D /Y /R /L /D /Y W G Q G S L V T  
TATTGTGCAAGAGAGAGATTATAGGCTTGACTACTGGGGTCAGGGCTCCCTCGTCACA  
BanII  
V S S Stop  
GTCTCCTCATAAGCTTCCTTACAACCTCTCTTCTATTTCAGCTTAA.....BamHI  
HindIII

b) Light chain (variable domain), Hu Vllys HindIII.....

G V H S D I Q N T Q S P S S L S A  
CTCTCCACAGGTGTCCACTCCGATATCCAGATGACCCAGAGCCCAAGCAGCCTGAGCGCC  
EcoRV  
S V G D R V T I T C R/ A/ S/ G/ N/ I/ H/ N/ Y/ L  
AGCGTGGGTGACAGGGTGACCATCACCTGTAGAGCCAGCGGTAACATCCACAACCTACCTG  
BstEII  
/A/ W Y Q Q K P G K A P K L L I Y /Y/ T/ T/ T  
GCTTGGTACCAGCAGAAGCCAGGTAAGGCTCCAAAGCTGCTGATCTACTACACCACCACC  
60 70  
/L/ A/ D G V P S R F S G S G S G T D F T F  
CTGGCTGACGGTGTGCCAAGCAGATTACGCGGTAGCGGTAGCGGTACCGACTTCACCTTC  
80 90 CDR3  
T I S S L Q P E D I A T Y Y C /Q /H /F /W /S  
ACCATCAGCAGCCTCCAGCCAGAGGACATCGCCACCTACTACTGCCAGCACTTCTGGAGC  
100  
/T /P /R /T F G Q G T K V E I K R..E STOP  
ACCCCAAGGACGTTTCGGCCAAGGTACCAAGGTGGAAATCAAACGTGAGTAGAATTAAAC  
KpnI  
TTTGCTTCCTCAGTTGGATCC  
BamHI

TAB. 5

The Antibody Expression Plasmid pFMT

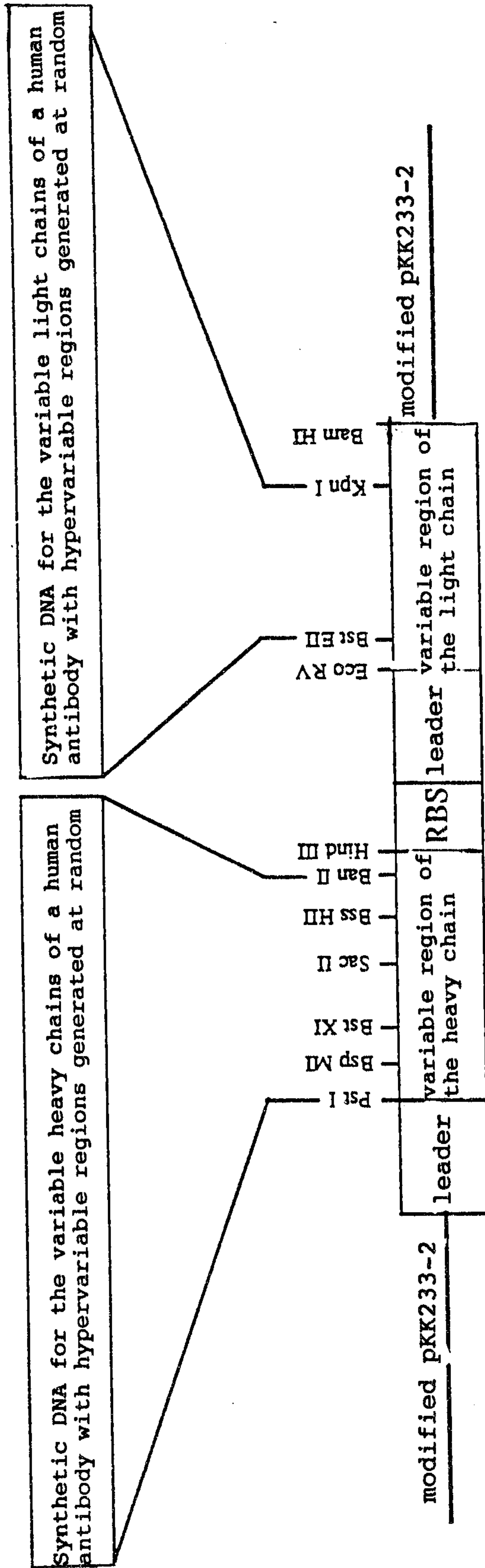


There is an RBS in the plasmid upstream of the heavy chain part but is not drawn in here.



TAB. 6

Insertion of the antibody libraries in the expression vector pFMT



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**THE EMBODIMENTS OF THE INVENTION IN WHICH AN EXCLUSIVE  
PROPERTY OR PRIVILEGE IS CLAIMED ARE DEFINED AS FOLLOWS:**

1. A synthetic human antibody-DNA library obtained by:
  - 5 (1) synthesizing almost random sequences for the antibody hypervariable regions, wherein (a) relatively conserved amino acids in the hypervariable regions have been taken into account in the choice of appropriate nucleotides during the oligonucleotide synthesis and (b) the ratio of the nucleotides used is chosen such that a nonsense codon is to be  
10 expected at most in every 89th position;
  - (2) inserting said almost random sequences for the hypervariable regions into a human antibody framework; and
  - (3) incorporating said almost random sequences combined with said human antibody framework into an expression vector.
- 15 2. The synthetic human antibody-DNA library as claimed in claim 1, wherein the synthesized almost random sequences for the hypervariable regions are amplified before the incorporation in the expression vector.
- 20 3. The synthetic human antibody-DNA library as claimed in claim 1, wherein said almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.
4. The synthetic human antibody-DNA library as claimed in claim 2,  
25 wherein said almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.
5. The synthetic human antibody-DNA library of claim 1 wherein the DNA is transfected into a microbial expression system.
- 30 6. The synthetic human antibody-DNA library of claim 2 wherein the DNA is transfected into a microbial expression system.

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7. The synthetic human antibody-DNA library of claim 3 wherein the DNA is transfected into a microbial expression system.

5 8. The synthetic human antibody-DNA library of claim 4 wherein the DNA is transfected into a microbial expression system.

9. The synthetic human antibody-DNA library of claim 5 wherein the microbial expression system is bacterial.

10

10. The synthetic human antibody-DNA library of claim 6 wherein the microbial expression system is bacterial.

11. The synthetic human antibody-DNA library of claim 7 wherein the  
15 microbial expression system is bacterial.

12. The synthetic human antibody-DNA library of claim 8 wherein the microbial expression system is bacterial.

20 13. The synthetic human antibody-DNA library of claim 9 wherein the bacteria are *E. coli*.

14. The synthetic human antibody-DNA library of claim 10 wherein the bacteria are *E. coli*.

25

15. The synthetic human antibody-DNA library of claim 11 wherein the bacteria are *E. coli*.

16. The synthetic human antibody-DNA library of claim 12 wherein the  
30 bacteria are *E. coli*.



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17. A process for preparing a synthetic human antibody-DNA library, which comprises:

(1) synthesizing almost random sequences for the antibody hypervariable regions, wherein

5 (a) relatively conserved amino acids in the hypervariable regions have been taken into account in the choice of appropriate nucleotides during the oligonucleotide synthesis and

(b) the ratio of the nucleotides used is chosen such that a nonsense codon is to be expected at most in every 89<sup>th</sup> position;

10 (2) inserting said almost random sequences for the hypervariable regions into a human antibody framework; and

(3) incorporating said almost random sequences combined with said human antibody framework into an expression vector.

15 18. The process for preparing a synthetic human antibody-DNA library as claimed in claim 17, wherein the synthesized random sequences are amplified before incorporation in an expression vector.

19. The process for preparing a synthetic human antibody-DNA library as  
20 claimed in claim 17, wherein the almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.

20. The process for preparing a synthetic human antibody-DNA library as  
25 claimed in claim 18, wherein the almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.

21. The process as claimed in claim 17, wherein said expression vector further comprises DNA coding for a marker peptide and the desired clones are identified using antibodies against the marker peptide.

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22. The process as claimed in claim 21, wherein said marker peptide is the TAG sequence and the desired clones are identified using the antibody YL 1/2.

5 23. The process as claimed in claim 18, wherein said expression vector further comprises DNA coding for a marker peptide and the desired clones are identified using antibodies against the marker peptide.

24. The process as claimed in claim 23, wherein said marker peptide is the TAG sequence and the desired clones are identified using the antibody YL1/2.

25. The process for preparing a synthetic human antibody-DNA library of claim 17 wherein the variable regions are incorporated into pFMT.

15 26. The process for preparing a synthetic human antibody-DNA library of claim 18 wherein the variable regions are incorporated into pFMT.

27. The process for preparing a synthetic human antibody-DNA library of claim 17 wherein the expression vector is transfected into a microbial expression system.

28. The process of claim 27 wherein the microbial expression system is bacterial.

25 29. The process of claim 28 wherein the bacteria are *E. coli*.

30. The process for preparing a synthetic human antibody-DNA library of claim 18 wherein the expression vector is transfected into a microbial expression system.

30

31. The process of claim 30 wherein the microbial expression system is bacterial.

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32. The process of claim 31 wherein the bacteria are *E. coli*.

33. A process for obtaining clones secreting specific human antibodies  
5 comprising the steps of:

(a) screening synthetic human antibody-DNA libraries using specific antigens, and

(b) isolating said clones which secrete the desired specific human antibodies,

10 wherein said specific human antibody-DNA library is obtained by:

(1) synthesizing almost random sequences for the antibody hypervariable regions, wherein

(a) relatively conserved amino acids in the hypervariable regions have been taken into account in the choice of appropriate nucleotides during the  
15 oligonucleotide synthesis and

(b) the ratio of the nucleotides used is chosen such that a nonsense codon is to be expected at most in every 89<sup>th</sup> position;

(2) inserting said almost random sequences for the hypervariable regions into a human antibody framework; and

20 (3) incorporating said almost random sequences combined with said human antibody framework into an expression vector.

34. The process for obtaining clones secreting specific human antibodies  
25 as claimed in claim 33, wherein the process of obtaining the specific human antibody-DNA library further comprises the step of amplifying the almost random sequences of the hypervariable regions before incorporation into the expression vector.

30 35. The process for obtaining clones secreting specific human antibodies as claimed in claim 33, wherein the almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.



- 25 -

36. The process for obtaining clones secreting specific human antibodies as claimed in claim 34, wherein the almost random sequences are incorporated into antibody variable regions derived from HuVhlys or HuVlls.

5

37. The process as claimed in claim 33, wherein said expression vector further comprises DNA coding for a marker peptide and the desired clones are identified using antibodies against the marker peptide.

10 38. The process as claimed in claim 37, wherein said marker peptide is the TAG sequence and the desired clones are identified using the antibody YL 1/2.

15 39. The process as claimed in claim 34, wherein said expression vector further comprises DNA coding for a marker peptide and the desired clones are identified using antibodies against the marker peptide.

20 40. The process as claimed in claim 39, wherein said marker peptide is the TAG sequence and the desired clones are identified using the antibody YL 1/2.

41. The process for obtaining clones secreting specific human antibodies of claim 33 wherein the variable regions are incorporated into pFMT.

25 42. The process for obtaining clones secreting specific human antibodies of claim 34 wherein the variable regions are incorporated into pFMT.

30 43. The process for obtaining clones secreting specific human antibodies of claim 33 wherein the expression vector is transfected into a microbial expression system.

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44. The process of claim 43 wherein the microbial expression system is bacterial.

45. The process of claim 44 wherein the bacteria are *E. coli*.

5

46. The process for obtaining clones secreting specific human antibodies of claim 34 wherein the expression vector is transfected into a microbial expression system.

10 47. The process of claim 46 wherein the microbial expression system is bacterial.

48. The process of claim 47 wherein the bacteria are *E. coli*.

15 49. The synthetic human antibody-DNA library as claimed in claim 1 wherein the expression vector further comprises DNA coding for a marker peptide.

20 50. The synthetic human antibody-DNA library as claimed in claim 49 wherein the marker peptide is the TAG sequence.

51. The synthetic human antibody-DNA library as claimed in claim 2 wherein the expression vector further comprises DNA coding for a marker peptide.

25

52. The synthetic human antibody-DNA library as claimed in claim 51 wherein the marker peptide is the TAG sequence.

30 53. The synthetic human antibody-DNA library as claimed in claim 2 wherein the variable regions are incorporated into pFMT.

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54. The synthetic human antibody-DNA library as claimed in claim 53 wherein the variable regions are incorporated into pFMT.



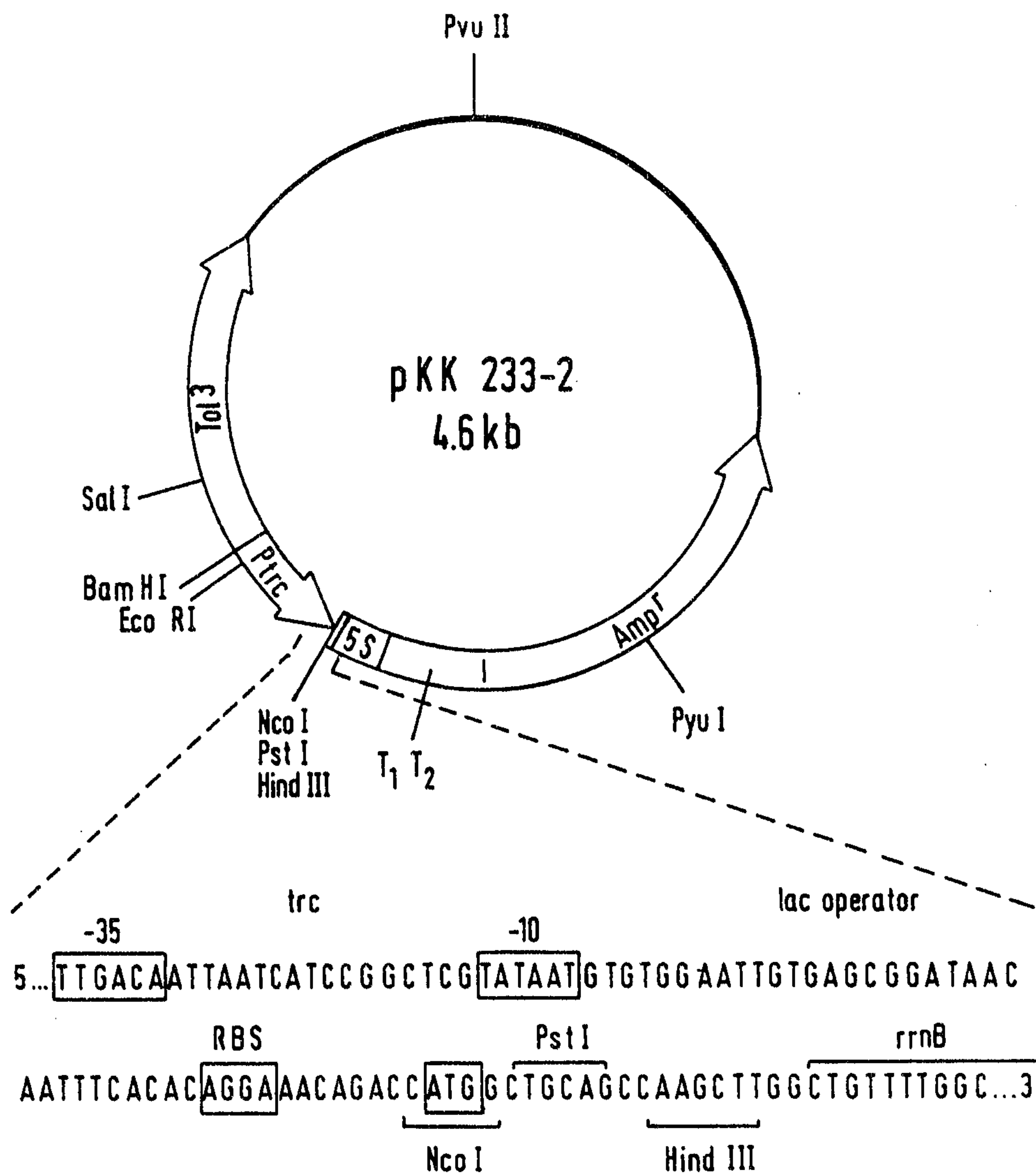


FIG. 1

By: Rogers, Breckin & Parr.