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(54) **POLYVALENT VACCINE**

**Related U.S. Application Data**

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*A61K 39/295* (2006.01)  
*A61P 37/04* (2006.01)  
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(52) **U.S. Cl.** ..... **424/188.1**; 530/324; 530/350; 536/23.72; 435/320.1; 424/208.1

(57) **ABSTRACT**

The present invention relates, in general, to an immunogenic composition (e.g., a vaccine) and, in particular, to a polyvalent immunogenic composition, such as a polyvalent HIV vaccine, and to methods of using same. The invention further relates to methods that use a genetic algorithm to create sets of polyvalent antigens suitable for use, for example, in vaccination strategies.

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(22) Filed: **Dec. 3, 2010**

Fig. 1A

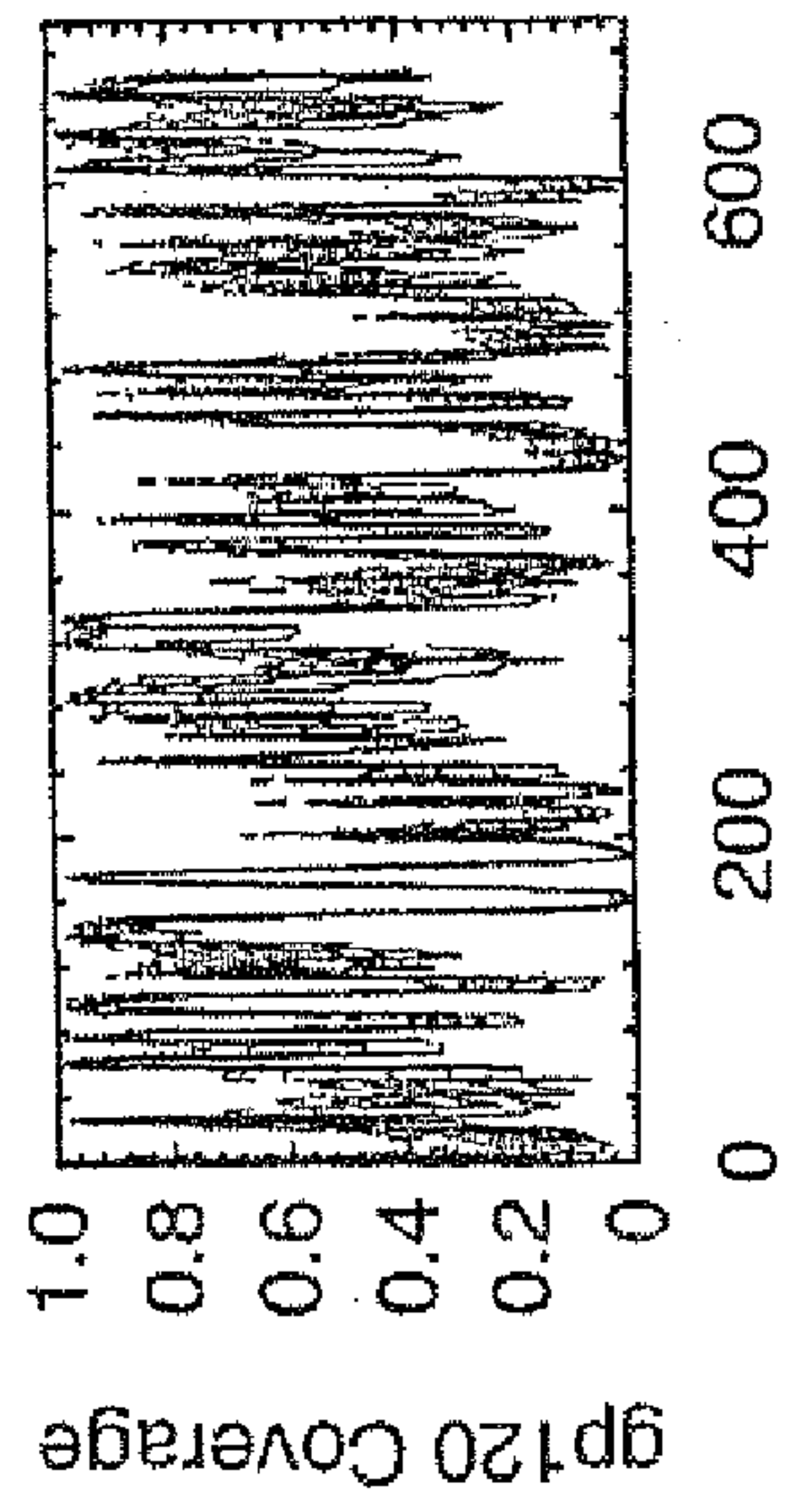


Fig. 1B

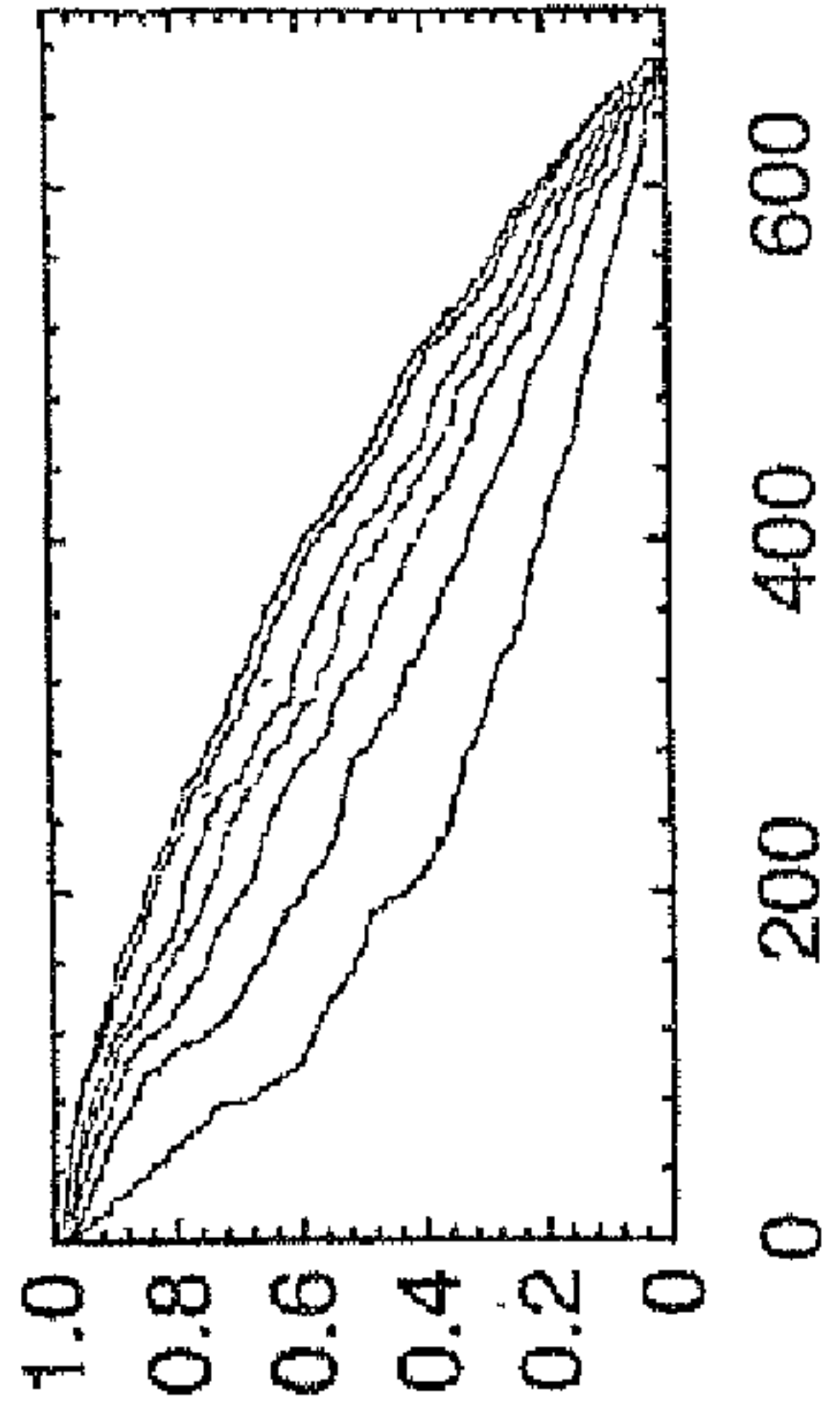


Fig. 1C

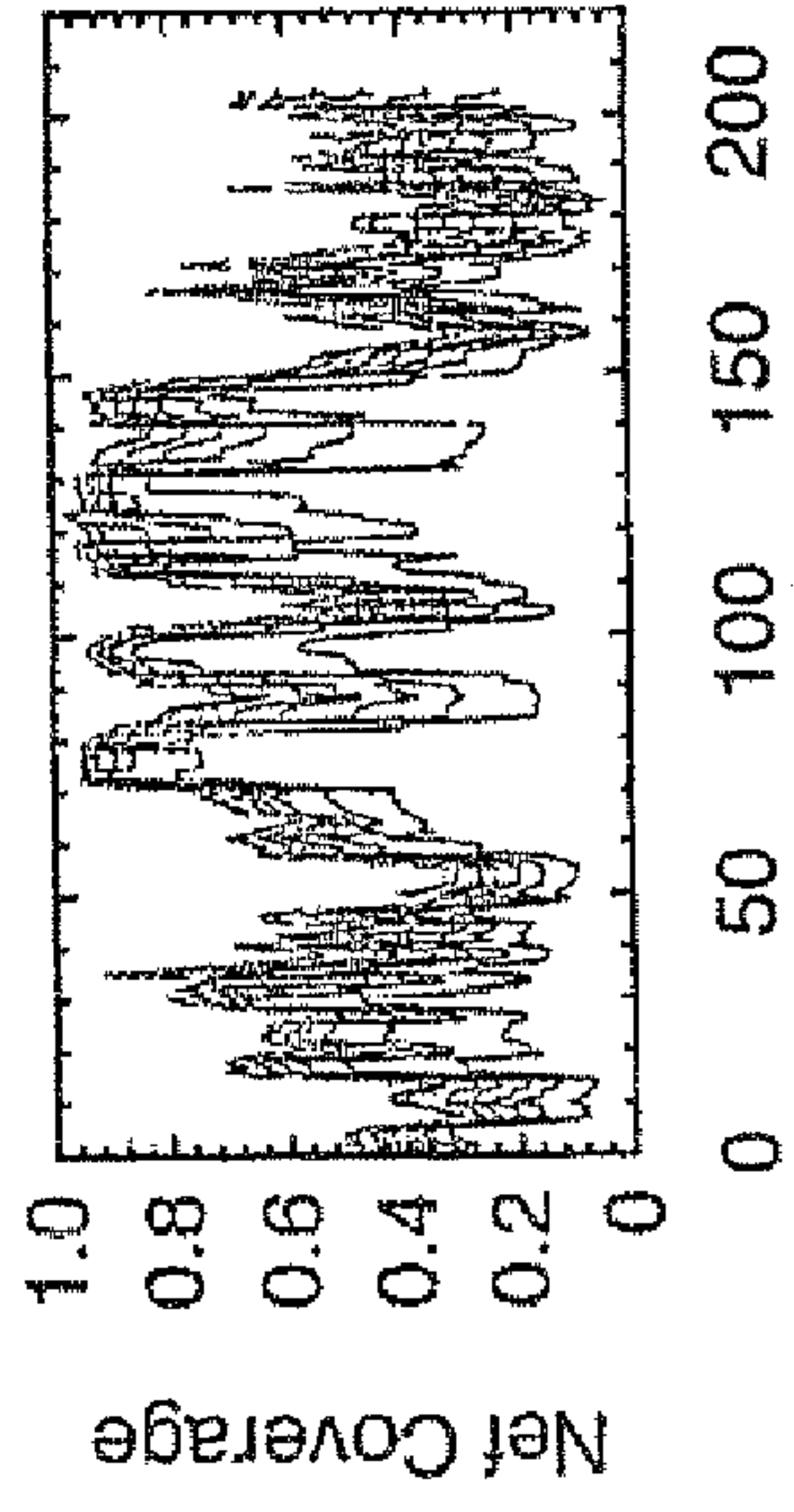


Fig. 1D

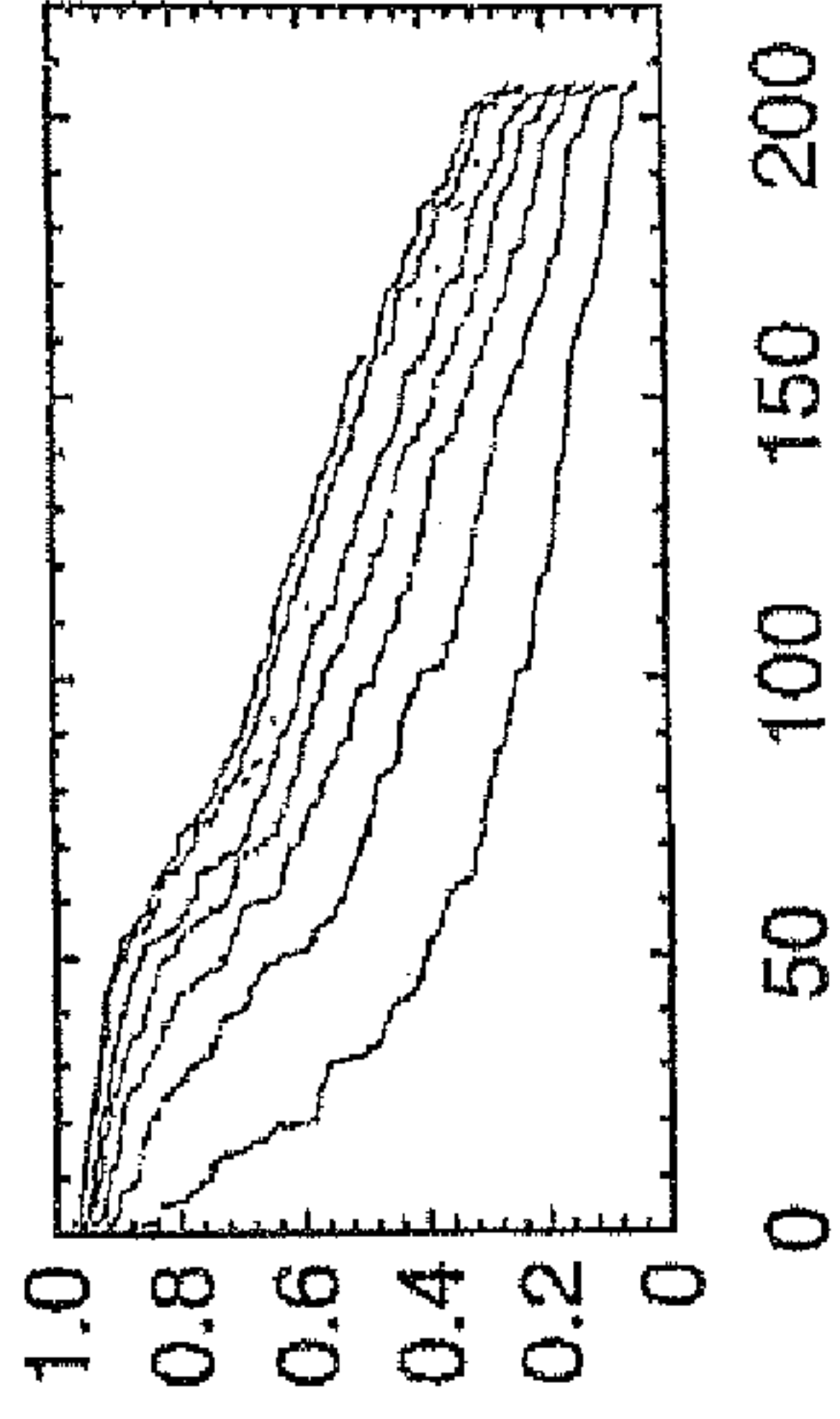


Fig. 1E

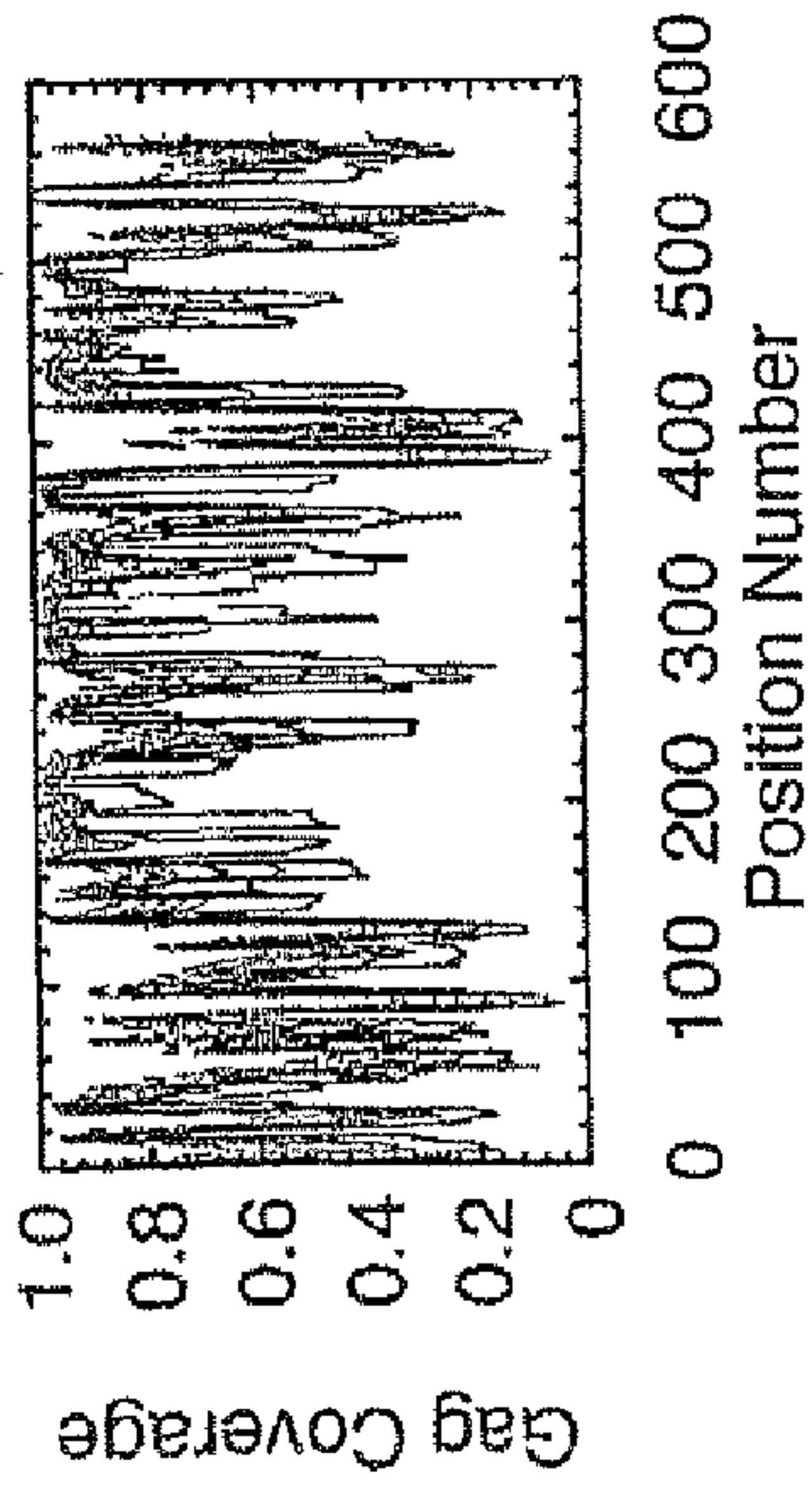


Fig. 1F

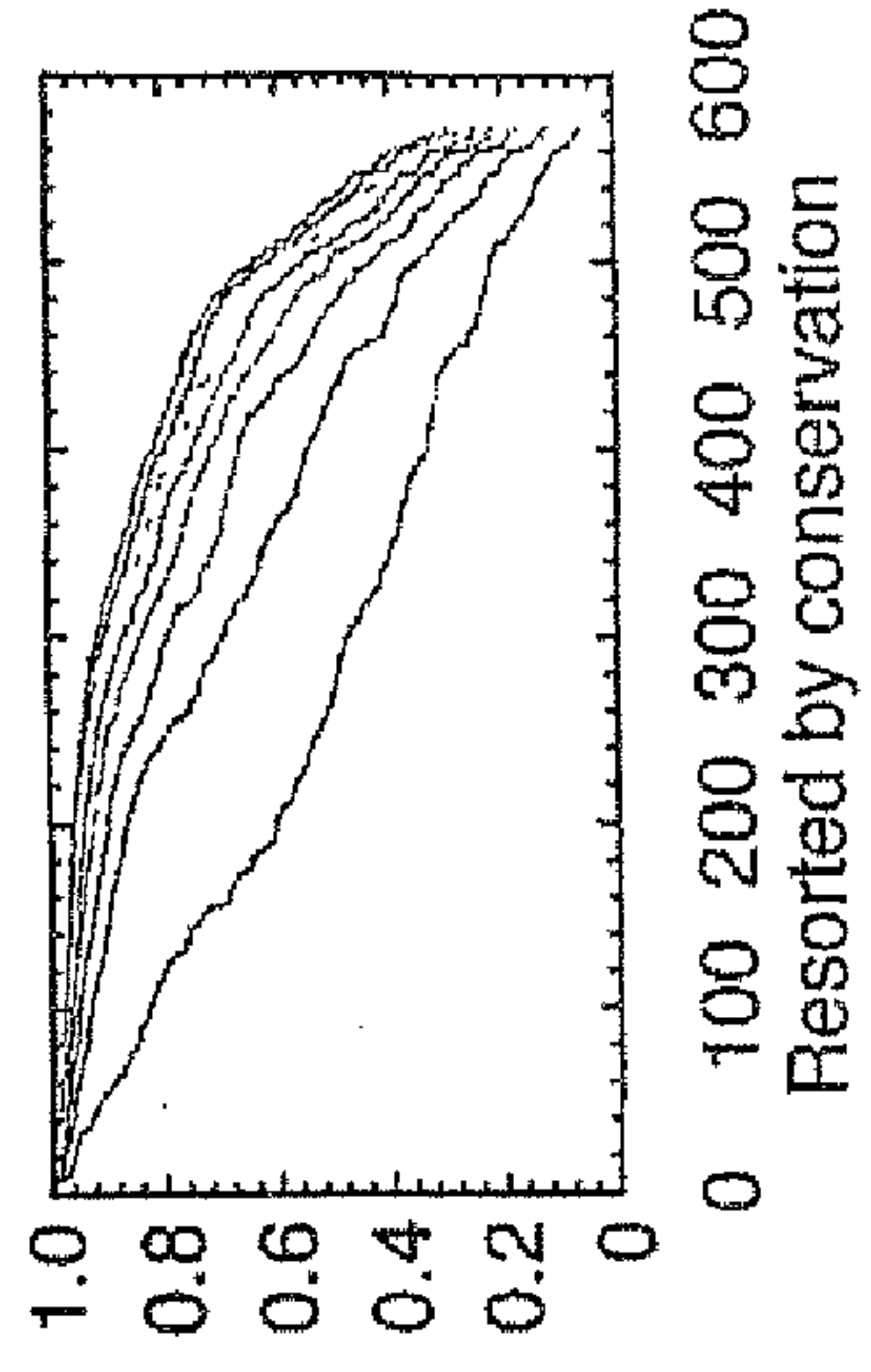




Fig. 2A

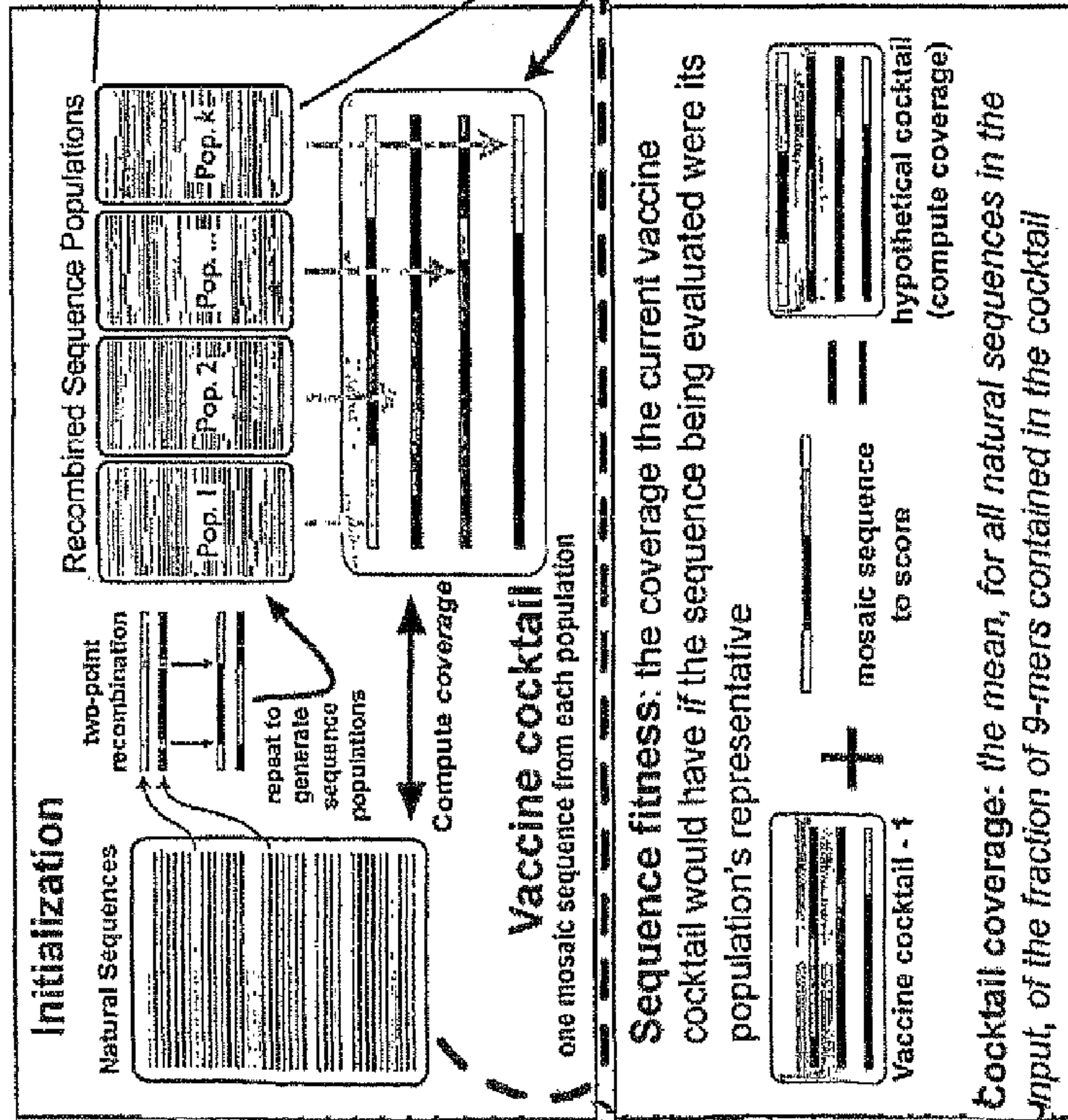


Fig. 2C

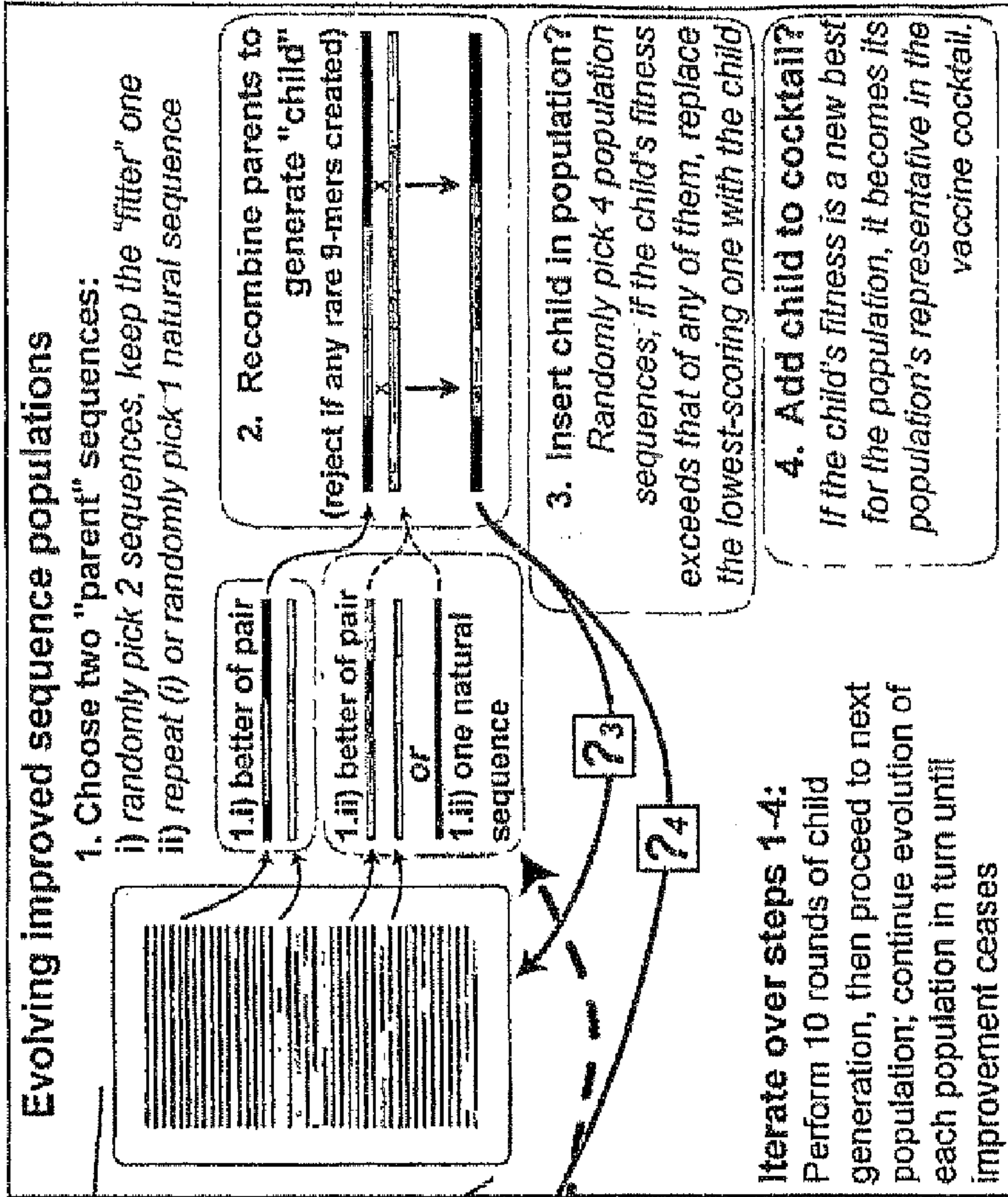


Fig. 2B

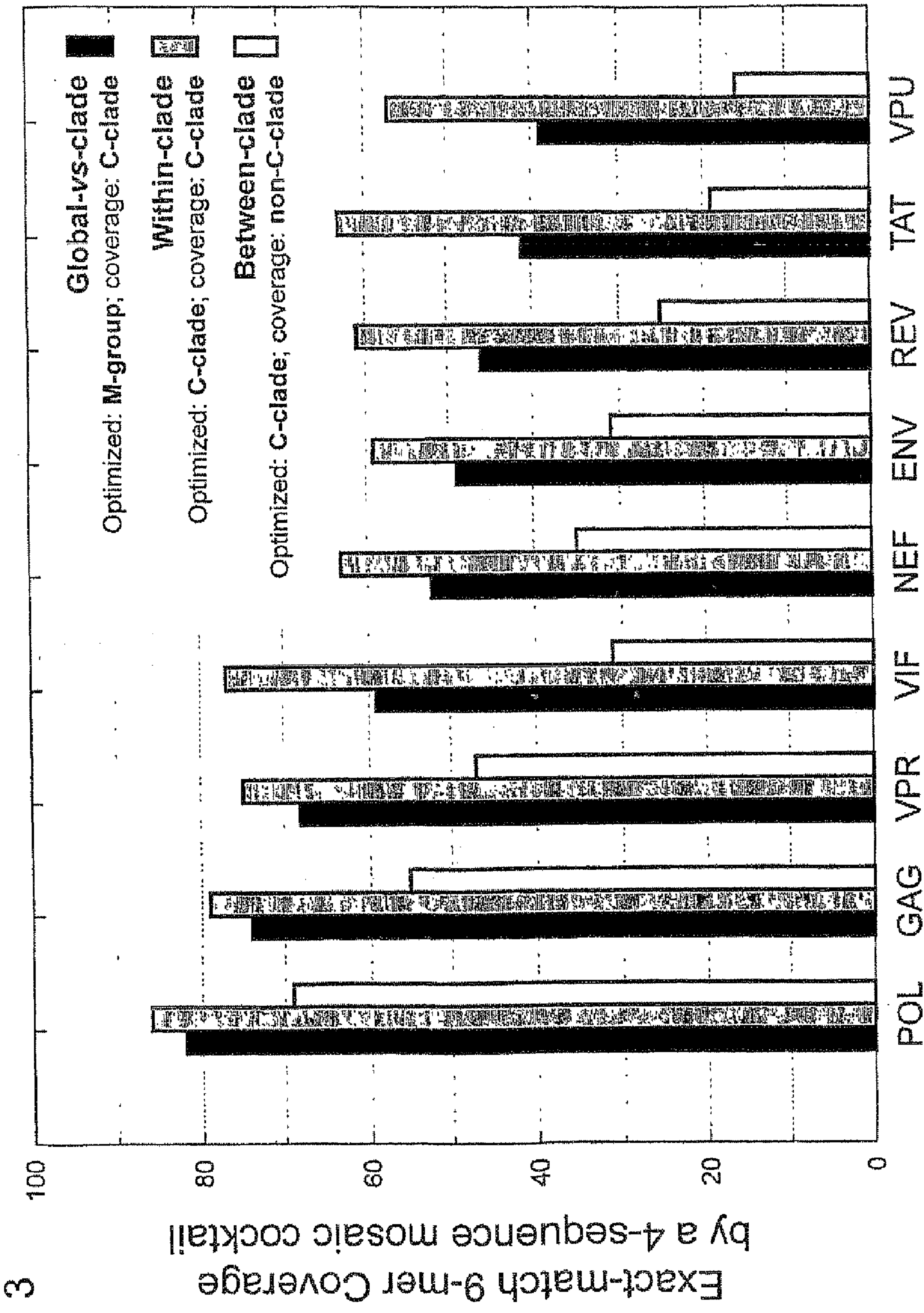


Fig. 3



Fig. 4A

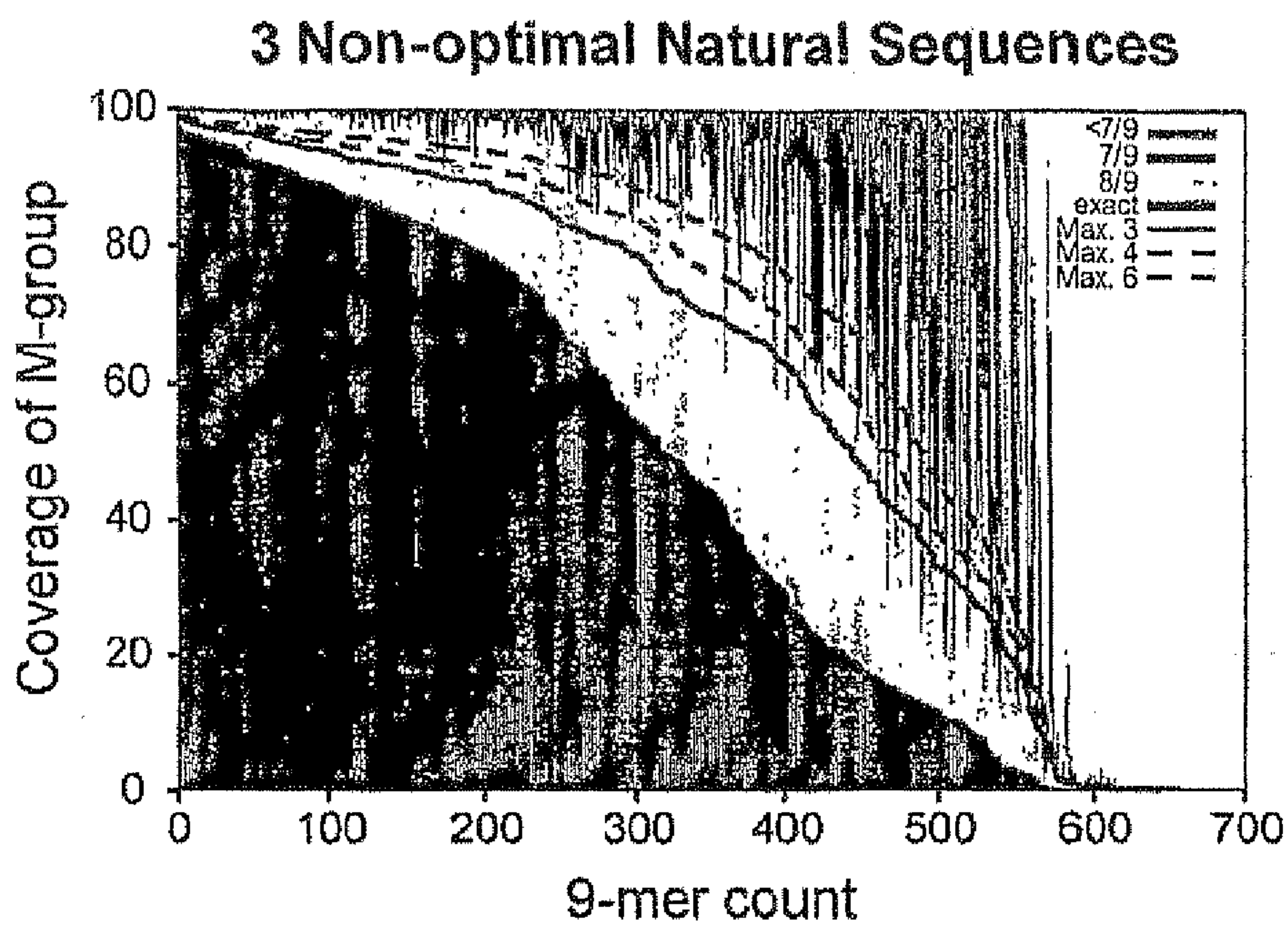


Fig. 4B

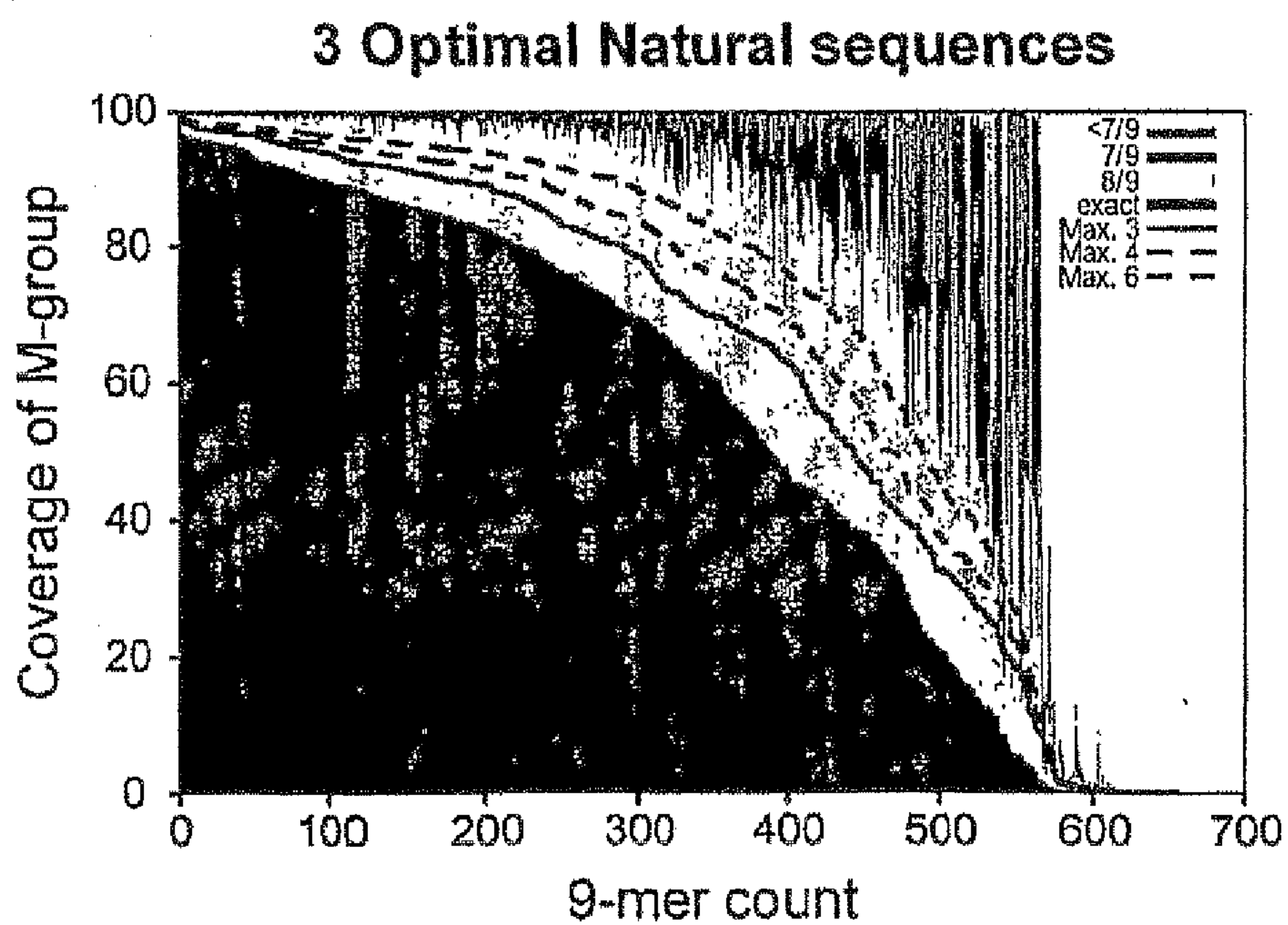


Fig. 4C

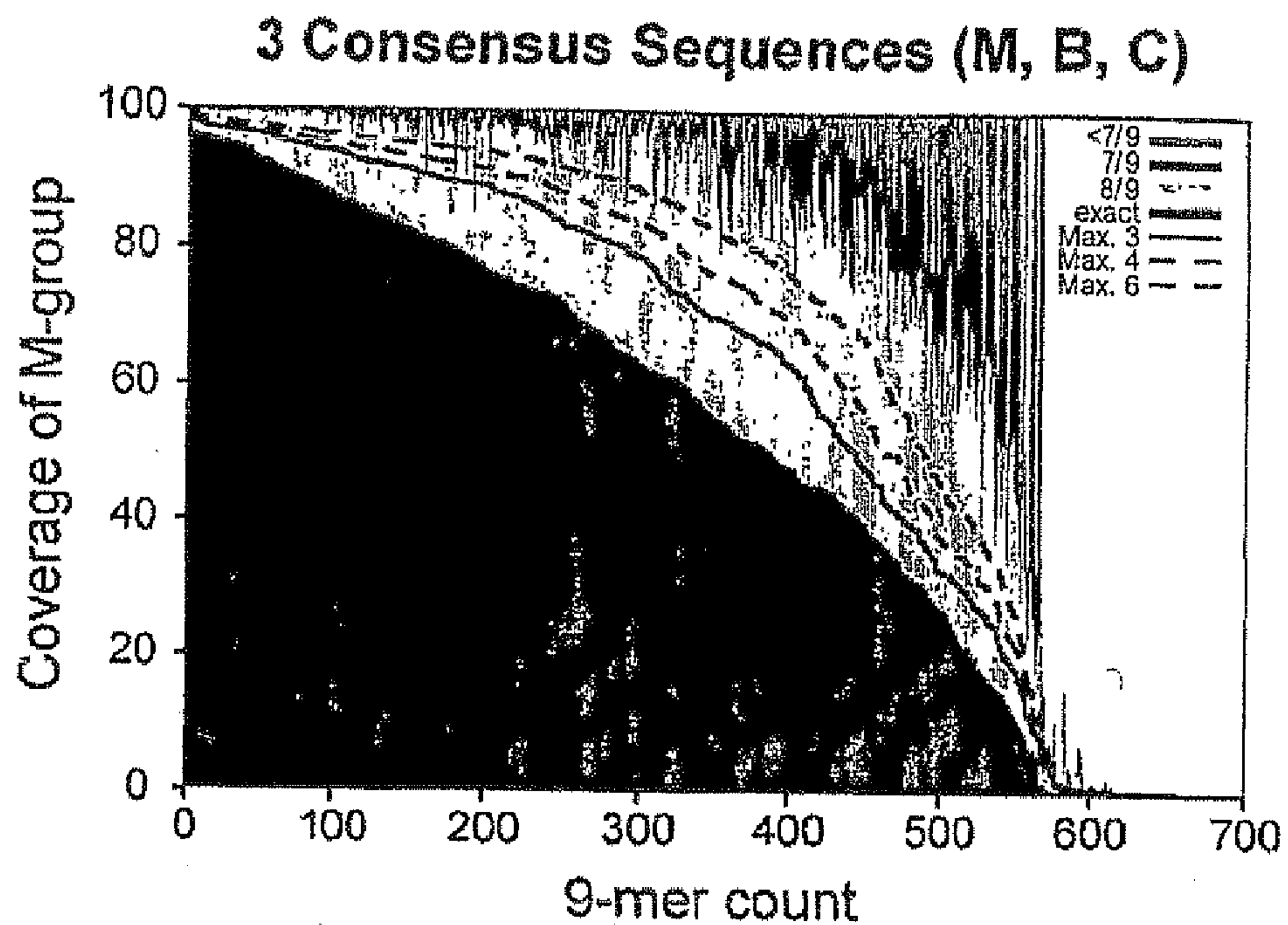


Fig. 4D

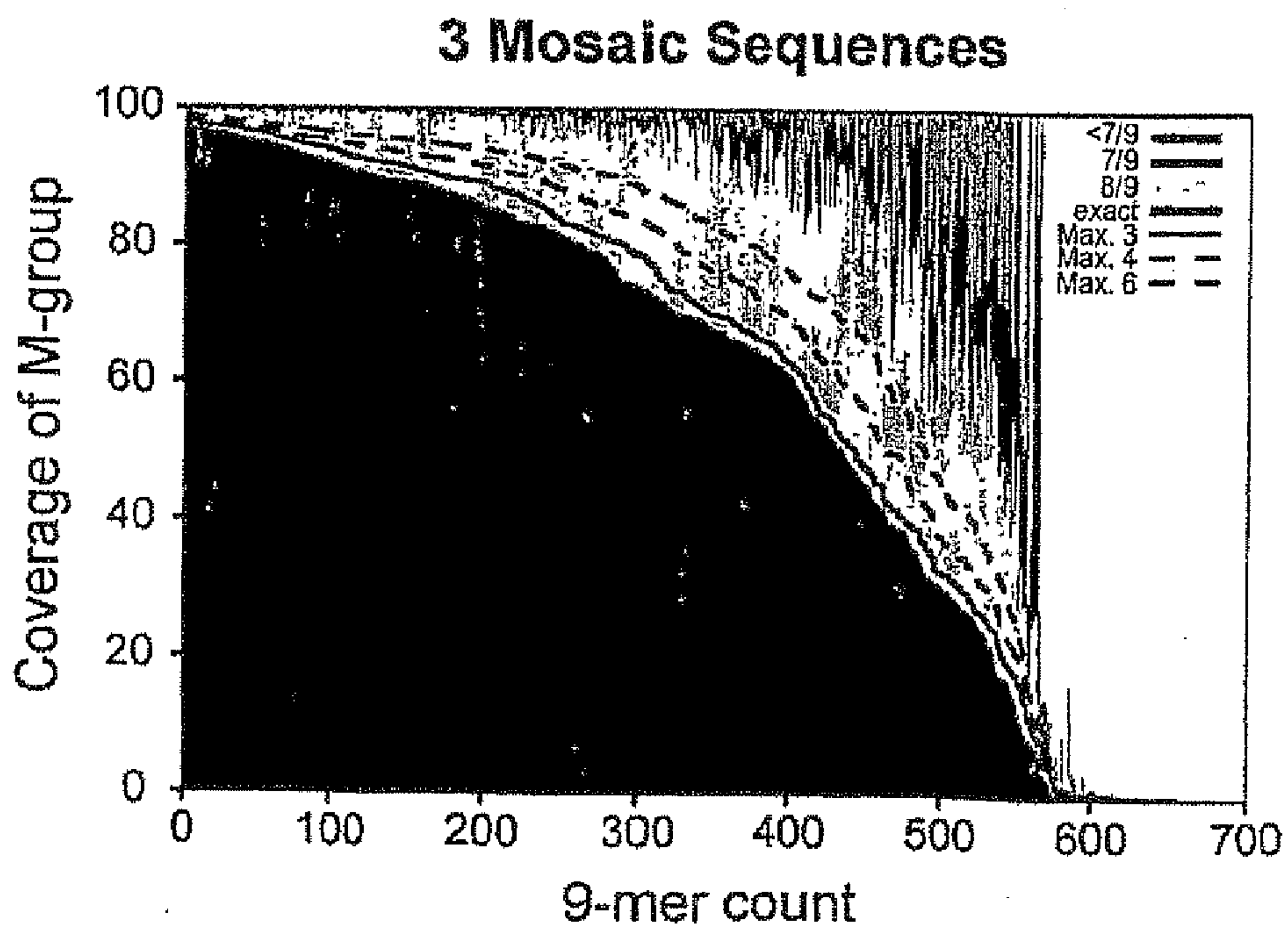




Fig. 4E

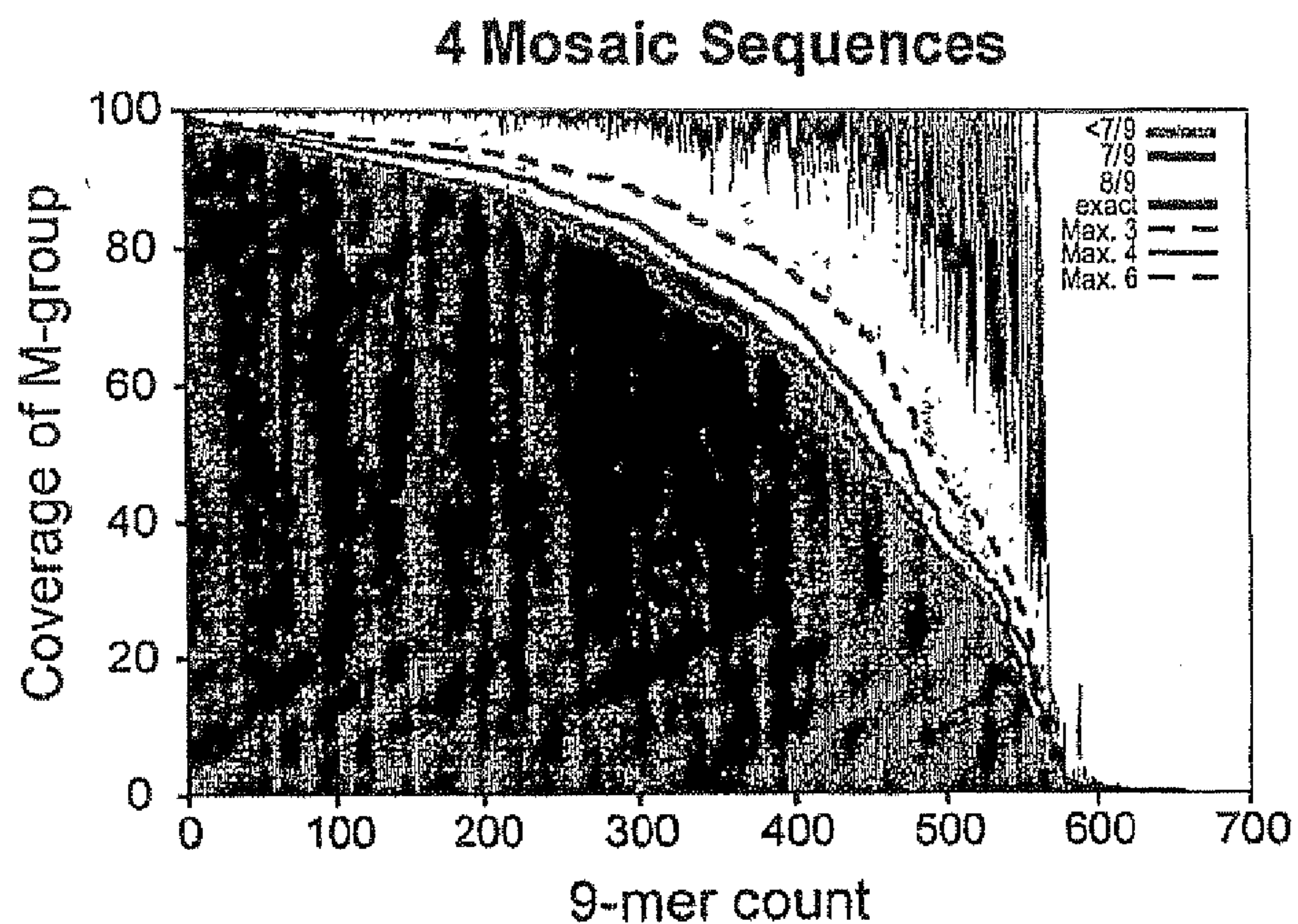


Fig. 4F

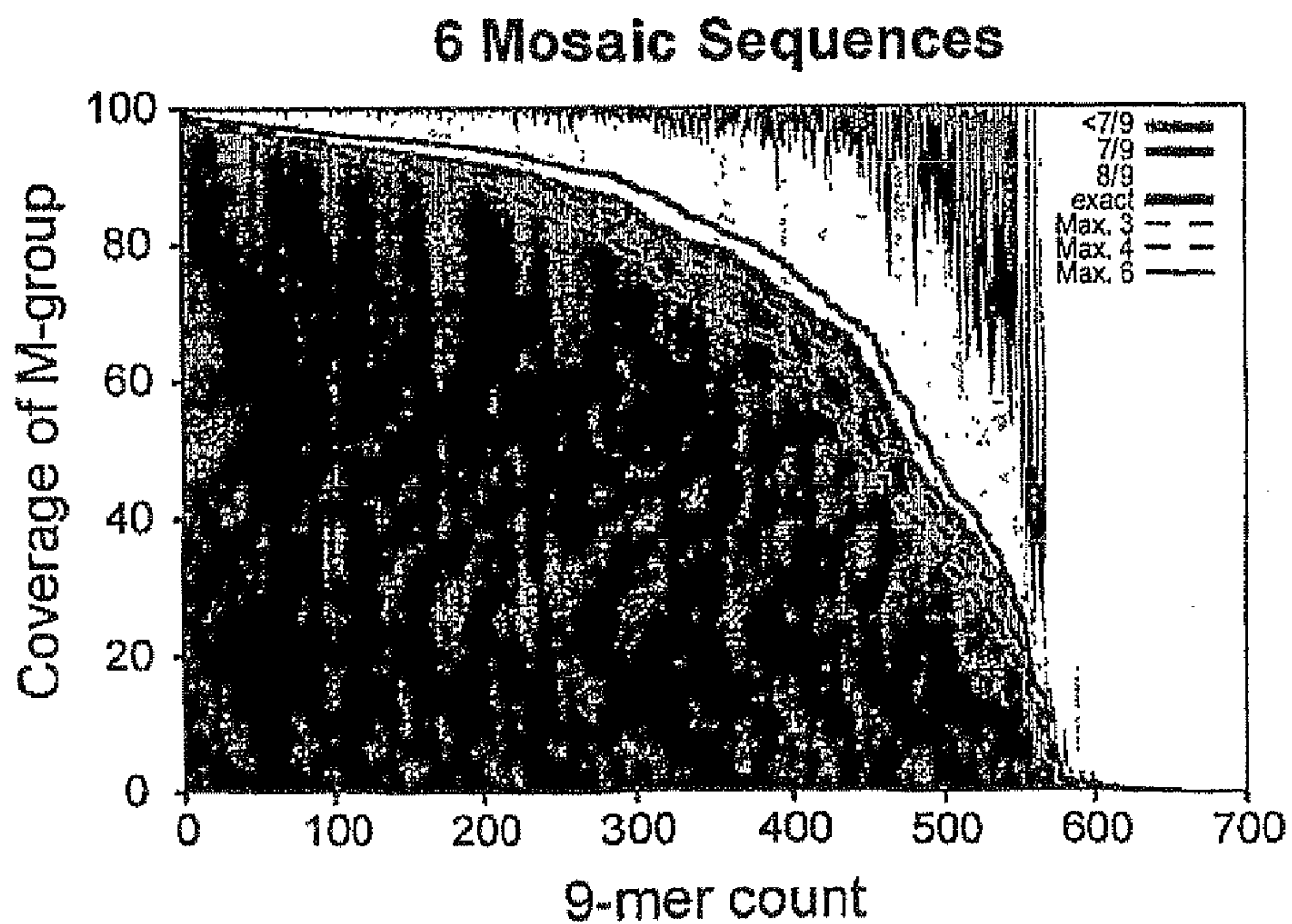




Fig. 5A

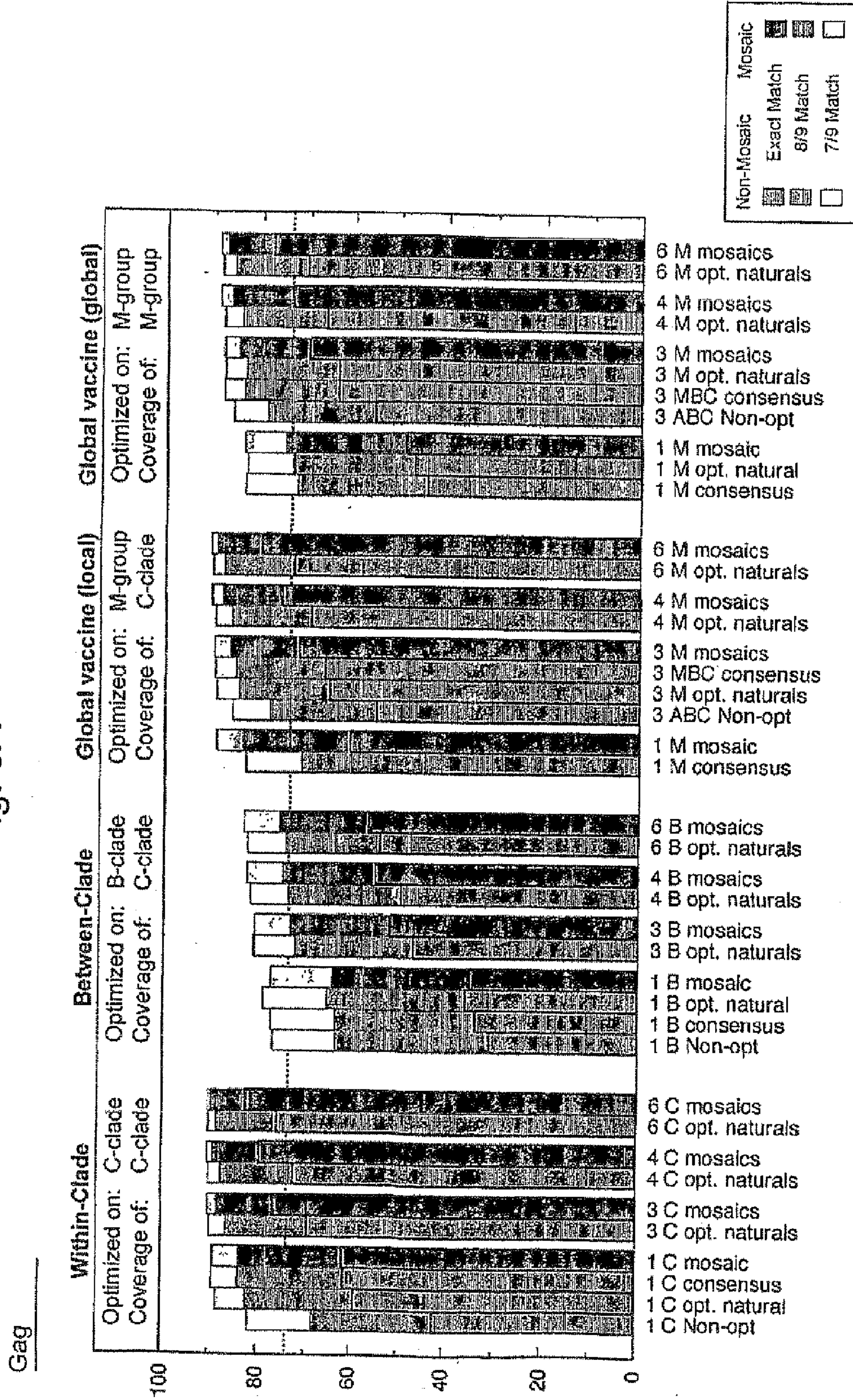




Fig. 5B

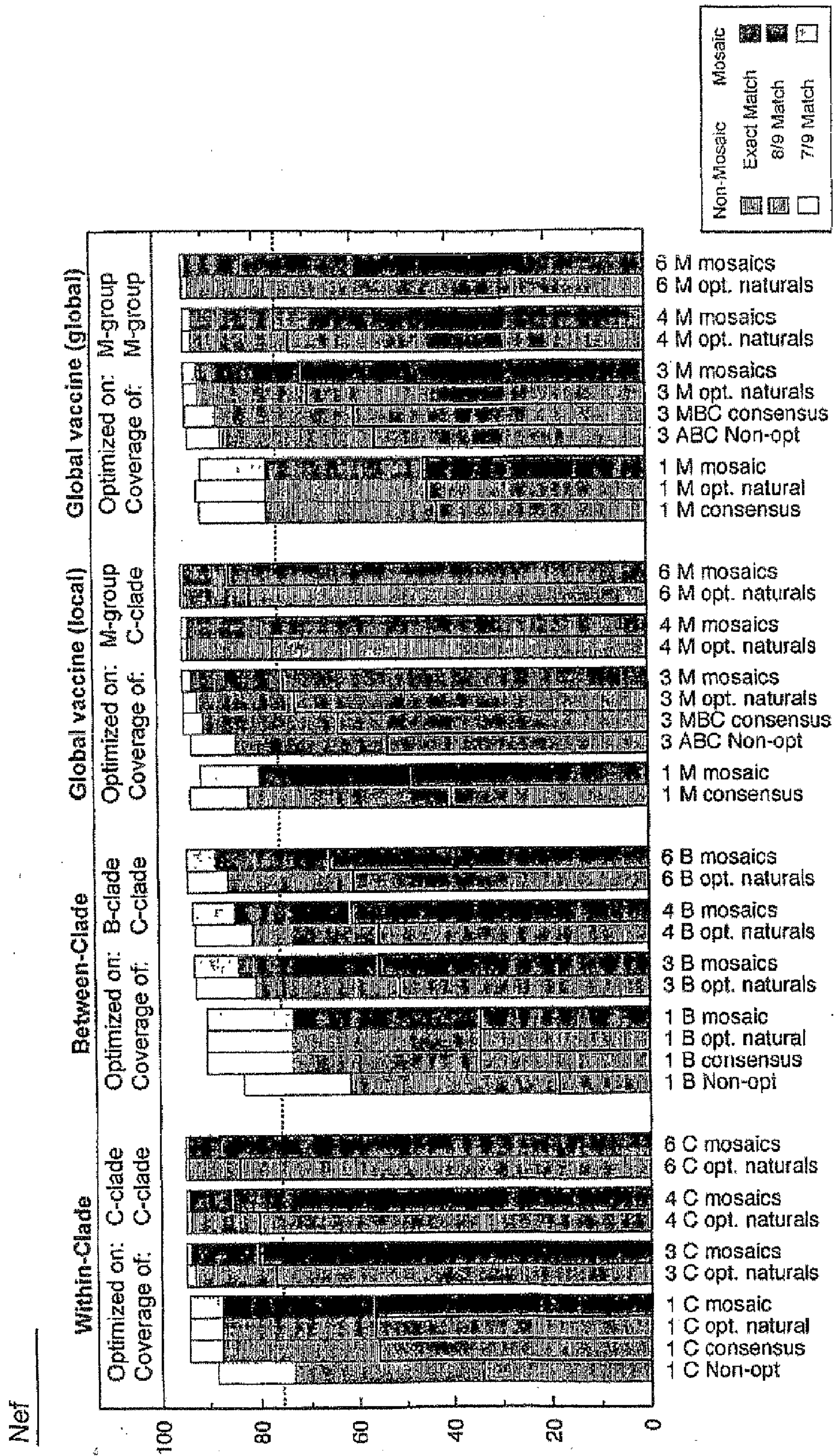




Fig. 6A

Gag coverage

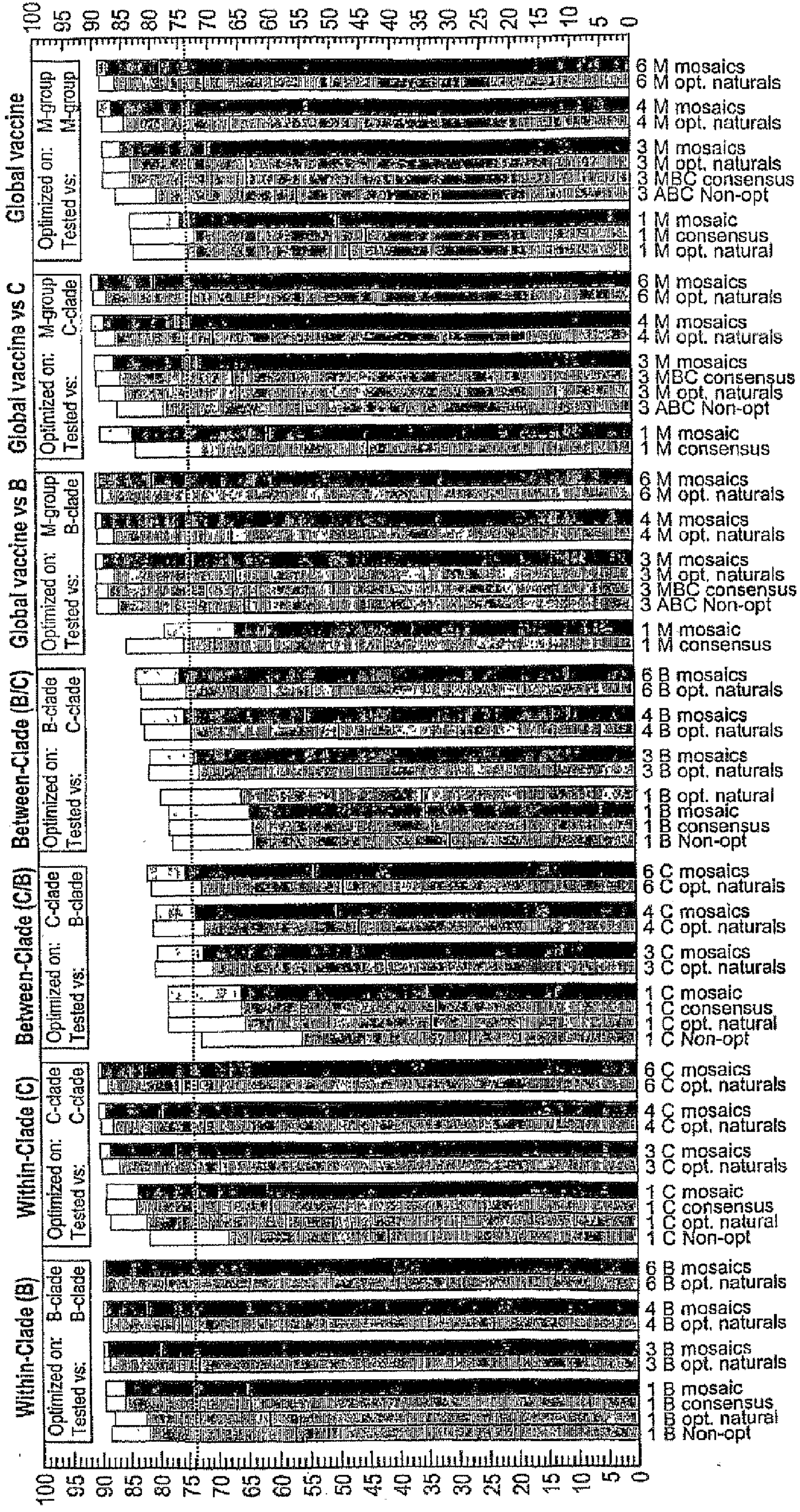




Fig. 6B

Nef coverage (central region)

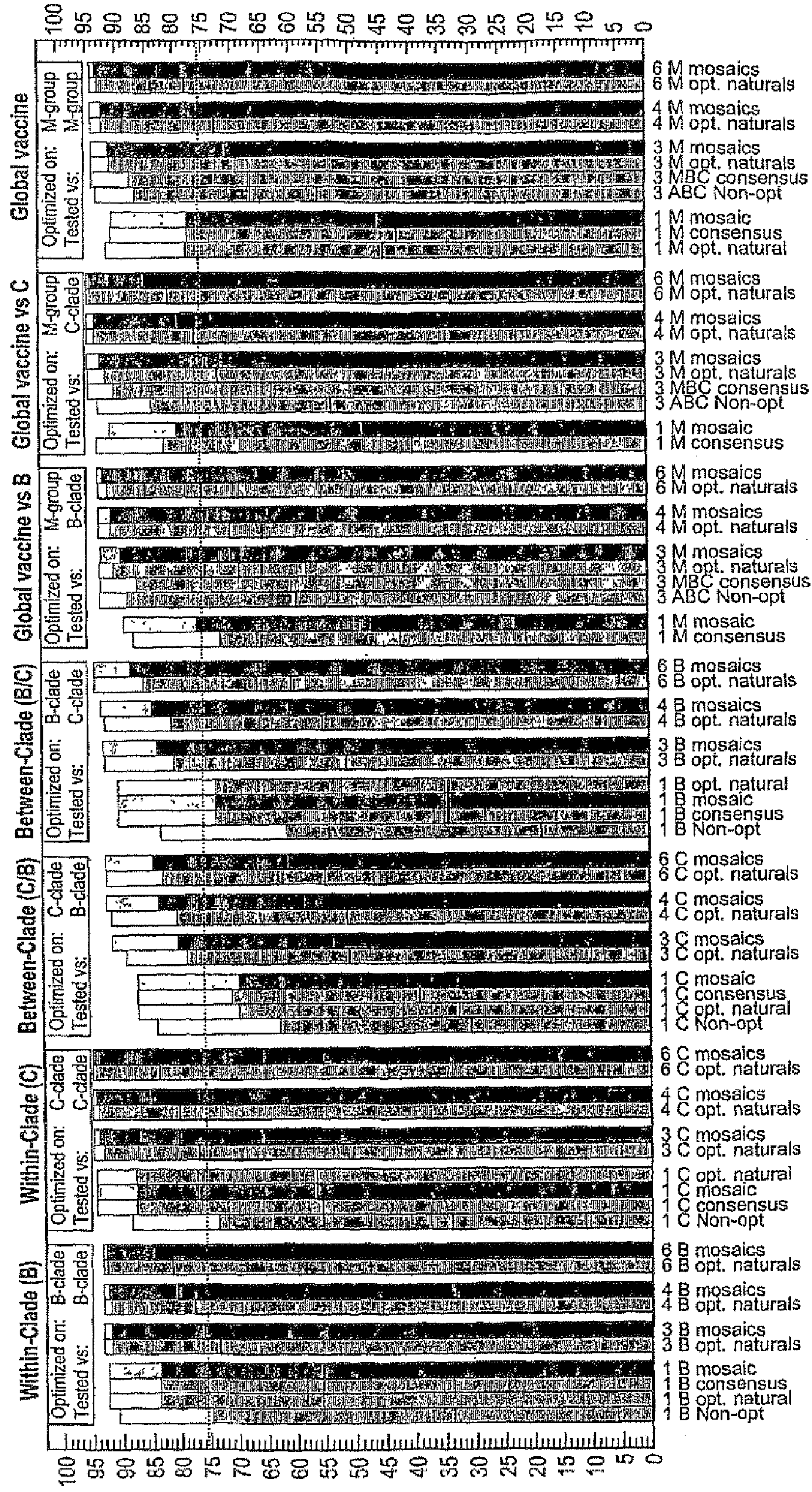




Fig. 7A

9-mer Frequencies (0-60%)

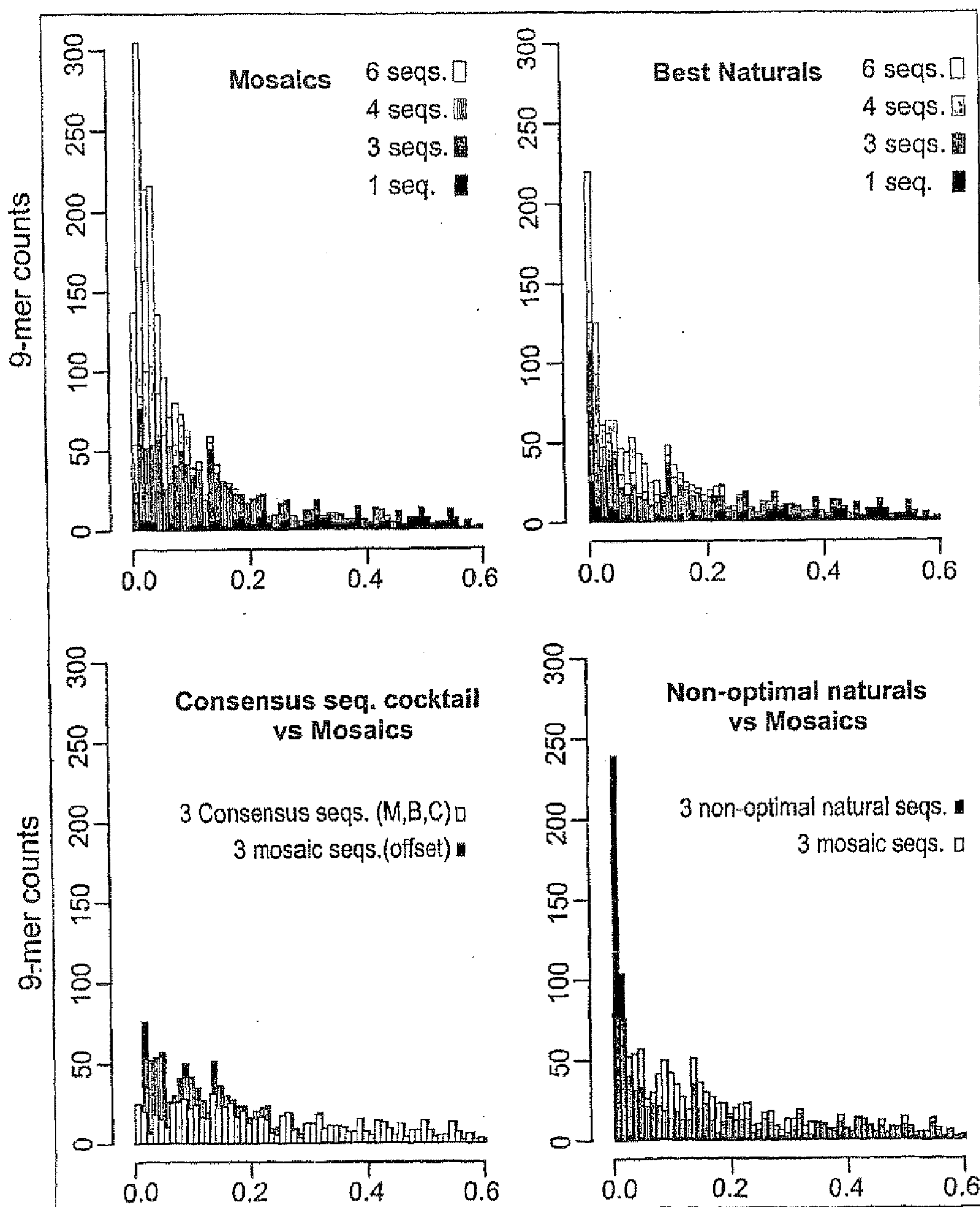
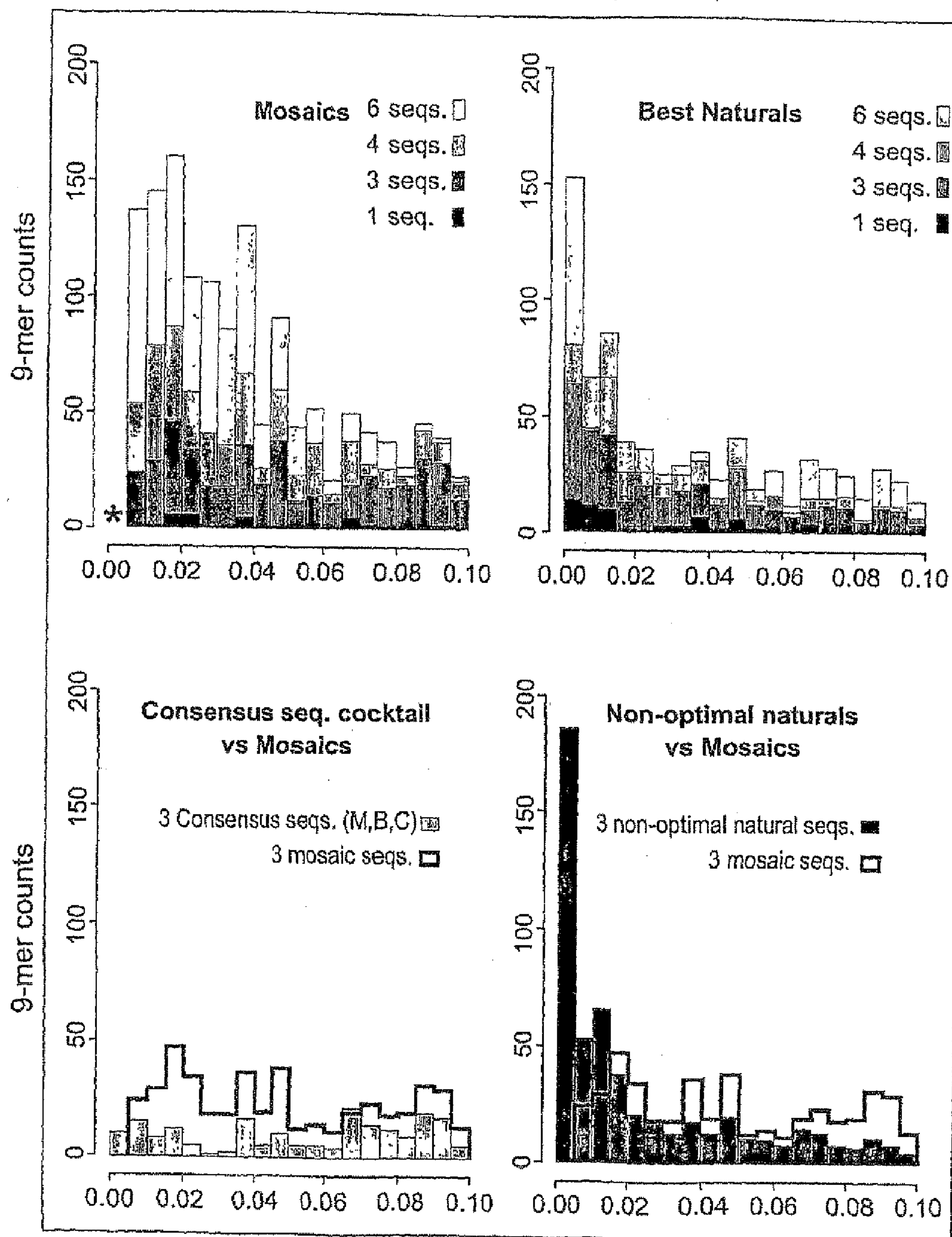
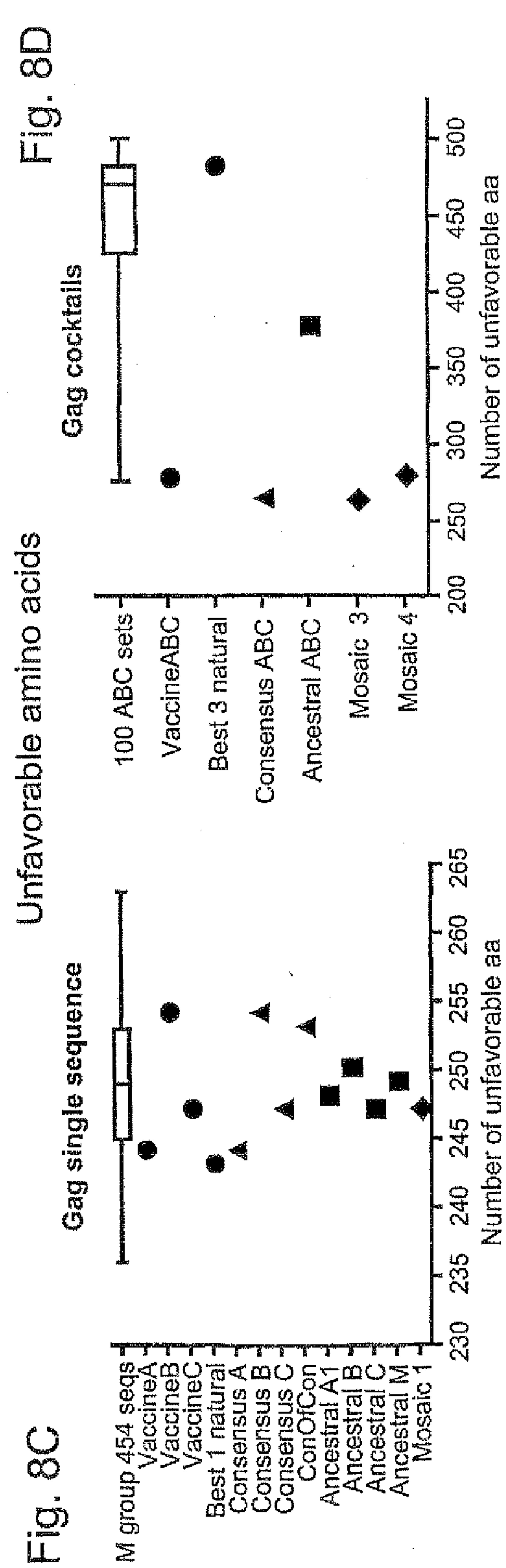
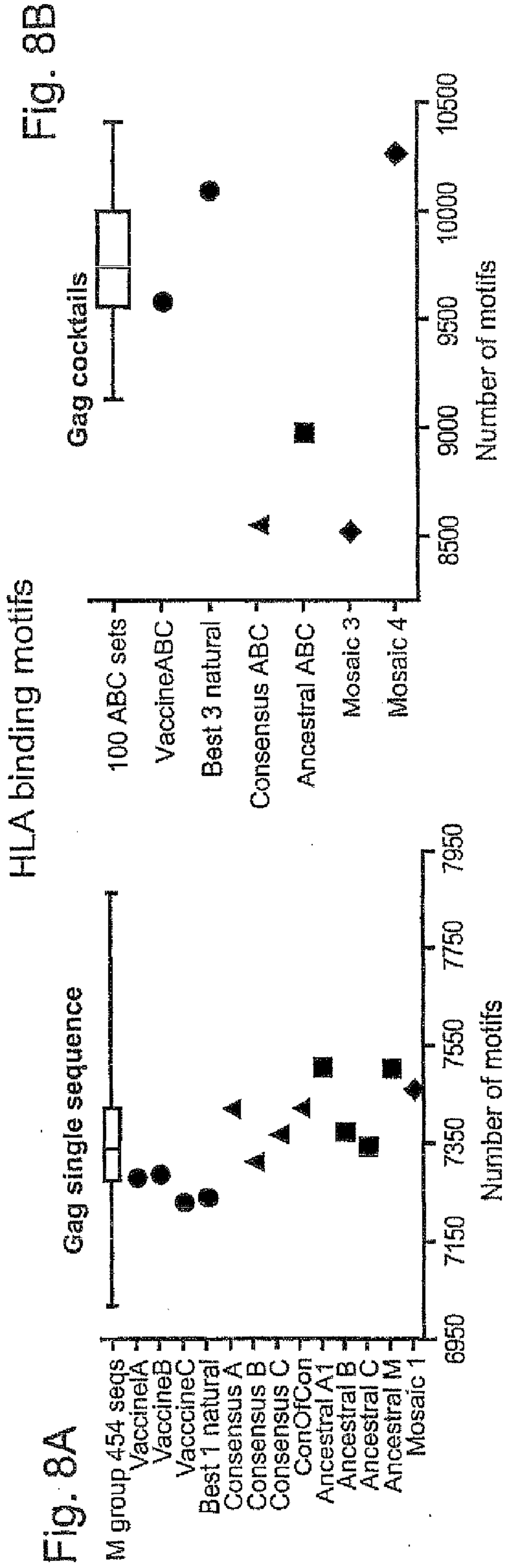




Fig. 7B

9-mer Frequencies (0-10%)





**Fig. 8A**

**Fig. 8C**

HLA binding motifs

Unfavorable amino acids

**Fig. 8B**

**Fig. 8D**



Fig. 9

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Fig. 9 cont'd-1

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Fig. 9 cont'd-2

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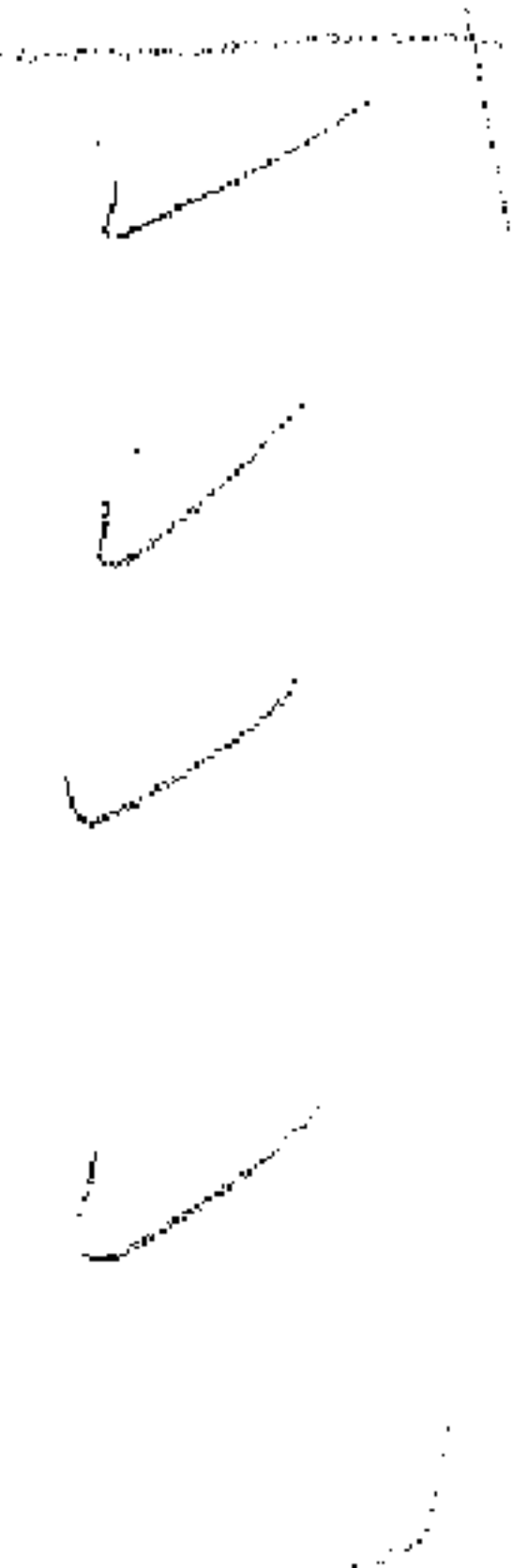


Fig. 9 cont'd-3

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Fig. 9 cont'd-4

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GTGNSSQVSNYPPIVQNMQGMVHQALS PRTLNAWVKVIEEKAFSPEVIMFSALSEGAT  
PQDLNTMLNTVGGHQAAMQMLKDTINEEAAEWDRVHPVHAGPIAPGQMRDPRGSDIAGTT  
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Fig. 9 cont'd-5

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Fig. 9 cont'd-6

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Fig. 9 cont'd-7

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STLQEQITWMTSNPPVPVGEIYKRWIILGLNKIVRMYSVPSILDIRQGPKEPFRDYVDRF  
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ADK--GKVSQNYPIVQNAQGGQMVHQALS PRTLNAWVKVVEEKAFSPEIIMFTALSEGAA  
PQDLNMTLNTVGGHQAAMQMLKDTINEEAAEWDRLHPVHAGPIAPGQIREPRGSDIAGTT  
STLQEQVAVMTSNPPVPVVDIYKRWIIVLGLNKIVRMYSVPSILDIKQGPKEPFRDYVDRF  
FKTLRAEQSSQEVKNWMTDTLLVQANANPDCKTILRALGPAASLEEMMTACQGVGGPSHKA  
RVLAEAMSQ-ANSTNIMMQRGNFKGPKRIVKCFNCGREGHIARNCRAPRKKGCWKCGQEG  
HQMKDC-IERQANFLGKIWPSHKG-RPGNFIQSRPE-----PTAPP-----EPTAP  
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Fig. 9 cont'd-8

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Fig. 9 cont'd-9

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RVLAEAMSQ-TNSA-ILMORSNFKGSKRIVKCFNCGKEGHIARNCRAPRKRKGCWKCQEG
HQMKDC-TERQVNFGLKIWPSNKG-RPGNFLQNRPE-----PTAPPA-----EPTAP
PAESFRFEE--TTPAPKQEPKDRE--PLTSLRSLFGNDPSSQ

>gagM.syn4.1

MGARASVLSGGELDRWEKIRLRPGGKKKYKLVWASRELERFAVNPGLLETSEGCRQI
LGQLQPSLQGTGSEELRSLYNTVAVLYCVHQRI DVKDTKEALEKIEEEQNKSOQKTQOAKA
ADG---KVSQNYPIVQNAOQGMVHQALSPRTLNAWVKVIEEKGFSPPEVI PMF'SALAEAT
PQDLNMTMLNTIGGHQAAMQMLKDTINDEAAEWDRLHPVQAGPVAPGQIREPRGSDIAGTT
SNLQEQIGWMTSNPPIPVGDIYKRWIIMGLNKIVRMYSPTSILDIRQGPKEPFRDYVDRF
FRTLRAEQASQEVKNWMTETLLVQNSNPDKTILKALGPAATLEEMMTACQGVGGPGHKA
RVLAEAMSQ-VQPNIMMQRGNFKGQKR- IKCFNCGREGHLARNCRAPRKKGCWKCQREG
HQMKDC-TESKANFLGKIWPSNKG-RPGNFLQSRP-----EPSAP
PAESFGFGEE-ITPSQKQEQKDKELYPLASLKSFLGNDPLSQ

>gagM.syn4.2

MGARASVLSGGKLDWEKIRLRPGGKKKYRLKHLVWASRELDRELFALNPGLLETAEAGCKQI
MKQLQPALKTGTTELKSLYNTVATLYCVHEKIDVRDTKEALDKIEEEQNKIQOKTQOAKE
ADG---KVSQNYPIVQNIQOQGMVHQALSPRTLNAWVKVVEEKAFSPPEVI PMFTALSDGAT
PQDLNSMLNAVGGHQAAMQILKDTINEEAAEWDRLHPVHAGPVAPGQMREPRGSDIAGTT
STLQEQIGWMTNPPPIPVGEIYKRWIILGLNKIVRMYSVPSILDIKQGPKEPFRDYVDRF
FKVLRAEQATQDVKNWMTDTLLIQNANPDCKSILRALGPGATLEEMMTACQGVGGPSHKA
RILAEAMSQVTNSATIMMQRGNFRNQRKTVKCFNCGKEGHLARNCKAPRKRKGCWKCQEG
HQMKEC-TERQANFLGKIWPSNKG-RPGNFPQSRP-----EPTAP
PEESFRFGEETTTSPQKQEPIDKELYPLASLRSFLGNDPSSQ

>gagM.syn4.3

MGARASILRGGKLDKWEKIRLRPGGKKRYMLKHLI WASRELERFALNPGLLETAEAGCQQI
IEQLQSTLKTGTGSEELKSLFNTVATLYCVHQRIEVKDTKEALDKVEEEQNKSKKKAQOAAA
DTGNSSQVSNYPIVQNLQOQGMVHQALS PRTLNAWVKVIEEKAFSPPEIIPMFTALSEGAT
PSDLNMTMLNTVGGHQAAMQMLKDTINEEAAEWDRVHPVHAGPIPPGQMREPRGSDIAGTT
SSLQEQIAWMTSNPPVPVGEIYKRWIVLGLNKIVRMYSVPSILDIRQGPKEPFRDYVDRF
FKTLRAEQASQDVKNWMTETLLVQANANPDCKTILRALGPGASLEEMMTACQGVGGPSHKA
RVLAEAMSQ-TNSA-ILMORSNFKGSKRIVKCFNCGKEGHIARNCRAPRKRKGCWKCQEG
HQMKDC-NERQANFLGKIWPSHKG-RPGNFLQNRPE-----PTAPP-----EPTAP
PAESFRFEE--TTPAPKQELKDRE--PLTSLKSLFGSDPLSQ

>gagM.syn4.4

MGARASVLRGEKLDKWERIRLRPGGKKHYMLKHLVWASRELEKRFALNPGLLETSEGCKQI
IKQLQPALQGTGTEELRSLFNTVATLYCVHAGIEVRDTKEALDKIEEI QNKSKQKTQOAAA
GTGSSSKVSQNYPIVQNLQOQGMVHQPLSPRTLNAWVKVVEEKGFNPEVI PMFSALSEGAT
PQDLNMMMLNIVGGHQAAMQMLKETINEEAAEWDRLHPVHAGPIAPGQMREPRGSDIAGST
STLQEQIAWMTGNPPVPVVDIYKRWIILGLNKIVKMSPTSILDIKQGPKEPFRDYVDRF
YKTLRAEQATQEVKNWMTDTLLVQANANPDCKSILKALGTGATLEEMMSACQGVGGPAHKA
RVLAEAMSQ-ANNTNIMMQRSNFKGPKRI IKCFNCGKEGHIAKNCRAPRKKGCWKCQEG
HQMKDC-TERQANFLGRIWPSNKG-RPGNFLQSRPE-----PTAPPA-----EPTAP
PAESFKFEE--TTPAPKQEPKDRE--PLTSLRSLFGSDPLLQ

>gagM.syn6.1

MGARASILSGGKLDWEKIRLRPGGRKHYMLKHLI WASRELERFALNPGLLETAEAGCQQI
IEQLQSTLKTGTGSEELKSLFNTVATLWCVHQRIEVKDTKEALDKLEEEQNKSOQKTQOAKA
ADG---KVSQNYPIVQNLQOQGMVHQSI SPRTLNAWVKVIEEKAFSPPEVI PMFSALAEAT
PQDLNMTMLNTIGGHQAAMQILKDTINEEAAEWDRIHPVHAGPVAPGQMRDPRGSDIAGTT
SNLQEQIAWMTSNPPVPVGEIYKRWIILGLDKIVRMYSVPSILDIRQGPKEPFRDYVDRF
FKTLRAEQATQEVKGWMTDTLLVQANANPDCKTILKALGPGATLEEMMSACQGVGGPGHKA



Fig. 9 cont'd-10

RVLAEAMSQ-ANNTNIMMOKSNFKGPKRI IKCFNCGKEGHLARNCRAPRKKGCWKCGQEG  
HOMKDC-TERQANFLGRIWPSHKG-RPGNFPQSRL-----EPTAP  
PAESFGFGEE-IAPSPKQEPKEKELYPLTSLKSLFGNDPLSQ

>gagM. syn6.2

MGARASILRGGKLDKWEKIRLRPGGKKKYKXKXIVWASRELEKFAFNPGLEETSEGCRQI  
LGQLQPSLQGTGSEELKSLYNTVATLYCVHQRIDVKDTKEALEKIEEEQNKSQOKTQOAAA  
DKG----VSQNYPIVQNLQGMVHQALSPRTLNAWVKVIEEKAFSPEIIPMFTALSEGAT  
PQDLTTLMLNTVGGHQAAMQMLKETINDEAAEWDRLHPVHAGPVAPGQLREPRGSDIAGST  
STLQEQIAWMTGNPPVPVGDYKRWIVLGLNKIVRMYSPTSILDIRQGPKEPFRDYVDRF  
YKTLRAEQASQDVKNWMTETLLVQANANPDCRTILKALGPAATLEEMMTACQGVGGPAHKA  
RVLAEAMSQVTPATIMMQRGNFRNQRKTVKCFNCGKEGHLAKNCRAPRKRGCWKCGKEG  
HOMKDC-NERQANFLGKIWPSNKG-RPGNFLQNR-----EPTAP  
PAESFRFGEEKTPSQKQEPIDKELYPLASLRSLEFGNDPLSQ

>gagM. syn6.3

MGARASVLRGEKLDKWERIRLRPGGKRYMLKHLI WASRELERFALNPSLLETSEGCKQI  
IQQLQPALKTGTEELRSLYNTVATLYCVHEKIEVRDTKEAVDKIEEEQNKSKKKAQOAAA  
DTGNSSQVSNYPIVQNIQGMVHQALSPRTLNAWVKVVEEKGFNPEVIPMFSALSEGAT  
PQDLNMLNIVGGHQAAMQMLKDTINEEAAEWDRLHPVQAGPVAPGOMREPRGSDIAGTT  
STLQEQITWMTSNPPIPVGEIYKRWIIMGLNKIVRMYSVPSILDIKQGPKEPFRDYVDRF  
FRTLRAEQASQEVKNWMTETLLIQANANPDCKTILRALGPAASLEEMMTACQGVGGPGHKA  
RVLAEAMSQ-TNSA-ILMORSNFKGSKRIVKCFNCGKEGHIARNCRAPRKKGCWRCGKEG  
HOMKDC-TESKANFLGKIWPSHKG-RPGNFLQNRPEPTAPPEPTAPPAEPTAPPAEPTAP  
PAESFKFEE--TTPAPKQELKDRE--PLISLKSLFGSDPLLQ

>gagM. syn6.4

MGARASILRGEKLDWEKIRLRPGGKKQYRLKXIVWASRELDRELFALNPSLLETAEGCKQI  
IKQLHPALQGTGTEELRSLFNTVATLYCVHAGIEVRDTKEALDKIEEEQNKIQOKTQOAKA  
ADE---KVSQNYPIVQNMQGMVHQPLSPRTLNAWVKVVEEKAFSPEVIPMFAALSEGAT  
PSDLNMLNIVGGHQAAMQMLKDTINDEAAEWDRLHPAQAGPIPPGQIREPRGSDIAGTT  
STPQEQIGWMTNPPPIPVGEIYKRWIVLGLNKIVRMYSPTSILDIRQGPKEPFRDYVDRF  
FKALRAEQATQEVKGWMTETLLVQNSNPDCKTILRALGPGASLEEMMTACQGVGGPSHKA  
RILAEAMSQ-ANS-NIMMORSNFKGPKRIVKCFNCGKEGHIARNCRAPRKKGCWKCGREG  
HOMKDC-IERQANFLGKIWPSQKG-RPGNFLQSRP-----EPSAP  
PAESFRFGE--TTPAPKQEPKDRE--PLTSLRSLEFGSDPLSQ

>gagM. syn6.5

MGARASVLSGGELDRWEKIRLRPGGKKYRLKHLVWASRELERFAINPGLLETSDGCKQI  
IKQLQPALQGTGSEELRSLYNTIATLYCVHQKIEVKDTKEALDKIEEIQNKSKOKTQOAAA  
GTGSSSKVSQNYPIVQNAQGMVHQSLSPRTLNAWVKVIEEKGFNPEVIPMFTALSEGAT  
PHDLNMLNIVGGHQAAMQMLKETINEEAAEWDRVHPVHAGPIPPGOMREPRGSDIAGST  
STLQEQIGWMTSNPPIPVGDYKRWIILGLNKIVRMYSPTSILDIRQGPKEPFRDYVDRF  
FKCLRAEQATQEVKNWMTDTLLIQANANPDCKSILRALGPGATLEEMMTACQGVGGPGHKA  
RILAEAMSQ-VQPNIMMQRGNFKGQKR-IKCFNCGREGHIARNCKAPRKKGCWKCGKEG  
HOMKDC-TERQVNFLGKIWPSYKG-RPGNFLQSRP-----EPTAP  
PEESFRFGEETTPSQKQETIDKELYPLASLKSLFGNDPSSQ

>gagM. syn6.6

MGARASVLSGGKLDWEKIRLRPGGKKHYMLKHLVWASRELERFAVNPGLLETSEGCKQI  
MKQLQPALQGTGTEELKSLYNTVAVLYCVHQRIETKDTKEALDKIEEIQNKCOOKTQOAKE  
ADG---KVSQNYPIVQNLQGMVHQPI SPRTLNAWVKVIEEKGFSPPEVIPMFTALSDGAT  
PQDLNSMLNAVGGHQAAMQMLKDTINEEAAEWDRLHPVHAGPIAPGOMREPRGSDIAGTT  
SSLQEQIAWMTNPPVPVGEIYRRWII LGLNKIVKMYSPTSILDIKQGPKEPFRDYVDRF  
FKVLRAEQATQDVKNWMTDTLLVQANANPDCKSILKALGTGATLEEMMTACQGVGGPSHKA  
RVLAEAMSQVTPNSATIMMOKGNFRNQRKIVKCFNCGREGHLARNCKAPRKRGCWKCGKEG  
HOMKEC-TERQANFLGKIWPSKSG-RPGNFPQSRP-----EPTAP  
PAESFRFEE--TTPAPKQESKDRE--PLTSLKSLFGSDPSSQ



Fig. 10

>ENV-B.syn1.1  
 MRVTGIRKKNYQHLWRWGTMLLWRWGTMLLGLMLICSATEKLWVTVYYGVPVWKEATTTLF  
 CASDAKAYDTEVHNVWATHACVPTDPNPQEVVLENTENFNMWKNMVEQMHEDIISLWD  
 QSLKPCVKLTPLCVTLNCTDDVRNVT--NNATNTNSSW--GEPMEKGEIKNCSFNITTSIRD  
 KVQKEYALFYKLDVVP--DNDSNNTN-----YRLISCNTSVITQACPKVSFEPIPIH  
 YCAPAGFAILKCNCKKFNGTGPTNVSTVQCTHGIRPVVSTQLLLNGLSLAEEVVIRSEN  
 FTNNAKTIMVQLNVSVEINCTRPNNNTRKSIHIGPGRAFYTTGDIIGDIRQAHCNISRAQ  
 WNNTLKHIVEKLGKQFGNNTIIVFNHSSGGDPEIVMHSFNCGGEFFYCNSSTKLFNSTWTR  
 N-NGTWTRN---DTERSINSTE---EHITLPCRICKQIINMWQEVGKAMYAPPIRGQIRCSS  
 NITGLLLTRDGGNDT-----SGTEIFRPGGGDMRDNRSELYKYKVVKIEPLGVAPTAK  
 RRVVQREKRAVG--IGAVFLGFLGAAGSTMGAASMTLTVQARQLLSGIVQQQNNLLRAIEA  
 QQHLLQLTVWGIKQLQARVLAVERYLRDQQLLGIWGCSSGKLICTTAVPWNASWSNKSLE  
 IWNMTWMEWEKEIDNYTNLIYNLLEESQNOQEKNEQELLELDKWASLWNWFDISNWLWY  
 IKIFIMIVGGLVGLRIVEFAVLSIVNRVRQGYSPFSFQTRLPAPRGPDRPEGIEEEGGGERD  
 RDRSVRLVDGFLALIWDLLRSLCLFSYHRLRDLILLIVTRIVELLG-----RRGWEALK  
 YWWNLLQYWSQELKNSAVSLLNATAIAVAEGTDRVIEALQRACRAILHIPRRIROGLERA  
 LL-

>ENV-B.syn3.1  
 MRVKETRKNYQHLWKWGTML-----LGMLMICSATEKLWVTVYYGVPVWRDANATLF  
 CASDAKAYDTEAHNVWATHACVPTDPNPQEVVELKNVTENFNMWKNMVEQMHEDIINLWD  
 QSLKPCVELTPLCVTLNCTDYVKNIT--NNATSTNSSW--GKPMKGEIKFCSFNITTSIRN  
 KVQKQYALFYKLDIVPI--DNDNTS-----YRLISCNTSTITQACPKVTFEPIPIH  
 YCAPAGFAILKCNKTFNGTGPTNVSTVQCTHGIRPVIISTQLLLNGLSLAEEVVIRSEN  
 FTNNAKTIMVQLNVSVEINCTRPNNNTRRSIPIGPRVFTTEDIIGDIRQAHCNISRAQ  
 WNNTLKHIVEKLGKQFGNNTIIVFNHSSGGDPEIVMHSFNCRGGEFFYCKSTKLFNSTWTR  
 N-NGTWTRN---DTERSINSTE---EHITLPCRICKQIINMWQEVGKAMYAPPIKGOISCS  
 NITGLLLTRDGGNDT-----SGTEIFRPGGGDMRDNRSELYKYKVVRIEPLGVAPTAK  
 RRVVQREKRAVG--IGAVFLGFLGAAGSTMGAASMTLTVQARLLLSGIVQQQSNLLRAIEA  
 QQHMLQLTVWGIKQLQARLLAVERYLRDQQLLGLWGCSSGKLICTTTPWNTSWSNKSLE  
 IWDNMTWQWEREIDNYTGLIYNLLEKSQNOQEKNEQELLELDKWASLWNWFDITNWLWY  
 IKIFIMIVGGLVGLRIVFTVLSIVNRVRKGYSPFSFQTRLPTRGPDRPGGIEEEGGEQD  
 RDRSGPLVNGFLALIWDLLRSLFLFSYHRLRDLILLIVARIVELLG-----RRGWEILK  
 YWWNLLLYWSQELKNSAVSLLNATAIAVAEGTDRVIEVVQRAFRAILHIPRRIROGFERA  
 LL-

>ENV-B.syn3.2  
 MRVTGIRKKNYQHLWRWGTMLLWRWGTMLLGLMICSAAAGKLWVTVYYGVPVWKEANTTLF  
 CASDAKAYDTEVHNVWATHACVPIIDPNPQEVVLENTENFNMWKNMVEQMHEDIISLWD  
 ESLKPCVKLTPICVTLNCTDDVRNVT--NNATNTNSSW--GEPMEKGEIKNCSFNITTSIRD  
 KVQKQYALFYKLDVVP--DNDSNNTN-----YRLISCNTSVITQACPKISFEPIPIH  
 FCAPAGFAILKCNCKKFNGTGPTNVSTVQCTHGIRPVVSTQLLLNGLSLAEEIVIRSEN  
 FTDNAKTIVQLNESVVINCTRPNNNTRKSIHIGPGRAFYATGETIIGDIRQAHCNLSRAK  
 WNDTLKQIVIKLREQFG--NKTIVFNQSSGGDPEIVMHSFNCGGEFFYCNTTQLFNSTWTW  
 N-NSTW--N---NTRKSNSTE---EITLPCRICKQIINMWQKVGKAMYAPPIRGQIRCSS  
 NITGILLTRDGGNNNET---NRTETFRPGGGNMKDNWRSELYKYKVVKIEPLGIAPTAK  
 RRVVQREKRAVGTIGAMFLGFLGAAGSTMGAASLTLTVQARQLLPGIVQQQNNLLKAIEA  
 QQHLLRLTVWGIKQLQARVLAVERYLRDQQLLGIWGCSSGKIICTTAVPWNASWSNKSLE  
 IWDNMTWMEWEKEIDNYTNLIYNLLEESQNOQEKNELELELDKWANLWNWFDISNWLWY  
 IKIFIMIIGGLVGLRIVFAVLSIVNRVRQGYSPFSFQTRLPAPRGPDRPEGIEEEGGGERD  
 RDRSVRLVDGFLALIWDLLRSLCLFLYHRLRDLILLIARIVELLG-----RRGWEALK  
 YWWNLLQYWSQELKNSAISLLNATAVAVAEGTDRVIEALQRACRAILHIPRRIROGLERL  
 LL-

>ENV-B.syn3.3  
 MRVKGIRKNCQHLWRWGTML-----LGMLMICSAAEQWVTVYYGVPVWKEATTTLF  
 CASDAKAYDKEVHNVWATHASVPTDPNPQEVVLENTENFNMWKNMVDQMHEDIISLWD  
 QSLKPCVKLTPLCVTLNCTD--LRNATNGNDTNTTSSS--REMMGGGEMKNCSFNVTTSIRD  
 KVQKEYALFYKLDVPI--DSRNNSNSTE---YNSYRLINCNTSVITQACPKVSFEPIPIH



Fig. 10 cont'd-1

YCTPAGFAILKCKDKKFNGTGPCTKVSTVQCTHGIRPVVSTQLLNGSLAE EEEVIIRSEN
FTNNAKTIIVQLKEAVEINCTRPSNNTRKSIPIGPGRAFYT TGDII GDIRKAHCNISRA N
WNN'LRQIVEKLG EQFGNNKTIIFKQSSGGDPEIVTHSFNCGGGEFFYCNSTQLFNSTWNF
--NGTWNKN---FNNTWNNTEGTNDTITLPCRIKQIINRWQEVGKAMYAPPISGQIRCSS
NITGLILTRDGGNNGNET--NGTEIFRPGGGMRDNRSELYRYKVVKIEPLGVAPTKAK
RRVVQREKRAVG-IGALFLGFLGAAGSTMGAASITLTVQARQLLSGIVQQONNLLRAIEA
QQHLLQLT VWGIKQLQARILAVERYLKDQQLLGIWGCSGKLICTTAVFWNTSWSNRSLNE
IWNNTWMEWEREIDNYTSLIYTLIEESQOQEKNEQELLELDK WASLWNWF SITNWLWY
IRIFIMIVGGLIGLRIVFAVLSVVRVROGYSPLSFQTHLPAQRGPDRPEGTEEEGGERD
RDRSGRLVDGFLAIIWVDLRSCLFSYHRLRDL LLIVTRIVELLG-----RRGWEVLK
YWWNLLQYWIQELKNSAVSLFNATAIAVAEGTDRIIEVVQRAYRAILHIPTIRQGLERA
LL-

>ENV-B.syn4.1

MRVTGIRKNYQHLWRWGTMLLWRWGTMLLGIILMICSAAAGKLWVTVYYGVPVWKDATTLEF
CASDAKAYDTEVHNVWATHASVPTDPNPQEVVLENTEDFNMWKNNMVDQM HEDII SLWD
QSLKPCVELTPLCVTLNCTDDVRNVT-NNATNTNSSW-GEPMEKGEIKFCSFNITTSIRN
KVQKQYALFYKLDV VPI-DNDSNNTN-----YRLISCNTSVITQACPKVTFEPIPIH
YCAPAGFAILKCNKTFNGTGPCTKVSTVQCTHGIRPVVSTHLLLNGSLAE EEEVIIRSEN
FTDNTKTIIVQLKEAVEINCTRPNNTRKGIHIGPGRAFYT TGEIIGDIRQAHCNLSRAK
WNDTLKQIVIKLREQFG-NKTIIFNQSSGGDPEIVMHTFNCGGGEFFYCNSTQLFNSTW--
-----QN---ETSGSINITDIGENITLPCRIKQIVNMWQKVGKAMYAPPIKQOISCSS
NITGLLLTRDGGNNGNET--NGTEIFRPGGGMKDNWRSELYRYKVVKIEPLGVAPTRAK
RRVVQREKRAVT-LGAMFLGFLGAAGSTMGAASMTLTVQARLLLSGIVQQOSNLLRAIEA
QQHLLRLTVWGIKQLQARILAVERYLQDQQLLGIWGCSGKLICTTAVPWNASWSNKSQDE
IWNNTWMEWEREIDNYTGLIYTLLEESQIQOQEKNEQELLELDK WASLWNWF'DITNWLWY
IKIFIMIVGGLIGLRIVFTVLSIVNRVRKGYSP LSFQTHLPAPRGPDRPGGIEEEGGEQD
RDRSGPLVNGFLALI IWVDLRSCLFSYHRLRDL LLIVARIVELLG-----RRGWEVLK
YWWNLLQYWSQELKNSAISLLNATAVAVAEGTDRVIEALQACRAILHIPRRIRQGLERL
LL-

>ENV-B.syn4.2

MRVKGIRKNCQHLWRWGILL-----LGMLMICSAAEQWLWVTVYYGVPVWRDANATLEF
CASDAKAYDTEAHNVWATHACVPTDPNPQEVVLEKNTENFNMWKNNMVEQM QEDIISLWD
QSLKPCVKLTPICVTLNCTDYVKNIT-NNATSTNSSW-GKPM EKGEIKNCSFNVTTSIRD
KVQKEYALFYRLDVVPI-DNDSNNDSTNTNYTNYRLISCNTSTITQACPKV SFQPIPIH
YCAPAGFALLKCNKDFNGTGPCKNVSTVQCTHGSIKPVVSTQLLNGSLAE EEEIVIRSEN
FTNNAKTIIVQLNESVVINCTRPNNTRKSIHIGPGRAFYATGDII GDIRKAHCNISRA N
WNN'LRQIVEKLG EQFG-NKTIVFNQSSGGDVEIVMHSFNCGGGEFFYCNSTQLFNSTWNF
--NGTWNKN---FNNTWNNTEGTNDTITLPCRIKQIINMWQGVGKAMYAPPISGQIRCSS
NITGLILTRDGGNN-NET--NRTETFRPGGDMRDNRSELYKYKVVKIEPLGIAPTKAR
RRVVQREKRAVGTIGAMFLGFLGTAGSTMGAASLTLTVQARLLLSGIVQQONNLLKAIEA
QQHLLQLT VWGIKQLQARLLAVERYLGDQQLLGLWGCSGKLICTTVPWNASWSNKS LDK
IWDNMTWMEWEREIDNYTGLIYNLLEKSQOQEKNELELLELDK WANLWNWF'DITKWLWY
IKIFIMIIGGLIGLRIVFAVLSVVRVROGYSPLSLQTRLPTQRGPDRPEGIAEEGGERD
RDRSGPLVDGFLAIIWVDLRSCLFSYHHLRDL LLIVTRIVELLG-----RRGWEALK
YWWNLLLYWSQELKNSAVNLLNTTAIAVAEGTDRIIEVLQRIYRAFLHIPRRIRQGFERA
LL-

>ENV-B.syn4.3

MRVKEIRKNYQHLWKWGTML-----LGMLMICSAAAGNLWVTVYYGVPVWKEANTTLEF
CASDAKAYETE VHNWATHACVPIDPNPQEVV LGNV TENFNMGKNNMVEQM HEDII SLWD
ESLKPCVKLTPLCVTLNCTDELKNATFRSNTT TNSSW--EKMEKGEIKNCSFNITTNMRD
KMQKEYALFYKLDVIFI-DSRNNSSNSTE--YNSYRLINCNTSVITQACPKISFEPIPIH
YCTPAGFAILKCKDKKFNGKGPCTNVSTVQCTHGIRPVVSTQLLNGSLAE KEVVIRSDN
FTNNAKTIMVQLNVSVEINCTRPNNTRRSIPIGPGRVFYTTEDIIGDIRQAHCNISRAQ
WNN'TLKHIVEKLGKQFGNNKTIIVFNHSSGGDPEIVMHSFNCRGGEFFYCKSTKLENSTWTR



Fig. 10 cont'd-2

N-NGTWTRN---DTERSNSSTE---EHITLPCRIKQIINMWQEVGKAMYAPPPIRGQIRCSS  
NITGILLTRDGGNDT-----SGTEIFRPGGDMKDNWRSELYKYKVVRIEPLGVAPTEAK  
RRVVQREKRAVG-IGAVFLGFLGAAGSTMGAAVTLTVQARLLLSGIVQQQNNLLRAIEA  
QORLLQLTVWGIKQLQARVLAVERYLRDQQLLGIWGC SGKI ICTTAVPWNTSWSNRS LNE  
IWNMTWMEWEKEIDNYTNLIYNLLEESQNQQEKNEQELLALDKWANLWNWFDISNWLWY  
IKIFIIVGGLVGLRIVFAVLSIVNRVRQGYSPLSFQTRLPA PRGPDRPEGIEEEGGGERD  
RDRSVRLVDGFLALIWDLLRSLCLFSYHRLRDL LLI-----VELLG-----RRGWEILK  
YWWNLLQYWGQELKNSAVSLLNATAIAVAEGTDRVIEVVQRAYRAI LHIPTRIROGLERA  
LL-

>ENV-B.syn4.4

MRVKETRKNYQHLWRWGIML-----LGMLMICSATEKLWVTVYYGVPVWKEATTTLF  
CASDAKAYDKEVHNWVATHACVPTDPSPQEVVLENTENFNMMWKNMVEQM HEDIINLWD  
QSLKPCVRLTPLCVTLNCTN-VNVTNLKNETNTKSSSGG EKMEEGEMKNCSFNITTSIRD  
KVQKQYALFYKLDVVPI-DNDNTS-----YRLISCNTSVIKQACPKVSFEPIPIE  
FCAPAGFAILKCNDDKFNGTGPCTNVSTVQCTHGIRPVI STQLLLNGLSLAEEVVIRSEN  
FTDNAKTIIVQLNETVEINCTRPSNNTRKSIPIGPGR AFYTTGDIIGDIRQAYCNISRAK  
WNNTLKQIVTKLREQFGNNKTIIFKQSSGGDPEIVTHSFNCGGEFFYCNSTKLFNSTWTW  
N-NSTW--N---NTRKRSNDTE---EIITLPCRIKQIINRWQEVGKAMYAPPIEGQIRCLS  
NITGLLLLTRDGGTNNT---NTNETFRPGGGMNRDNWRSELYKYKVVQIEPLGVAPTKAK  
RRVVQREKRAVG-IGALFLGFLGAAGSTMGAASITLTVQARQLLSGIVQQQNNLLRAIEA  
QQHMLQLTVWGIKQLRARVLAVERYLKDQQLLGIWGC SGRLICTTNVPWNTSWSNKS LNE  
IWDNMTWMEWEREIDNYTSLIYTLIEESQNQQEKNEQD LLDKWA SLWNWFSITNWLWY  
IRIFIMIVGGLVGLRIVFTVISIVTRVRQGYSPLSFQTRLPTPRGPDRPEGTEEEGGGERD  
RDRSGRLVDGFLALFWDDLRSCLFLYHRLRDL LLI AARIVEL LG-----RRGWELLK  
YWWNLLQYWIQELKNSAVSLENAIAIAVAEGTDWVIEISQRAFR AVLHIPPVRIROGLERA  
LQ-

>ENV-B.syn6.1

MRVTGIRKNYQHLWRWGTM LLLWRWGTM L LGILMICS AAGKLWVTVYYGVPVWKDATTTLF  
CASDAKAYDTEAHNVWATHACVPI DPNPQEVVLENTENFN AWKNNMVEQM HEDMISLWD  
QSLQPCVRLTPLCVTLNCTDDVRN-----ATSTNSSW-GKPM EKGEIKNCSFNITTSIRD  
KVQKQYALFYKLDVVPI-DNDSNNTN-----YRLISCNTS IITQACPKITFEPIPIH  
YCTPAGFALLKCNDDKFNGTGPCTKVSTVQCTHGIRPVVSTHLLLNGLSLAEEVVIIRSEN  
FTNNAKTIMVQLNVSVEINCTRPSNNTRKSIHIGPGR AFYTTGDIIGDIRKAHCNISRAN  
WNNTLRQIVEKLGEQFGNNKTIIVFNHSSGGDLEIVTHSFICGGEFFYCNSTKLFNSTWTW  
N-NSTW--N---NTRKRSNDTE---EIITLPCRIKQIINMWQEVGKAMYAPPPIRGKIRCSS  
NITGLLLLTRDGGTNNT---NTNETFRPGGDMRDNWRNELYKYKVVRIEPLGIAPTEAK  
RRVVQREKRAVG-IGAMFLGFLGTAGSTMGAASVALTVQARQLLP GIVQQQNNLLRAIDA  
QQHLLQLTVWGIKQLQARILAVERYLKDQQLLGFWGC SGKLICTTNVPWNTSWSNKSYSQ  
IWENMTWMEWEREINNYTGLIYNLLEKSQNQQEKNEQEL LELDKWASLWSWFDISNWLWY  
IKIFIIVGGLVGLRIVFAVLSIINRVRQGYSPLSFQTHLPAPRGPDRPEGIAEEGGGERD  
RDRSGRLVNGFLALIWDVLRSLCLFSYHHLRDL LLI-----VELLG-----RRGWEVLK  
YWWNLLLYWSQELKNSAISLLNATAVAVAEGTDRVIEALQ RACRAILHI PRRIROGLERL  
LL-

>ENV-B.syn6.2

MRVKETRKNYQHLWKWGTM L-----LGILMICSATENLWVTVYYGVPVWKEATTTLF  
CASDAKAYDKEVHNWVATHACVPTDPNPQEVVELKNVTENFNMMWKNMVEQM QEDIISLWD  
QSLKPCVRLTPLCVTLNCTD-LRNATNGNDTNTTSSS-REMMGGGEMKNCSFNITTNIRD  
KVQKEYALFYKLDIVPI-DNDNTN-----YRLISCNTSVVTQACPKVSFEPIPIH  
FCAPAGFAILKCNDDKFNKG PCTNVSTVQCTHGIRPVI STQLLLNGLSLAEEVVIRSEN



Fig. 10 cont'd-3

FTNNVKTIIIVQLNETVEINCTRFNNNTRRSIPIGPGRVFTTEDIIGDIRQAHCNLSRTQ
WNNLTKQIVTKLREQFG-NKTIIIFNQSSGGDPEIVMHTFNCGGEFFYCNTTKLFNSTW--
--NDTTINR----TEGSNNTR----NITLPCRIKQIINLWQEVGKAMYAPPIQGQISCS
NITGLLLLTRDGGNN-NET--NRTETFRPGGNNMRDNWRSELYKYKVVKIEPLGVAPTKAK
RRVVQREKRAVG-IGALFLGFLGAAGSTMGAASVTLTVQARQLLSGIVQQRNLLRAIEA
QQRMLQLTVWGIKQLRARVLAVERYLKDQQLMGIWGCSGKLICTTTPVWNASWSNKSLE
IWDNMTWMEWEREIDNYTSLIYTLIEESQNOQEKNELELLELDKWASLWNWFSITNWLWY
IRLFIMIVGGLVGLRIVFTVISIVTRVRQGYSPLSFQTRLPTPRGPDRPGGIEEGGEQD
RDRSIRLVDGFLALIWDLLRSLCLFSYHRLRDLWI----VELLG-----RRGWEALK
YLWNLQYWSQELKKSASVSLFNATAIAVAEGTDWVIEVIQRAFRAFIHIPTRVROGLERA
LQ-

>ENV-B.syn6.3

MRVKGIRKNCQHLWRWGILL-----LGMLMICSATEKLWVTVYYGVPVWKETTTTLF
CASDAKAYVAEKHNWATHACVPTDPNPREVVMGNVTEEFNIWNNSMVEQMHEDIISLWE
QSLKPCVKLTPLCVSLKCTDL-----KNDTNTNSSSGRMIMEKGEIKNCSFNITTGIRG
KVQ-EYSLFYKLDVVQM-DEDNTS-----YRLINCNTSVITQACPKVSFQPIPIH
YCAPAGFAILKCKDKKFNGTGCKNVSTVQCTHGIRPVISTQLLNGSLAEGEVVIRSEN
FTDNAKTIIIVQLKDFKINCTRFNNNTRKSIPIGPGRAFATGDIIGDIRQAHCNISTTK
WNKTLGQVVKLREQFK-NKTIVFKQSSGGDPEVVMHSFNCGGEFFYCNTSOLFNSTW--
-----N---STSLFNSTN---GTITLQCRIKQIINRWQEVGKAMYAPPIEGQIRCLS
NITGLLLVRDGGINVTNN--TGTEVERPGGGDMRDNRSELYKYKVIKIEPLGVAPTRAK
RRVVQREKRAVG-LGAMFLGFLGAAGSTMGAASITLTVQARQLLSGIVQQRNLLRAIEA
QQHMLQLTVWGIKQLQARVLAVERYLQDQQLLGIWGCSSGKLICTTTPVWNASWSNKSLE
IWDNMTWMEWEREIDNYTGLIYTLIEESQNOQEKNEHELELLELDKWASLWNWFSITNWLWY
IKIFIMIIGGLIGLRIVFAVLSIVNRVRQGYSPISFQTRLPAAPRGPDRPDGIEEGGDRD
RDRSGRLVDGFLALIWDLLRSLCLFSYHRLRDLLLIAARIVELG-----HRGWEALK
YWWNLQYWIQELKNSAVNLNNTAIAVAEGTDRVIEVVQRAYRAILNIPTRIROGFERA
LL-

>ENV-B.syn6.4

MRVKEIRKNCQRLWRWGTM-----LGMLMICSAAEQWVTVYYGVPVWRDANATLE
CASDAKAYDTEVHNWATHASVPTDPNPQEVVLGNVTENFNMMWKNMVEQMHEDEVISLWD
QSLKPCVKLTPICVTLNCTDYVKNIT--NNATSTNSSW--GEPMEKGEIKNCSFNITTSMD
KVQKYALFYKLDVVPI-DNDSNNNDSTNTNYRNLISCNTSVIKQACPKVSFDPIPIH
YCTPAGFAILKCRDKKFNGTGCKNVSTVQCTHGIRPVVPTQLLLNGSLAEEDVIRSEN
FSDNAKTIIIVHLNESVEINCTRLNNNTRKSIHMGPGRAFATGEIIGDIRQAHCNISRAK
WNNLTKQIAIKLREQFGNKTIIIFKQSSGGDPEIVTHSFNCGGEFFYCNTSOLFNSTWNF
--NGTWNKN---FNNTWNTEGTNDTITLPCRIKQIINMWQKVGKAMYAPPIISGQIRCTS
NITGLLLTRDGGN---DT--SGTEIFRPGGNNMKDNWRSELYKYKVVQIEPLGVAPTEAK
RRVVQREKRAVG-IGAVFLGFLGAAGSTMGAAVTLTVQARQLLSGIVQQRNLLKAIEA
QQHLLRLTVWGIKQLQARVLAVERYLGDQQLLGLWGCSSGKLICTTAVPWNTSWSNRSLE
IWNMTWMEWEREIDNYTNLIYNLLEESQNOQEKNEKELLELDKWANLWNWFDISNWLWY
IRIFIMIVGGLIGLRIVFIVLSVNRVRQGYSPLSLQTRLPTQRPDRPEGTEEGGERD
RDTSGRLVDGFLALIWDLLRSLCLFSYHRLRDLLLIVTRIVELG-----RRGWEILK
YWWNLQYWGQELKNSAVSLLNATAITVAEGTDRVIEVLQAGRAILNIPTRIROGLERI
LL-

>ENV-B.syn6.5

MRVKGIRRNQHLWRWGIML-----LGMLMICSATEQLWVTVYYGVPVWKEANTTLE
CASDAKAYKTEAHNVWATHASVPTDPNPQEVVLGNVTENFNMMWKNMMAEQMHEDIINLWD
QSLKPCVELTFLCVTLNCTDELKNATFRSNTTNSW--EKMEKGEIKNCSFNVTTSIRD
KMQKEYALFYRLDVVPI-DNDNTS-----YRLISCNTSVITQACPKISFEPIPIH
YCVFAGFAILKCNKTFNGTGCTNVSTVQCTHGIRPVVSTQLLLNGSLAEEDVIRSEN
FTDNKTIIIVQLKEAVEINCTRFNNNTRKGIHIGPGRAFATGDIIGDIRQAHCNLSRAK
WNDTLKQIVIKLREQFG-NKTIVFNQSSGGDVEIVMHSFNCGGEFFYCNTSOLFNSTW--
--NANDIRN---VTRGSNRTTGGNDTLLPCRIKQIVNMWQEVGKAMYAPPIKQIKCSS



Fig. 10 cont'd-4

NITGLLLTRDGGNNGNET--NGTEIFRPGGGDMRNNWRSELYKYKVVRIEPLGVAPT KAR  
RRVVQREKRAVT-LGAMFLGFLGAAGSTMGAASMTLTVQARLLLSGIVQQQS NLLRAIEA  
QORLLQLTVWGIKQLQARILAIERYLKDQQLLGIWGC SGKI ICTTAVPWNASWSNKSQDE  
IWNMTW MQWEREIDNYTGLIYNLIEESQNQQEKNEQELLALDKWANLWNWFDITKWLWY  
IKIFIMIVGGLIGLRIVFTVLSIVNRVRKGYSPLSFQTRLPAQRGPDRPEGIEEGGERD  
RDRSGPLVDGFLAIFWVDLRSFLFSYRHLRDL LLIVARIVELLG-----RRGWELLK  
YWWNLLQYWSQELKSSAVSLLNATAI AVAEGTDRI LEVLQRAYRAILHIPVRIROGLERA  
LL-

>ENV-B.syn6.6

MRVKGIRKNYQHLWRWGMMML-----FGMLMICSAAGNLWVTVYYGVPVWREATTLE  
CASDAKAYETE VHNWATHACVPTDPS PQEVVLENTEDFNMWKNNMVDQM HEDIISLWD  
ESLKPCVKLTPLCVTLNCTDDVRNVT-NNATNTNSSW-CEPMEKGEIKFC SFNITTSIRN  
KVQKQYALFYKLDVIPI-DSRNNNSNSTE--YNSYRLINCNSSTITQACPKVTFEPIPIH  
YCAPAGFAILKCNKKFNGTGPCNNVSTVQCTHGIRPVVSTQ LLLNGSLAEKEVVIRSDN  
FTNNAKTIIVQLNESVVINCTRPNNNTRKRISMGPGRVYYTTGEIIGDIRRAHCNISRAQ  
WNNTLKHIVEKLGKQFGNNKTI-FNHSSGGDPEIVMHSFNCRGEFFYCKSTKLFNSTWTR  
N-NGTWTRN---DTERS NSTE---EHITLPCR IKQIINMWQGVGKAMYAPP IRGQIRCSS  
NITGLILTRDGGNNDT----RGTEIFRPGGGDMKDNWRSELYRYKVVKIEPLGIAPTKAK  
RRVVQREKRAVGTIGAMFLGFLGTAGSTMGAASLTLTVQARLLLSGIVQQQNNLLRAIEA  
QQHLLQLTVWGIKQLQAKVLAVERYLRDQQLLGIWGC SGR LICTTNVPWNASWSNKS LDK  
IWNMTWMEWDREINNYTSLIYSLIEESQNQQEKNEQDLLALDKWASLWNWFDITNWLWY  
IKIFIMVVGGLVGLRIIFAVLSIVNKVRQGYSPLSLQTHLPARRGPDRPEGIEEGGERD  
RDRSVRLVDGFLALFWDDLRSCLFLYHRLRDL LLIVTRVVELLG-----RRGWEALK  
YCWNLLQYWSQELKNSAVSLFNAIAI AVAEGTDRIIEVVQRICRAIRHIPRRIROGF ERA  
LL-

>ENV-C.syn1.1

MRVRGIQRNWPQWWIWGILG-----FWMLMICNVVGNLWVTVYYGVPVWKEAKTTLE  
CASDAKAYEKEVHNWATHACVPTDNPQEI VLENTENFNMWKNDMVDQM HEDIISLWD  
QSLKPCVKLTPLCVTLNCTDVKNATSNGT TTYNNSI-DS--MNGEIKNCSFNITTEIRD  
KKKQVYALFYRLDIVPL-DNNSSE-----YRLINCNTSTITQACPKVSFDPIPIH  
YCAPAGYAILKCNKTFNGTGPCNNVSTVQCTHG IKPVVSTQ LLLNGSLAE EEEIIRSEN  
LTNNAKTIIVHLNESVEIVCTRPNNNTRKSIRIGPGQTFYATGDIIGDIRQAHCNISEKQ  
WDQTLYRVSEKLEHFP-NKTIKFAPSSGGDLEITTHSFNCRGEFFYCNTSKLFNSTY--  
--NSTQMHN---DTGS--NST-----ITLPCR IKQIINMWQEVGRAMYAPPIAGNITCKS  
NITGLLLTRDGGTNN-----NNTETFRPGGGDMRDNWRSELYKYKVVVEIKPLGIAPTKAK  
RRVVEREKRAVG-IGAVFLGFLGAAGSTMGAASITLTVQARQLLSGIVQQQS NLLRAIEA  
QQHMLQLTVWGIKQLQTRVLA IERYLKDQQLLGIWGC SGKLICTTAVPWNSSWSNKSQTD  
IWDNMTW MQWDREISNYTDTIYRLL EDSQNQQEKNEKDLLALDSWKNLWNWFDITNWLWY  
IKIFIMIVGGLIGLRIIFAVLSIVNRVRQGYSPLSFQTLTPNPRGPDR LGRIEEGGEQD  
RDRSIRLVSGELALAWDDLRSCLFLSYHRLRDFILVTARAVELLGRSSLRGLQRGWEALK  
YLGSLVQYWGLELKKS AISLLDPIAIAVAEGTDRIIE LIQRICRAIRNIPRRIROGF EAA  
LL-

>ENV-C.syn3.1

MRVMGIQRNCQQWWIWGSLG-----FWMLMIYNVMGNLWVTVYYGVPVWKEAKTTLE  
CASDAKAYDTEVHNWATYACVPTDNPQEMVLENTENFNMWKNNMVDQM HEDIISLWD  
QSLKPCVKMTPLCVTLNCSNAK KD-----NTTI-DNE-MKGEIKNCSFNITTELRD  
KKKQVYALFYKLDIVPL-NSNSSE-----YRLINCNTSAITQACPKVSFDPIPIH  
YCAPAGYAILKCNNETFNGTGPCNNVSTVQCTHGIRPVVSTQ LLLNGSLAEKEIIRSEN  
LTDNVKTIIVHLNESVEINCTRPNNNTRRSIRIGPGQAFYATGEIIGDIRQAYCNISGEK  
WNETLQRVGKKLEHFP-NKTIKFAPSSGGDLEITTHSFNCRREFFYCNTSGLFN GTY--  
--NGNGTYN---GTGTDNST-----ITIPCR IKQIINMWQEVGRAMYAPPIEGNITCKS  
NITGLLLVRDGGTENNTET-NNTETFRPGGGDMRDNWRSELYRYRVVEIKPLGIAPTKAK  
RRVVERGKRAVG-IGAVFLGFLGVAGSTMGAASITLTVQARQVLSGIVQQQS NLLRAIEA  
QQHLLQLTVWGIKQLQTRVLA IERYLKDQQLLGIWGYSGKLICTTAVPWNSSWSNRSQED



Fig. 10 cont'd-5

IWNNMTWMQWDREINNYTNTIYRLLEDSQNQQEKNEQDLLALDSWKNLWNWFEDITNWLWY  
IRIFIMIVGGLIGLRIIFAVLSIVNVRVQGYSPLSLQTLTPNPRELDRLGRIEEGGGEQD  
RDRSIRLVSGFLALAWDDLRSCLCFSYHRLRDFILIAARAAELLGRSSLKGLQRCWEILK  
YLGSLIQYWGLELKKSAINLLDTIAIVVAEGTDRIIEFIQRICRAICNI PRRIROGFEEA  
LQ-

>ENV-C.syn3.2

MRVRGILRNWQQWIIWGILG-----FWMLMICNVVGNLWVTVYYGVPVWREAKTTLF  
CASDAKAYEREVHNWATHACVPTDPNFQELVLENTENFNMMWKNMVDQMHQDIISLWD  
ESLKPCVKLTPLCVTLKCVNVTKNS--NATVNSNATV--NNNTMGEEIKNCSEFNATTEIRD  
KKQNVYALFYRLDIVPL--NENNDNSS-----YRLINCNTSTITQACPKVTFDPIPIH  
YCTPAGYAILKCNKTFNGTGPCHNVSTVQCTHGKIPVISTQLLNGSLABEEIIRSEN  
LTNNVKTIIVHLNKSVEIVCTRPGNNTKRSVRIGPGQTFYATGDIIGDIRQAHCNISRTA  
WNKTLQEVGKKLAHF--NKTIEFKPSSGGDLEVTTHSFNCRGEFFYCNTSKLFNSTYNS  
TYNSTYNSN---STNSNSNST-----ITLQCRKQIINMWQKVGRAIYAPPIAGNITCRS  
NITGLLLLTRDGGNNDTGNNDTEIFRPGGGDMKDNWRNELYKYKVVVEVKPLGIAPTGA  
RRVVEREKRAVG--LGAVFLGFLGAAGSTMGAASMTLTVQARQLLSGIVQQNNLLRAIEA  
QQHMWQVTVWGIKQLQARVLALERYLKDQQLLGLWGCSGKLICTTNVPWNSSWSNKS LTD  
IWENMTWMQWDKEISNYTDTIYRLLEVSQNQQEKNEKDLLALDSWNNLWNWFSITKWLWY  
IKIFIMIVGGLIGLRIIFGVLSIVKRVVQGYSPLSFQTLTPNPRGPDRLGRIEEGGGEQD  
KDRSIRLVNGFLALAWDDLRLNLCFSYHQLRDFILIVARAVELLGHSSLRGLQRCWEALK  
YLGSLVQYWGLELKRSAISLLDTIAIIVAEGTDRIIEVIQRICRAIRNIPTRIROGFEEA  
LLQ

>ENV-C.syn3.3

MRVRGIQRNWPQWIIWGILG-----FWMIIICRVVGNLWVTVYYGVPVWTEAKATLF  
CASDAKAYEKEVHNWATHSCVPTDPNPQEIIVLGNVTENFNMWENDMVDQMHEDVISLWD  
QSLKPCVKLTPLCVTLNCT-----NANVTVNATSDGS--IKEEIKNCSEFNTTTEIRD  
KKQKVYALFYRPDIVPLSGSNSSE-----YILINCNTSTVTQACPKVSFEPIPIH  
YCAPASYAILKCNKTFNGTGPCQNVSTVQCTHGKIPVVSTQLLNGSLAEGEIIRSEN  
LTNNAKTIIVHLNESIEIVCTRPNNTKRSIRIGPGQTFATGDIIGNIRQAHCNISEEK  
WNKTLQEVSRKLEHFP--NKTIIFNSSSGDLEITTHSFNCGGEFFYCNTTKLFNDS---  
-----ALSAFNKTS--NETITLPCRIKQIINMWQGVGRAMYAPPIAGNITCNS  
SITGLLLLTRDGGT-----NTEIFRPGGGNMKDNWRSELYKYKVVVEIKPLGVAPTEAK  
RRVVEREKRAVG--IGAVLLGFLGAAGSTMGAASITLTAQARQLLSGIVQQSNLLKAIEA  
QQHMLQTVWGIKQLQARVLAIERYLQDQQLLGIWGCSSGKLICTTTVPWNSSWSNKSQTD  
IWDNMTWMQWDREISNYTNTIYRLLEESQNQQEQNEKDLLALDKWQNLWSWFSITNWLWY  
IKIFIIIVGGLIGLRIILGVLSIVRRVVRVQGYSPLSFQTLIPNPRGPDRLGGIEEGGEQD  
RDRSVRLVSGFLSLAWDDLRSCLCFSYHQLRDFILVTARAVELLGRSSLRGLQKWEALK  
YLGSLVQYWGLELKRSAISLLDTIAIIVAEGTDRIIEFIQRICRAIRNIPTRIROGLEAA  
LQ-

>ENV-C.syn4.1

MRVRGILRNYQQWIIWGSLG-----FWMLMIYNVGGNLWVTVYYGVPVWKEAKTTLF  
CASDAKAYDTEVHNWATHACVPTDPDPQEIIVLGNVTENFNMWENDMVDQMHEDIISLWD  
ESLKPCVKLTPLCVTLKCTNVTST---GNTTRGNNTS--EN---REEMKNCSFNNTTTEIRD  
KKQKVYALFYKPDVVPL--KENSSE-----YILINCNTSTVTQACPKVSFDPIPIH  
YCAPAGFAILKCNKTFNGTGPCNNVSTVQCTHGKIPVVSTQLLNGSLAEEEIIRSEN  
LTDNAKTIIVHLNESIEIVCTRPGNNTKRSIRIGPGQAFYATGDIIGDIRQAYCNISKAT  
WNKTLQEVGKELAKHFP--NKTINFNSSSGDLEITTHSFNCGGEFFYCNTTKLFNNSL--  
-----LNNTADNST---STITLQCRKQIINMWQGVGQAMYAPPIAGNITCKS  
NITGLLLLTRDGGDTST---NGTEIFRPGGGNMKDNWRSELYKYKVVVEVKPLGIAPTGA  
RRVVEREKRAVG--IGAVLLGFLGAAGSTMGAASITLTAQARQVLSGTVQQSNLLRAVEA  
QQHMLQTVWGIKQLQTRVLAIERYLKDQQLLGIWGCSSGKLICTTNVPWNSSWSNKSQEE  
IWENMTWMQWDREISNYTGTIYRLLEESQNQQEKNEQDLLALDSWKNLWNWFDISNWLWY  
IKIFIIIVGGLIGLRIIFGVLSIVKRVVQGYSPLSFQTLIPNPRGPDRLERIEEGGEQD  
RGRSIRLVSGFLAIAWDDLRSCLCFSYHQLRDFILIAVRAVELLGHSSLRGLQRCWEALK  
YLGSLVQYWGLELKRSAISLLDTIAIIVAEGTDRIIEFIQRICRAIRNIPTRIROGFEEA  
LQ-



Fig. 10 cont'd-6

>ENV-C.syn4.2  
 MRVMGIQRNCQQWWIWGILG-----FWILMICNVMGNLWVTVYYGVPVWKEAKATLF  
 CASDAKAYEKEVHNIWATHACVPTDPNPQELVLENTENFNMWDMNDMVDQMHDIIISLWD  
 QSLKPCVKLAPLCVTLNCTNATVTATRNGSDIMNTTS-ND----GEMKNCSFNVTTELDRD  
 KKKKEYALFYRLDIVPL-NEGSGNANQNNSNYSDYRLINCNTSAITQACPKVTFDPIPIH  
 YCTPAGYAILKCNKTFNGTGPCHNVSTVQCTHGIRPVVSTQLLNGLSLAEGEIMIRSEN  
 LTNNAKTIIVHLNKSVEIVCTRPNNNTRKSVRIGPGQTFYATNDIIGDIRQAHCNISEEK  
 WNKTLLQQVGGKLAHFPP-NKTIEFKPSSGGDLEVTTHSFNCRGEFFYCNTSGLFNFTF--  
 --DGT-----ESNSTSNAT-----ITIPCRKQIINMWQKVGRAIYAPPIAGNITCRS  
 NITGLLLVRDGGNDNKT---NDTETFRPGGGDMRDNRSELYKYKVVEVKPLGVAPTAKK  
 RRVVQREKRAVG-IGAVFLGFLGVAGSTMGAASMTLVQARQVLSGIVQQQSNLLRAIEA  
 QQHLLQLTVWGIKQLQARVLALERYLRDQQLLGMWGCSSGKLICTTAVPWNSSWSNKSQED  
 IWGNMTWMQWDKEISNYTNTIYRLLEDSQNQQERNEKDLLALDSWKNLWSWEDITNWLWY  
 IKIFIMIIGGLIGLRIFAVLSIVNRVRQGYSPLSLQTLTPNPRGPDRLGRIEEEGGEQD  
 KDRSIRLVNGFLALAWDDLRLNCLFSYHRLRDFILIVARAVELLGRNSLRGLQRGWETLK  
 YLGSLLIQYWGLELKKSAISLLDTTAVAEAGTDRIIELIQRICRAICNIIPRRIRQGLEAA  
 LQ-

>ENV-C.syn4.3  
 MRVRGIQRNWPQWWIWGILG-----FWMIICRVVGNLWVTVYYGVPVWREAKTTLF  
 CASNAKAYEKEVHNWVATHACVPTDPNPQELVLENTENFNMWKNDMVDQMHDVIVSLWD  
 QSLKPCVKMTPLCVTLNCTDVKNATSNGT'TYNNNSI-DS--MNGEIKNCSFNVTTELDRD  
 KKQKAYALFYRPDIVPLPGKDNSKDNSSEYEE--YILINCNSSTITQACPKVSFEPIPIH  
 YCAPASYAILKCNNETFNGTGPCKNVSTVQCTHGKIPVISTQLLNGLSLAEKEIIRSEN  
 LTNNVKTIIIVHLKESVEINCTRPNNNTRKSVRIGPGQTFYATGDIIGNIRQAHCNISREK  
 WNTTLKRVKEKLEHFP-NKTIKFAFPSSGGDLEITHTFNCRGEFFYCNTSKLENSTYV-  
 --NRTDMND---D--TGNNST-----ITLPCRKQIINMWQEVGRAMYAPPIAGNITCNS  
 SITGLLLTRDGGNNT-----ENTETFRPGGGNMKDNWRNELYKYKVVEIKPLGVAPTEAK  
 RRVVEREKRAVG-LGAVFLGFLGAAGSTMGAASITLVQARQLLSGIVQQQSNLLRAIEA  
 QQHMLQLAVWGIKQLQARVLAIERYLQDQQLLGIWGCSSGKLICTTSPVWNSSWSNRSQED  
 IWNNMTWMQWDREISNYTDTIYRLLEVSQNQQEQNEKDLLALDKWQNLWSWFSITNWLWY  
 IRIFIMIVGGLIGLRIFAVLSLVNRVRQGYSPLSFQTLTPSPRGPDRLGRIEEEGGEQD  
 RDRSIRLVSGFLSLAWDDLRLSLCLFSYHRLRDFILIAARAAELLGRSSLRGLQRGWEILK  
 YLGSLLAQYWGLELKKSAINLLDTTAVAEAGTDRIIEVIQRICRAIYNIIPRRIRQGFEEA  
 LL-

>ENV-C.syn4.4  
 MRVRGIQRNWPQWWIWGILG-----FWMLMICNVVGNLWVTVYYGVPVWTEAKTTLF  
 CASDAKAYEREVHNWVATYACVPTDPNPQEMVLENTENFNMWKNDMVEQMHDIIISLWD  
 QGLKPCVKLTPLCVTLNCSNAKKD-----NTTI-DNE-MRGEIKNCSFNITTELDRD  
 KKQQVYALFYKLDIVPL-NSNSSE-----YRLINCNTSTITQACPKVNFDPPIPIH  
 YCAPAGYAILKCNKTFNGTGPQNVSTVQCTHRIKPVVSTQLLINGSLAEGEIIRSEN  
 LTDNVKTIIIVHLNESVEIVCTRPNNNTRKSMRIGPGQTFYATGEIIGDIRQAHCNISKEK  
 WNTTLQEVREKLEHFP-NKTIKFAHSGGDPEITTHSFNCRGEFFYCNTSOLFENSTY--  
 --NSTQMHN---DTGS--NST-----ITLPCRKQIINMWQGVGRAMYAPPIEGNITCTS  
 NITGLLLTRDGGT-----NNTETFRPGGGDMRNNWRSELYKYKVVEIKPLGIAPTAKK  
 RRVVERGKRAVG-IGAVFLGFLGAAGSTMGAASIALTAQARQLLSGIVQQQSNLLKAIEA  
 QQHMWQVTVWGIKQLQARVLAMERYLKDQQLLGLWGCSSGKLICTTTPVWNSSWSNKSQTD  
 IWDNMTWMQWDREINNYTNTIYKLLLEDSQNQQEKNEKDLLALDSWNNLWNWFSITKWLWY  
 IKIFIMIVGGLIGLRILGVLSIVRRVRQGYSPLSFQTLTPNPRELDRLGRIEEEGGEQD  
 RDRSVRLVSGFLALAWDDLRLSLCLFCYHRLRDFILVTRAVELLGRSSLKGLQRGWEALK  
 YLGNLVQYWGLELKKSAISLFDITAVAEAGTDRIIELVQRICRAIRNIIPRRIRQGFEEA  
 LL-

>ENV-C.syn6.1  
 MRVRGIQRNWPQWWIWGILG-----FWIIMCRVMGNMWVTVYYGVPVWREAKTTLF  
 CASDAKGYEKEVHNAWATHACVPTGPNPQEMVLENTENFNMWKNNMVDQMHDIIINLWD  
 QSLKPCVRLTPLCVTLKCVNVTKNS--NATVNSNATV--NNNTMGEEIKNCSFNATTEIRD  
 KKQKAYALFYRPDIVPL-NENSSSENNSSE----YILINCNTSTITQACPKVSFDPIPIH  
 YCAPASYAILKCNNETFNGTGPQNVSTVQCTHGKIPVISTQLLNGLSLAEEDIIIRSEN



Fig. 10 cont'd-7

LTNNAKTIIVHLNQSVEIVCTRPGNNTRKSMRIGPGQTFYATNDIIGNIRQAHCNISEGK
WNETLLRVKKLEEHFP-NKTIKFEPPSSGGDLEITHTFNCRGEEFFYCDTSTLFNHTY--
--VSAYMNTDVSADRKNDTQ-SNSTITLPCRIRQIINMWQEVGRAIYAPPIAGNITCRS
NITGLLLVRDGGNTT-----NSTETFRPEGGNMKDNWRSELYKYKVVEIRPLGIAPTGA
RRVVEREKRAVG-IGAVFLGFLGVAGSTMGAASMTLTVQARQVLSGVVQQQSNLLQAIEA
QQHLLQLTVWGIKQLQTRVLALERYLRDQQLLGIWGC SGKI ICTTAVPWNTSWSNKSQED
IWNNTWMOQWDREINNYTNTIYKLLSESONQOEKNEQD LLDLSDWNSLWNWFSITKWLWY
IRIFIIIVGSLIGLRIIFGVLSIVKRVRQGYSP LLSQTLTPNPREPDRLGRIEEGGGEQD
RDRSVRLVNGFLALVWDDLRLSLCLFCYHRLRDFILVTARVV ELLGRSSLRGLQKGWEALK
YLGSLVQYWGLELKKSAINLLDTIAIAVGEGETDRIIEVIQRICRAIYNI PRRIROGFEAS
LL-

>ENV-C.syn6.2

MRVRGBILRNYQQWWIWGSLG-----FWMLMIYNVGGNLWVTVYYGVPVWTDKATTLF
CASDAKAYDKEVHNVWATHACVPTDPNPQELVLENTENFNMWKNDMVNQMHEDIISLWD
ESLKPCVKLTPLCVTLNCTNATVTATRNGSDIMNTTS-ND----GEMKNCSFNITTEL RD
KKRKEYALFYRLDIVPL-DENNSSEKSSSENSSEYRLINCNTSAITQACPKVTFDPIPLH
YCAPAGYAILKCKDKTFNGTGPCSNVSTVQCTHGIKPVVSTRLLNGSLAEGEIIIRSEN
LTNNVKTIIIVHLKEPVEINCTRPNNTRESIRIGPGQTFYATGDIIGDIRQAHCNISREK
WNKTLQEVGKKLAEHFP-NKTIKFAPHSGGDLEITMHSFNCRGEEFFYCN TSGLENGTY--
--MPTYMPN---GTESNSNST-----ITIPCRIKQIINMWQEVGRAMYAPPIEGNITCNS
NITGLLLVRDGGINKT-----NNTETFRPGGGDMRNNWRSELYKYKVVEIKPLGVAPTEAK
RRVVEREKRA-A-LGAMFLGFLGAAGSNMGAASITLTAQARQLLSGIVQQRSNLLRAIEA
QQHLLQLTVWGVKQLQARVLAMERYLKDQQLLGLWGC SGKLICTTSVPWNSSWSNRSQEE
IWNNTWMEWDREISNYTNTIYRLLEDSQNQOEKNEKDLLALDSWKNLWSWFDITNWLWY
IKIFIMIIGGLIGLRIVFAVLSIVNRVRQGYSP LSFQTLTPSPRGPDR LGRIEEEGGEQD
KDRSVRLVSGFLSLAWDDLRLSLCLFSYHRLRDLILIAARAV ELLGHSSLRGLQRGWEILK
YLGSLAQYWGLELKRSAISLLDTIAITVAEGTDRIIEI IQRICRAICNI PRRIROGFETA
LL-

>ENV-C.syn6.3

MRVMGILRNCQQWWIWGVLG-----FWMLMICNVVGNLWVTVYYGVPVWKEAKTTLF
CASNAKAYEREVHNIWATHACVPTDPNPQEMVLKNVTENFNMWKNDMVDQMHEDVISLWD
QSLKPCVKLAPLVCVTLNCTNVTVNDTLHQNF-----DMKNCSFNVTTEL RD
KKQKVYALFYRLDVVPL-GDNSS-----YRLINCNTSTIAQACPKVNFDP IPIH
YCTPAGYAILKCNDKTFNGTGPCKNVSTVQCTHEIKPVVSTQ LLLNGSLAEEGIIRSEN
LTDNAKTIIVHLNESVEINCTRPGNNTRQSIRIGPGQAFYATGAIIGDIRQAHCNISKDE
WEKTLKRVSEKLEHFP-NKTIEFKPSSGGDLEVTTHSFNCRREFFYCN TSKLFNSTY--
--NSTQMHN---DTGS--NST-----ITLPCIKQIINMWQGVGQAMYAPPIKGNITCKS
NITGILLTRDGGNLT-----NGTETFRPGGGDMKDNWRSELYRYRVVEIKPLGIAPT KAK
RRVVQREKRAVG-IGALFLGFLGTAGSTMGAASLTLTVQARQLLS SIVQQQSNLLRAIEA
QQHMLQLTIWGIKQLQTRVLAVERYLKDQQLLGMWGC SGKLICTTAVPWNASWSNKSQEE
IWGNMTWMOQWDREISNYTDIIYRLLEESQNQOERNEKDLLALDSWNNLWNWFNITNWLWY
IKIFIMIVGGVIGLRIIFAVLSLVNRVRQGYSP LSFQTLTPNPRELDRLGRIEEEGGEQD
RDRSIRLVNGFLAIAWDDLRLSLCLFSYRRLRDFILIAARA AELLGRSSLRGLQRGWETLK
YLGSLIQYWGLELKKSAISLFDTIAIAVAEGTDRIIEI IQRICRAIRNI PRRIROGLEAA
LQ-

>ENV-C.syn6.4

MRVMGIQRNCQQWWIWGILG-----FWMLMIYNVVG NLWVTIYYGVPVWKEAKATLF
CASDAKAYDTEVHNVWATHACVPTDPDPQEMVLGNVTENFN MWKNDMADQMHEDIISLWD
QGLKPCVKLTPLCVTLHCTN-----TNITNENRTI-GDKLNE-EMKNCSFNITTEL RD
KKQQVYALFYKPDVVPL-NGGEHNETGE-----YILINCNSSTITQACPKVSFEPIPIH
YCAPAGFAILKCNKTFNGTGPCHNVSTVQCTHGI RFPVVSTQ LLLNGSLAEEEIIRSEN
LTDNVKTIIIVHLNKSVEIVCTRPNNNTRKSIRIGPGQTFYATNDIIGDIRQAYCNISAEK
WNKTLERVEEKLKEHFP-NKTIKFNSSSGGDLEITTHSFNCRGEEFFYCN TSNLENGTY--
--HGTQSTN---ST---NST-----ITLQCRIKQIINMWQKVGRAMYAPPIAGNITCKS
NITGLLLLRDGGTEN-----NDTETFRPGGGNM RDNWRSELYKYKVVEVKPLGIAPTTAK
RRVVERDKRAVG-IGAVLLGFLGAAGSTMGAASMALTVQARQLLSGIVQQQSNLLRAVEA



Fig. 10 cont'd-8

QQHMLQLTWVGKIQQLQARVLALERYLKDQQLLGIWGC SGRLICTTAVPWNS SWSNKTQGE  
I WENMTWMQWDKEINNYTNTIYRLLEESQTQQEQNEKDLLALDSWKNLWNWFDTKWLWY  
IKIFIMVVGGLIGLRIIFAVLSIVNSVRQGYSPLSLQTLTPNPRGPDRLERIEEGGEQD  
RNRSIRLVNGFLALAWDDLRSCLFSYHHLRDFILVTARAVELLGRSSLKGLQRGWEALK  
YLGNLVQYWGLELKKSAISLLDTTAVAEAGTDRIIELVQRICRAILNIPTRIRQGFEEA  
LQ-

>ENV-C. syn6.5

MRVRGIPRNWPQWWTWGILG-----FWMIIICRVVGNLWVTVVYGVVPVWTEAKTTLF  
CASDAKAYEREVHNVWATHSCVPTDPNPQEIIVLGNVTENEFNMWENDMVDQMHQDIISLWD  
QSLKPCVKMTPLCVTLNCSNAKDD-----NTTI-DNE-MKGEIKNCSFNITTEIRD  
KKQKVHALFYRLDIVPL-NEGSGNANQNSNSYDYRLINCNTSTVTQACPKVTFDPIPIH  
YCAPARYAILKCNNTFNGTGPCNNVSTVQCTHGKIPVSTQLLLSGSLAE EIVIRSEN  
LTNNAKIIIVHLNESVEIVCTRPNNTRRSIRIGPGQTFYATGEIIGDIRQAHCNISAKQ  
WNTTLERVKEKLEHFP-NKTIKFEPHSGGDPEITTHSFNCGGEFFYCNTSOLFENSTY--  
--NSTYMSN---NTGENSNET-----ITLPCRKQIINMWQVGRAMYAPPIAGNITCNS  
SITGLLLTRDGGNNNDTGNNDTEEFRPGGGDMRDNRSELYKYKVVVELKPLGIAPTEAK  
RRVVKREKRAVG-IGAVLFGFLGAAGSTMGAASIALTAQARQVLSGIVQQNNLLRAIEA  
QQHVLQLTWVGKIQQLQTRVLAIERYLKDQQLLSLWGC SGKLICTTVPWNS SWSNKS LTD  
I WDNMTWMQWDREISNYTGTIYRLLEDSQSQQEKNEKDLLELDKWNLWNWFDISNWLWY  
IKIFIIVGGLIGLRIIFAVLSIINRVRQGYSPLLFQTLTPNPRGLDRLGRIIEEGGEQD  
KDRSIRLVNGFLALAWEDLRSCLFSYHQLRDFILIVARAVELLG-----RRGWEALK  
YLGNLVLYWGLELKKSAVSLLDTIAI AVAGGTDRIEVVQRICRAIRNIPTRIRQGLEAA  
LL-

>ENV-C. syn6.6

MRVRGILRNWQQWVIWGILG-----FWMVMICNVMGNLWVTVVYGVVPVWQEAKTTLF  
CASDAKAYEKEVHNVWATHACVPTDPSQEIIVLENVTENEFNMWKNDMVEQM HEDIISIWD  
QSLKPCVTLTPLCVTLNCTDVKNVATSNGTTTYNN SI-DS--MNGEIKNCSFNITTEIRD  
KKKQVYALFYKLDIVPL-NSNSSE-----YRLINCNTSAVTQACPKVSWDPIPIH  
YCAPAGYAILKCNKTFNGTGPCNTVSTVQCTHRIKPVVTTQLLNGSLAEKEIIRSEN  
LTNNIKTIIIVHLNESIEIVCTRPNNTNRKSVRIGPGQTFATGDIIGDIRKAHCNISEDK  
WNETLQRVGKKLVEHFP-NKTIKFA PSSGGDLEVTTHSFNCKGGEFFYCNTTKLFD-----  
-----DSERINTT-----TIIILPCRKQFINMWQVGRAMYAPPIAGNITCTS  
NITGLLLTRDGGT-----NNTTEIFRPGGGNMKDNWRNELYKYKVVVEVKPLGVAPT KAK  
RRVVEREKRAVG-LGAVFLGFLGAAGSTMGAASITLTVQARQLLEFGIVQQQSNLLKAIEA  
QQHMWQVTVWVGKIQQLQARVLAIERYLQDQQLLGIWGC SGKLICTTNVPWNS SWSNKS QTD  
I WDNMTWMQWDKEISNYTDTIYRLLEVSQNOQEEKDLLALDKWQNLWNWF SITNWLWY  
IRIFIMIVGGLIGLRIILGVLSIVRRVRQGYSPLSFQTLIPNPRGPDRLGGIEEGGEQD  
RDRSIRLVSGFLALAWDDLRLNCLFSYHRLRDFILIVRRAVELLGRNSLRGLQRGWEALK  
YLGSLGQYWGLEIKKSAISLLDTIAIVVAEGTDRIIEFIQRFCAIRNLPRRIRQGFEEA  
LL-

>ENV-M. syn1.1

MRVTGIRKNYQHLWRWGTMLLWRWGTMLLGLMLICSAAGNLWVTVVYGVVPVWKEATTTLF  
CASDAKAYDTEVHNVWATHACVPTDPNPQEVVLENVTENEFNMWKNNMVEQM HEDIISLWD  
QSLKPCVKLTPLCVTLNCTDDVRNVT-NNATNTNS SW-GEPM EKGEIKNCSFNITTSIRD  
KVQKEYALFYKLDVVP I-DNDSNNTN-----YRLISCNTSVITQACPKVSFEPIPIH  
YCAPAGFAILKCNDDKKNFTNGTGPCNTVSTVQCTHGKIPVSTQLLNGSLAE EEVVIRSEN  
FTNNAKTIIVQLNESVEINCTRPNNTNRKSVRIGPGQTFYATGDIIGDIRQAHCNISRAQ  
WNNTLKHIVEKLGKQFGNKTIVFNHSSGGDPEIVMHSFNCGGEFFYCNTTKLFNSTWTR  
N-NGTWTRN---DTERS NSTE---EHITLPCRKQIINMWQEVGKAMYAPPIRGQIRCSS  
NITGLLLTRDGGNDT-----SGTEIFRPGGGDMRDNRSELYKYKVVKIEPLGVAPT KAK  
RRVVQREKRAVG-IGAVFLGFLGAAGSTMGAASITLTVQARQLLSGIVQQQNNLLRAIEA  
QQHLLQLTWVGKIQQLQARVLAVERYLKDQQLLGIWGC SGKLICTTAVPWNS SWSNKS LNE  
I WNNMTWMEWEREIDNYTGLIYTLIEESQNOQEKNEQELLELDKWA SLWNWFDISNWLWY  
IKIFIMIVGGLIGLRIVFAVLSIVNRVRQGYSPLSFQTRLPA PRGPD RPEGIEEGGERD  
RDRSIRLVSGFLALAWDDLRSCLFSYHRLRDL LLIVTRIVELLG-----RRGWEALK



Fig. 10 cont'd-9

YWWNLLQYWSQELKNSAVSLLNATAIAVAEGTDRVIEALQRACRAILHIPRRIROGLERA  
LL-

>ENV-M.syn3.1

MRVRGIQRNWPQWWIWGILG-----FWMLMICNVVGNLWVTVYYGVPVWKEAKTTLF  
CASDAKAYEKEVHNVWATYACVPTDPNPQEIHLNVTEEFNMWKNMVDQMHEDEIISLWD  
QSLKPCVQLTPLCVTLNCTN-VNVTNLKNETNTKSSSGGKMEEGEMKNCSFNITTSIRD  
KVQKEYALFYKLDVVPIDNDSNNTN-----YRLISCNTSVITQACPKVTFEPIPIH  
YCTPAGFAILKCKDKKFNGTGPKNVSTVQCTHGKIPVISTQLLNGSLAEDEIIRSEN  
ITNNAKTIIVQLNESVEINCTRPNNTRKSVRIGPGQTFYATGEIIGDIRQAHCNLSRAK  
WNDTLKQIVIKLREQFG-NKTIVFNQSSGGDPEIVMHSFNCGGEFFYCNTTQLFNSTW--  
-----N---STSLFNSTN---GTITLQCRIKQIINMWQEVGKAMYAPPIEGNITCKS  
NITGLLLVRDGGT---EP--NDTETFRPGGGNMKDNWRSELYKYKVVKIEPLGVAPTKAK  
RRVVQREKRAVGTIGAMFLGFLGAAGSTMGAASMTLTVQARLLLSGIVQQQNNLLRAIEA  
QQHLLQLTVWGIKQLQARILAVERYLKDQQLLGLWGCSGKLICTTAVPWNTSWSNKSQTD  
IWDNMTWMEWEREIDNYTGLIYTLIEESQNOQEKNEQELLELDKWASLWNWFDITKWLWY  
IKIFIMIVGGLVGLRIVFAVLSIVNRVRKGYSPLSFQTLTPNPRGPDRLGRIEEEGGEQD  
RDRSIRLVSGFLALAWDDLRSCLFSYHQLRDFILIVARAVELLGRSSLRGLQRGWALK  
YLGSLVQYWGLELKKSAISLLDTIAIAVAEGTDRIIEVIQRICRAIRNIPRRIROGFERA  
LL-

>ENV-M.syn3.2

MRVTGIRKNYQHLWRWGTMLLWRWGTMLLGIILMICSAAAGKLWVTVYYGVPVWRDAETTLF  
CASDAKAHETEVEHNIWATHACVPTDPNPQEVVLGNVTENFNMWKNMVEQMHEDEIISLWD  
ESLKPCVKLTPICVTLNCTDDVRNVT--NNATNTNSSW-GEPMEKGEIKNCSFNMTTELRO  
KKQKVHALFYKLDIVPL-NSNSSE-----YRLINCNTSAITQACPKVSFEPIPIH  
YCAPAGFAILKCNKDFNGTGPCNTVSTVQCTHGIRPVVSTQLLNGSLAEDEEVVIRSEN  
FTNNAKTIMVQLNVSVEINCTRPNNTRKSIHIGPGRAFYTGDIIIGDIRQAHCNISRAQ  
WNNTLKHIVEKLGKQFGNKTIVFNHSSGGDPEITTHSFNCGGEFFYCNSTKLFNSTWTR  
N-NGTWTRN---DTERSNSSTE---EHITLPCRIKQIVNMWQRVGQAMYAPPIRGQIRCSS  
NITGLLLTRDGGNDT-----SGTEIFRPGGGDMRNNWRNELYKYKVVRIEPLGVAPTRAK  
RRVVEREKRAVG-LGAVFLGFLGAAGSTMGAASLTLTVQARQVLSGIVQQQSNLLKAIEA  
QQHLLKLTWGIKQLQARVLAVERYLRDQQLLGIWGCSSGKLICTTTVPWNASWSNKSLE  
IWNMTWMEWEKEIDNYTNLIYNLLEESQNOQEKNEQDLLALDKWANLWNWFDISNWLWY  
IKIFIIIVGGLIGLRIVFAVLSIINRVROGYSPLSLQTLIPNPRGPDRPGGIEEEEGGEQG  
RDRSIRLVNGFLALAWDDLRLNCLFSYHRLRDLILLIVTRIVELLG-----RRGWALK  
YWWNLLQYWSQELKNSAISLLNATAVAVAEGTDRVIEALQRACRAILHIPRRIROGLERL  
LL-

>ENV-M.syn3.3

MRVKETRKNYQHLWKWGTML-----LGMLMICSAAEQWLWVTVYYGVPVWKEATTTLF  
CASDAKAYDTEVHNVWATHACVPTDPSQEVVLENVTENFNMWKNMVEQMHTDIISLWD  
QSLKPCVKLTPICVTLNCTDYVKNIT--NNATSTNSSW-GKPMKGEIKNCSFNITTSIRN  
KVQKQYALFYKLDIVPI--DNDNTS-----YRLINCNTSTITQACPKVSFDPIPIH  
YCAPAGYAILKCNKTFNGTGPCNNVSTVQCTHGKIPVVSTQLLNGSLAEDEIIRSEN  
LTNNAKTIIVHLNKSVEINCTRPNNTRKSIHIGPQAFYATGDIIIGDIRKAHCNISGTK  
WNHTLEQVMEELKKHFP-NKTIKFNSSSGGDLEITTHSFNCRGEFFYCNTSGLFNSTW--  
--NDTTINR---TEGSNNTR---NITLPCRIKQIINMWQGVGRAMYAPPIAGNITCKS  
NITGILLTRDGGNNN-----STNETFRPGGGDMRDNWRSELYKYKVVVEIKPLGIAPTKAK  
RRVVEREKRAVG-IGAVFLGFLGTAGSTMGAASITLTVQARQLLSGIVQQQSNLLRAIEA  
QQHMLQLTVWGIKQLQTRVLAERYLKDQQLLGIWGCSSGKLICTTNVPWNSSWSNKSQSE  
IWDNMTWMMQWDREISNYTDTIYRLLEDSONQOQEKNEKDLLALDSWKNLWNWFDITNWLWY  
IRIFIMIVGGLIGLRIFAVLSIVNRVRQGYSPLSFQTRLPAAPRGPDRPEGIEEEEGGERD  
RDRSVRLVDGFLALIWDLLRSCLFSYHRLRDFILIAARTVELLGHSSLKGLRLGWGLK  
YLWNLLQYWIQELKNSAVSLLNATAIAVAEGTDRVIEVVQRAYRAILHIPTRIROGLERA  
LL-



Fig. 10 cont'd-10

>ENV-M.syn4.1  
 MRVKETRKNYQHLWKWGTML-----LGMLMICSAAEQWVTVVYGVVWKEATTTLF  
 CASDAKAHETEVEHNIWATHACVPTDPNPQEVVLENTENFNMWKNMVEQMHTDIISLWD  
 QSLKPCVELTFLCVTLHCTDL-----GNATNT-----MEKGEIKNCSFNMTTELDR  
 KKQKVYALFYRLDIVPI-DNDNTS-----YRLINCNTSVIKQACPKVTFEPIPIH  
 YCTPAGFAILKCNCKNFNGTGPCKNVSTVQCTHGIRPVVSTQLLLNGSLAEKEIIRSEN  
 LTDNAKTIIVHLNKSVEINCTRPSNNTRKSVRIGPGQTFYATGDIIGDIRQAHCNISRAK  
 WNNTLQKQIVTKLREQFK-NKTIVFNQSSGGDLEITTHSFNCRGEEFFYCNTTQLENSTW--  
 -----KN---DTEVSNNTK-GNDTITLPCRKQIVNMWQEVGRAMYAPPIEGNITCNS  
 NITGILLTRDGGNNGNET--NGTEIFRPGGGNMRDMWRNELYKYKVVVEIKPLGVAPTEAK  
 RRVVEREKRAVG-IGAVFLGFLGAAGSTMGAASITLTVQARQLLTGIVQQQSNLLRAIEA  
 QQHMLQLTVWGIKQLQTRVLAIERYLKQDQQLLGLWGCSGKLICTTAVPWNSWSNKTynd  
 IWDNMTWMOVDREISNYTDTIYRLLEDSQNOQEKNEKDLLALDSWKNLWNWFDITNWLWY  
 IKIFIMIVGGLIGLRIFAVLSIVNRVRQGYSPLSFQTLTPNPRGPDRPGGIEEEGGEGG  
 RDRSIRLVNGFLALAWDDLRLNCLFSYHQLRDFILIVARAVELLGRSSLRGLQRGWEALK  
 YLGSLVQYWGLELKKSAISLLDTIAIAVAEGTDRVIEVVQRAYRAILHIPTRIRQGLERL  
 LL-

>ENV-M.syn4.2  
 MRVIRGIQRNWPQWIIWGIIG-----FWMLMICNVVGNLWVTVVYGVVWKEAKTTLF  
 CASDAKAYEKEVEHNVWATHACVPTDPSQEVVLENTENFDMWKNMVEQMVEDVISLWD  
 QSLKPCVKLAPLCVTLNCTDYVKNIT-NNATSTNSSW-GKPMKGEIKFCSEFNITTSIRM  
 KVQKQYALFYKLDVVQM-DEDNTS-----YRLISCNTSTITQACPKVTFDPIPIH  
 YCAPAGFAILKCNKNTFNGTGPCNTVSTVQCTHGIRPVVSTQLLLNGSLAEKEIIRSEN  
 LTNNAKTIIVHLNESVEIVCTRPNNNTRKSIHIGPGRFYATGEIIGDIRQAHCNLSRAK  
 WNDTLKQIVIKLREQFG-NKTIIIFNQSSGGDPEITTHSFNCGGEEFFYCNS'TQLENSTWNF  
 --NGTWNKN---FNNTWNTEGTNDTITLPCCKIKQIINMWQVGOAMYAPPISGQIRCSS  
 NITGLILTRDGGN---DT--SGTEIFRPGGGDMRDNRSELYKYKVVKIEPLGVAPTKAK  
 RRVVQREKRAVG-IGALFLGFLGAAGSTMGAASMTLTVQARQLLSGIVQQQSNLLKAIEA  
 QQHLLQLTVWGIKQLQARILAVERYLKDQQLLGIWGCSSGKLICTTTPWNASWSNKSLSNE  
 IWDNMTWMEWEREIDNYTGLIYNLIEESQTQOQEKNEQELLELDKASLWNWFDITKWLWY  
 IKIFIMIIGGLIGLRIVFTVLSIVNRVRKGYSPLSFQTLTHHQREPDRPERIEEGGEGQD  
 RDRSGRLVDGFLAIWVDLRSCLFSYHRLRDLILLIVPRIVELLG-----RRGWEVLK  
 YWWNLLQYWSQELKNSAVSLLNATAIAVAEGTDRIIEVIQRICRAIRNIPRIRQGLERA  
 LL-

>ENV-M.syn4.3  
 MRVKETQMNWPNLWKWGTLI-----LGLVIICSASDNLWVTVVYGVVWKAETTLF  
 CASDAKAYDTEVHNWATYACVPTDPNPQEIHLNVTENFNMWKNMVDQMHEDIISLWD  
 ESLKPCVKLTPLCVTLNCTDEVRNVT-NNATNTNSSW-GEPMEKGEIKNCSFNITTSIRD  
 KVQKEYALFYKLDVPI-DNDSNNTN-----YRLISCNTSVITQACPKVSFEPIPIH  
 YCAPAGYAILKCNCKNFNGTGPCNNVSTVQCTHGIRPVVTTQLLLNGSLAEKEIIRSEN  
 ITNNAKTIIVQLNESVVINCTRPNNNTRKSIIRIGPGQAFYATGDIIGNIRQAHCNISRAK  
 WNNTLRQIVEKLGEQFGNNKTIVFNHSSGGDPEIVTHSFNCAGEFFYCNTTKLENSTWTR  
 N-NGTWTRN---DTERSNSTE---EHITLPCRKQIINMWQEVGKAMYAPPIRGQIRCSS  
 NITGLLLLTRDGGNNN-----STNETFRPGGGNMKDNWRSELYKYKVVQIEPLGIAPTAK  
 RRVVEREKRAVG-LGAVFLGFLGTAGSTMGAASLTLTVQARQVLSGIVQQQSNLLRAIEA  
 QQHLLKLTWGIKQLQARVLAIERYLQDQQLLGMWGCSSGKLICTTNVPWNSWSNKSQTD  
 IWDNMTWLQWDKEISNYTSLIYTLIEESQNOQEKNEQDLLALDKASLWSWFDISNWLWY  
 IKIFIIIVGGLIGLRIVFAVLSIINRVRQGYSPLSLQTLIPNPRGPDRLGRIEEEGGEGQD  
 RDRSIRLVSGFLALAWDDLRLSLCIFSYHRLRDFILIAARTVELLGHSSLKGLRLGWGLK  
 YLGNLLLYWGQELKNSAINLLDTIAIAVAGWTDRIEIGQRAGRAILNIPRIRQGFERA  
 LL-

>ENV-M.syn4.4  
 MRVTGIRKNYQHLWRWGTMLLWRWGTMLLGLMICSAAAGKLWVTVVYGVVWRDADTTLF  
 CASDAKAYDTEAHNVWATEASVPTDPNPQEIIVLENTENFNMWKNMVEQMHEDIISLWD  
 QSLKPCVQLTFLCVTLNCTN-VNVTNLKNETNTKSSSGGKMEEGEMKNCSFNITTEIRD  
 KKQKVHALFYKLDIVPL-NSNSSE-----YRLINCNTSAITQACPKVSFDPIPIH



Fig. 10 cont'd-11

YCTPAGYAILKCNKKFNGTGPCKNVSSVQCTHGIKPVISTQLLNGSLAE EEVVIRSEN
FTNNAKTIMVQLNVSVEINCTRPNNNTRRSIPIGPGRAFYTTGDIIGDIRKAHCNISRAQ
WNNTLKHIVEKLGKQFGNNKTIIFKPSSGGDPEIVMHSFNCGGEFFYCNTSGLFNSTW--
-----N---STSLFNSTN---GTITLQCRIKQIINMWQGVGRAMYAPPIAGNITCKS
NITGLLLVVRDGGT---EP--NDTETFRPGGGDMKDNWRSELYKYKVVRIEPLGVAPTRAK
RRVVEREKRAIG-LGAMFLGFLGAAGSTMGAASVTLTVQARLLLSGIVQQONLLRAIEA
QQHLRLTVWGIKQLQARVLAVERYLRDQQLLGIWGC SGKI ICTTAVPWNTSWSNRS LNE
IWNMTWMEWEKEIDNYTNLIYNLLEESQNQQEKNEQEL LALDKWANLWNWEDI SNWLWY
IRIFIMIVGGLVGLRIVFAVLSIVKRV RQYSPLSFQTRLPAPRGPDRPEGIEEEGGGERD
RDRSVRLVDGFLALIWDLLRSLCLFSYHHLRDL LLIVARIVELLG-----RRGWEALK
YWWNLLQYWIQELKNSAISLLNATAVAVAEGTDRVIEALQRACRAILHIPRRIROGFEAA
LL-

>ENV-M. syn6.1

MRVMGIQRNCQQWWIWGILG-----FWMLMICNVMGNLWVTVYYGVPVWKEANTTLF
CASDAKAYEREVHNWATHASVPTDPNPQEVVLENTEDFNMMWKNMVEQMVEDVISLWD
QSLQPCVKLTPLCVTLNCTD-LRNATNGNDTNTTSSS-REMMGGGEMKNCSFNITTEIRD
KKQKVYALFYKLDVVP I-DNDSNNTN-----YRLISCN TSAVTQACPKVTFDPIPIH
YCTPAGFAILKCRDKKFN GTGPCKNVSTVQCTHGIKPVVTTQLLNGSLAE EIVIRSEN
FTDNAKTIIVQLKEAVEINCTRPNNNTRKGIHIGPGRAFYATGEIIGDIRQAHCNVSRSF
WNKTLQQVATQLRKHF--NKTIIFNSSSGDLEITTHSEFNCRGEFFYCNTSGLFNSTW--
--NDTTINR----TEGSNNTR----NITLPCRIKQFINMWQEVGRAMYAPPIAGNITCRS
NITGLLLTRDGGNNGNET--NGTEIFRPGGGDMRNNWRSELYKYKVV EIKPLGIAPTKAR
RRVQREKRAVG-IGAVFLGFLSAAGSTMGAASITLTVQARQLLTGIVQQOSNLLKAIEA
QQHMLQLTVWGVKQLQARVLAVERYLRDQQLLGIWGC SGRLICTTAVPWNTSWSNKS LNE
IWDNMTWMEWEREIDNYTSLIYTLIEESQNQQEKNEQELLELDKWANLWNWFSITNWLWY
IRIFIMIVGGLIGLRIIFGVLSIVKRV RQYSPLSFQTRLPAPRGPDRPEGIEEEGGGERD
RDRSGRLVDGFLALIWDLLRSLCLFSYHRLRDL LILIAARIVELLGHSSLKGLRLGWEALK
YLWNLLLYWGQELKNSAISLLNTTAIVVAEGTDRVIEVLQ RAGRAILNIPRRIROGFEAA
LL-

>ENV-M. syn6.2

MRVTGIRKNYQHLWRWGTMLLWRWGTMLL GILMICSAAAGKLWVTVYYGVPVWREAKTTLF
CASDAKAYEKEVHNWATYACVPTDPNPQEMVLENTENFNMMWKNMVDQM HEDIISLWD
ESLKPCVKLTPLCVTLHCTDL-----GNATNT-----MEKGEIKNCSFNITTEIRD
KKQKVHALFYRLDVVPI-DNDNTS-----YTLINCNTSVITQACPKVTFEPIPIH
YCAPAGFAILKCNKKFNGTGPCTNVSTVQCTHGIKPVVSTQLLNGSLAE GEIIRSEN
LTDNAKTIIVHLNESVEIVCTRPNNNTRKSVRIGPGQTFYATGAIIGDIRQAYCNISRAK
WNNTLKQIVTKLREQFGNNKTIIFKPSSGGDLEITMHHFNCRGEFFYCNTTQLFNSTWNF
--NGTWNKN---FNNTWNNTEGTNDTITLPCRIKQIINMWQGVGRAMYAPPIISGQIRCSS
NITGLLLTRDGGT-----NNT EIFRPGGGNMRDNWRSELYKYKVVKIEPLGVAPTKAK
RRVQREKRAVG-IGALFLGFLGAAGSTMGAASMTLTVQARQVLSGIVQQORNLLRAIEA
QQHLLQLTVWGIKQLQARILAVERYLKDQKFLGLWGC SGKI ICTTAVPWNASWSNKS LDD
IWNMTWMOWEREIDNYTGLIYSLIEESQTQQEKNEQELLQLDKWASLWNWFDITNWLWY
IRLFIMIVGGLVGLRIVFTVLSIVNRV RKGYSPLSFQTLTHHQREPDRPERIEEGGGEQG
RDRSVRLVSGFLALEWDDLRSCLFCYHRLRDFELIAARTVELLGHSSLKGLRRGW EGLK
YLWNLLQYWIQELKNSAISLLNATAVAVAEGTDRVIEALQRACRAILHIPRRIROGLERL
LL-

>ENV-M. syn6.3

MRVRGIQRNWPQWWIWGILG-----FWMIICRVVGNLWVTVYYGVPVWKDAETTLF
CASDAKSYETEAHNIWATHACVPTDPS PQEVVLGNVTENFNMMWKNDMVEQM HEDIISLWD
QSLKPCVELTPLCVILNCTDDVRNVT-NNATNTNSSW-GEPMEKGEIKNCSFNITTSIRN
KVQKQYALFYKLDVQI-DDNNTSNTS-----YRLINCNTSAITQACPKVSFDPIPIH
YCAPAGYAILKCNKTFNGTGPCHNVSTVQCTHGIKPVISTQLLNGSLAE EEVVIRSEN
FTNNAKTIMVQLNVSVEINCTRPNNNTRKSIHIGPGRAFYTTGDIIGDIRQAHCNISRAQ
WNNTLKHIVEKLGKQFGNNKTIIFKPSSGGDPEIVMHSFNCRGEFFYCNTSKLFNSTWTR
N-NGTWTRN---DTERSNSTE---EHITLPCRIKQIINMWQRVGQAMYAPPIAGNITCNS
SITGLLLTRDGGN---DT--SGTEIFRPGGGNIKDNWRSELYKYKVVQIEPLGVAPTRAK



Fig. 10 cont'd-12

RRVVEREKRAVG--IGAMIFGFLGAAGSTMGAASMLTVQARQLLSGIVQQSNLLMAIEA  
QOHLKLTVWGIKQLRARVLAVERYLKDQQLLGIWGCSGKHICTTNVPWNSSWSNKSLE  
IWNMTWIEWEREINNYTGLIYNLLEKSONQOEKNEQDLLALDKWASLWSWFDISNWLWY  
IKIFIIIVGGLIGLRIVFAVLSLVNRVRQGYSPLSLQTLPTFRGPDRPEGTEEEGGEQG  
RDRSIRLVSGFLALAWDDLRSCLRFSYHRLRDFILIVARTVELLGRSSLKGLRLGWGLK  
YLGNNLLYWGQELKISALSLLDTTATAVAGWTDRIEIQORLCRAIRNIPRIRQGAERA  
LQ-

>ENV-M.syn6.4

MRVKETQMNWPNLWKWGTLI-----LGLVIICSASDNLWVTVYYGVPVWRDADTTLF  
CASDAKAHETEVEHNVWATHACVPTDPNPQEIHLNVTEEFNMWKNMVEQMHTDIIISLWD  
QSLKPCVRLTPLCVTLNCTDELKNATFRSNTTNSW--EKMERGEIKNCSFNITTSIRD  
KVQKEYALFYKLDIVPL-NSNSSE-----YRLINCNTSVIKQACPKISFDPPIPIH  
YCAPAGFAILKCKDKKFNGTGPCQNVSTVQCTHRIKPVVSTQFLLNGSLAEEDIIIRSEN  
ITNNAKTIIVQLNESVVINCTRPNNNTRRSIPIGPRVFTTEDIIGDIRQAHCNLSRAK  
WNTDLKQIVIKLREQFG-NKTIVFNQSSGGDLEIVMHSFNCGGEFFYCNSTQLFNSTWF-  
--NSTW-----STEGSNTE-GSDTITLPCRIKQIVNMWQGVGKAMYAPPIRQIRCSS  
NITGILLTRDGGTNGT----NETETFRPGGDMRDNWRSELYRYKVVKIEPLGIAPTAK  
RRVVEREKRAIG-LGAMFLGFLGTAGSTMGAASLTLTVQARQLMSGIVQQNNLLRAIEA  
QOHLKLTVWGIKQLQARVLAVERYLKDQQLLGLWGCSGKLICTTTPWNSSWSNKSQTD  
IWDNMTWQWDREISNYTNTIYRLLEDSONQOEKNEKDLLALDSWKNLWNWFDITKWLWY  
IKIFIMIVGGLIGLKIIVFAVLSIINVRVQGYSPLSFQTLIPNPRGPDRPGGIEEGGEQD  
RDRSIRLVNGFLALIWVDLRSCLFSYHRLRDLIIIVTRIVELLG-----RRGWEALK  
YWWNLLQYWSQELKNSAINLLDTTATAVAEGTDRIIEVIQORICRAIRNIPRIRQGLERA  
LL-

>ENV-M.syn6.5

MRVKGIRKNYQHLWKWGTM-----LGMLMICSATEKLWVTVYYGVPVWKEATTTLF  
CASDAKAYDTEVHNVWATYACVPTDPNPQELVLENVTENFDMWKNMVEQMHEIDIINLWD  
QSLKPCVKLTPICVTLNCTDYVKNIT-NNATSTNSW-GKPMKGEIKNCSFNMTTEL RD  
KKQKVYSLFYKLDVVQM-DEDNTS-----YRLISCNTSVITQACPKISFEPIPIH  
YCTPAGYAILKCNKFNNGTGPCKNVSSVQCTHGKIPVISTQLLLNGSLAEEDIIIRSEN  
LTNNVKTIIIVHLNKSVEINCTRPNSNTRTSIRIGPGQAFYATGDIIGDIRKAHCNISRAN  
WNNTLRQIVEKLGEQFGNKTIVFNHSSGGDPEITTHSFNCGGEFFYCNTTKLFNSTWTW  
N-NSTW--N---NTRKSDTE---EITLPCRIKQIINMWQEVGKAMYAPPIQGVIRCES  
NITGLIILTRDGGNNN-----STNETFRPGGDMRDNWRSELYKYKVVRIEPLGVAPTEAK  
RRVVQREKRAVGTIGAMFLGFLGAAGSTMGAASVTLTVQARLLLSGIVQQNNLLKAIEA  
QOHLRLTVWGIKQLQARVLAVERYLQDQQLLGIWGCSGKLICTTNVPWNSSWSNRSLNE  
IWNMTWMEWEKEIDNYTNLIYNLLEESQIQOEKNEQELLALDKWANLWNWFDISNWLWY  
IRIFIIIVGGLVGLRIVFAVLSIVNKVRQGYSPLSFQTHLPAQRGPDRPEGIEEGGEQD  
RDRSVRLVDGFLAIWVDLRSCLFSYHRLRDLIIIVARIVELLG-----RRGWEVLK  
YWWNLLKYWSQELKNSAVSLLNATAVAEGTDRIIEVIQORICRAICNIPRIRQGFERA  
LL-

>ENV-M.syn6.6

MRVKETRKNYQHLWRWGIML-----LGMLMICSAAEQWVTVYYGVPVWKEAKTTLF  
CASNAKAYDTEAHNVWATHACIPTDPNPQEIIVLENVTESFNMWKNMVDQMHEVDVISLWD  
QSLKPCVQLTPLCVTLNCTN-VNVTLKNETNTKSSSGGKMEEGEMKNCSFNVTTEL RD  
KKKKEYALFYRLDIVPL-NEGNNSNSSY-----YRLINCNTSTITQACPKVSFEPIPIH  
FCAPAGFAILKCNKFNNGTGPCKNVSTVQCTHGKIPVISTQLLLNGSLAEKEIIIRSEN  
LTNNAKIIIVQLNESVEINCTRPNNTRKSIRIGPGQTFYATGDIIGNIRQAHCNISRTQ  
WNNTLKQIAIKLREQFG-NKTIIIFNQSSGGDPEIVTHSFNCGGEFFYCKSTKLFNSTW--  
-----N---STSLFNSTN---GTITLQCRKQIINRWQEVGKAMYAPPIEGNITCKS  
NITGLLLVRDGGINVTNN--TGTEVFRPGGDMRDNWRNELYKYKVVVEIKPLGVAPTRAR  
RRVVEREKRAVG-LGAVFLGFLGAAGSTMGAASVTLTVQARQLLFGIVQQSNLLRAIEA  
QQRMLQLTVWGIKQLQTRVLAVERYLKDQQLLGMWGCSGKLICTTAVPWNSSWSNKTYND  
IWDNMTWLQWDKEISNYTDTIYRLLEESQNERNEKDLLELDKWASLWNWFNITNLWY  
IKIFIMIIGGLIGLRIIFAVLSIVNRVQGYSPLSFQTLTPNPRGPDRLGRIEIEEGGEQD  
KDRSIRLVNGFSALIWDDLRLNCLFSYHQLRDFILVTARAVELLGRSSIRGLQRGWEALK  
YLGSLVQYWGLELKKSAISLLDTTATAVANWTDRIEIVVQRAYRAILHIPRIRQGFEEA  
LQ-



Fig. 10 cont'd-13

>POL-B.syn1.1

FFRENLAFFPQKAREFSSEQTRANSPTRR-----ELQVWGRDNNSLSEAGA
DR----QGTVS-FSFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMNLPGRWKPK
MIGGIGGFIVRQYDQIPIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPISPIE
TVPVKLKPMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNRRTQDFWEVQLGIPHPAGLKKKKSVTVLDVGDAYFSVPLDKDFRK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGSFAIFQS SMTKILEPFRKQNPDIYIYQYMD
LYVGSdleIGQHRTKIEELRQHLLRWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPIVL
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVKQLCKLLRGTALTEVIPLTEEALELAEN
REILKEPVHGVYDPSKDLIAEIQKQGGQWTYQIYQEPFKNLKTGKYARMRGAHTNDVK
QLTEAVQKIATESIVIWGKTPKFKLPIQKETWETWTEYQATWIPEWEFVNTPPVLKLV
YQLEKEPIVGAETFYVDGAANRETCLGKAGYVDRGRQKVVSLDITTNQKTELQAIHLAL
QDSGLEVNIVTDSQYALGI IQAQPDKSESELVSQIIEQLIKKEKVYLAWVPAHKGIGNE
QVDKLVSAGIRKVLFLDGDIDKAQEEHEKYHSNWRAMASDFNLPPVVAKEIVASCDKCQLK
GEAMHGQVDCSPGIWQLDCTHLEGKIIILVAVHVASGYIEAEVI PAETGQETAYFLLKLAG
RWPVKTHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQSQGVVESMNKELKKIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDIIATDIQTKELQKQITKIQNFRVY
YRDSRDPLWKGPAKLLWKGEAVVIQDNSDIKVVPRRKAKIIRDYGKQ MAGDDCVASRQD
ED-

>POL-B.syn3.1

FFRENLAFFPQKAREFPSEQTRANSPTR-----ELQVWGGDNNSLSEAGD
DR----QGTVS-FSFPQITLWQRPIVTIKIGGQKKEALLDTGADDTVLEEMNLPGRWKPK
IIGGIGGFIVKQYDQILIEICGHKAIGTVLVGPTPANIIGRNLLTQIGCTLNFPISPIE
TVPVRLKPGMDGPKVKQWPLTEEKIKALIEICTELENEGKISKIGPENPYNTPVFAIKKK
DGTKWRKLVDFRELNRRTQDFWEVQLGIPHPAGLKKKKSVTVLDVGDAYFSVPLDKDFRK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGSFSIFQSSMTKILEPFRKQNPDIYIYQYMD
LYVGSdleIEQHRTKIEELRQHLLRWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPIVL
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVKQLCRLLRGTALTEVVPLTEEALELAEN
REILKEPVHGVYDPSKDLVAEIQKQGLGQWTYQIYQEPYKNLKTGKYAKMRGAHTNDVK
QLTEAVQKIATESIVIWGKTPKFKLPIQKETWEAWMEYQATWIPEWEFVNTPPVLKLV
YQLEKEPIVGAETFYVDGAANRDTKLGKAGYVDRGRQKVVSLDITTNQKTELQAIHLAL
QDSGLEVNIVTDSQYALGI IQAQPDRSESELVSQIIEQLIKKEKVYLAWVPAHKGIGNE
QVDKLVSSGIRKVLFLDGDIDKAQEDHEKYHSNWKAMASDFNLPPVVAKEIVACCDKCQLK
GEAMHGQVDCSPGIWQLDCTHLEGKIVILVAVHVASGYIEAEVI PAETGQETAYFILKLAG
RWPVTTHTDNGSNFTSTAVKAACWWAGIKQEFGI PYNPQSQGVVESINKELKKIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERMIDIATDIQTTELQKQITKLQNFRVY
FRDSRDPLWKGPAKLLWKGEAVVIQDNSEIKVVPRRKAKIIRDYGKQ MAGDDCVASRQD
ED-

>POL-B.syn3.2

FFREDLAFLOQKAREFSSEQTRANSPTRG-----ELQVWGRDNNSLSEAGA
DR----QGTVS-LSFPQITLWQRPLVTVKIGGQLKEALLDTGADDTVLEEMSLPGRWKPK
MIGGIGGFIVRQYDQILVEICGHKAIGTVLVGPTPVNIIGRDLLTQIGCTLNFPISPID
TVPVKLKPMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPIFAIAIKKK
DSTKWRKLVDFRELNRRTQDFWEVQLGIPHPAGLQKKSVTVLDVGDAYFSVPLDEDERK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGSFAIFQSSMTRILEPFRKQNPDLVIYQYMD
LYVGSdleIGQHRTKIEELREHLLRWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPIML
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVKQLCKLLRGAKALTEVIPLTKEALELAEN
REILKEPVHGAYYDPSKDLIAEIQKQGGQWTYQIYQDPFKNLKTGKYARMRGAHTNDVR
QLTEAVQKITTESIVIWGKIPKFKLPIQKETWETWTEYQATWIPEWEFVNTPPVLKLV
YQLEKEPIIGAETFYVDGAANRETCLGKAGYVTNKGROKVVSIITDTTNQKTELQAILLAL
QDSGLEVNIVTDSQYALGI IQAQPDKSESELVSQIIEELIKKEKVYLTWVPAHKGIGNE
QIDKLVSAGIRKVLFLDGDIDQAQEEHEKYHSNWRAMASDFNI PPVVAKEIVASCDKCQLK
GEAIIHGQVDCSPGIWQLDCTHLEGKIIILVAVHVASGYIEAEI IPTETGQETAYFLLKLAG
RWPVKTHTDNGSNFTSTTVKAACWWAGVQKQEFGI PYNPQSQGVVESMNNELKKIIGQIR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDIIATDIQTKELQKQITKIQNFRVY
YRDNRDPLWKGPAKLLWKGEAVVIQENS DIKVVPRRKVKIIRDYGKQ MAGDDCVASGQD
ED-



Fig. 10 cont'd-14

>POL-B.syn3.3

FFREDLAF PQGEAREFSSEQTRANSPTRR-----ELQVWGRDSNSLSEAGA
DR----QGTVS-FNEFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEDMNLP GKWKPK
MIGGIGGF IKVRQYDQIPIEICGHKAVGTVLVGPTFVNIIGRNLLTQLGCTLNFPISPIE
TVPVKLKP GMDGPKVKQWPLTEEKIKALVEICTELEKEGKISKIGPENPYNTPVF AIKKK
DSTKWRKVVD FRELNKKTQDFWEVQLGIPHP SGLKKKKS VTVLDVGDAYFSVPLDKDFRK
YTAFTIPSVNNETPGVRYQYNVLPQG WKGSPAIFQCSMTKILEPFRKQNP DIVIYQYMDD
LYVGS DLEIGQHRAKIEELRQHLLRWGFTTPDKKHQNEPPFLW MGYELHPDKWTVQPIEL
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVRQLCKLLRGTKALTEVIPLTEEA ELELAEN
REILREP VHG VYYDPTKDLIAEIQKQEQGQW TYQIYQEPFKNLKTGKYARTRGAHTNDVK
QLTEAVQKVATESI VIWGT PKFKLP IQKETWEAWWTEY WQATWIPEW EFWNT PPLVKLW
YQLEKEPIEGAETFYVDGASNRET KL GKAGYVTNRGRQKV VPLTDTTNQKTELQAIYLAL
QDSGSEVNI VTD SQYALGI IQAQPDKSESELVNQI IEQLIKKEKIYLA WVP AHKGIGGNE
QVDKLV SAGIRKILFLDGI DKAQEEHEKYHNNWRAMASDFNL PPIVAKEIVASCDKCQLK
GEA IHGQVDCSPGIWQLDCTHLEGKVILVAVHVASGYIEAEVI PAETGOETAYFILKLAG
RWPVKTIHTDNGSNFTSATVKAACWWAGIKQEFGI PYNPQSOGV VESMNKELKKIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGEYSAGERI IDIIATDIQTRELQKQITKI QNFRVY
YRDSRDPLWKGP AKLLWKGEGAVVIQDNSDIKV VPRRKAKI IRDYGKQ MAGDDCVAGRQD
ED-

>POL-B.syn4.1

FFRENLA FPQGEAREFSSEQNRANSPTRR-----ELQVWGGDNNSLSEAGA
DR----QGTVS-LSFPQITLWQRPLVTIRIGGQLKEALLDTGADDTVLEEMSLPGRWKPK
MIGGIGGF IKVRQYDQILIEICGHKAIGTVLIGPTPVNIIGRNLLTQLGCTLNFPISPID
TVPVKLKP GMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPIFAIKKK
DSTKWRKLVDFREJNRRTQDFWEVQLGIPHP SGLKKKKS VTVLDVGDAYFSVPLDKDFRK
YTAFTIP SINNETPGVRYQYNVLPQG WKGSPAIFQSSMTKILEPFRKQNPDMVIYQYMDD
LYVGS DLEIGQHRTKIEELRQHLLRWGLTTPDKKHQKEPPFLW MGYELHPDKWTVQPIKL
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVRQLCKLLRGAKALTEVIPLTAEAELELAEN
REILREP VHG VYYDPSKDLIAEIQKQGGQGW TYQIYQDPFKNLKTGKYAKMRGAHTNDVK
QLTEAVQKVATESI VIWGT PKFERLP IQKETWEAWWTEY WQATWIPEW EFWNT PPLVKLW
YQLEKEPIEGAETFYVDGAANRDTKL GKAGYVTD RGRQKV VSLTDTTNQKTELQAIHLAL
QDSGLEVNI VTD SQYALGI IQAQPDKSESELVSQI IEELIKKEKVYLA WVP AHKGIGGNE
QIDKLV SAGIRRVLF DGI DQAQEEHEKYHSNWRAMASDFNL PPIVAKEIVASCDKCQLK
GEA IHGQVDCSPGIWQLDCTHLEGKILLVAVHVASGYIEAEVI PAETGOETAYFLLKLAG
RWPVKTIHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQSOGV VESMNKELKKIIEQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGEYSAGERI IDIIATDIQTRELQKQITKI QNFRVY
YRDNRDPLWKGP AKLLWKGEGAVVIQDNSDIKV VPRRKAKI IRDYGKQ MAGDDCVAGRQD
ED-

>POL-B.syn4.2

FFRENLA FPQ GKAREFPSEQTRANSPTSR-----ELQVWGRDNNSLSEAGD
DR----CGTVS-FSFPQITLWQRPLVTVKIGGQLKEALLDTGADDTVLEEMNLPGRWKPK
MIGGIGGF IKVRQYDQIPIEICGHKAVGTVLVGPTFVNIIGRDLLTQIGCTLNFPISPIE
TVPVKLKP GMDGPKVKQWPLTEEKIKALIEICTELENEGKISKIGPENPYNTPVF AIKKK
DGTKWRKLVDFRELNKRTQDFWEVQLGIPHPAGLQKKS VTVLDVGDAYFSVPLDKEFRK
YTAFTIPSTNNETPGIRYQYNVLPQG WKGSPAIFQCSMTKILEPFRKQNPDLVIYQYMDD
LYVGS DLEIEQHRTKIEELREHLLKWGFTTPDKKHQNEPPFLW MGYELHPDKWTVQPIML
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVKQLCKLLRGTKALTEVVPLTEEA ELELAEN
REILKVPVHG VYYDPSKDLVAEIQKQGLGQW TYQIYQEPFKNLKTGKYARTRGAHTNDVR
QLTEAVQKIATESI VIWGT PKFKLP IQKETWEAWWMEY WQATWIPEW EFWNT PPLVKLW
YQLEKEPIVGAETFYVDGAANRET KL GKAGYVTD KGRQKV VPLTDTTNQKTELQAINLAL
QDSGSEVNI VTD SQYAIGI IQAQPDRSESELVSQI IEQLINKEKVYLA WVP AHKGIGGNE
QVDKLV SSGIRKVLFLDGI DKAQEDHEKYHSNWRAMAGDFNL PPVVAKEIVACCDKCQLK
GEA MHGQVDCSPGIWQLDCTHLEGKVILVAVHVASGYIEAEVI PAETGOETAYFILKLAG
RWPVKTVHTDNGSNFISTTVKAACWWAGVKQEFGI PYNPQSOGV VESMNNELKKIIGQVR
DQAEHLKTAVQMAVVFVHNFKRKGGIGGYTAGERI VDIIASDIQTKELQKQITKI QNFRVY



Fig. 10 cont'd-15

YRDSRDPLWKGPakllWkGEGAVVIQDNSEIKVVPRRKAKIIRDYgkQmAGDDCVASRQn  
ED-

>POL-B.syn4.3

FFREDLAFLOgkAREFSSEQTRANSPTRR-----ELQVWGRDNNSPSEAGA  
DR----QGTVS-FNFPQITLWQRPIVVIKIGGQLKEALLDTGADDTVLEDMNLPgkWKPK  
MIGGIGGFikVRQYDQILVEICGHKAIGTVLVGPTPANIIGRNLLTQIGCTLNFPISPIE  
TVPVklKSGMDGPKVKQWPLTEEKIKALIEICTEMEKEGKISKIGPENPYNTPVFAIKKK  
DSTKWRklVDFRELNKRtQDFWEVQLGIPHPSGLKKKKSvtVLDVGDAYFSVPLDEDFRK  
YTAFTIpsVNNETPGIRYQYNVLPQGWKGS PAIFQSSMTRILEPFRKQNPDI VIYQYMDD  
LYVGSdleIGQhRAKIEELRQHLLRWGFTTPDKKHQKEPFLWmGYELHPDKWTVQPIEL  
PEKDSWTVNDIQklVGKLNWASQIYPGIKVRQLCKLLRGTKALTEVIPLTEEALELAEN  
REILKEPvHGvYYDPSKELIAEIQKQEQGQWTYQIYQEPFKNLKTGKYARMRGTHTNDVK  
QLTEAVQKITTESIViWGRTPKFKLPIQKETWESWTEYwQATWIPewEFVNTpPLVklW  
YQLEREPIAGAETfYVDGASNRETKLgKAGYVTNRGRQKVVS LpDTTNQKTELQAIYLAL  
QDSGLEVNIvTDSQYAIGIIQAQPDkSESELVNOIIEQLIKKEKIYLAWVPAHKGIGGNE  
QVDKLVsNGIRKI LFLDGIDKAQDEHEKYHSNWKAMASDFNLPPVAKEIVACCDKcQLK  
GEAMHGQVDCSPGIWQLDCTHLEGKIILVAVHVASGYIEAEVI PAETGQETAYFLLKLAG  
RWPVKIiHTDNGSNFTSTAVKAACWWAGIKQEFGI PYNPQSQGVVESINKELKKI IKQVR  
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERMIDI IATDIQTTELQKQITKLQNFVY  
FRDSRDPLWKGPakllWkGEGAVVIQDNNDIKVVPRRKVKIIRDYgkQmAGDDCVASGQD  
ED-

>POL-B.syn4.4

FFREDLAFPOgkARELSSEQTRANSPTRG-----ELQVWGRDSNSLSEAGA  
DR----PGTVS-FSFPQITLWQRPLVTIKIGGQKEALLDTGADDTVLEEINLPGRWKPK  
IIGGIGGFikVKQYDQIPIEICGHKVIgTVLVGPTPANIIGRNLLTQIGCTLNFPISPIE  
TVPVRLKPGMDGPKVKQWPLTEEKIKALVEICTELEKEGKISKIGPENPYNTPVFAIKKK  
DSTKWRkVVDFRELNKKtQDFWEVQLGIPHAGLKKKKSvtVLDVGDAYFSVPLDENFRK  
YTAFTIpsINNETPGIRYQYNVLPQGWKGSPSIFQSSMTKILEPFRKQNP EIVIYQYMDD  
LYVGSdleLGQhRTKIEELRQHLLKwGFYTPDKKHQKEPFLWmGYELHPDKWTVQPIVL  
PEKDSWTVNDIQklVGKLNWASQIYPGIKVKQLCRLLRGAKALTEVIPLTKEAELELAEN  
REILKEPvHGAYDPTKDLIAEIQKQEGEQWTYQIYQEPYKNLKTGKYARMRGAHTNDVK  
QLTETVQKITTESIViWGKIPKFKLPIQKETWETWWTEYwQATWIPewEFVNTpPLVklW  
YQLEKEPIIGAETfYVDGAASRETKLgKAGYVTNKGRQKVVSITD TTNQKTELQAILLAL  
QDSGLEVNIvTDSQYAIIGIIQAQPDkSESEIVSQIIEQLIKKEKVYLTWVPAHKGIGGNE  
QVDKLVsAGIRKVLFLDGIDKAQEEHEKYHNNWRAMASDFNI PPVAKEIVASCDKcQLK  
GEAMHGQVDCSPGIWQLDCTHLEGKVI L VAVHVASGYIEAEI IPTETGQETAYFILKLAG  
RWPVTTIHTDNGSNFTSATVKAACWWAGVKQEFGI PYNPQSQGVIESMNKELKKI IGQIR  
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDI IATDIQTKELQNCITKIQNFVY  
YRDSRDPLWKGPakllWkGEGAVVIQENS DIKVVPRRKVKIIRDYgkQmAGDDCVASRQD  
ED-

>POL-B.syn6.1

FFREDLAFPOGAEAREFCSEQTRANSPATR-----ELQVWGRDNTSLSEAGA  
DR----PGTVS-FSFPQITLWQRPIVTVKIEGQLKEALLDTGADDTVLEEMNLPgkWKPK  
MIGGIGGFikVRQYDQVSIEICGHKAIGTVLVGPTPANIIGRNLLTQIGCTLNFPISPIE  
TVPVklKPGMDGPKVKQWPLTEEKIKALVEICTELEKEGKISKIGPENPYNTPVFAIKKK  
DSTKWRkVVDFRELNKRtQDFWEVQLGIPHPSGLKKKKSvtVLDVGDAYFSVPLDENFRK  
YTAFTIpsINNETPGIRYQYNVLPQGWKGS PAIFQSSMTKILEPFRKQNPDI IYQYMDD  
LYVGSdleIGQhRAKIEELRQHLLKwGFYTPDKKHQKEPFLWmGYELHPDKWTVQPIEL  
PEKDSWTVNDIQklVGKLNWASQIYAGIKVKELCKLLRGTKALTEVVPLTEEALELAEN  
REILKEPvHGvYYDPSKDLIAELQKQGGQWTYQIYQEPYKNLKTGKYARTRGAHTNDVR  
QLTEAVQKIATEGIViWGKTPKFKLPIQKETWEAWTEYwQATWIPewEFVNTpPLVklW  
YQLEKEPIlGAETfYVDGASNRETKLgKAGYVTDGRGRQKVVS LpDTTNQKTELQAINLAL  
QDSGLEVNIvTDSQYALGIIQAQPDkRSESELVSQIIEQLINKEKVYLA WVPAHKGIGGNE  
QVDKLVSTGIRRVFLDGIDKAQEEHEKYHSNWRAMASDFNLPPIVAKEIVASCDKcQLK  
GEAIHGQVDCSPGIWQLDCTHLEGKVI L VAVHVASGYIEAEVI PAETGQETAYFILKLAG



Fig. 10 cont'd-16

RWPVKTIHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQS QGVV ESMNKELKKIIEQVR  
DQAEHLKTAVQMAV FVHNFKRKG GIGEYSAGERI VDI IATDI QTKELQKHITKI QNFRVY  
YRDSRDELWKGPAKLLWKGE GAVVIQDNSEIKV VPRRKAKI IRDYGKQ MAGDDCVASRQD  
ED-

>POL-B.syn6.2

FFREDLAFPQ GKARELSSEQTRANSPTS PTRG-----ELQVWGRDSNSLSEAGA  
DR----QGPVS--FSFPQITLWQRPIVTIKIGGQLKEALLDTGADDTVLEDMNLPGRWKPK  
MIGGIGGFIVKQYDEILVEICGHKAIGTVLIGPTPVNIIGRNLLTQLGCTLNFPISPIE  
TVPVKLKSMDGPKVKQWPLTEEKIKALVEICTELEKEGKISKIGPENPYNTPIFAIAKKK  
DSTKWRKLVDFRELNRKTQDFWEVQLGIPHPAGLKKKSVTVLDVGDAYFSVPLDKDFRK  
YTAFTI P SVNNETPGIRYQYNVLPQGWKGS PAIFQCSMTKILEPFRKQNPDMVIYQYMDD  
LYVGS DLEIGQHRIKIEELREHLLKWF TTPDKKHQNEPPFLWMGYELHPDKWTVQPIVL  
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVRQLCKLLRGTKALTEVIPLTEEALELAEN  
REILREP VHG VYYDPTKDLIAEIQKQGQWTYQIYQEPFKNLKTGKYARMRGAHTNDVK  
QLTEAVQKITTESIVIWGKIPKFRLP IQKETWEAWWIEYWQATWIPEWEFVNT PPLVKLW  
YQLEREPIAGAE TFYVDGAANRETKLGKAGYVTNRGRQKVSITDTTNQKTELQAILLAL  
QDSGLEVNIVTDSQYALGI IQAQPDKSESELVNIIEQLIKKEKIYLA WVP AHKGIGGNE  
QIDKLV SAGIRKVLFLDGI DKAQDEHEKYHSNWRAMAGDFNLPPVVAKEIVACCDKCQLK  
GEAMHGQVDCSPGIWQLDCTHLEGKII LVAHVHVASGYIEAEVIPAETGQETAYFLLKLAG  
RWPVTTIHTDNGSNFTSATVKAACWWAGVKQEFGI PYNPQS QGVV ESMNKELKKIIEQVR  
DQAEHLKTAVQMAVFIHNFKRKG GIGEYSAGERI IIDI IATDI QTKELQKHITKI QNFRVY  
YRDSRDP IWKGP AKLLWKGE GAVVIQDNSEIKV VPRRKAKI IRDYGKQ MAGDDCVAGRQD  
ED-

>POL-B.syn6.3

FFRENLAFFPQGEAREFSSEQTRANSPTRG-----ELQVWGRDSNSLSEAGD  
DR----QGTVS--FSFPQITLWQRPLVTIKIGGQKKEALLDTGADDTVLEEMNLPGRWKPK  
IIGGIGGFIVKQYDQIPIEICGHKAVGTVLVGPTPVNIIGRDLLTQIGCTLNFPISPIE  
TVPVKLKSMDGPKVKQWPLTEEKIKALIEICTEMEKEGKISKIGPENPYNTPVFAIAKKK  
DGTKWRKLVDFRELNRKTQDFWEVQLGIPHPAGLKKKSVTVLDVGDAYFSVPLDREFRK  
YTAFTI PSLNNETPGIRYQYNVLPQGWKGS P SIFQSSMTKILEPFRKQNPDLVIYQYMDD  
LYVGS DLELGQHRTKIEELRQHLLKWFYTPDKKHQKEPPFLWMGYELHPDKWTVQPIKL  
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVKQLCRLLRGAKALTEVVPLTKEAELELAEN  
REILKEPVH GAYYDPTKDLIAEVQKQELGQWTYQIYQEPFKNLKTGKYARMKGAHTNDVK  
QLTETVQKITTESIVIWGKTPKFRLP IQKETWESWWT EYWQATWIPEWEFVNT PPLVKLW  
YQLEKEPI TGAETFYVDGAANRETKIGKAGYVTDKGRQKVVSLPDTTNQKTELQAIHLAL  
QDSGSEVNIVTDSQYAIGI IQAQPDRSESEVNIIEQLIKKEKVYLA WVP AHKGIGGNE  
QVDKLV SNGIRKILFLDGI DKAQEEH ERYHSNWKAMASDFNLPPVVAKEIVACCDKCQLK  
GEA IHGQVDCSPGIWQLDCTHLEGKVI LVAHVHVASGYIEAEVIPAETGQETAYFILKLAG  
RWPVKI IHTDNGSNFTSTAVKAACWWAGIKQEFGI PYNPQS QGVV ESMNKELKKIIEQVR  
DQAEHLKTAVQMAVFIHNFKRKG GIGYSAGERI IIDI IASDI QTKELQKHITKI QNFRVY  
YRDSRDP VWKGP AKLLWKGE GAVVIQDNSEIKV VPRRKAKI IRDYGKQ MAGDDCVASRQD  
ED-

>POL-B.syn6.4

FFRENLAFFPQRKAREFSSEQTRANSPTRR-----ELQVWGGDNNSLSEAGA  
DR----QGTVS--LSFPQITLWQRPLVTIKVGGQLKEALLDTGADDTVLEEINLPGRWKPK  
MIGGIGGFIVRQYDQILVEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPISPIE  
TVPVKLKP GMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPIFAIAKKK  
DSTKWRKLVDFRELNRRTQDFWEVQLGIPHPAGLKKKSVTVLDVGDAYFSVPLDEDFRK  
YTAFTI P S INNETPGVRYQYNVLPQGWKGS PAIFQASMTKILEPFRKQNPDI VIYQYMDD  
LYVGS DLEIGQHRIKIEELRQHLLRWGLTTPDKKHQKEPPFLWMGYELHPDKWTVQPIML  
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVRQLCKLLRGAKALTEVIPLTAEAELELAEN  
REILKVPVHG VYYDFSKELIAEIQKQEQGQWTYQIYQDPFKNLKTGKYARMRGHTNDVR  
QLTEAVQKITTESIVIWGKIPKFKLP IQKETWETWWT EYWQATWIPEWEFVNT PPLVKLW  
YQLEKEPI IGAETFYVDGAASRETKLGKAGYVTDGRQKVISLTDTTNQKTELQAIHLAL  
QDSGVEVNIVTDSQYALGI IQAQPDKSESEIVSQIIEQLIKKEKVYLTWVPAHKGIGGNE



Fig. 10 cont'd-17

QVDKLVSTGIRKVLFLDGDQAQEEHEKYHSNWRMTMASDFNLPPIVAKEIVASCDKCQLK
GEAMHGQVDCSPGIWQLDCTHLEGKVILVAVHVASGYIEAEVIPAETGQETAYFLLKLAG
RWPVKTHTDNGPNEISTTVKAACWWAGIKQEFGI PYNPQSOGVVE SMNRELKKIIGQVR
EQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERMIDI IATDIQTTELQKQITKIQNFRVY
FRDSRDPLWKGP AKLLWKGEGAVVIQENSDIKV VPRRKAKI IRDYGKQ MAGDDCVAGRQD
ED-

>POL-B.syn6.5

FFRENLAFFPQ GKAREFPSEQTRANSPTS R-----ELQVWGRDNNSLSEAGA
NR----QGTVS-FSFPQITLWQRPLVTVKIGGQLKEALLDTGADDTVLEMDLPGRWKPK
MIGGIGGFIVRQYDQIPIEICGHKVI GTVLVGP TPANI IGRNLLTQIGCTLNFPISP I E
TVPVRLKPGMDGPKVKQWPLTEEKIKALVEICTELEKEGKISKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNKKTQDFWEVQLGIPHPSGLK KKKSVTVLDVGDAYFSVPLDKEFRK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGS PAIFQASMTKILEPFRKQNP EIVYQYMDD
LYVGS DLEIEQHRTKIEELRQHLLRWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPIVL
PDKDSWTVNDIQKLVGKLNWASQIYPGIKIRQLCKLLRGAKALTEVIPLTKEAELELAEN
REILKEPVHGVYDPSKDLIAEIQKQGGQWTYQIYQEPFKNLKTGKYAKMRGAHTNDVK
QLTEAVQKVATESIVIWGKTPKFKLPIQKETWEAWWMEYWQATWIPEWEFVNT PPLVKLW
YQLEKEPIEGAETFYVDGAANRDTKLGKAGYVTKGRQKVVTLTDTTNQKTELQAIHLAL
QDSGLEVNIVTDSQYALGIIQAQPKSESEIVNQIIEQLIKKEKVYLAWVPAHKGIGGNE
QVDKLVSSGIRKVLFLDGDIDKAQEDHEKYHSNWRAMANDENLPPVVAKEIVACCDKCQLK
GEAMHGQVDCSPGIWQLDCTHLEGKVILVAVHVASGYIEAEVIPAETGQETAYFILKLAG
RWPVKT VHTDNGSNFTSNTVKAACWWAGIKQEFGI PYNPQSOGVVE SMNKQLKQIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDI IATDIQTRELQKQITKIQNFRVY
YRDSREPLWKGP AKLLWKGEGAVVIQDNSDIK VVPRRKAKI IRDYGKQ MAGDDCVASGQD
ED-

>POL-B.syn6.6

FFREDLAFLQ GKAREFSSEQTRAISPTRR-----ELQVWGRDNNSPSEAGA
DR----QGTVS-FNFPQITLWQRPLVTIRIGGQLKEALLDTGADDTVLEEMSLPGRWKPK
MIGGIGGFIVRQYDQILIEICGHKAVGTVLIGPTPVNI IGRNLLTQIGCTLNFPISP ID
TVPVKLPGMDGPKVKQWPLTEEKIKALIEICTELENEGKISKIGPENPYNTPIFAIKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPSGLK KKKSVTVLDVGDAYFSIPLDEDFRK
YTAFTIPSINNETPGTRYQYNVLPQGWKGS PAIFQS SMTRILEPFRKQNP DIVIYQYVDD
LYVGS DLEIGQHRTKIEELREHLLRWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPI TL
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVKQLCKLLRGTKSLTEVVPLTAEAELELAEN
REILKEPVHGAYDPSKDLVAEIQKQGLGQWTYQIYQEPFKNLKTGKYAKMRGTHTNDVK
QLTEAVQKIATESIVIWGRTPKFKLPIQKETWDAAWTEYWQATWIPEWEFVNT PPLVKLW
YQLEKEPIVGAETFYVDGAANRETRLGKAGYVTDGRQKVVPLTDTTNQKTELQAIYLAL
QDSGLEVNIVTDSQYALGIIQAQPKSESELVSQIIEELIKKEKVYLAWVPAHKGIGGNE
QVDKLVSAGIRRVLFLDGDIDKAQEEHEKYHNNWRAMASDFNI PPVVAKEIVASCDKCQLK
GEA IHGQVDCSPGIWQLDCTHLEGKIIILVAVHVASGYIEAEI IPTETGQETAYFLLKLAG
RWPVKTHTDNGRNFTSNSVKAACWWAGIKQEFGI PYNPQSOGVVE SMNRELKKIIGQIR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDI IASDIQTKELQKQITKIQNFRVY
YRDNRDPLWKGP AKLLWKGEGAVVIQDNNDIK VVPRRKVKI IRDYGKQ MAGDDCVASRQD
ED-

>POL-C.syn1.1

FFRENLAFFPQGEAREFPSEQTRANSPTS RANSPTS R-----ELQV--RGDNPRSEAGA
ER----QGT---LNFPQITLWQRPLVSIKVGGOIKEALLDTGADDTVLEEINLPGKWKPK
MIGGIGGFIVRQYDQILIEICGKKAIGTVLVGP TPVNI IGRNMLTQLGCTLNFPISP I E
TVPVKLPGMDGPKVKQWPLTEEKIKALTAICEEMEKEGKITKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHAGLKK KKKSVTVLDVGDAYFSVPLDEGFRK
YTAFTIPSINNETPGIRYQYNVLPQGWKGS PAIFQSSMTKILEPFRAQNP EIVYQYMDD
LYVGS DLEIGQHRAKIEELREHLLKWGFTTPDKKHQKEPPFLWMGYELHPDKWTVQPIQL
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVRQLCKLLRGAKALTDIVPLTEEALELAEN
REILKEPVHGVYDPSKDLIAEIQKQGHQWTYQIYQEPFKNLKTGKYAKMRTAHTNDVK



Fig. 10 cont'd-18

QLTEAVQKIAMESIIVIWGKTPKFRLLPIQKETWETWWTDYWQATWIPWEFVNTPPPLVKLW
YOLEKEPIAGAETFYVDGAANRETKIGKAGYVTDGRGRQKIVSLTETTNOKTELQAIQLAL
QDSGSEVNIIVTDSQYALGIIQAQPKSESELVNQIIEQLIKKERVYLSWVPAHKGIGGNE
QVDKLVSSGIRKVLFLDGDIDKAQEEHEKYHSNWRAMASEFNLPPIVAKEIVASCDKQCQLK
GEAIHGQVDCSPGIWQLDCTHLEGKIILVAVHVASGYIEAEVI PAETGQETAYYILKLAG
RWPVKVIHTDNGSNFTSAAVKAACWWAGIQQEFGIPYNPQSOGVVE SMNKELKKIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIIDIIATDIQTKELQKQITKIQNFRVY
YRDSRDPIWKGPAKLLWKGE GAVVIQDNSDIKVVPRRKVKI IKDYGKQ MAGADCVAGRQD
EDQ

>POL-C.syn3.1

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ER----QGT---FNFPQITLWQRPLVTIKVGGQVKEALLDTGADDTVLEEINLPGKWKPK
MIGGIGGFIVRQYDQIVIEICGKKAIGTVLIGTPVNIIGRNMLTQLGCTLNFPISPIE
TVPVKLPGMDGPKIKQWPLTEEKIKALTAICDEMEKEGKIEKIGPENPYNTPIFAIKKK
DSTKWRKLVDFKELNKRTQDFWEVQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDESFRK
YTAFTIPSINNETPGIRYQYNVLPQGWKGSFAIFQSSMTRILEPFRAKNPEIIVYQYMDD
LYIGSDLEIGQHRAKVEELREHLLRWGFTTPDKKHQKEPPFLWMGYELHDPKWTVQPIQL
PEKDSWTVNDIQRLVGLNWSAQIYPGIKVRQLCKLLRGTALTDIVPLTEEALELAEN
REILKEPVHGVYDPSKDLIAEIQKQGGDQWTYQIYQESFKNLKTGKYAKMRSHTNDVK
QLTEAVQKIALESIIVIWGKAPKFRLLPIQKETWEIWWTDYWQATWIPDWEFVNTPPPLVKLW
YOLEKEPIAGAETFYVDGAANRETKIGKAGYVTDGRGRQKVVTLTETTNOKTELQAIQLAL
QDSGLEVNIIVTDSQYALGIIQAQPKSESELVNQIIEELIKKERVYLSWVPAHKGIGENE
QVDKLVSSNGIRKVLFLDGDIDKAQEEHEKYHSNWRAMANEFNLPVVAKEIVASCDKQCQLK
GEAIHGQVDCSPGMWQLDCTHLEGKIVLAVHVASGYVEAEVI PAETGQETAYFILKLAG
RWPVKI IHTDNGSNFTSNAVKAACWWAGIQQEFGIPYNPQSOGVVE SMNKELKKIIGQVR
EQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIIDIIASDIQTKELQKQITKIQNFRVY
YRDSRDPVWKGPAKLLWKGE GAVVIQDNGDIKVVPRRKAKI IKDYGKQ MAGDDCVAGRQD
EDQ

>POL-C.syn3.2

FFRENLAFOQGEAREFPSEQTRANSPTSRSNSPTSRTNSPTSRELQV--RGDNPRSEAGV
ER----QGT---LNFPQITLWQRPLVSIKVGGOIKEALLDTGADDTVLEDINLPGKWKPR
MIGGIGGFIVRQYDQIPIEICGKKAIGTVLVGTPVNIIRRNMLTQLRCTLNFPISPIK
TVPVKLPGMDGPKVKQWPLTEEKIKALTEICEEMEKEGKITKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPGSLKKKKSVTVLDVEDAYFSVPLDEGFRK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGSFAIFQCSMTKILEPFRTQNPDIVIYQYMDD
LYVGSLEIGQHRAKIEELRAHLLKWGLTTPDKKHQKEPPFLWMGYELHDPKWTVQPIKL
PEKDSWTVNDIQKIVGKLNWSAQIYPGIKVKQLCKLLRGAKALTDIIPLTEEALELAEN
REILKEPVHGAYDPSKDLVAEIQKQGHQWTYQIYQEPYKNLKTGKYAKMRTAHTNDVR
QLTEAVQKIAQESIIVIWGKTPKFRLLPIQKETWETWWTDYWQATWIPWEFINTPPPLVKLW
YOLEKEPIVGAETFYVDGAANRETKMGKAGYVTDKGRQKIVSLTETTNOKTELQAIQLAL
QDSGPEVNIIVTDSQYALGIIQAQPKDRSESELVNQIIEQLINKERIYLSWVPAHKGIGGNE
QVDKLVSSGIRKVLFLDGDIDKAQEDHEKYHNNWRAMASDFNLPPIVAREIVASCDKQCQLK
GEAMHGQVDCSPGVWQLDCTHLEGKIILVAVHVASGYMEAEVI PAETGQETAYYILKLAG
RWPVKVIHTDNGSNFTSTAVKAACWWAGIKQEFGIPYNPQSOGVVEAMNKELKKIIEQVR
DQAEHLKTAVQMAVLIHNFKRKGGIGGYSAGERIVDIIATDIQTRRELQKQIIQIQNFRVY
YRDSRDPVWKGPAKLLWKGE GAVVIQDNSDIKVVPRRKVKI IKDYGKQ MAGADCVASRQD
ED-

>POL-C.syn3.3

FFRENLAFPQKAREFPSEQARANSPTSRSNSPTS-----ELQV--RRDNPRSEAGA
ER----QGT---LNCQITLWQRPLVSIKIGGQTRREALDTGADDTVLEEISLPGKWKPK
MIGGIGGFIVRQYDQILIEICGKKAIGSVLVGTPVNIIGRNLLTQLGCTLNFPISPIE
TIPVKLPGMDGPRVKQWPLTEEKIKALTAICEEMEKEGKISKIGPENPYNTPIFAIKKK
DSTKWRKLVDFRELNKRTQDFWEIQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDEDFRK
YTAFTIPSINNATPGIRYQYNVLPQGWKGSPSIFQSSMTKILEPFRAQNPEIIVYQYMDD
LYVGSLEIEQHRAKIEELREHLLKWGFTTPDKKHQKEPPFLWMGCELHDPKWTVQPIQL



Fig. 10 cont'd-19

PEKESWTVNDIQKLVGKLNWASQIYAGIKVKQLCRLLRGAKALTDIVPLTAEAELELAEN
REILREFVHGVYYDPSKELIAEIQKQGQDQWTYQIYQEPFKNLKTGKYAKRRTAHTNDVK
QLAEAVQKIAMESIVIWGKIPKFRLP IQKETWEAWWTDYWQATWIPWEFVNT PPLVKLW
YQLEKDP IAGVETFYVDGAANRET KLGKAGYVTD RGRQKIVSLSETTNQKTELHAIQLAL
QDSGSEVNIIVTDSQYALRI IQAQPKSESEIVNQIIEQLIKKERVYLA WVP AHKGI GENE
QVDKLVSKGIRKVLFLD GIEKAQEEHERYHSNWRAMASEFNLPPIVAKEIVASCDKCQLK
GEATHGQVDCSPGIWQLDCTHLEGKVIIVAVHVASGYIEAEVI PAETGQETAYFLKLAG
RWPVKTIHTDNGSNFTSAAVKAACWWAGIHQEFGI PYNPQSOGVVE SMNKELKKIIGQVR
DQAEHLKTAVLMAVFIHNFKRKGGIGDYSAGERIIDIIATDIQTKELQKQI I KIQNERVY
YRDNRPDIWKGP AKLLWKGEGAVVLQDNSDIKVI PRRKAKIIRDY GKQ MAGADCVAGRQD
ENQ

>POL-C.syn4.1

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ER----QGT---LNL P QITLWQRPLVSIKVG GQIKEALLDTGADDTVLEDINLPGKWKPK
MIGGIGGF I KVRQYDQIPIEICGKKAIGTVLVGPTPVNII GRNMLTQLGCTLNFPISPIE
TIPVKLKPGMDGPKVKQWPLTEEKIKALTEICKEMEKEGKIEKIGPENPYNTPVFALKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPSGLKKKKS VTVL DVEDAYFSVPLDENFRK
YTAFTIPSVNNETPGIRYQYNVLPQGWKGS PAIFQC SMTKILEPFRTQNP EIVIYQYMDD
LYVGS DLEIGQHRAKIEKLRHLLKWGFTTPDKKHQKEPPFLW MGYELHPDKWTVQPIKL
PEKDSWTVNDIQRLV GKLNWASQIYAGIKVRQLCKLLRGAKALTDIVPLTKEAELELAEN
REILKEPFVHGVYYDPSKELIAEIQKQGQDQWTYQIYQEPFKNLKTGKYAKRRTAHTNDVK
QLAEAVQKITMESIVIWGRTPKFRLP IQKETWEAWWTDYWQATWIPWEFVNT PPLVKLW
YQLEKEPIAEAEETFYVDGAANRET KMGKAGYVTD KGRQKIVSLTETT NQKTELHAIQLAL
QDSGPEVNIIVTDSQYALGI IQAQPKSESESELVSQIIEQLINKERIYLSWVPAHKGI GNE
QVDKLVSKGIRKVLFLD GIDKAQEEHERYHSNWRAMANEFNLPPIVAREIVASCDKCQLK
GEATHGQVDCSPGVWQLDCTHLEGKIIIVAVHVASGYVEAEVI PAETGQETAYFLKLAG
RWPVKTIHTDNGSNFTSAAVKAACWWAGVQQEFGI PYNPQSOGVVE SMNKELKKIIGQIR
DQAEHLKTAVQMAVFIHNFKRKGGIGDYSAGERIIDIIATDIQTRELOKQI I QIQNERVY
YRDSRDPVWKGP AKLLWKGEGAVVLQDNSDIKVVPRRKVKI IKDY GKQ MAGADCVAGRQD
ENQ

>POL-C.syn4.2

FFRENLAFFPEGEAREFPSEQTRANSPT-RANSPTS R-----KLQV--RGDNPRSEAGV
ER----QGT---LNF P QITLWQRPLVSIKIGGQTR EALLDTGADDTVLEEIKLPGNWKPK
MIGGIGGF I KVRQYDQILIEICGKKAIGTVLIGPTPVNII GRNLLTQLGCTLNFPISPIK
TVPVKLKPGMDGPKVKQWPLSEEKIKALTEICEEMEKEGKISKIGPENPYNTPVFALKKK
DSTKWRKLVDFRELNKRTQDFWEIQLGIPHAGLKKKKS VTVL DVGDAYFSVPLDEDFRK
YTAFTIP S INNATPGIRYQYNVLPQGWKGS PSIFQSSMTKILEPFRAKNPEIVIYQYMDD
LYVGS DLEIGQHRAKVEELREHLLKWGLTTPDKKHQKEPPFLW MGYELHPDKWTVQPIQL
PDKDSWTVNDIQKLV GKLNWASQIYPGIKVKQLCKLLRGTKALTDIVPLTAEAELELAEN
REILREFVHGVYYDPSKDLVAEIQKQGNDQWTYQIYQEPYKNLKTGKYAKMRS AHTNDVK
QLTEAVQKIALESIVIWGKAPKFRLP IQKETWEIWWTDYWQATWIPDWEFVNT PPLVKLW
YQLEKDP IAGVETFYVDGAANRET KLGKAGYVTD RGRQKIVSLSETTNQKTELQAIQLAL
QDSGLEVNIIVTDSQYALGI IQAQPKSESESELVNQIIEELIKKEKVYLSWVPAHKGI GENE
QVDKLVSSGIRKVLFLD GIEKAQEEHEKYHNNWRAMASEFNLPPIVAKEIVASCDKCQLK
GEATHGQVDCSPGMWQLDCTHLEGKIIIVAVHVASGYLEAEVI PAETGQDTAYYILKLAG
RWPVKVIHTDNGTNFTSAAVKAACWWAGIQQEFGI PYNPQSOGVVE SMNKELKKIIGQVR
EQAEHLKTAVLMAVFIHNFKRKGGIGEYSAGERIIDMIATDIQTKELQNQITKIQNERVY
YRDSRDPDIWKGP AKLLWKGEGAVVIQDNGDIKVVPRRKVKIIRDY GKQ MAGDDCVAGRQD
EDQ

>POL-C.syn4.3

FFRENLAFFPQGEAREFPPEQTRANSPTSRTNSPTS R-----ELQV--RGDNPHSEAGA
ERQGTLOGT---LNC P QITLWQRPLVSI R VGGQIKEALLDTGADDTVLEEISLPGKWKPK
MIGGIGGF I KVRQYDQIVIEICGKKAIGSVLVGPTPVNII RRNMLTQLRCTLNFPIS SIE
TVPVKLKPGMDGPRVKQWPLTEEKIKALTAICEEMEKEGKITKIGPDNPNYNTPVFALKKK



Fig. 10 cont'd-20

DSTKWRKLVDFKELNKRRTQDFWEVQLGIPHPAGLKRKKSVTVLDVGDAYFSVPLDEGFRK
YTAFTIPS'TNNETPGIRYQYNVLPQGWKGSFAIFQSSMIKILEPFRAQNPDIYIYQYMDD
LYIGSDLEIEQHRAKIEELRAHLLKWGFTT'PDKKHQKEPPFLWMGCELHPDKWTVQPIQL
PEKESWTVNDIQRLVGKLNWASQIYSGIKVRQLCKLLRGVKALTDIVPLTEEALELELAEN
REILKETVHGAYYDPSKDLIAEIQKQGHQDQWYQIYQEPFKNLKTGKYAKMRTAHTNDIK
QLTEAVQKIAMESIVIWGKTPKFRLLPIQKETWETWWTWTDYQATWIPEWEIFINTPPLVKLW
YQLEKEPIVGAETFYVDGAANRDTKLGKAGYITDRGRQKVVTLTETTNQKAEIQAIQLAL
QDSGSKVNIIVTDSQYALGIIQAQPDSESELVNCIIEQLIKKERVYLSWVPAHKGIGGNE
QIDKLVSSGIRRVFLDGDIDKAQEDHEKYHSNWRAMASDFNLPPIVAKEIIASCDKCQK
GEAMHGQVDCSPGIWQLDCTHLEGKVIIVAVHVASGYIEAEVI PAETGQETAYYILKLAG
RWPVKI IHTDNGSNFTSNAVKAACWWAGIHQEFGIPYNPQSQGVVEAMNKELKKIIGQVR
DQAEHLKTAVLMAVFIHNFKRGGGIGGYSAGERIIDIIASDIQTKELQKQITKIQNFRVY
YRDSRDFIWKGPAKLLWKGEGAVVIQDNSDIKVI PRRKAKI IKDYGKQ MAGADCVAGGQD
EN-

>POL-C.syn4.4

FFRENLAFFQGEAREFPSEQTRAISPTSR-----ELQV--RGDNPCEAGA
ER----QGT---FNFPQITLWQRPLVTIKVGGQVKEALLDTGADDTVLEEINLPGKWKPR
MIGGIGGFIVRQYEQILIEICGKRAIGTVLVGPTPINIIGRNMLTQIGCTLNFPISSIE
TVPVKLKPMDGPKIKQWPLTEEKIKALTAICDEMEKEGKITKIGPENPYNTPIFAIKKK
DSTKWRKLVDFRELNKRRTQDFWEVQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDESFRK
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LYVGSDEIEQHRAKIEELREHLLRWGFTT'PDKKHQKEPPFLWMGYELHPDKWTVQPIQL
PEKDSWTVNDIQKIVGKLNWASQIYPGIKVRQLCRLLRGAKALTDIIPLTEEALELELAEN
REILKEPVHGAYYDPSKDLIAEIQKQGDQWYQIYQESFKNLKTGKYAKMRTAHTNDVR
QLTEAVQKIAQESIVIWGKIPKFRLLPIQKDTWETWWTWTDYQATWIPDWEFVNTPPLVKLW
YQLEKEPIAGAETFYVDGAANRETKIGKAGYVTDGRGRQKVI TLTETTNQKTELQAIHLAL
QDSGSEVNIIVTDSQYALRI IQAOPDKSESEIVNQCIEQLINKERVYLSWVPAHKGIGGNE
QVDKLYSNGIRKVLFLDGDIDKAQEEHEKYHSNWRAMASEFNLPVVAKEIVASCDKCQOK
GEAIHGQVDCSPRIWQLDCTHLEGKVIILVAVHVASGYMEAEVI PAETGQETAYFILKLAG
RWPVKVIHTDNGSNFTSTAVKAACWWAGIKQEFGIPYNPQSQGVVESMNKELKKIIEQVR
DQAEHLKTAVQMAVLIHNFKRGGGIGGYSAGERIVDIIATDIQTKELQKQIILKIQNFRVY
YRDNRPDIWKGPAKLLWKGEGAVVIQDNSEIKVVPRRKAKI IRDYGKQ MAGADCVASRQD
ED-

>POL-C.syn6.1

FFRENLAFFQGEAREFPSEQTRAISPTSR-----ELQV--RGNNPRSEAGA
ER----QGT---LNLFPQITLWQRPLVSIKIGGQTRREALLDTGADDTVLEEIKLPGNWKPK
MIGGIGGFIVRQYDQILIEICGKRAIGTVLVGPTPVNIIIGRNMLTQIGCTLNFPISSIE
TVPVKLKPMDGPKVKQWPLTEEKIKALTEICEEMEKEGKITKIGPDNPYNTPVFAIKKK
DSTKWRKLVDFKELNKRRTQDFWEVQLGIPHPAGLKKKNSVTVLDVGDAYFSVPLDESFRK
YTAFTIPSINNETPGIRYQYNVLPQGWKGSFAIFQSSMTRILEPFRAQNPDIYIYQYMDD
LYVGSDEIEQHRAKIEELREHLLRWGFTT'PDKKHQKEPPFLWMGCELHPDKWTVQPIQL
PEKDSWTVNDIQKLVGKLNWASQIYSGIKVRQLCKLLRGVKALTDIVPLTKEAELELELAEN
REILREPVGHYDPAKDLIAEIQKQGDQWYQIYQEPFKNLKTGKYAKRRRTAHTNDVK
QLTEAVQKIATESIVIWGKIPKFRLLPIQKDTWETWWTWTDYQATWIPDWEFVNTPPLVKLW
YQLEKEPIAEAEETFYVDGAASRETKMGKAGYVTDGRGRQKVITLTETTNQKTELQAIKLAL
QDSGSEVNVVTVDSQYALGIIQAQPDSESEIVNQCIEQLINKERVYLSWVPAHKGIGGNE
QVDKLVSRGIRKVLFLDGDIDKAQDEHEKYHSNWRAMASEFNLPVIVAREIVASCDKCQK
GEATHGQVDCSPGIWQLDCTHLEGKVIILVAVHVASGYIEAEVIPTETGQETAYYILKLAG
RWPVKI IHTDNGSNFTSSAVKAACWWAGIQEFGIPYNPQSQGVVESMNKELKKIIGQVG
DQAEHLKTAVQMAVFIHNFKRGGGIGGYSAGERIIDIIATDIQTRRELQKQIILKIQNFRVY
YRDSRDFIWKGPAKLLWKGEGAVVIQDNSEIKVVPRRKVKI IKDYGKQ MAGADCMASRQD
ED-

>POL-C.syn6.2

FFRENLAFFQGEARELPSEQTRANGPTSR-----ELQV--RGDNPCEAGA
ER----QGT---FNFPQITLWQRPLVTIKVGGQVKEALLDTGADDTVLEEINLPGKWKPK



Fig. 10 cont'd-21

MIGGIGGFIVRQYDQIPIEICGKRAIGTVLVGPTPINIIGRNMLTQLGCTLNFPISPIE  
TVPVQLKPGMDGPRVKQWPLTEEKIKALTEICKEMEKEGKISKIGPENPYNTPIFAIKKK  
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDEDFRK  
YTAFTIP SINNETPGVRYQYNVLPQGWKGSPIFQSSMTKILEPFRTQNP EIV IYQYMDD  
LYIGSDLEIGQHREKIEELREHLLKWGFTT PDKKHQKEPPFLWMGYELHPDKWTVQPIKL  
PEKDSWTVNDIQRLVGKLNWASQIYAGIKVRQLCKLLKGAKALTDIVTLTEEALELELAEN  
REILKEPVYGVYDPSKDLVAEIQKQGNQWTYQIYQESFKNLKTGKYAKMRTAHTNDIK  
QLTEAVQKIAQESIVIWGKTPKFRLP IQKETWEAWWTDYWQATWIPDWEFVNT PPLVKLW  
YQLEKEPMAGVETFYVDGAANRETKIGKAGYVTDGRGRQKVVTITETTNOKTELQAIYLAL  
QDSGSKVNIIVTDSQYALGIIQAQPKSESELVSIIEQLINKEKIYLSWVPAHKGIGGNE  
QVDKLVSSGIRKVLFLDGDIDKAQEDHERYHSNWRAMASDFNLPPTIVAKEIVASCDCQCLK  
GEAMHGQVDCSPGMWQLDCTHLEGKIIIVAVHVASGYIEAEVISAETGQETAYYILKLAG  
RWPVKVHTDNGSNFTSAAVKAACWWAGVQOEFGI PYNPQSOGVVE SMNKELKRIIGQVR  
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIIDMIATDIQTKELQKQI IQIQNFRVY  
YRDSRDP IWKGPAKLLWKGEGAVVIQDKGDIKVVPRRKAKI IRDYGKQ MAGADCMAGRQD  
EDQ

>POL-C.syn6.3

FFREDLAFPQGEARKFPPEQTRANSPTSRSR-----KLQV--RGDNPRSEAGV  
ER----CGT---LNFPQITLWQRPLVSIKVGGOIREALLDTGADDTVLEEMSLPGKWKPK  
MIGGIGGFIVKQYEQILIEICGKKAIGSVLVGPTPVNIIGRNLLTQLGCTLNFPISPIE  
TVPVKLPGMDGPKVKQWPLTEEKIKALTAICDEMEKEGKITKIGPENPYNTPVFAIKKK  
DSTKWRKLVDFKELNKRTQDFWEVQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDENFRK  
YTAFTIPSRNNETPGIRYQYNVLPQGWKGSPIFQASMTKILEPFRAKNPEIV IYQYMDD  
LYVGS DLEIGQHRAKIEELRDHLLKWGFTT PDKKHQKEPPFLWMGYELHPDKWTVQPIEL  
PEKDSWTVNDIQKLVGKLNWASQIYPGIQVKQLCKLLRGAKALTDVVPLTEEALELELAEN  
REILKEPVHGAYYDPSKDLIAEIQKQGHQWTYQIYQEPYKNLKTGKYAKRRAAHTNDVK  
QLTEAVQKIAMESIVIWGKTPKFRLP IQKETWETWWTEYWQATWIPEWEFVNT PPLVKLW  
YQLEKEPIAGAETFYVDGAANRETKMGKAGYITDRGRQKIISLTETTNOKTELHAIQLAL  
QDSGSEANIVTDSQYALGIIQAQFDRSESELVNQIIEQLIKKERVYLAWVPAHKGIGENE  
QVDKLVSSGIRKILFLDGDIDKAQEEHEKYHSNWKAMASEFNLPVAREIVASCDCQCLK  
GEAMHGQVDCSPRIWQLDCTHLERKVLVAVHVASGYMEAEVIPAETGQETAYFILKLAG  
RWPVKVIHTDNGSNFTSNAVKAACWWAGIHQOEFGI PYNPQSOGVVE SMNKELKKIIEQVR  
DQAEHLKTAVQMAVLIHNFKRKGGIGGYTAGERIIDI IATDIQTKELQKQITKIQNFRVY  
YRDNRP IWKGPAKLLWKGEGAVVLQDNSDIKVVPRRKVKI IRDYGKQ MAGADCVAGRQD  
ED-

>POL-C.syn6.4

FFRKNLAFPQGEAREFPPEQTRANSPTSRSR-----ELQV--RGDNPLSEAGA  
ERQGTLQGT---LNCPQITLWQRPLVSIKVGGOIKEALLDTGADDTVLEEISLPGKWKPK  
MIGGIGGFIVRQYDQIVIEICGKKAIGAVLVGPTPVNI IRRNMLTQLRCTLNFPISPIK  
TVPVKLPGMDGPKVKQWPLSEEKIKALTAICEDMEKEGKITKIGPENPYNTPIFAIKKK  
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPAGLKKKRSVTVLDVEDAYFSVPLDEGFRK  
YTAFTIP SINNATPGIRYQYNVLPQGWKGSPIFQSSMTKILEPFRTKNPDIV IYQYMDD  
LYVGS DLEIGQHRAKIEKLREHLLRWGLTTPDKKHQKEPPFLWMGYELHPDKWTVQPIQL  
PDKDSWTVNDIQKLVGKLNWASQIYAGIKVKQLCRLLRGAKALTDIIPLTEEALELELAEN  
REILKEPVHGAYYDPSKDLIAEIQKQGGQWTYQIYQEPYKNLKTGKYAKMRTAHTNDVK  
QLAEAVQKITMESIVIWGRTPKFRLP IQKETWETWWTDYWQATWIPEWEFINT PPLVKLW  
YQLEKEPIVGAETFYVDGAANRETKLGKAGYVTDKGRQKIVSLTETTNOKTELQAIHLAL  
QDSGSEVNIIVTDSQYALRI IQAOPDKSESELVNQIIEQLINKERIYLSWVPAHKGIGGNE  
QVDKLVSNIRKVLFLDGDIEKAQEEHERYHSNWRAMANEFNLPPIVAREIVASCDCQCIK  
GEAMHGQVDCSPGVWQLDCTHLEGKVIIVAVHVASGYVEAEVIPAETGQEAAYFILKLAG  
RWPVKTIHTDNGSNFTSTAVKAACWWAGIKOEFGI PYNPQSOGVVE SMNKELKKIIGQVR  
DQAEHLKTAVQMAVLIHNFKRKGGIGDYSAGERIIDIIATDMQTKELQKQIIKVNFRVY  
YRDSRDP IWKGPAKLLWKGEGAVVIQDNGDIKVVPRRKVKI IKDYGRQ MAGADCVASRQD  
ED-



Fig. 10 cont'd-22

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>POL-C.syn6.5
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MIGGIGGFIVRQYDQITIEICGKKAIGTVLVGPTPINTIGRNMLTQIGCTLNFPISPIE
TVPVKLKPMDGPKIKQWPLTEEKIKALKAI CEEMEKEGKIEKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNKRTQDFWEIQLGIPHPAGLKRKKSVTVLDVGDAYFSVPLYEDFRK
YTAFTIPSVNNETPGIRYQYNVLPQGWKGS PAIFQCSMTKILEPFRAQNPEI VIYQYMD
LYVGS DLEIGQHRAKVEELREHLLKWGLTTPDKKHQKEPPFLWMGYELHDPKWTVQPIQL
PEKESWTVNDIQKIVGKLNWASQIYPGIKVRQLCRLLRGAKALTDIVPLTAEAELELAEN
REILKEPVHGVYYDPSKELIAEIQKQGDQWYQIYQEPFKNLKTGKYAKMRS AHTNDVK
QLTEAVQKIALESIVIWGKAPKFR LPIQKETWEIWWTDYWQATWIPEWEFVNT PPLVKLW
YQLEKEPIAGVETFYVDGAANRDTKIGKAGYVTD RGRQKIVSLSETTNQKTELQAIQLAL
QDSGLEVNIVTDSQYALGIIQAQPDNSESELVNQIIEELIKKERVYLSWVPAHKGIGGNE
QVDKLVSKGIRKVLFLDGIDKAQEEHEKYHNNWRAMASDFNLPPVVAKEIVACCDK CQLK
GEA IHGQVDCSPGIWQLDCTHLEGKVIIVAVHVASGYLEAEVI PAETGQETAYFLLKLAG
RWPVKVIHTDNGPNFTSAAVKAACWWAGINQEFGI PYNPQSQGVVESMNKELKKIIGQVR
EQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERIVDI IATDIQTKELQNQI I KIQNFRVY
YRDSRDP IWKGP AKLLWKGE GAVVIQENS DIKVVPRRKAKI IKDYGKQ MAGDDCVAGRQD
EDQ

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>POL-C.syn6.6
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MIGGIGGFIVRQYEQIPIEICGKKAIGTVLIGPTPVNIIGRNLLTQLGCTLNFPISPIE
TIPVKLKPMDGPKVKQWPLTEEKIKALTAI CEEMEKEGKITKIGPDNPYNTPVFAIKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPAGLKKKKSVTVLDVGDAYFSVPLDKDFRK
YTAFTIPSTNNETPGIRYQYNVLPQGWKGS PAIFQSSMIKILEPFRAKNPELVIYQYMD
LYVGS DLEIMQHRAKIEELRAHLLKWGFTTPDKKHQKEPPFLWMGYELHDPKWTVQPIVL
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVKQLCKLLRGTALTDIVPLTEEAELELAEN
REILKETVHGAYYDPSKDLIAEIQKQGYDQWYQIYQEPFKNLKTGKYAKKRTAHTNDVR
QLTEAVQKIAIESIVIWGKTPKFR LPIQKETWETWWADYWQATWIPEWEFVNT PPLVKLW
YQLEKDP IAGAETFYVDGAANRET KKGKAGYVTDKGRQKVVTLETETNQAELQAIQLAL
QDSGPEVNIVTDSQYALRIIQAQPDKSESEGLVNQIIEQLIKKEKVYLSWVPAHKGIGGNE
QIDKLVSSGIRRVLFLDGIDKAQEDHEKYHSNWRAMAGEFNLPPVVAKEIVASCDK CQQK
GEA IHGQVDCSPGIWQLDCTHLEGKIIIVAVHVASGYIEAEVI PAETGQDTAYYILKLAG
RWPVKVIHTDNGTNFTSAAVKAACWWASIQQEFGI PYNPQSQGVVEAMN KELKKIIGQIR
DQAEHLKTAVLMAVFIHNFKRKGGIGGEYSAGERI IDI IASDIQTKELQKQITKI QNFRVY
YRDSRDPVWKGP AKLLWKGE GAVVIQDNS DIKVI PRRKAKI IRDYGKQ MAGADCVAGGQD
ED-

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>POL-M.syn1.1
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ER----QGT---LNFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEDINLPGKWKPK
MIGGIGGFIVRQYDQILIEICGKKAIGTVLVGPTPVNIIGRNMLTQIGCTLNFPISPIE
TVPVKLKPMDGPKVKQWPLTEEKIKALTEICTEMEKEGKISKIGPENPYNTPVFAIKKK
DSTKWRKLVDFRELNKRTQDFWEVQLGIPHPAGLKKKKSVTVLDVGDAYFSVPLDESFRK
YTAFTIP SINNETPGIRYQYNVLPQGWKGS PAIFQSSMTKILEPFRKQNPDI VIYQYMD
LYVGS DLEIGQHRAKIEELREHLLKWGFTTPDKKHQKEPPFLWMGYELHDPKWTVQPIQL
PEKDSWTVNDIQKLVGKLNWASQIYPGIKVKQLCKLLRGAKALTDIVPLTEEAELELAEN
REILKEPVHGVYYDPSKDLIAEIQKQGDQWYQIYQEPFKNLKTGKYAKMRTAHTNDVK
QLTEAVQKIATESIVIWGKTPKFR LPIQKETWETWWTDYWQATWIPEWEFVNT PPLVKLW
YQLEKEPIVGAETFYVDGAANRET KLGKAGYVTD RGRQKVVSLETETNQKTELQAIQLAL
QDSGSEVNIVTDSQYALGIIQAQPDKSESELVNQIIEQLIKKEKVYLSWVPAHKGIGGNE
QVDKLVSSGIRKVLFLDGIDKAQEEHEKYHSNWRAMASDFNLPPIVAKEIVASCDK CQLK
GEA MHGQVDCSPGIWQLDCTHLEGKVIIVAVHVASGYIEAEVI PAETGQETAYFILKLAG
RWPVKVIHTDNGSNFTSAAVKAACWWAGIQQEFGI PYNPQSQGVVESMNKELKKIIGQVR
DQAEHLKTAVQMAVFIHNFKRKGGIGGYSAGERI IDI IATDIQTKELQKQITKI QNFRVY

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Fig. 10 cont'd-23

YRDSRDPiWKGPakLLWkGEGAVVIQDNsDIKVVPRRkAKIIRDYgKQMaGDDCVaGRQD  
EDQ

>POL-M.syn3.1

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MIGGIGGFikVRQYDQIPiEiCGKRAIGTVLVGPTPINIIGRNMLTQlGCTLNFPiSPiK  
TVPVklKPGMDGPKVQWPLTEEKIKALTAICEEMEKEGKITKIGPENPYNTPVFAIKKK  
DGTkWRKLVDFRELnkRTQDFWEVQLGIPHPsGLKkKkSVSVLDVGDAYFSVPLDESFRK  
YTAFTiPSINNETPGIRYCYNVLPQGWKGS PAIFQCSMTKILEPFRAQNPEIVIYQYMDD  
LYIGSDLEIGQHRAKIEELREHLLRWGFTTPDKKHQKEPFLWMGCELHPDKWTVQPIQL  
PEKDSWTVNDIQKIVGKLNWASQIYPGIKVRQLCKLLRGAKALTDIVPLTEEAELELAEN  
REILREPvHGvYDPSKDLVAEIQKQGDQWTYQIYQEPFKNLKTGKYAKMRTAHTNDVK  
QLTEAVQKIALESIVIWGKIPKfRLPIQKETWEAWWMEYWOATWIPeWefINTPPLVklW  
YQLEKEPIAGAETfYVDGAANRETKIGKAGYVTDGRGRQKIVSLTETTNOkAELQAIQLAL  
QDSGPEVNIvTDSQYALGIIQAHPDKSESELVNQIIEQLIKKERVYLSWVPAHKGI GENE  
QVDKLVsNGIRKILFLDGDIDKAQEEHEKYHSNWRAMASEFNLPPIVAKeIVASCNKcQLK  
GEALHGQVDCSPGMWQLDCTHLEGKVIIVAVHVASGYMEAEVIPAETGQETAYYTLKLAG  
RWPVKVvHTDNGSNFTSTAVKAACWWAGIQEFGIPYNPQSOGVIESMNKELKKIIGQIR  
DQAEHLKTAVQMAVFIHNFKRKGGIGeYSAGERIIDLIATDIQTRELQKQIKIQNFRVY  
YRDSRDPiWKGPakLLWkGEGAVVLQDNsDIKVVPRRkVKIKDYgKQMaGADCVaGRQD  
ENQ

>POL-M.syn3.2

FFRENLAfQGEARKfSSEQTGAnsPTr-----ELRV--RRGDNPLSEAGA  
ER----RGTVPsLSfPQITLWQRPLVTVKIGGQlIEALLDTGADDTVLEdINLPGKWKPR  
MIGGIGGFikVKQYDQILiEiCGKKAIGTVLVGPTPVNIIGRNMLTQlGCTLNFPiSPiD  
TVPVTLKPGMDGPKIKQWPLTEEKIKALTEICTEMEKEGKISRIGPENPYNTPIFAIKKK  
DSTkWRKLVDFRELnkKTQDFWEVQLGIPHPAGLkKkRSVTVLDVGDAYFSVPLDKDFRK  
YTAFTiPSTNNETPGIRYQYNVLPQGWKGS PSiFQSSMTRI LEPFRAKNPEIVIYQYMDD  
LYVGSdLEIEQHRTKIEELRQHLLRWGLTTPDKKHQKEPFLWMGYELHPDRWTVQPIEL  
PEKESWTVNDIQKLVGKLNWASQIYAGIKVKQLCKLLRGTKALTEVVPLTEEAEL ELEEN  
REILKDPVHGAYYDPSKDLIAEIQKQGHQWTYQIYQEQYKNLKTGKYARKRSaHTNDVR  
QLTEAVQKIATESIVIWGKTPKfRLPIQRETWETWWTDYWOATWIPeWefVNTPPLVklW  
YQLEKDPiAGVETfYVDGASNRETKKGGKAGYVTDKGRQKVSLTETTNOkTELHAIHLAL  
QDSGSEVNIvTDSQYALGIIQAQPDRSESELVNQIIEELIKKEKVYLSWVPAHKGI GNE  
QVDKLVSSGIRKVLFLDRIDKAQEEHeryHSNWRtMASDFNLPPIVAKeIVANCDKcQLK  
GEAMHGQVDCSPGIWQIDCTHLEGKVIIVAVHVASGYIEAEVIPAETGQETAYFLKLAG  
RWPVKVIHTDNGSNFTSAAVKAACWWANVTQEFGI PYNPQSOGVVESMNKELKKIIGQVR  
EQAEHLKTAVQMAVLIHNFKRKGGIGGYSAGERIIDIIASDIQTKELQKQIKVQNFRVY  
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EDQ

>POL-M.syn3.3

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DR----QGTvS-fSFPQITLWQRPLVTIKIGGQlKEALLDTGADDTVLEEMNLPGRWKPK  
MIGGIGGFikVRQYDQILiEiCGHKAIGTVLIGPTPVNIIGRNLLTQlGCTLNFPiSPiE  
TVPVklKPGMDGPRVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPVFAIKKK  
DSTRWRKLVDFRELnkRTQDFWEIQLGIPHPAGLkKkKSVTVLDVGDAYFSVPLDEGFRK  
YTAFTiPSVNNETPGVRYQYNVLPQGWKGS PAIFQSSMTKILEPFKQNPDIVIYQYMDD  
LYVGSdLEIGQHRTKIEELREHLLKwGFTTPDKKHQNEPFLWMGYELHPDKWTVQPIVL  
PEKDSWTVNDLQKLVGKLNWASQIYPGIKVKQLCRLLRGAKALTEVIPLTKEAELELAEN  
REILKEPvHGvYDPSKELIAEIQKQGGQWTYQIYQEPYKNLKTGKYARMRGaHTNDVK  
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YQLEKEPIVGAETfYVDGAANRETKLGKAGYVTDGRGRQKVSLTDTTNOkTELQAIHLAL  
QDSGLEVNIvTDSQYAIgIIQAQPDKSESELVSQIIEQLIKKEKVYLAWVPAHKGI GNE  
QVDKLVsAGIRKVLFLDGDIDKAQEDHEKYHNNWRAMASDFNLPPVVAKeIVASCDKcQLK



Fig. 10 cont'd-24

GEA IHGQVDCSPGIWQLDCTHLEGKIILVAVHVASGYLEAEVIPAETGQETAYFILKLAG  
RWPVKTIHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQSQGVVESINKELKKIIGQVR  
DQAEHLKTAVLMAVFIHNFKRRGGIGGYSAGERIVDI IATDIQTKELQKQITKI QNERVY  
YRDSRDPLWKGP AKLLWKGEGAVVIQDNSEIKV VPRRKAKI IRDYGKQ MAGDDCVASRQD  
ED-

>POL-M.syn4.1

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MIGGIGGFIVKQYDQILIEICGKKAIGTVLVGPTPVNI IGRNMLTQIGCTLNFPISPID  
TVPVTLKPGMDGPRIKQWPLTEEEKIKALTEICKEMEEEGKISKIGPENPYNTPIFAIKKK  
NSTRWRKLVDFRELNKKTQDFWEVQLGIPHPAGLKRKKS SVTVLDVEDAYFSVPLDESFRK  
YTAFTIIP SINNETPGVRYQYNVLPQGWKGS PAIFQCSMTKILEPFR IKNPEMVIYQYMDD  
LYVGS DLEIGQHRIKIEELRAHLLSWGFTTPDKKHQKDP PFLWMGYELHPDRWTVQPIEL  
PEKDSWTVNDIQKLVKLNWASQIYSGIKVRQLCRLLRGAKALTDIVPLTEEALELAEN  
REILKEPVHGAYYDPSKDLVAEIQKQGDQWTYQIYQEPFKNLKTGKYARKRSAHTNDVK  
QLTEVVQKIATESIVIWGKTPKFRLP IQRETWETWWTEYWQATWIPWEFVNTPPPLVKLW  
YQLEKDP IAGVETFYVDGAASRETKLGKAGYVTD RGRQKVSLTETTNOKTELHAIHLAL  
QDSGSEVNI VTD SQYVLGIIQAQPD RSESELVNQIIEELIKKEKVYLSWVPAHKGIGGNE  
QVDKLVSSGIRKVLFLNGIDKAQEEHERYHSNWRTMASDFNLPPIVAKEIVANCDKCQLK  
GEAMHGQVDCSPGVWQLDCTHLEGKVIIVAVHVASGYIEAEVIPAETGQEAAYFILKLAG  
RWPVKVHTDNGSNFTSAAVKAACWWANVRQEFGI PYNPQSQGVVESMNNELKKIIGQIR  
DQAEHLKTAVLMAVFIHNFKRRGGIGEYSAGERIIDI IATDIQTR ELQKQITKI QNFRVY  
FRDSRDPIWKGP AKLLWKGEGAVVIQDNSEIKV VPRRKVKI IRDYGKQ MAGDDCVAGRQD  
EN-

>POL-M.syn4.2

FFRENLAFPQGEAREFPSEQARANSPTS RANSPTS R-----DLWDGGRDNL P--SEAGA  
ER----QGT---LNFPQITLWQRPLVTVRIGGQLREALLD TGADDTVLEDIDLPGKWKPK  
IIGGIGGFIVRQYEQIPIEICGHKAIGTVLVGPTPINI IGRNMLTQLGCTLNFPISPIK  
TVPVKLPGMDGPRVKQWPLTEEEKIKALTAICEEMEKEGKITKIGPENPYNTPVFAIKKK  
DSTRWRKLVDFRELNRRTQDFWEVQLGIPHPAGLKKKRS SVTVLDVGDAYFSVPLDEGFRK  
YTAFTIIP SVNNETPGIRYQYNVLPQGWKGS PSI FQSSMTRILEPFR AKNPEIVIYQYIDD  
LYVRS DLEIGQHRAKIEELREHLLRWGFTTPDKKHQKEP PFLWMGCELHPDKWTVQPIQL  
PEKDSWTINDIQKLVGKLNWASQIYPGIKVRQLCKLLRGTKALTDIVTLTEEALELAEN  
REILKDPVHG VYYDPSKELIAEIQKQGDQWTYQIYQEQYKNLKTGKYAKRRTAHTNDVR  
QLTEAVQKIALESIVIWGKIPKFRLP IQKETWEAWWMEYWQATWIPWEYVNTPPPLVKLW  
YQLEKEPIIGAE TFYVDGAANRETKLGKAGYV TNRGRQKVSLTDTTNQKTELQAIQLAL  
QDSGSEVNVVTD SQYALGIIQAHPDKSESELVNQIIEQLIKKERVYLSWVPAHKGIGGNE  
QVDKLV SAGIRKVLFLDGDIDKAQEDHERYHSNWRAMASDFNLPPVVAKEIVASCNKCQLK  
GEA IHGQVDCSPGIWQLDCTHLEGKIIIVAVHVASGYMEAEVIPAETGQETAYFILKLAG  
RWPVKI IHTDNGSNFTSATVKAACWWANVTQEFGI PYNPQSQGVVESINKELKKIIGQVR  
DQAEHLRTAVQMAVFIHNFKRRGGIGGYSAGERIVDI IATDIQTKELQKQITKI QKFRVY  
YRDSRDPLWKGP AKLLWKGEGAVVIQDN S DIKVI PRRKAKI IKDYGKQ MAGADCVAGRQD  
EDQ

>POL-M.syn4.3

FFRENLAFPQ GKAREFPSEQTRANSPTS RANSPTS R-----ELQV--RGDNPRSEAGA  
ER----QGT---FNFPQITLWQRPLVSIKVG GQIKEALLDTGADDTVLEEINLPGKWKPR  
MIGGIGGFIVRQYDQILIEICGKRAIGTVLVGPTPANI IGRNLLTQLGCTLNFPISPTE  
TVPVKLPGMDGPKIKQWPLTEEEKIKALTEICTEMEKEGKISKIGPENPYNTPIFAIKKK  
DSTKWRKLVDFKELNKR TQDFWEVQLGIPHPSGLK KKS SVTVLDVGDAYFSVPLDEDFRK  
YTAFTIIP STNNETPGIRYQYNVLPQGWKGS PAIFQASMTKILEPFR AQNPEIVIYQYMDD  
LYVGS DLEIEQRRAKVEELREHLLK WGFTTPDKKHQNEP PFLWMGYELHPDKWTVQPIKL  
PEKESWTVNDIQKLVGKLNWASQIYPGIKVKQLCRLLRGTKALTEVIPLTKEAELELAEN  
REILREP VHG VYYDPTKDLIAEIQKQGHQWTYQIYQEPHKNLKTGKYAKMRTAHTNDVK  
QLAEAVQKIAMESIVIWGKIPKFKLPIQKETWETWWT DYWQATWIPDWEFVNTPPPLVKLW



Fig. 10 cont'd-25

YQLEKEPIAGAETFYVDGAANRETKIGKAGYVTDGRGRQKIVSLTETTNQKAELOAIQLAL
QDSGPEVNIIVTDSQYALGIIQAQPKSESEIVNQIIEKLEKDKVYLSWVPAHKGIGGNE
QIDKLVSNNGIRKVLFLDGIKAQEEHEKYHSNWRAMASEFNLPPIVAKEIVASCDKCOLK
GEATHGQVDCSPGMWQLDCTHLEGKIIILVAVHVASGYIEAEVIPTETGQETAYYILKLAG
RWPVKVIHTDNGSNFTSTAVKAACWWAGIQQEFGI PYNPQGGVVE SMNKELKKIIGQVR
EQAEHLKTAVQMAVLIHNFKRKGGIGGYSAGERIIDIIASDIQTKELQKQI IKVQNFVY
YRDSRDP IWKGP AKLLWK GEGAVVLQDNSDIKVVPRRKVKI IKDYGKQ MAGADCVASRQD
EN-

>POL-M.syn4.4

FFREDLAFPQ GKAREFSSEQTRANSPTRR-----ELQVWGRDNNSLSEAGA
DR----QGTVS-FSFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMNLPGRWKPK
MIGGIGGF I KVRQYDQIPIEICGKKAIGTVLIGPTPVNII GRNLLTQIGCTLNFPISPIE
TIPVVKLPGMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISRIGPENPYNTPVF AIKKK
DGTKWRKLVDFRELNKRTQDFWEIQLGIPHPAGLKKKKS SVSVLDVGDAYFSVPLDKDFRK
YTAFTIP SINNETPGIRYCYNVLPQGWKGS PAIFQSSMTKILEPFRKQNPDI VIYQYMD
LYIGSDLEIGQHRTKIEELRQHLLRWGLTTPDKKHQNEPPFLWMGYELHHPDKWTVQPIVL
PEKDSWTVNDLQKLVGKLNWASQIYAGIKVKQLCKLLRGAKALTEVVPLTEAELELEEN
REILKEPVHGVYYDPSKDLIAEIQKQGQGWTYQIYQEPYKNLKTGKYARMRGAHTNDVK
QLTEAVQKIAQECIVIWGKTPKFKLPIQKETWETWWM DYWQATWIPEWEFINTPPLVKLW
YQLEKEPIVGAETFYVDGASNRETKK GKAGYVTDKGRQKVVTLTETTNQKTELQAIHLAL
QDSGLEVNIIVTDSQYAIGI IQAQPDKSESELVSQIIEQLIKKEKVYLA WVPAHKGIGGNE
QVDKLVSNNGIRKILFLDGI DKAQEEHEKYHNNWRAMASDFNI PPVVAKEIVACCDKCOLK
GEALHGQVDCSPRIWQLDCTHLEGKVILVAVHVASGYLEAEVI PAETGQETAYFLLKLAG
RWPVKTIHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQSQGVIESM NKELKKIIEQVR
DQAEHLKTAVQMAV FVHNFKRKGGIGDYSAGERIIDII STDIQTRELOKQI IKIQNFVY
YRDSRDPVWKGP AKLLWK GEGAVVIQDNNEIKVVPRRKAKI IRDYGKQ MAGDDCVASRQD
ED-

>POL-M.syn6.1

FFREDLAFPQGEARKFPSEQTRANSPTRG-----ELQVWGRDNNSLSEAGD
DR----QGTVS-FNLPQITLWQRPLVTVRIGGQLIEALLDTGADDTVLEDMNLP GKWKPK
MIGGIGGF I KVRQYEQIPIEICGHKAIGTVLIGPTPVNII GRNLLTQIGCTLNFPISPID
TVPVTLKPGMDGPKIKQWPLTEEKIKALTEICTEMEKEGKISRIGPENPYNTPVF AIKKK
NSTRWRKLVDFRELNKRTQDFCEVQLGIPHPAGLKKKRSVTVLDVGDAYFSVPLDENFRK
YTAFTIP SINNETPGVRYQYNVLPQGWKGS PAIFQASMTKILEPFRTKNPELVIYQYMD
LYVGS DLEIEQHRTKIEELRAHLLSWGFTTPDKKHQKEPPFLWMGYELHHPDKWTVQPIEL
PEKDSWTVNDIQKIVGKLNWASQIYPGIKVRQLCRLLRGFKALTDIVPLTAEAELELAEN
REILREP VHG VYYDPSKELIAEIQKQGHQWTYQIYQDPFKNLKTGKYARKRSAHTNDVR
QLTEAVQKITTESIVIWGKTPKFR LPIQRETWEAWWMEYWQATWIPEWEFINTPPLVKLW
YQLEKDP I VGAETFYVDGAASRETKLGKAGYVTNKGROKVVSLNETTNQKTELHAIHLAL
QDSGSEANIVTDSQYALGIIQAQPD RSESEVVNQIIEELIKKEKVYLSWVPAHKGIGGNE
QVDKLVSSGIRKVLFLDGI DKAQEDHERYHSNWRMTASDFNLPPIVAREIVASCDKCOQK
GEAMHGQVDCGPGIWQLDCTHLERKVILVAVHVASGYIEAEVI PAETGQETAYFVLKLAG
RWPVKVIHTDNGSNFTSAAVKAACWWANVTQEFGI PYNPQSQGVVE SMNNE LKKIIGQVR
EQAEHLKTAVLMAVFIHNFKRKRGIGGYSAGERIVDIIASDIQTKELQNOITKI QNFVY
FRDSRDP IWKGP AKLLWK GEGAVVIQDNNDIKVVPRRKVKI IRDYGKQ MAGDDCVAGRQD
EN-

>POL-M.syn6.2

FFREDLAFQQGEARKFSSEQTRANSPTSR-----ELRVWG-GDNTLSETGA
ER----QGT---LNFQITLWQRPLVTIKVGGQIKEALLDTGADDTVLEDINLP GKWKPK
MIGGIGGF I KVRQYDQIPIEICGKKAIGSVLVGPTPVNII GRNMLTQLGCTLNFPISPIK
TVPVKLPGMDGPKVKQWPLSEEKIKALTAICDEMEKEGKITKIGPDNPYNTPVF AIKKK
DGTKWRKLVDFKELNKRTQDFWEIQLGIPHPAGLKKKKS SVSVLDVGDAYFSVPLDESFRK
YTAFTIP SLNNETPGIRYCYNVLPQGWKGS PAIFQSSMTKILEPFRAQNF EIVIYQYIDD
LYVRS DLEIGQHRAKIEELREHLLKWGLTTPDKKHQKEPPFLWMGYELHHPDRWTVQPIQL



Fig. 10 cont'd-26

PKDSWTVNDLQKLVGKLNWASQIYPGIRVKQLCKLLKGAKALTDIVTLTEEALELAEN  
REILKNPVHGVYDPAKDLIAEIQKQGNDQWTYQIYQEPHKNLKTGKYAKMRTAHTNDVK  
QLTEVVQKIAMESIVIWGKVPKFRLPQKETWETWWTDYWQATWIPDWEFVNTPLVVKLW  
YQLEKEPIAGAETFYVDGAANRETKMGKAGYVTDGRGRQKVVSLETTNQTTELQAIQLAL  
QDSGPEVNIIVTDSQYAIGI IQAQPDKSESEIVNQIIEQLIKKERVYLSWVPAHKGIGENE  
QVDKLVSTGIRRVLFLDGIDKAQEEHERYHSNWRAMASEFNLPPIVAKEIVASCDQCQLK  
GEAMHGQVDCSPGVWQLDCTHLEGKIIIVAVHVASGYMEAEVI PAETGQETAYFILKLA  
RWPVKVIHTDNGPNFTSATVKAACWWANITQEFGI PYNPQGGQGVVESMNKELKKI IKQVR  
DQAEHLKTAVQMAVLIHNFKRKGGIGGYSAGERIIDIIASDIQTKELQKQI IKIQNFQVY  
YRDSRDPIWKGPAKLLWKGEGAVVLQDNSDIKVVPRRKVKI IKDYGKQMAGADCVAGGQD  
ED-

>POL-M. syn6.3

FFRENLAFFPQKAREFFSEQTRAISPTSR-----ELQVWGGDNNLSLSEAGA  
ER----QGTVS-FSFPQITLWQRPIVTIKIGGQLREALLDTGADDTVLEEMNLPGRWPKPK  
MIGGIGGFIVKQYDNILIEICGHKAVGSTVLVGPFPANI IGRNLLTQLGCTLNFPISPIE  
TVPVKLKPGLDGPVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPYNTPIFAIKKK  
DSTRWRKLVDFRELNRRRTQDFWEVQLGIPHPAGLKRKKSVTVLDVGDAYFSVPLDKEFRK  
YTAFTIPSINNETPGIRYQYNVLPQGWKGPSIFQSSMTKILEPFRKKNPEMVIYQYMD  
LYIGSDLEIGQHRIKIEELREHLLKWGF TTPDKKHQKEPPFLWMGCELHPDKWTVQPIML  
PEKDSWTVNDIQKLVGKLNWASQIYAGIKVRQLCKLLRGTALTEVVPLTEEALELAEN  
REILKEPVHGVYDPSKDLIAEVQKQGQDQWTYQIYQEPFKNLKTGKYAKKRSHTNDVK  
QLTEAVQKIALESIVIWGKAPKFRLPQKETWEAWWTEYWQATWVPEWEFVNTPLVVKLW  
YQLETEPIAGAETYYVDGAANRETKLKGAGYVTDNRGRQKVVSLETTNQTTELQAIHLAL  
QDSGLEVNIIVTDSQYALGI IHAQPDKSESELVNQIIEQLINKERIYLSWVPAHKGIGENE  
QVDKLVSKGIRKVLFLDGI EKAQEEHEKYHSNWKAMASEFNLPVVAKEIVACCDKQCQLK  
GEALHGQVDCSPGMWQLDCTHLEGKIIIVAVHVASGYIEAEVIPTETGQETAYFLLKLAG  
RWPVKTIHTDNGSNFTSTTVKAACWWAGIKQEFGI PYNPQSQGVVESINKELKKI IGQIR  
DQAEHLKTAVLMAVFIHNFKRKGGIGGYTAGERIVDIIATDIQTKELQKQITKVQNFVY  
YRDSREPLWKGPAKLLWKGEGAVVIQDNNEIKVVPRRKAKILRDYGKQMAGADCVASRQD  
EN-

>POL-M. syn6.4

FFRENLAFFQGEAREFSSEQTRTNSPTSR-----ELWDGGRDNLPS-SEAGA  
ER----RGTVPSLSFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEINLPGKWKPK  
LIGGIGGFIVKQYDQILIEICGKKAIGTVLVGPTPINI IGRNMLTQIGCTLNFPISPIE  
TIPVKLKPMDGPRVKQWPLTEEKIKALIEICTEMEKEGKISRVPENPYNTPIFAIKKK  
NSNRWRKLVDFRELNKRTQDFWEVQLGIPHPGGLKKKKSVTILDVGDAYFSVPLDEDFRK  
YTAFTIPSINNATPGIRYQYNVLPQGWKGPSAIFQCSMTKILEPFRKQNPETIIYQYMD  
LYVRSLEIGQHRTKIEELRQHLLKWGFYTPDKKHQKEPPFLWMGYELHPDKWTVQPIKL  
PEKESWTVNDIQKLVKLNWASQIYPGIKVKQLCRLLRGAKALTEVIPLTEEALELEEN  
REILKDPVHGVYDPTKDLIAEIQKQGDQWTYQIYQEPYKNLKTGKYAKRRTAHTNDVR  
QLTEVVQKVATESIVIWGKIPKFKLPQKETWEIWWTDYWQATWIPWEFVNTPHLVVKLW  
YQLEKEPIIGAETFYVDGASNRETKKKGAGYVTDGRGRQKIVSLETTNQTTELQAIQLAL  
QDSGSEVNIIVTDSQYALGI IQAHPDKSESELVSQIIEQLIKKEKVYLAWVPAHKGIGGNE  
QIDKLVSNKIRKILFLDGI DKAQEEHEKYHNNWRAMASDFNLPPVVAKEIVASCNKQCQLK  
GEA IHGQVDCSPRIWQLDCTHLEGKVIIVAVHVASGYVEAEVI PAETGQDTAYFILKLAG  
RWPVKVHTDNGSNFTSAAFKAACWWANVQOEFGI PYNPQSQGVVEAMNKELKKI IEQVR  
DQAEHLKTAVQMAVVFVHNFKRKGGIGDY SAGERIIDIIATDIQTRRELQKQI IKIQNFVY  
YRDNRPDIWKGPAKLLWKGEGAVVIQDNSDIKVI PRRKAKI IRDYGKQMAGDDCMAGRQD  
EDQ

>POL-M. syn6.5

FFREDLAFLOQKAREFSSEQTRANSPTRR-----ELQVWGRDSNSLSEAGA  
DR----QGTVS-FNFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEDIDLPGKWKPK  
IIGGIGGFIVKQYDQILIEICGKRAIGTVLVGPTPVNI IGRNILTQIGCTLNFPISPID  
TVPVKLKPMDGPRIKQWPLTEEKIKALTEICKEMEEEGKISKIGPENPYNTPVFAIKKK



Fig. 10 cont'd-27

DSTKWRKVVDRELNKGTQDFWEVQLGIPHPAGLKQKKSVTVLDVEDAYFSVPLDKDFRK  
YTAFTIPSVNNETPGIRYQYNVLPQGWKGS PAIFQS SMTRILEPFRKQNPDIYQYMDD  
LYVGS DLEIGQHRTKVEELRQHLLRWGFTTPDKKHQKDPFLWMGYELHPDKWTVQPIVL  
PEKDSWTINDIQKLVGKLNWASQIYSGIKVRQLCKCLRGTALTEVIPLTKEAELELAEN  
KEILKEPVHGVYDPSKDLVAEIQKQGQGWYQIYQEQYKNLKTGKYARMRGAHTNDVK  
QLAEAVQKIATESIWIWGKIPKFRLP IQRETWETWWTEYWQATWIPEWEYVNT PPLVKLW  
YQLEKEPIVGAETFYVDGAANRDTKLGKAGYVTDGRGRQKVPLTDTTNQKTELQAINLAL  
QDSGSKVNIIVTDSQYVLGT IQAQ PDRSESEIVNQIIEKLIKDKVYLSWVPAHKGIGNE  
QVDKLV SAGIRKVLFLDGDIDKAQDEHEKYHSNWRAMASDFNLPPIVAKEIVASCDKCQLK  
GEATHGQVDCSPGIWQLDCTHLEGKVIIVAVHVASGYIEAEVISAETGQETAYYILKLAG  
RWPVKI IHTDNGSNFTSTAVKAACWWAGIQQEFGI PYS PQSQGVV ESMNKQLKQIIGQVR  
DQAEQLKTAVQMAVFIHNFKRKGGIGEYSAGERIIDII STDIQTRELOKQITKI QNFRVY  
YRDSRDPVWKGP AKLLWK GEGAVVIQDNSEIKV VPRRKAKI IRHYGKQ MAGDDCVASRQD  
EDQ

>POL-M.syn6.6

FFRENLAFPQGEAREFPSEQARANSPTS SRANSPTS R-----ELQV--RGDNPRSEAGA  
ERQGT LQGT---LNC PQITLWQRPLVSIKVG GQVKEALLDTGADDTVLEEMSLPGKWKPK  
MVG GIGGF I KVRQYDQILVEICGHKAIGTVLVGPTPVNI IRRNMLTQLRCTLNFPISPIE  
TVPVTLKPGMDGPKVRQWPLTEEKIKALTAICEEMEKEGKITKIGPENPYNTPVFAIRKK  
DSTKWRKLVDFRELNKKTQDFWEVQLGIPHP SGLK KKSVTVLDVGDAYFSVPLDEGFRK  
YTAFTIPSTNNETPGIRYQYNVLPQGWKGS PAIFQSSMIKILEPFRAKNPEIYIYQYMDD  
LYVGS DLEIGQHRAKVEELREHLLRWGFTTPDKKHQNEPPFLWMGYELHPDKWTVQPIQL  
PEKDSWTVNDIQRLVGKLNWASQIYAGIKVKQLCKLLRGAKALTDIVPLTEEAELELAEN  
REILKTPVHGVYDPSKDLIAEIQKQGQDQWSYQIYQEPFKNLKTGKYARTRGAHTNDVR  
QLTEAVQKIAQECIWIWGKTPKFKLPIQKDTWETWWM DYWQATWIPKWEFVNT PPLVKLW  
YQLEKDPIAGVETFYVDGAANRETKIGKAGYVTDKGRQKVVTLET TTNQKTELHAIYLAL  
QDSGSEVNVV TDSQYALGI IQAQ PDRSESELVNQIIEKLI GKD VYLSWVPAHKGIGENE  
QVDKLV SNGIRKVLFLDGDIDKAQEDHEKYHSNWRAMANEFNLPPIVAKEIVANCDKCQLK  
GEAMHGQVDCSPGIWQIDCTHLEGKVIIVAVHVASGYLEAEVIPAETGQEAAYF I LKLAG  
RWPVKTVHTDNGSNFTSNAVKAACWWANVRQEFGI PYNPQSQGVIESMNKELKKIIGQVR  
DQAEHLRTAVQMAVFIHNFKRKGGIGGYSAGERMIDI IATDIQTTELOKQITKI QKFRVY  
YRDSRDPVWKGP AKLLWK GEGAVVIQENS DIKVVPRRKAKI IKDYGKQVAGADCVAGRQD  
EDQ



## POLYVALENT VACCINE

**[0001]** This application is a continuation of U.S. application Ser. No. 11/990,222, filed Apr. 20, 2009, which is the U.S. national phase International Application No. PCT/US2006/032907, filed Aug. 23, 2006, which designated the U.S. and claims priority from U.S. Provisional Application No. 60/710,154, filed Aug. 23, 2005, and U.S. Provisional Application No. 60/739,413, filed Nov. 25, 2005, the entire contents of which are incorporated herein by reference.

**[0002]** This invention was made with Government support under Contract No. DE-AC52-06NA25396 awarded by the U.S. Department of Energy. The Government has certain rights in the invention.

**[0003]** The content of the ASCII text file submitted with this application on Dec. 3, 2010 named Sequence\_Listing.txt created Dec. 3, 2010, which is 870 KB, is also incorporated herein by reference.

**[0004]** The Sequence Listing filed Apr. 20, 2009 in U.S. application Ser. No. 11/990,222 is incorporated herein by reference.

## TECHNICAL FIELD

**[0005]** The present invention relates, in general, to an immunogenic composition (e.g., a vaccine) and, in particular, to a polyvalent immunogenic composition, such as a polyvalent HIV vaccine, and to methods of using same. The invention further relates to methods that use a genetic algorithm to create sets of polyvalent antigens suitable for use, for example, in vaccination strategies.

## BACKGROUND

**[0006]** Designing an effective HIV vaccine is a many-faceted challenge. The vaccine preferably elicits an immune response capable of either preventing infection or, minimally, controlling viral replication if infection occurs, despite the failure of immune responses to natural infection to eliminate the virus (Nabel, *Vaccine* 20:1945-1947 (2002)) or to protect from superinfection (Altfeld et al, *Nature* 420:434-439 (2002)). Potent vaccines are needed, with optimized vectors, immunization protocols, and adjuvants (Nabel, *Vaccine* 20:1945-1947 (2002)), combined with antigens that can stimulate cross-reactive responses against the diverse spectrum of circulating viruses (Gaschen et al, *Science* 296:2354-2360 (2002), Korber et al, *Br. Med. Bull.* 58:19-42 (2001)). The problems that influenza vaccinologists have confronted for decades highlight the challenge posed by HIV-1: human influenza strains undergoing antigenic drift diverge from one another by around 1-2% per year, yet vaccine antigens often fail to elicit cross-reactive B-cell responses from one year to the next, requiring that contemporary strains be continuously monitored and vaccines be updated every few years (Korber et al, *Br. Med. Bull.* 58:19-42 (2001)). In contrast, co-circulating individual HIV-1 strains can differ from one another by 20% or more in relatively conserved proteins, and up to 35% in the Envelope protein (Gaschen et al, *Science* 296:2354-2360 (2002), Korber et al, *Br. Med. Bull.* 58:19-42 (2001)).

**[0007]** Different degrees of viral diversity in regional HIV-1 epidemics provide a potentially useful hierarchy for vaccine design strategies. Some geographic regions recapitulate global diversity, with a majority of known HIV-1 subtypes, or clades, co-circulating (e.g., the Democratic Repub-

lic of the Congo (Mokili & Korber, *J. Neurovirol* 11(Suppl. 1):66-75 (2005)); others are dominated by two subtypes and their recombinants (e.g., Uganda (Barugahare et al, *J. Virol.* 79:4132-4139 (2005)), and others by a single subtype (e.g., South Africa (Williamson et al, *AIDS Res. Hum. Retroviruses* 19:133-144 (2003)). Even areas with predominantly single-subtype epidemics must address extensive within-clade diversity (Williamson et al, *AIDS Res. Hum. Retroviruses* 19:133-44 (2003)) but, since international travel can be expected to further blur geographic distinctions, all nations would benefit from a global vaccine.

**[0008]** Presented herein is the design of polyvalent vaccine antigen sets focusing on T lymphocyte responses, optimized for either the common B and C subtypes, or all HIV-1 variants in global circulation [the HIV-1 Main (M) group]. Cytotoxic T-lymphocytes (CTL) directly kill infected, virus-producing host cells, recognizing them via viral protein fragments (epitopes) presented on infected cell surfaces by human leukocyte antigen (HLA) molecules. Helper T-cell responses control varied aspects of the immune response through the release of cytokines. Both are likely to be crucial for an HIV-1 vaccine: CTL responses have been implicated in slowing disease progression (Oxenius et al, *J. Infect. Dis.* 189:1199-208 (2004)); vaccine-elicited cellular immune responses in nonhuman primates help control pathogenic SIV or SHIV, reducing the likelihood of disease after challenge (Barouch et al, *Science* 290:486-92 (2000)); and experimental depletion of CD8+ T-cells results in increased viremia in SIV infected rhesus macaques Schmitz et al, *Science* 283:857-60 (1999)). Furthermore, CTL escape mutations are associated with disease progression (Barouch et al, *J. Virol.* 77:7367-75 (2003)), thus vaccine-stimulated memory responses that block potential escape routes may be valuable.

**[0009]** The highly variable Env protein is the primary target for neutralizing antibodies against HIV; since immune protection will likely require both B-cell and T-cell responses (Moore and Burton, *Nat. Med.* 10:769-71 (2004)), Env vaccine antigens will also need to be optimized separately to elicit antibody responses. T-cell-directed vaccine components, in contrast, can target the more conserved proteins, but even the most conserved HIV-1 proteins are diverse enough that variation is an issue. Artificial central-sequence vaccine approaches (e.g., consensus sequences, in which every amino acid is found in a plurality of sequences, or maximum likelihood reconstructions of ancestral sequences (Gaschen et al, *Science* 296:2354-60 (2002), Gao et al, *J. Virol.* 79:1154-63 (2005), Doria-Rose et al, *J. Virol.* 79:11214-24 (2005), Weaver et al, *J. Virol.*, in press)) are promising; nevertheless, even centralized strains provide limited coverage of HIV-1 variants, and consensus-based reagents fail to detect many autologous T-cell responses (Altfeld et al, *J. Virol.* 77:7330-40 (2003)).

**[0010]** Single amino acid changes can allow an epitope to escape T-cell surveillance; since many T-cell epitopes differ between HIV-1 strains at one or more positions, potential responses to any single vaccine antigen are limited. Whether a particular mutation results in escape depends upon the specific epitope/T-cell combination, although some changes broadly affect between-subtype cross-reactivity (Norris et al, *AIDS Res. Hum. Retroviruses* 20:315-25 (2004)). Including multiple variants in a polyvalent vaccine could enable responses to a broader range of circulating variants, and could also prime the immune system against common escape mutants (Jones et al, *J. Exp. Med.* 200:1243-56 (2004)).



Escape from one T-cell receptor may create a variant that is susceptible to another (Allen et al, *J. Virol*, 79:12952-60 (2005), Feeney et al, *J. Immunol.* 174:7524-30 (2005)), so stimulating polyclonal responses to epitope variants may be beneficial (Killian et al, *Aids* 19:887-96 (2005)). Escape mutations that inhibit processing (Milicic et al, *J. Immunol.* 175:4618-26 (2005)) or HLA binding (Ammaranond et al, *AIDS Res. Hum. Retroviruses* 21:395-7 (2005)) cannot be directly countered by a T-cell with a different specificity, but responses to overlapping epitopes may block even some of these escape routes.

**[0011]** The present invention relates to a polyvalent vaccine comprising several "mosaic" proteins (or genes encoding these proteins). The candidate vaccine antigens can be cocktails of  $k$  composite proteins ( $k$  being the number of sequence variants in the cocktail), optimized to include the maximum number of potential T-cell epitopes in an input set of viral proteins. The mosaics are generated from natural sequences: they resemble natural proteins and include the most common forms of potential epitopes. Since CD8+ epitopes are contiguous and typically nine amino-acids long, sets of mosaics can be scored by "coverage" of nonamers (9-mers) in the natural sequences (fragments of similar lengths are also well represented). 9-Mers not found at least three times can be excluded. This strategy provides the level of diversity coverage achieved by a massively polyvalent multiple-peptide vaccine but with important advantages: it allows vaccine delivery as intact proteins or genes, excludes low-frequency or unnatural epitopes that are not relevant to circulating strains, and its intact protein antigens are more likely to be processed as in a natural infection.

#### SUMMARY OF THE INVENTION

**[0012]** In general, the present invention relates to an immunogenic composition. More specifically, the invention relates to a polyvalent immunogenic composition (e.g., an HIV vaccine), and to methods of using same. The invention further relates to methods that involve the use of a genetic algorithm to design sets of polyvalent antigens suitable for use as vaccines.

**[0013]** Objects and advantages of the present invention will be clear from the description that follows.

#### BRIEF DESCRIPTION OF THE DRAWINGS

**[0014]** FIGS. 1A-1F. The upper bound of potential epitope coverage of the HIV-1 M group. The upper bound for population coverage of 9-mers for increasing numbers of variants is shown, for  $k=1-8$  variants. A sliding window of length nine was applied across aligned sequences, moving down by one position. Different colors denote results for different numbers of sequences. At each window, the coverage given by the  $k$  most common 9-mers is plotted for Gag (FIGS. 1A and 1B), Nef (FIGS. 1C and 1D) and Env gp120 (FIGS. 1E and 1F). Gaps inserted to maintain the alignment are treated as characters. The diminishing returns of adding more variants are evident, since, as  $k$  increases, increasingly rare forms are added. In FIGS. 1A, 1C and 1E, the scores for each consecutive 9-mer are plotted in their natural order to show how diversity varies in different protein regions; both p24 in the center of Gag and the central region of Nef are particularly highly conserved. In FIGS. 1B, 1D and 1F, the scores for each 9-mer are reordered by coverage (a strategy also used in FIG. 4), to provide a sense of the overall population coverage of a given protein. Coverage of gp120, even with 8 variant 9-mers, is particularly poor (FIGS. 1E and 1F).

**[0015]** FIGS. 2A-2C. Mosaic initialization, scoring, and optimization. FIG. 2A) A set of  $k$  populations is generated by random 2-point recombination of natural sequences (1-6 populations of 50-500 sequences each have been tested). One sequence from each population is chosen (initially at random) for the mosaic cocktail, which is subsequently optimized. The cocktail sequences are scored by computing coverage (defined as the mean fraction of natural-sequence 9-mers included in the cocktail, averaged over all natural sequences in the input data set). Any new sequence that covers more epitopes will increase the score of the whole cocktail. FIG. 2B) The fitness score of any individual sequence is the coverage of a cocktail containing that sequence plus the current representatives from other populations. FIG. 2C) Optimization: 1) two "parents" are chosen: the higher-scoring of a randomly chosen pair of recombined sequences, and either (with 50% probability) the higher-scoring sequence of a second random pair, or a randomly chosen natural sequence. 2) Two-point recombination between the two parents is used to generate a "child" sequence. If the child contains unnatural or rare 9-mers, it is immediately rejected, otherwise it is scored (Gaschen et al, *Science* 296:2354-2360 (2002)). If the score is higher than that of any of four randomly-selected population members, the child is inserted in the population in place of the weakest of the four, thus evolving an improved population; 4) if its score is a new high score, the new child replaces the current cocktail member from its population. Ten cycles of child generation are repeated for each population in turn, and the process iterates until improvement stalls.

**[0016]** FIG. 3. Mosaic strain coverage for all HIV proteins. The level of 9-mer coverage achieved by sets of four mosaic proteins for each HIV protein is shown, with mosaics optimized using either the M group or the C subtype. The fraction of C subtype sequence 9-mers covered by mosaics optimized on the C subtype (within-clade optimization) is shown in gray. Coverage of 9-mers found in non-C subtype M-group sequences by subtype-C-optimized mosaics (between-clade coverage) is shown in white. Coverage of subtype C sequences by M-group optimized mosaics is shown in black. B clade comparisons gave comparable results (data not shown).

**[0017]** FIGS. 4A-4F. Coverage of M group sequences by different vaccine candidates, nine-mer by nine-mer. Each plot presents site-by-site coverage (i.e., for each nine-mer) of an M-group natural-sequence alignment by a single tri-valent vaccine candidate. Bars along the x-axis represent the proportion of sequences matched by the vaccine candidate for a given alignment position: 9/9 matches (in red), 8/9 (yellow), 7/9 (blue). Aligned 9-mers are sorted along the x-axis by exact-match coverage value. 656 positions include both the complete Gag and the central region of Nef. For each alignment position, the maximum possible matching value (i.e. the proportion of aligned sequences without gaps in that nine-mer) is shown in gray. FIG. 4A) Non-optimal natural sequences selected from among strains being used in vaccine studies (Kong et al, *J. Virol.* 77:12764-72 (2003)) including an individual clade A, B, and C viral sequences (Gag: GenBank accession numbers AF004885, K03455, and U52953; Nef core: AF069670, K02083, and U52953). FIG. 4B) Optimum set of natural sequences [isolates US2 (subtype B, USA), 70177 (subtype C, India), and 99TH.R2399 (subtype CRF15\_01B, Thailand); accession numbers AY173953, AF533131, and AF530576] selected by choosing the single sequence with maximum coverage, followed by the sequence



that had the best coverage when combined with the first (i.e. the best complement), and so on, selected for M group coverage FIG. 4C) Consensus sequence cocktail (M group, B- and C-subtypes). FIG. 4D) 3 mosaic sequences, FIG. 4E) 4 mosaic sequences, FIG. 4F) 6 mosaic sequences. FIGS. 4D-4F were all optimized for M group coverage.

**[0018]** FIGS. 5A and 5B. Overall coverage of vaccine candidates: coverage of 9-mers in C clade sequences using different input data sets for mosaic optimization, allowing different numbers of antigens, and comparing to different candidate vaccines. Exact (blue), 8/9 (one-off; red), and 7/9 (two-off; yellow) coverage was computed for mono- and polyvalent vaccine candidates for Gag (FIG. 5A) and Nef (core) (FIG. 5B) for four test situations: within-clade (C-clade-optimized candidates scored for C-clade coverage), between-clade (B-clade-optimized candidates scored for C-clade coverage), global-against-single-subtype (M-group-optimized candidates scored for C-clade coverage), global-against-global (M-group-optimized candidates scored for global coverage). Within each set of results, vaccine candidates are grouped by number of sequences in the cocktail (1-6); mosaic sequences are plotted with darker colors. “Non-opt” refers to one set of sequences moving into vaccine trials (Kong et al, *J. Virol.* 77:12764-72 (2003)); “mosaic” denotes sequences generated by the genetic algorithm; “opt. natural” denotes intact natural sequences selected for maximum 9-mer coverage; “MBC consensus” denotes a cocktail of 3 consensus sequences, for M-group, B-subtype, and C-subtype. For ease of comparison, a dashed line marks the coverage of a 4-sequence set of M-group mosaics (73.7-75.6%). Over 150 combinations of mosaic-number, virus subset, protein region, and optimization and test sets were tested. The C clade/B clade/M group comparisons illustrated in this figure are generally representative of within-clade, between-clade, and M group coverage. In particular, levels of mosaic coverage for B and C clade were very similar, despite there being many more C clade sequences in the Gag collection, and many more B clade sequences in the Nef collection (see FIG. 6 for a full B and C clade comparison). There were relatively few A and G clade sequences in the alignments (24 Gag, 75 Nef), and while 9-mer coverage by M-group optimized mosaics was not as high as for subtypes for B and C clades (4-mosaic coverage for A and G subtypes was 63% for Gag, 74% for Nef), it was much better than a non-optimal cocktail (52% Gag, 52% for Nef).

**[0019]** FIGS. 6A and 6B. Overall coverage of vaccine candidates: coverage of 9-mers in B-clade, C-clade, and M-group sequences using different input data sets for mosaic, optimization, allowing different numbers of antigens, and comparing to different candidate vaccines. Exact (blue), 8/9 (one-off; red), and 7/9 (two-off; yellow) coverage was computed for mono- and polyvalent vaccine candidates for Gag (FIG. 6A) and Nef (core) (FIG. 6B) for seven test situations: within-clade (B- or C-clade-optimized candidates scored against the same clade), between-clade (B- or C-clade-optimized candidates scored against the other clade), global vaccine against single subtype (M-group-optimized candidates scored against B- or C-clade), global vaccine against global viruses (M-group-optimized candidates scored against all M-group sequences). Within each set of results, vaccine candidates are grouped by number of sequences in the cocktail (1-6); mosaic sequences are plotted with darker colors. “Non-opt” refers to a particular set of natural sequences previously proposed for a vaccine (Kong, W. P. et al. *J Virol* 77, 12764-72 (2003)); “mosaic” denotes sequences generated by the genetic algorithm; “opt. natural” denotes intact natural sequences selected for maximum 9-mer coverage; “MBC consensus”

denotes a cocktail of 3 consensus sequences, for M-group, B-subtype, and C-subtype. A dashed line is shown at the level of exact-match M-group coverage for a 4-valent mosaic set optimized on the M-group.

**[0020]** FIGS. 7A and 7B. The distribution of 9-mers by frequency of occurrence in natural, consensus, and mosaic sequences. Occurrence counts (y-axis) for different 9-mer frequencies (x-axis) for vaccine cocktails produced by several methods. FIG. 7A: frequencies from 0-60% (for 9-mer frequencies >60%, the distributions are equivalent for all methods). FIG. 7B: Details of low-frequency 9-mers. Natural sequences have large numbers of rare or unique-to-isolate 9-mers (bottom right, FIGS. 7A and 7B); these are unlikely to induce useful vaccine responses. Selecting optimal natural sequences does select for more common 9-mers, but rare and unique 9-mers are still included (top right, FIGS. 7A and 7B). Consensus cocktails, in contrast, under-represent uncommon 9-mers, especially below 20% frequency (bottom left, FIGS. 7A and 7B). For mosaic sequences, the number of lower-frequency 9-mers monotonically increases with the number of sequences (top left, each panel), but unique-to-isolate 9-mers are completely excluded (top left of right panel: \* marks the absence of 9-mers with frequencies <0.005).

**[0021]** FIGS. 8A-8D. HLA binding potential of vaccine candidates. FIGS. 8A and 8B) HLA binding motif counts. FIGS. 8C and 8D) number of unfavorable amino acids. In all graphs: natural sequences are marked with black circles ( $\lambda$ ); consensus sequences with blue triangles ( $\sigma$ ); inferred ancestral sequences with green squares ( $\nu$ ); and mosaic sequences with red diamonds ( $\heartsuit$ ). Left panel (FIGS. 8A and 8C) shows HLA-binding-motif counts (FIG. 8A) and counts of unfavorable amino acids (FIG. 8C) calculated for individual sequences; Right panel (FIGS. 8B and 8D) shows HLA binding motifs counts (FIG. 8B) and counts of unfavorable amino acids (FIG. 8D) calculated for sequence cocktails. The top portion of each graph (box-and-whiskers graph) shows the distribution of respective counts (motif counts or counts of unfavorable amino acids) based either on alignment of M group sequences (for individual sequences, FIGS. 8A and 8C) or on 100 randomly composed cocktails of three sequences, one from each A, B and C subtypes (for sequence cocktails, FIGS. 8B and 8D). The alignment was downloaded from the Los Alamos HIV database. The box extends from the 25 percentile to the 75 percentile, with the line at the median. The whiskers extending outside the box show the highest and lowest values. Amino acids that are very rarely found as C-terminal anchor residues are G, S, T, P, N, Q, D, E, and H, and tend to be small, polar, or negatively charged (Yusim et al, *J. Virol.* 76:8757-8768 (2002)). Results are shown for Gag, but the same qualitative results hold for Nef core and complete Nef. The same procedure was done for supertype motifs with results qualitatively similar to the results for HLA binding motifs (data not shown).

**[0022]** FIG. 9. Mosaic protein sets limited to 4 sequences (k=4), spanning Gag and the central region of Nef: optimized for subtype B, subtype C, and the M group. Figure discloses SEQ ID NOS 1-84, respectively, in order of appearance.

**[0023]** FIG. 10. Mosaic sets for Env and Pol. Figure discloses SEQ ID NOS 85-168, respectively, in order of appearance.

#### DETAILED DESCRIPTION OF THE INVENTION

**[0024]** The present invention results from the realization that a polyvalent set of antigens comprising synthetic viral proteins, the sequences of which provide maximum coverage of non-rare short stretches of circulating viral sequences, constitutes a good vaccine candidate. The invention provides



a “genetic algorithm” strategy to create such sets of polyvalent antigens as mosaic blends of fragments of an arbitrary set of natural protein sequences provided as inputs. In the context of HIV, the proteins Gag and the inner core (but not the whole) of Nef are ideal candidates for such antigens. The invention further provides optimized sets for these proteins.

**[0025]** The genetic algorithm strategy of the invention uses unaligned protein sequences from the general population as an input data set, and thus has the virtue of being “alignment independent”. It creates artificial mosaic proteins that resemble proteins found in nature—the success of the consensus antigens in small animals models suggest this works well. 9 Mers are the focus of the in studies described herein, however, different length peptides can be selected depending on the intended target. In accordance with the present approach, 9 mers (for example) that do not exist in nature or that are very rare can be excluded—this is an improvement relative to consensus sequences since the latter can contain some 9 mers (for example) that have not been found in nature, and relative to natural strains that almost invariably contain some 9 mers (for example) that are unique to that strain. The definition of fitness used for the genetic algorithm is that the most “fit” polyvalent cocktail is the combination of mosaic strains that gives the best coverage (highest fraction of perfect matches) of all of the 9 mers in the population and is subject to the constraint that no 9 mer is absent or rare in the population.

**[0026]** The mosaics protein sets of the invention can be optimized with respect to different input data sets—this allows use of current data to assess virtues of a subtype or region specific vaccines from a T cell perspective. By way of example, options that have been compared include:

**[0027]** 1) Optimal polyvalent mosaic sets based on M group, B clade and C clade. The question presented was how much better is intra-clade coverage than inter-clade or global.

**[0028]** 2) Different numbers of antigens: 1, 3, 4, 6

**[0029]** 3) Natural strains currently in use for vaccine protocols just to exemplify “typical” strains (Merck, VRC)

**[0030]** 4) Natural strains selected to give the best coverage of 9-mers in a population

**[0031]** 5) Sets of consensus: A+B+C.

**[0032]** 6) Optimized cocktails that include one “given” strain in a polyvalent antigen, one ancestral+3 mosaic strains, one consensus+3 mosaic strains.

**[0033]** 7) Coverage of 9 mers that were perfectly matched was compared with those that match 8/9, 7/9, and 6/9 or less.

This is a computationally difficult problem, as the best set to cover one 9-mer may not be the best set to cover overlapping 9-mers.

**[0034]** It will be appreciated from a reading of this disclosure that the approach described herein can be used to design peptide reagents to test HIV immune responses, and be applied to other variable pathogens as well. For example, the present approach can be adapted to the highly variable virus Hepatitis C.

**[0035]** The proteins/polypeptides/peptides (“immunogens”) of the invention can be formulated into compositions with a pharmaceutically acceptable carrier and/or adjuvant using techniques well known in the art. Suitable routes of administration include systemic (e.g. intramuscular or subcutaneous), oral, intravaginal, intrarectal and intranasal.

**[0036]** The immunogens of the invention can be chemically synthesized and purified using methods which are well known to the ordinarily skilled artisan. The immunogens can also be synthesized by well-known recombinant DNA techniques.

**[0037]** Nucleic acids encoding the immunogens of the invention can be used as components of, for example, a DNA vaccine wherein the encoding sequence is administered as naked DNA or, for example, a minigene encoding the immunogen can be present in a viral vector. The encoding sequences can be expressed, for example, in mycobacterium, in a recombinant chimeric adenovirus, or in a recombinant attenuated vesicular stomatitis virus. The encoding sequence can also be present, for example, in a replicating or non-replicating adenoviral vector, an adeno-associated virus vector, an attenuated mycobacterium tuberculosis vector, a Bacillus Calmette Guerin (BCG) vector, a vaccinia or Modified Vaccinia Ankara (MVA) vector, another pox virus vector, recombinant polio and other enteric virus vector, Salmonella species bacterial vector, Shigella species bacterial vector, Venezuelan Equine Encephalitis Virus (VEE) vector, a Semliki Forest Virus vector, or a Tobacco Mosaic Virus vector. The encoding sequence, can also be expressed as a DNA plasmid with, for example, an active promoter such as a CMV promoter. Other live vectors can also be used to express the sequences of the invention. Expression of the immunogen of the invention can be induced in a patient’s own cells, by introduction into those cells of nucleic acids that encode the immunogen, preferably using codons and promoters that optimize expression in human cells. Examples of methods of making and using DNA vaccines are disclosed in U.S. Pat. Nos. 5,580,859, 5,589,466, and 5,703,055.

**[0038]** It will be appreciated that adjuvants can be included in the compositions of the invention (or otherwise administered to enhance the immunogenic effect). Examples of suitable adjuvants include TRL-9 agonists, TRL-4 agonists, and TRL-7, 8 and 9 agonist combinations (as well as alum). Adjuvants can take the form of oil and water emulsions. Squalene adjuvants can also be used.

**[0039]** The composition of the invention comprises an immunologically effective amount of the immunogen of this invention, or nucleic acid sequence encoding same, in a pharmaceutically acceptable delivery system. The compositions can be used for prevention and/or treatment of virus infection (e.g. HIV infection). As indicated above, the compositions of the invention can be formulated using adjuvants, emulsifiers, pharmaceutically-acceptable carriers or other ingredients routinely provided in vaccine compositions. Optimum formulations can be readily designed by one of ordinary skill in the art and can include formulations for immediate release and/or for sustained release, and for induction of systemic immunity and/or induction of localized mucosal immunity (e.g. the formulation can be designed for intranasal, intravaginal or intrarectal administration). As noted above, the present compositions can be administered by any convenient route including subcutaneous, intranasal, oral, intramuscular, or other parenteral or enteral route. The immunogens can be administered as a single dose or multiple doses. Optimum immunization schedules can be readily determined by the ordinarily skilled artisan and can vary with the patient, the composition and the effect sought.

**[0040]** The invention contemplates the direct use of both the immunogen of the invention and/or nucleic acids encoding same and/or the immunogen expressed as as indicated above. For example, a minigene encoding the immunogen can be used as a prime and/or boost.



**[0041]** The invention includes any and all amino acid sequences disclosed herein, as well as nucleic acid sequences encoding same (and nucleic acids complementary to such encoding sequences).

**[0042]** Specifically disclosed herein are vaccine antigen sets optimized for single B or C subtypes, targeting regional epidemics, as well as for all HIV-1 variants in global circulation [the HIV-1 Main (M) group]. In the study described in the Example that follows, the focus is on designing polyvalent vaccines specifically for T-cell responses. HIV-1 specific T-cells are likely to be crucial to an HIV-1-specific vaccine response: CTL responses are correlated with slow disease progression in humans (Oxenius et al, *J. Infect. Dis.* 189: 1199-1208 (2004)), and the importance of CTL responses in non-human primate vaccination models is well-established. Vaccine elicited cellular immune responses help control pathogenic SIV or SHIV, and reduce the likelihood of disease after challenge with pathogenic virus (Barouch et al, *Science* 290:486-492 (2000)). Temporary depletion of CD8+ T cells results in increased viremia in SIV-infected rhesus macaques (Schmitz et al, *Science* 283:857-860 (1999)). Furthermore, the evolution of escape mutations has been associated with disease progression, indicating that CTL responses help constrain viral replication in vivo (Barouch et al, *J. Virol.* 77:7367-7375 (2003)), and so vaccine-stimulated memory responses that could block potential escape routes may be of value. While the highly variable Envelope (Env) is the primary target for neutralizing antibodies against HIV, and vaccine antigens will also need to be tailored to elicit these antibody responses (Moore & Burton, *Nat. Med.* 10:769-771 (2004)), T-cell vaccine components can target more conserved proteins to trigger responses that are more likely to cross-react. But even the most conserved HIV-1 proteins are diverse enough that variation will be an issue. Artificial central-sequence vaccine approaches, consensus and ancestral sequences (Gaschen et al, *Science* 296:2354-2360 (2002), Gao et al, *J. Virol.* 79:1154-1163 (2005), Doria-Rose et al, *J. Virol.* 79:11214-11224 (2005)), which essentially “split the differences” between strains, show promise, stimulating responses with enhanced cross-reactivity compared to natural strain vaccines (Gao et al, *J. Virol.* 79:1154-1163 (2005)) (Liao et al. and Weaver et al., submitted.) Nevertheless, even central strains cover the spectrum of HIV diversity to a very limited extent, and consensus-based peptide reagents fail to detect many autologous CD8+ T-cell responses (Altfeld et al, *J. Virol.* 77:7330-7340 (2003)).

**[0043]** A single amino acid substitution can mediate T-cell escape, and as one or more amino acids in many T-cell epitopes differ between HIV-1 strains, the potential effectiveness of responses to any one vaccine antigen is limited. Whether a particular mutation will diminish T-cell cross-reactivity is epitope- and T-cell-specific, although some changes can broadly affect between-clade cross-reactivity (Norris et al, *AIDS Res. Hum. Retroviruses* 20:315-325 (2004)). Including more variants in a polyvalent vaccine could enable responses to a broader range of circulating variants. It could also prime the immune system against common escape variants (Jones et al, *J. Exp. Med.* 200:1243-1256 (2004)); escape from one T-cell receptor might create a variant that is susceptible to another (Lee et al, *J. Exp. Med.* 200:1455-1466 (2004)), thus stimulating polyclonal responses to epitope variants may be beneficial (Killian et al, *AIDS* 19:887-896 (2005)). Immune escape involving avenues that inhibit processing (Milicic et al, *J. Immunol.* 175:4618-4626 (2005)) or HLA binding (Ammaranond et al, *AIDS Res. Hum. Retroviruses* 21:395-397 (2005)) prevent epitope presentation, and in such cases the escape variant

could not be countered by a T-cell with a different specificity. However, it is possible the presence of T-cells that recognize overlapping epitopes may in some cases block these even escape routes.

**[0044]** Certain aspects of the invention can be described in greater detail in the non-limiting Example that follows.

## EXAMPLE

### Experimental Details

**[0045]** HIV-1 sequence data. The reference alignments from the 2005 HIV sequence database (<http://hiv.lanl.gov>), which contain one sequence per person, were used, supplemented by additional recently available C subtype Gag and Nef sequences from Durban, South Africa (GenBank accession numbers AY856956-AY857186) (Kiepiela et al, *Nature* 432:769-75 (2004)). This set contained 551 Gag and 1,131 NefM group sequences from throughout the globe; recombinant sequences were included as well as pure subtype sequences for exploring M group diversity. The subsets of these alignments that contained 18 A, 102 B, 228 C, and 6 G subtype (Gag), and 62 A, 454 B, 284 C, and 13 G subtype (Nef) sequences were used for within- and between-single-clade optimizations and comparisons.

**[0046]** The genetic algorithm. GAs are computational analogues of biological processes (evolution, populations, selection, recombination) used to find solutions to problems that are difficult to solve analytically (Holland, *Adaptation in Natural and Artificial Systems: An Introductory Analysis with Applications to Biology, Control, and Artificial Intelligence*, (M.I.T. Press, Cambridge, Mass. (1992))). Solutions for a given input are “evolved” through a process of random modification and selection according to a “fitness” (optimality) criterion. GAs come in many flavors; a “steady-state co-evolutionary multi-population” GA was implemented. “Steady-state” refers to generating one new candidate solution at a time, rather than a whole new population at once; and “co-evolutionary” refers to simultaneously evolving several distinct populations that work together to form a complete solution. The input is an unaligned set of natural sequences; a candidate solution is a set of  $k$  pseudo-natural “mosaic” sequences, each of which is formed by concatenating sections of natural sequences. The fitness criterion is population coverage, defined as the proportion of all 9-amino-acid sequence fragments (potential epitopes) in the input sequences that are found in the cocktail.

**[0047]** To initialize the GA (FIG. 2),  $k$  populations of  $n$  initial candidate sequences are generated by 2-point recombination between randomly selected natural sequences. Because the input natural sequences are not aligned, “homologous” crossover is used: crossover points in each sequence are selected by searching for short matching strings in both sequences; strings of  $c-1=8$ , were used where a typical epitope length is  $c=9$ . This ensures that the recombined sequences resemble natural proteins: the boundaries between sections of sequence derived from different strains are seamless, the local sequences spanning the boundaries are always found in nature, and the mosaics are prevented from acquiring large insertions/deletions or unnatural combinations of amino acids. Mosaic sequence lengths fall within the distribution of natural sequence lengths as a consequence of mosaic construction: recombination is only allowed at identical regions, reinforced by an explicit software prohibition against excessive lengths to prevent reduplication of repeat regions. (Such “in frame” insertion of reduplicated epitopes could provide another way of increasing coverage without generating unnatural 9-mers, but their inclusion would create “unnatu-



ral” proteins.) Initially, the cocktail contains one randomly chosen “winner” from each population. The fitness score for any individual sequence in a population is the coverage value for the cocktail consisting of that sequence plus the current winners from the other populations. The individual fitness of any sequence in a population therefore depends dynamically upon the best sequences found in the other populations.

**[0048]** Optimization proceeds one population at a time. For each iteration, two “parent” sequences are chosen. The first parent is chosen using “2-tournament” selection: two sequences are picked at random from the current population, scored, and the better one is chosen. This selects parents with a probability inversely proportional to their fitness rank within the population, without the need to actually compute the fitness of all individuals. The second parent is chosen in the same way (50% of the time), or is selected at random from the set of natural sequences. 2-point homologous crossover between the parents is then used to generate a “child” sequence. Any child containing a 9-mer that was very rare in the natural population (found less than 3 times) is rejected immediately. Otherwise, the new sequence is scored, and its fitness is compared with the fitnesses of four randomly chosen sequences from the same population. If any of the four randomly chosen sequences has a score lower than that of the new sequence, it is replaced in the population by the new sequence. Whenever a sequence is encountered that yields a better score than the current population “winner”, that sequence becomes the winner for the current population and so is subsequently used in the cocktail to evaluate sequences in other populations. A few such optimization cycles (typically 10) are applied to each population in turn, and this process continues cycling through the populations until evolution stalls (i.e., no improvement has been made for a defined number of generations). At this point, the entire procedure is restarted using newly generated random starting populations, and the restarts are continued until no further improvement is seen. The GA was run on each data set with  $n=50$  or  $500$ ; each run was continued until no further improvement occurred for 12-24 hours on a 2 GHz Pentium processor. Cocktails were generated having  $k=1, 3, 4,$  or  $6$  mosaic sequences.

**[0049]** The GA also enables optional inclusion of one or more fixed sequences of interest (for example, a consensus) in the cocktail and will evolve the other elements of the cocktail in order to optimally complement that fixed strain. As these solutions were suboptimal, they are not included here. An additional program selects from the input file the  $k$  best natural strains that in combination provide the best population coverage.

**[0050]** Comparison with other polyvalent vaccine candidates. Population coverage scores were computed for other potential mono- or polyvalent vaccines to make direct comparisons with the mosaic-sequence vaccines, tracking identities with population 9-mers, as well as similarities of 8/9 and 7/9 amino acids. Potential vaccine candidates based on natural strains include single strains (for example, a single C strain for a vaccine for southern Africa (Williamson et al, AIDS Res. Hum. Retroviruses 19:133-44 (2003))) or combinations of natural strains (for example, one each of subtype A, B, and C (Kong et al, J. Virol. 77:12764-72 (2003))). To date, natural-strain vaccine candidates have not been systematically selected to maximize potential T-cell epitope coverage; vaccine candidates were picked from the literature to be representative of what could be expected from unselected vaccine candidates. An upper bound for coverage was also determined using only intact natural strains: optimal natural-sequence cocktails were generated by selecting the single sequence with the best coverage of the dataset, and then

successively adding the most complementary sequences up to a given  $k$ . The comparisons included optimal natural-sequence cocktails of various sizes, as well as consensus sequences, alone or in combination (Gaschen et al, Science 296:2354-60 (2002)), to represent the concept of central, synthetic vaccines. Finally, using the fixed-sequence option in the GA, consensus-plus-mosaic combinations in the comparisons; these scores were essentially equivalent to all-mosaic combinations were included for a given  $k$  (data not shown). The code used for performing these analyses are available at: <ftp://ftp-t10/pub/btk/mosaics>.

## Results

**[0051]** Protein Variation. In conserved HIV-1 proteins, most positions are essentially invariant, and most variable positions have only two to three amino acids that occur at appreciable frequencies, and variable positions are generally well dispersed between conserved positions. Therefore, within the boundaries of a CD8+ T-cell epitope (8-12 amino acids, typically nine), most of the population diversity can be covered with very few variants. FIG. 1 shows an upper bound for population coverage of 9-mers (stretches of nine contiguous amino acids) comparing Gag, Nef, and Env for increasing numbers of variants, sequentially adding variants that provide the best coverage. In conserved regions, a high degree of population coverage is achieved with 2-4 variants. By contrast, in variable regions like Env, limited population coverage is possible even with eight variants. Since each new addition is rarer, the relative benefits of each addition diminish as the number of variants increases.

**[0052]** Vaccine design optimization strategies. FIG. 1 shows an idealized level of 9-mer coverage. In reality, high-frequency 9-mers often conflict: because of local co-variation, the optimal amino acid for one 9-mer may differ from that for an overlapping 9-mer. To design mosaic protein sets that optimize population coverage, the relative benefits of each amino acid must be evaluated in combination with nearby variants. For example, Alanine (Ala) and Glutamate (Glu) might each frequently occur in adjacent positions, but if the Ala-Glu combination is never observed in nature, it should be excluded from the vaccine. Several optimization strategies were investigated: a greedy algorithm, a semi-automated compatible-9mer assembly strategy, an alignment-based genetic algorithm (GA), and an alignment-independent GA.

**[0053]** The alignment-independent GA generated mosaics with the best population coverage. This GA generates a user-specified number of mosaic sequences from a set of unaligned protein sequences, explicitly excluding rare or unnatural epitope-length fragments (potentially introduced at recombination breakpoints) that could induce non-protective vaccine-antigen-specific responses. These candidate vaccine sequences resemble natural proteins, but are assembled from frequency-weighted fragments of database sequences recombined at homologous breakpoints (FIG. 2); they approach maximal coverage of 9-mers for the input population.

**[0054]** Selecting HIV protein regions for an initial mosaic vaccine. The initial design focused on protein regions meeting specific criteria: i) relatively low variability, ii) high levels of recognition in natural infection, iii) a high density of known epitopes and iv) either early responses upon infection or CD8+ T-cell responses associated with good outcomes in infected patients. First, an assessment was made of the level of 9-mer coverage achieved by mosaics for different HIV proteins (FIG. 3). For each protein, a set of four mosaics was generated using either the M group or the B- and C-subtypes alone; coverage was scored on the C subtype. Several results are notable: i) within-subtype optimization provides the best



within-subtype coverage, but substantially poorer between-subtype coverage—nevertheless, B-subtype-optimized mosaics provide better C-subtype coverage than a single natural B subtype protein (Kong et al, *J. Virol.* 77:12764-72 (2003)); ii) Pal and Gag have the most potential to elicit broadly cross-reactive responses, whereas Rev, Tat, and Vpu have even fewer conserved 9-mers than the highly variable Env protein, iii) within-subtype coverage of M-group-optimized mosaic sets approached coverage of within-subtype optimized sets, particularly for more conserved proteins.

**[0055]** Gag and the central region of Nef meet the four criteria listed above. Nef is the HIV protein most frequently recognized by T-cells (Frahm et al, *J. Virol.* 78:2187-200 (2004)) and the target for the earliest response in natural infection (Lichterfeld et al, *Aids* 18:1383-92 (2004)). While overall it is variable (FIG. 3), its central region is as conserved as Gag (FIG. 1). It is not yet clear what optimum proteins for inclusion in a vaccine might be, and mosaics could be designed to maximize the potential coverage of even the most variable proteins (FIG. 3), but the prospects for global coverage are better for conserved proteins. Improved vaccine protection in macaques has been demonstrated by adding Rev, Tat, and Nef to a vaccine containing Gag, Pol, and Env (Hel et al, *J. Immunol.* 176:85-96 (2006)), but this was in the context of homologous challenge, where variability was not an issue. The extreme variability of regulatory proteins in circulating virus populations may preclude cross-reactive responses; in terms of conservation, Pol, Gag (particularly p24) and the central region of Nef (HXB2 positions 65-149) are promising potential immunogens (FIGS. 1,3). Pol, however, is infrequently recognized during natural infection (Frahm et al, *J. Virol.* 78:2187-200 (2004)), so it was not included in the initial immunogen design. The conserved portion of Nef that were included contains the most highly recognized peptides in HIV-1 (Frahm et al, *J. Virol.* 78:2187-200 (2004)), but as a protein fragment, would not allow Nef's immune inhibitory functions (e.g. HLA class I down-regulation (Blagoveshchenskaya, *Cell* 111:853-66 (2002))). Both Gag and Nef are densely packed with overlapping well-characterized CD8+ and CD4+ T-cell epitopes, presented by many different HLA molecules (<http://www.hiv.lanl.gov/content/immunology/maps/maps.html>), and Gag-specific CD8+ (Masemola et al, *J. Virol.* 78:3233-43 (2004)) and CD4+ (Oxenius et al, *J. Infect. Dis.* 189:1199-208 (2004)) T-cell responses have been associated with low viral set points in infected individuals (Masemola et al, *J. Virol.* 78:3233-43 (2004)).

**[0056]** To examine the potential impact of geographic variation and input sample size, a limited test was done using published subtype C sequences. The subtype C Gag data were divided into three sets of comparable size—two South African sets (Kiepiela et al, *Nature* 432:769-75 (2004)), and one non-South-African subtype C set. Mosaics were optimized independently on each of the sets, and the resulting mosaics were tested against all three sets. The coverage of 9-mers was slightly better for identical training and test sets (77-79% 9/9 coverage), but essentially equivalent when the training and test sets were the two different South African data sets (73-75%), or either of the South African sets and the non-South African C subtype sequences (74-76%). Thus between- and within-country coverage approximated within-clade coverage, and in this case no advantage to a country-specific C subtype mosaic design was found.

**[0057]** Designing mosaics for Gag and Nef and comparing vaccine strategies. To evaluate within- and between-subtype cross-reactivity for various vaccine design strategies, a calculation was made of the coverage they provided for natural

M-Group sequences. The fraction of all 9-mers in the natural sequences that were perfectly matched by 9-mers in the vaccine antigens were computed, as well as those having 8/9 or 7/9 matching amino acids, since single (and sometimes double) substitutions within epitopes may retain cross-reactivity. FIG. 4 shows M group coverage per 9-mer in Gag and the central region of Nef for cocktails designed by various strategies: a) three non-optimal natural strains from the A, B, and C subtypes that have been used as vaccine antigens (Kong et al, *J. Virol.* 77:12764-72 (2003)); b) three natural strains that were computationally selected to give the best M group coverage; c) M group, B subtype, and C subtype consensus sequences; and, d,e,f) three, four and six mosaic proteins. For cocktails of multiple strains, sets of k=3, k=4, and k=6, the mosaics clearly perform the best, and coverage approaches the upper bound for k strains. They are followed by optimally selected natural strains, the consensus protein cocktail, and finally, non-optimal natural strains. Allowing more antigens provides greater coverage, but gains for each addition are reduced as k increases (FIGS. 1 and 4).

**[0058]** FIG. 5 summarizes total coverage for the different vaccine design strategies, from single proteins through combinations of mosaic proteins, and compares within-subtype optimization to M group optimization. The performance of a single mosaic is comparable to the best single natural strain or a consensus sequence. Although a single consensus sequence out-performs a single best natural strain, the optimized natural-sequence cocktail does better than the consensus cocktail: the consensus sequences are more similar to each other than are natural strains, and are therefore somewhat redundant. Including even just two mosaic variants, however, markedly increases coverage, and four and six mosaic proteins give progressively better coverage than polyvalent cocktails of natural or consensus strains. Within-subtype optimized mosaics perform best—with four mosaic antigens 80-85% of the 9-mers are perfectly matched—but between-subtype coverage of these sets falls off dramatically, to 50-60%. In contrast, mosaic proteins optimized using the full M group give coverage of approximately 75-80% for individual subtypes, comparable to the coverage of the M group as a whole (FIGS. 5 and 6). If imperfect 8/9 matches are allowed, both M group optimized and within-subtype optimized mosaics approach 90% coverage.

**[0059]** Since coverage is increased by adding progressively rarer 9-mers, and rare epitopes may be problematic (e.g., by inducing vaccine-specific immunodominant responses), an investigation was made of the frequency distribution of 9-mers in the vaccine constructs relative to the natural sequences from which they were generated. Most additional epitopes in a cocktail compared to a k=4 cocktail are low-frequency (<0.1, FIG. 7). Despite enhancing coverage, these epitopes are relatively rare, and thus responses they induce might draw away from vaccine responses to more common, thus more useful, epitopes. Natural-sequence cocktails actually have fewer occurrences of moderately low-frequency epitopes than mosaics, which accrue some lower frequency 9-mers as coverage is optimized. On the other hand, the mosaics exclude unique or very rare 9-mers, while natural strains generally contain 9-mers present in no other sequence. For example, natural M group Gag sequences had a median of 35 (range 0-148) unique 9-mers per sequence. Retention of HLA-anchor motifs was also explored, and anchor motif frequencies were found to be comparable between four mosaics and three natural strains. Natural antigens did exhibit an increase in number of motifs per antigen, possibly due to inclusion of strain-specific motifs (FIG. 8).



**[0060]** The increase in ever-rarer epitopes with increasing  $k$ , coupled with concerns about vaccination-point dilution and reagent development costs, resulted in the initial production of mosaic protein sets limited to 4 sequences ( $k=4$ ), spanning Gag and the central region of Nef, optimized for subtype B, subtype C, and the M group (these sequences are included in FIG. 9; mosaic sets for Env and Pol are set forth in FIG. 10). Synthesis of various four-sequence Gag-Nef mosaics and initial antigenicity studies are underway. In the initial mosaic vaccine, targeted are just Gag and the center of the Nef protein, which are conserved enough to provide excellent global population coverage, and have the desirable properties described above in terms of natural responses (Bansal et al, *Aids* 19:241-50 (2005)). Additionally, including B subtype p24 variants in Elispot peptide mixtures to detect natural CTL responses to infection significantly enhanced both the number and the magnitude of responses detected supporting the idea that including variants of even the most conserved proteins will be useful. Finally, cocktails of proteins in a polyvalent HIV-1 vaccine given to rhesus macaques did not interfere with the development of robust responses to each antigen (Seaman et al, *J. Virol.* 79:2956-63 (2005)), and antigen cocktails did not produce antagonistic responses in murine models (Singh et al, *J. Immunol.* 169:6779-86 (2002)), indicating that antigenic mixtures are appropriate for T-cell vaccines.

**[0061]** Even with mosaics, variable proteins like Env have limited coverage of 9-mers, although mosaics improve coverage relative to natural strains. For example three M group natural proteins, one each selected from the A, B, and C clades, and currently under study for vaccine design (Seaman et al, *J. Virol.* 79:2956-63 (2005)) perfectly match only 39% of the 9-mers in M group proteins, and 65% have at least 8/9 matches. In contrast, three M group Env mosaics match 47% of 9-mers perfectly, and 70% have at least an 8/9 match. The code written to design polyvalent mosaic antigens is available, and could readily be applied to any input set of variable proteins, optimized for any desired number of antigens. The code also allows selection of optimal combinations of  $k$  natural strains, enabling rational selection of natural antigens for polyvalent vaccines. Included in Table 1 are the best natural strains for Gag and Nef population coverage of current database alignments.

TABLE 1

Natural sequence cocktails having the best available 9-mer coverage for different genes, subtype sets, and numbers of sequences
Gag, B-subtype, 1 natural sequence
B.US.86.AD87_AF004394 Gag, B-subtype, 3 natural sequences
B.US.86.AD87_AF004394 B.US.97.Ac_06_AY247251 B.US.88.WR27_AF286365 Gag, B-subtype, 4 natural sequences
B.US.86.AD87_AF004394 B.US.97.Ac_06_AY247251 B.US._R3_PDC1_AY206652 B.US.88.WR27_AF286365 Gag, B-subtype, 6 natural sequences
B.CN._CNHN24_AY180905 B.US.86.AD87_AF004394 B.US.97.Ac_06_AY247251 B.US._P2_AY206654 B.US._R3_PDC1_AY206652 B.US.88.WR27_AF286365

TABLE 1-continued

Natural sequence cocktails having the best available 9-mer coverage for different genes, subtype sets, and numbers of sequences
Gag, C-subtype, 1 natural sequence
C.IN._70177_AF533131 Gag, C-subtype, 3 natural sequences
C.ZA.97.97ZA012 C.ZA.x.04ZASK161B1 C.IN.-.70177_AF533131 Gag, C-subtype, 4 natural sequences
C.ZA.97.97ZA012 C.ZA.x.04ZASK142B1 C.ZA.x.04ZASK161B1 C.IN._70177_AF533131 Gag, C-subtype, 6 natural sequences
C.ZA.97.97ZA012 C.ZA.x.04ZASK142B1 C.ZA.x.04ZASK161B1 C.BW.99.99BWMC168_AF443087 C.IN._70177_AF533131 C.IN._MYA1_AF533139 Gag, M-group, 1 natural sequence
C.IN._70177_AF533131 Gag, M-group, 3 natural sequences
B.US.90.US2_AY173953 C.IN.-.70177_AF533131 15_01B.TH.99.99TH_R2399_AF530576 Gag, M-group, 4 natural sequences
B.US.90.US2_AY173953 C.IN._70177_AF533131 C.IN.93.93IN999_AF067154 15_01B.TH.99.99TH_R2399_AF530576 Gag, M-group, 6 natural sequences
C.ZA.x.04ZASK138B1 B.US.90.US2_AY173953 B.US._WT1_PDC1_AY206656 C.IN._70177_AF533131 C.IN.93.93IN999_AF067154 15_01B.TH.99.99TH_R2399_AF530576 Nef (central region), B-subtype, 1 natural sequence
B.GB.94.028jh_94_1_NP_AF129346 Nef (central region), B-subtype, 3 natural sequences
B.GB.94.028jh_94_1_NP_AF129346 B.KR.96.96KCS4_AY121471 B.FR.83.HXB2_K03455 Nef (central region), B-subtype, 4 natural sequences
B.GB.94.028jh_94_1_NP_AF129346 B.KR.96.96KCS4_AY121471 B.US.90.E90NEF_U43108 B.FR.83.HXB2_K03455 Nef (central region), B-subtype, 6 natural sequences
B.GB.94.028jh_94_1_NP_AF129346 B.KR.02.02HYJ3_AY121454 B.KR.96.96KCS4_AY121471 B.CN._RL42_U71182 B.US.90.E90NEF_U43108 B.FR.83.HXB2_K03455 Nef (central region), C-subtype, 1 natural sequence
C.ZA.04.04ZASK139B1 Nef (central region), C-subtype, 3 natural sequences
C.ZA.04.04ZASK180B1 C.ZA.04.04ZASK139B1 C.ZA._ZASW15_AF397568



TABLE 1-continued

Natural sequence cocktails having the best available 9-mer coverage for different genes, subtype sets, and numbers of sequences
Nef (central region), C-subtype, 4 natural sequences
C.ZA.97.ZA97004_AF529682 C.ZA.04.04ZASK180B1 C.ZA.04.04ZASK139B1 C.ZA._ZASW15_AF397568
Nef (central region), C-subtype, 6 natural sequences
C.ZA.97.ZA97004_AF529682 C.ZA.00.1192M3M C.ZA.04.04ZASK180B1 C.ZA.04.04ZASK139B1 C.04ZASK184B1 C.ZA._ZASW15_AF397568
Nef (central region), M-group, 1 natural sequence
B.GB.94.028jh_94_1_NP_AF129346
Nef (central region), M-group, 3 natural sequences
02_AG.CM._98CM1390_AY265107 C.ZA.03.03ZASK020B2 B.GB.94.028jh_94_1_NP_AF129346
Nef (central region), M-group, 4 natural sequences
02_AG.CM._98CM1390_AY265107 01A1.MM.99.mCSW105_AB097872 C.ZA.03.03ZASK020B2 B.GB.94.028jh_94_1_NP_AF129346
Nef (central region), M-group, 6 natural sequences
02_AG.CM._98CM1390_AY265107 01A1.MM.99.mCSW105_AB097872 C.ZA.03.03ZASK020B2 C.03ZASK111B1 B.GB.94.028jh_94_1_NP_AF129346 B.KR.01.01CWS2_AF462757

**[0063]** A centralized (consensus or ancestral) gene and protein strategy has been proposed previously to address HIV diversity (Gaschen et al, Science 296:2354-2360 (2002)). Proof-of-concept for the use of artificial genes as immunogens has been demonstrated by the induction of both T and B cell responses to wild-type HIV-1 strains by group M consensus immunogens (Gaschen et al, Science 296:2354-2360 (2002), Gao et al, J. Virol. 79:1154-63 (2005), Doria-Rose et al, J. Virol. 79:11214-24 (2005), Weaver et al, J. Virol., in press)). The mosaic protein design improves on consensus or natural immunogen design by co-optimizing reagents for a polyclonal vaccine, excluding rare CD8+ T-cell epitopes, and incorporating variants that, by virtue of their frequency at the population level, are likely to be involved in escape pathways.

**[0064]** The mosaic antigens maximize the number of epitope-length variants that are present in a small, practical number of vaccine antigens. The decision was made to use multiple antigens that resemble native proteins, rather than linking sets of concatenated epitopes in a poly-epitope pseudo-protein (Hanke et al, Vaccine 16:426-35 (1998)), reasoning that in vivo processing of native-like vaccine antigens will more closely resemble processing in natural infection, and will also allow expanded coverage of overlapping epitopes. T-cell mosaic antigens would be best employed in the context of a strong polyvalent immune response; improvements in other areas of vaccine design and a combination of the best strategies, incorporating mosaic antigens to cover diversity, may ultimately enable an effective cross-reactive vaccine-induced immune response against HIV-1.

**[0065]** All documents and other information sources cited above are hereby incorporated in their entirety by reference.

## SEQUENCE LISTING

The patent application contains a lengthy "Sequence Listing" section. A copy of the "Sequence Listing" is available in electronic form from the USPTO web site (<http://seqdata.uspto.gov/?pageRequest=docDetail&DocID=US20110150915A1>). An electronic copy of the "Sequence Listing" will also be available from the USPTO upon request and payment of the fee set forth in 37 CFR 1.19(b)(3).

**[0062]** Summarizing, the above-described study focuses on the design of T-cell vaccine components to counter HIV diversity at the moment of infection, and to block viral escape routes and thereby minimize disease progression in infected individuals. The polyvalent mosaic protein strategy developed here for HIV-1 vaccine design could be applied to any variable protein, to other pathogens, and to other immunological problems. For example, incorporating a minimal number of variant peptides into T-cell response assays could markedly increase sensitivity without excessive cost: a set of k mosaic proteins provides the maximum coverage possible for k antigens.

1. A polypeptide or protein comprising at least one sequence of amino acids set forth in FIG. 9 or FIG. 10.

2. The polypeptide or protein according to claim 1 wherein said polypeptide or protein comprises at least one sequence of amino acids set forth in FIG. 9.

3. The polypeptide or protein according to claim 1 wherein said polypeptide or protein comprises at least one sequence of amino acids set forth in FIG. 10.

4. A nucleic acid comprising a nucleotide sequence that encodes the polypeptide or protein according to claim 1.

5. The nucleic acid according to claim 4 wherein said nucleic acid encodes at least one sequence of amino acids set forth in FIG. 9.



6. The nucleic acid according to claim 4 wherein said nucleic acid encodes at least one sequence of amino acids set forth in FIG. 10.

7. A vector comprising the nucleic acid according to claim 4.

8. The vector according to claim 7 wherein said vector is a viral vector.

9. A composition comprising at least one polypeptide or protein according to claim 1 and a carrier.

10. A composition comprising at least one nucleic acid according to claim 4 and a carrier.

11. A method of inducing an immune response in a mammal comprising administering to said mammal an amount of at least one polypeptide or protein according to claim 1 sufficient to effect said induction.

12. A method of inducing an immune response in a mammal comprising administering to said mammal an amount of at least one nucleic acid according to claim 4 sufficient to effect said induction.

13. A polyvalent vaccine comprising a multiplicity of HIV-1 polypeptides, each of said polypeptides comprising a computationally-designed homologous recombinant mosaic blend of epitopes of natural HIV Gag, Nef, Env or Pol protein sequences,

wherein the length of each of said epitopes is about 9 amino acids,

wherein the length of each of said polypeptides is within the range of natural HIV protein lengths,

wherein the epitopes are linked within each polypeptide via homologous crossover so the unnatural combinations of amino acids are not present,

wherein said epitopes are natural, commonly occurring T-cell epitopes, and

wherein about 3 to about 6 different types of polypeptides are present in said vaccine.

\* \* \* \* \*