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(54) Title: RECOMBINATION OF INSERTION MODIFIED NUCLEIC ACIDS

N-terminal shuffled extein
N-terminal intein
C-terminal intein
C-terminal shuffled extein
Mature shuffled protein

(57) Abstract

Methods of modulating, tuning and improving hybridization properties and recombination properties of molecules for use in nucleic acid shuffling procedures, relates recombination mixtures and methods of modulating, tuning, improving and evolving splicing of RNAs and proteins are provided.
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RECOMBINATION OF INSERTION MODIFIED NUCLEIC ACIDS

CROSS-REFERENCES TO RELATED APPLICATIONS
This application is related to USSN 60/122,943 "RECOMBINATION OF INSERTION MODIFIED NUCLEIC ACIDS" by Patten et al., filed March 5, 1999, the disclosure of which is incorporated herein by reference. This application is also related to USSN 60/142,299 "RECOMBINATION OF INSERTION MODIFIED NUCLEIC ACIDS" by Patten et al., filed July 02, 1999, the disclosure of which is incorporated herein by reference. This application is also related to USSN 60/164,617 "RECOMBINATION OF INSERTION MODIFIED NUCLEIC ACIDS" by Patten et al., filed November 10, 1999, the disclosure of which is incorporated herein by reference. This case is also related to Patten et al. "ENCRYPTION OF TRAITS USING SPLIT GENE SEQUENCES AND ENGINEERED GENETIC ELEMENTS" USSN 60/164,618, Filed November 10, 1999.

The present application claims priority to and the benefit of each of these prior applications, pursuant to 35. U.S.C. 119(e).

FIELD OF THE INVENTION
The present invention relates to molecular shuffling, and to splicing of nucleic acids and proteins.

BACKGROUND OF THE INVENTION
Nucleic acid shuffling provides for the rapid evolution of nucleic acids, in vitro and in vivo. Rapid evolution provides for the commercial production of encoded molecules (e.g., nucleic acids and proteins) with new and/or improved properties. Proteins and nucleic acids of industrial, agricultural and therapeutic value can be created or improved through shuffling procedures. A number of publications by the inventors and their co-workers describe nucleic acid shuffling and applications of this technology. For example, Stemmer et al. (1994) "Rapid Evolution of a Protein" Nature 370:389-391; Stemmer (1994) "DNA Shuffling by Random Fragmentation and Reassembly: in vitro Recombination for Molecular Evolution," Proc. Natl. Acad. USA 91:10747-10751; Stemmer U.S. Patent No. 5,603,793 METHODS FOR IN VITRO RECOMBINATION; Stemmer et al. U.S. Pat. No. 5,830,721 DNA MUTAGENESIS BY RANDOM FRAGMENTATION AND REASSEMBLY; Stemmer et al., U.S. Pat. No. 5,811,238
METHODS FOR GENERATING POLYNUCLEOTIDES HAVING DESIRED
CHARACTERISTICS BY ITERATIVE SELECTION AND RECOMBINATION describe,
e.g., in vivo and in vitro nucleic acid, DNA and protein shuffling in a variety of formats,
e.g., by repeated cycles of mutagenesis, shuffling and selection, as well as methods of
generating libraries of displayed peptides and antibodies.

Applications of DNA shuffling technology have also been developed by the
inventors and their co-workers. In addition to the publications noted above, Minshull et al.,
U.S. Pat. No. 5,837,458 METHODS AND COMPOSITIONS FOR CELLULAR AND
METABOLIC ENGINEERING provides for the evolution of metabolic pathways and the
enhancement of bioprocessing through recursive shuffling techniques. Crameri et al. (1996),
“Construction And Evolution Of Antibody-Phage Libraries By DNA Shuffling” Nature
Medicine 2(1):100-103 describe, e.g., antibody shuffling for antibody phage libraries.
Additional details regarding DNA Shuffling can also be found in WO95/22625, WO97/
20078, WO96/33207, WO97/33957, WO98/27230, WO97/35966, WO98/31837,

Physical nucleic acid shuffling techniques (as opposed, e.g., to “in silico”
methods which are performed, at least in part, by manipulation of character strings in a
computer) rely upon actual recombination between physical nucleic acids, whether the
format is an in vitro or an in vivo format. Recombination occurs at a relatively high
frequency, e.g., where there are complementary nucleic acids between strands to be
recombined. Thus, nucleic acids to be recombined are typically e.g., about 70%
identical/complementary in sequence over regions of, e.g., about 30-40 nucleotides. It
would be desirable to be able to recombine low homology, or even non-homologous
sequences, thereby increasing access to the potential sequence space encoded by
recombinant nucleic acids resulting from shuffling methods. For example, for proteins
which are commercially valuable, it would be desirable to be able to gain access to a
recombination/mutation spectrum which is different than that of the native protein to
provide for greater diversity in products produced by the various available shuffling
strategies.

Similarly, nucleic acid recombination generally can be difficult to modulate,
resulting in regions of high or low crossover frequency between two different targets for
recombination. The crossover frequency for a particular pairing of sequences on two
different targets is one feature that mediates the recombinant nucleic acids that result from recombination methods. Improved methods of modulating the recombination frequency at potential recombination sites would be desirable to weight/bias recombination product outcomes.

In general, new techniques which faciltate, improve or add levels of control to recombination methods are highly desirable. In particular, techniques which permit shuffling of divergent nucleic acids, or which provide for modulation and tuning of shuffling rates are desirable. The present invention provides such significant new recombination protocols, as well as other features which will be apparent upon complete review of this disclosure.

**SUMMARY OF THE INVENTION**

The present invention provides a number of new nucleic acid recombination formats for nucleic acid shuffling. In the methods, a number of insertion sequences are inserted into one or more parental nucleic acid to provide a modified target nucleic acid substrate for recombination and subsequent mutation. The number, type and placement of such insertion sequences provides for the ability to shuffle nucleic acids with little or no homology other than the insertion sequences. In addition, these insertion sequences provide for the ability to modulate or "tune" recombination frequencies between target nucleic acids. The methods typically take advantage of self-splicing, trans-splicing or use cellular machinery to remove the insertion sequences from final coded nucleic acids or proteins, e.g., where the insertion sequences are introns, inteins, proteolyzed polypeptide sequences or the like. The insertion sequences can also comprise markers, molecular tags, or the like, e.g., for purification of encoded molecules or can serve to allow for expression of otherwise toxic proteins (e.g., RNases, Dnases, restriction enzymes, proteases, lipases, recombinases, ligases, polymerases, etc.) e.g., in a form where an intein is excised in vivo. Similarly, in vitro expression of insertion modified sequences can result in the production of these and other proteins in vitro, e.g., using in vitro expression systems.

Methods of shuffling two target nucleic acids (i.e., a first and a second target nucleic acid) are provided. In the methods, a first and a second target nucleic acid are provided, e.g., by cloning, PCR amplification, synthesis, isolation from an environmental source (soil, air, water, etc.), or other methods. At least one of the first and second target nucleic acids (and typically both) have a plurality of homologous or non-homologous
insertion nucleic acid sequences, such as one or more intron (e.g., self-splicing bacterial, eukaryotic or trans-splicing intron), intein, subsequence removed by site specific recombination (e.g., similar to V-D-J recombination for antibody production), or the like, optionally including intron splicing enhancers or the like. The target nucleic acids are recombined, producing a shuffled recombinant nucleic acid.

In addition to providing for new recombination methods per se, the invention also provides methods of producing selected proteins and RNAs, for any of the purposes that such proteins and RNAs are ordinarily produced. For example, in one aspect, a first shuffled nucleic acid subsequence encoding a first portion of the selected protein and a second nucleic acid subsequence encoding a second portion of the selected protein is provided. The nucleic acids can be on the same strand (as in cis-mediated reactions) or on different strands (as in trans mediated reactions). The first and second subsequences are expressed to produce a first protein subsequence and a second protein subsequence, which are spliced to produce the selected protein. Commonly, more than two subsequences are spliced, e.g., 3, 4, 5, 6, 7, 8, 9, 10 or more sequences, as set forth herein. The splicing reaction can be in cis or in trans (or both) and can be in vitro or in vivo (or both). Splicing can occur by spontaneous or controlled mechanisms.

Similarly, in RNA production methods, a first shuffled nucleic acid subsequence encoding a first portion of the selected RNA is provided and a second nucleic acid subsequence encoding a second portion of the selected RNA is also provided. Again, these subsequences can be on the same or on different molecules (depending on whether cis or trans splicing is employed). The first and second nucleic acid subsequences, or RNA copies thereof, are spliced to produce the selected RNA, which can encode a useful RNA (e.g., an antisense, or sense molecule or ribozyme) or the RNA can encode a protein. The intein and RNA shuffling/production methods are combinable, i.e., the spliced RNA molecules can encode intein-extein sequences which are spliced at the protein level to produce a useful protein.

In general, a parental nucleic acid can be broken into several exons or exteins by incorporation of a number of introns or inteins into the sequence of the parental nucleic acid. For example, the target nucleic acid resulting from incorporation of insertion sequences into the parental nucleic acid can have, e.g., about 5, 10, 15, 20, 30, 50, 100 or
more “mini exons,” or “mini exteins” separated by a corresponding number of insertion sequences.

In shuffling reactions, first and second target nucleic acids are optionally derived from a first and second parental nucleic acid which are sufficiently different in sequence that they do not substantially hybridize in solution. For example, the first and second target nucleic acids can be derived by integration of a plurality of insertion sequences into the first and second parental nucleic acid. The first and second parental nucleic acid can be, e.g., less than 50%, or less than e.g., 40%, or less than e.g., 30%, or less than e.g., 25%, or less than e.g., 15% identical over the full length of the first and second parental nucleic acid, when the first and second nucleic acids are aligned for maximum identity.

The insertion nucleic acid sequences can modulate a recombination frequency between the first and second target nucleic acid. For example, by placing an intron into a parental sequence, the recombination efficiency of nucleic acid subsequences to either side of the intron can be decreased. Similarly, placing homologous mini introns within the parental sequences provides sites for recombination within the resulting targets, e.g., where the targets display regions of low similarity in non-intronic sequences.

Insertion sequences can also modulate expression in one or more cell type, e.g., where the insertion sequences comprise one or more enhancer or other regulatory sequence. Similarly, insertion sequences optionally comprise splicing enhancer sequences (e.g., ISEs, such as the chicken cardiac troponin T (cTNT) ISE) to facilitate splicing.

Essentially any nucleic acid can be a parental nucleic acid with which insertion sequences can be combined to produce a target nucleic acid for splicing. Example sequences include parental nucleic acids corresponding a gene or cDNA encoding EPO, a gene or cDNA encoding an insulin protein, a gene or cDNA encoding a peptide hormone, a gene or cDNA encoding a cytokine, a gene or cDNA encoding an epidermal growth factor, a gene or cDNA encoding a fibroblast growth factor, a gene or cDNA encoding a hepatocyte growth factor, a gene or cDNA encoding insulin-like growth factor, a gene or cDNA encoding an interferon, a gene or cDNA encoding an interleukin, a gene or cDNA encoding a keratinocyte growth factor, a gene or cDNA encoding a leukemia inhibitory factor, a gene or cDNA encoding oncostatin M, a gene or cDNA encoding PD-ECSF, a gene or cDNA encoding PDGF, a gene or cDNA encoding pleiotropin, a gene or cDNA
encoding SCF, a gene or cDNA encoding c-kit ligand, a gene or cDNA encoding VEGF, a
gene or cDNA encoding G-CSF, a gene or cDNA encoding an oncogene, a gene or cDNA
encoding a tumor suppressor, a gene or cDNA encoding a steroid hormone receptor, a gene
or cDNA encoding a plant hormone, a gene or cDNA encoding a disease resistance gene, a
5 gene or cDNA encoding an herbicide resistance gene, a gene or cDNA encoding a bacterial
gene, a gene or cDNA encoding a monoxygenase, a gene or cDNA encoding a protease, a
gene or cDNA encoding a nuclease, an antibody, a peptide ligand, an angiogenesis inhibitor,
a gene or cDNA encoding a lipase, a gene or cDNA encoding a C-X-C chemokine, a gene
or cDNA encoding a C-C chemokine, a gene or cDNA encoding an antibody V gene, a gene
10 or cDNA encoding a cysteine knot protein such as TGFβ, NGF, PDGFβ or the like, a gene or
cDNA encoding a TNKα family member, a gene or cDNA encoding CNTF, a gene or
cDNA encoding 4F, and/or gene or cDNA encoding an RNase.

The methods herein are amenable to both physical recombination of nucleic acids and to virtual or “in silico” recombination of character strings representing nucleic
15 acids, e.g., in a computer. Following complete or partial sequence recombination in silico,
target nucleic acids, or nucleic acids derived from the target nucleic acids can be
synthesized. Such synthetic nucleic acids can be recombined, cloned, selected or otherwise
manipulated in the same manner as any other nucleic acid.

A variety of techniques can be used to produce target nucleic acids
20 comprising insertion sequences. Such methods include chemical synthesis, PCR
concatemerization, in silico character string formation or generation, and the like. For
example, in one embodiment, insertion of the plurality of insertion nucleic acid sequences
into one or more of the first and second parental nucleic acid sequences is performed by
physically joining a plurality of subsequences of the first or second parental nucleic acid
25 sequences to the plurality of insertion nucleic acid sequences.

As noted, the addition of insertion sequences to parental nucleic acids can
modify or modulate the recombination of resulting target nucleic acids. Similarly, the
addition of insertion sequences can alter the hybridization properties of resulting target
sequences. For example, even non-homologous parental nucleic acids can be made to
30 hybridize by the addition of a sufficient number and appropriate arrangement of insertion
sequences. Similarly, a target nucleic acid derived from a parental sequence can be made
which does not hybridize under a selected set of conditions (e.g., stringent hybridization
conditions) to the parental nucleic acid. As noted above, such insertion sequences can be used to tune recombination rates between selected regions of a target nucleic acid, e.g., where a particular region is targeted for an increased or decreased recombination rate.

The target and parental nucleic acids can have dramatically different hybridization properties as a result of the insertion sequences being present in the target nucleic acids. The target nucleic acids can be prevented from hybridizing to the parents by inclusion of the target sequences, or, conversely, one or more target sequence can even be made to hybridize to one or more parent, thereby controlling the recombination properties of resulting nucleic acid shuffling reactions. Thus, in one embodiment, the first and second parental nucleic acid sequences hybridize under stringent conditions, and the first and second target nucleic acids do not hybridize under stringent conditions. Similarly, in another embodiment, the first and second parental nucleic acid sequences do not hybridize under stringent conditions, while the first and second target nucleic acids hybridize under stringent conditions. In yet another embodiment, the first and second nucleic target nucleic acid hybridize under stringent conditions, while the first target nucleic acid does not hybridize under stringent conditions to the second parental nucleic acid, or wherein the second target nucleic acid does not hybridize under stringent conditions to the first parental nucleic acid. Similarly, in one embodiment, the first or second parental nucleic acid hybridizes to a third nucleic acid under stringent conditions, where the first and second target nucleic acids do not hybridize under stringent conditions to the third nucleic acid. A variety of other modifications in hybridization due to the number and arrangement of insertion sequences will be apparent upon complete review.

Recombinant nucleic acids generated by recombining nucleic acid sequences comprising insertion subsequences can, of course, be recombined or shuffled, cloned, amplified, expressed in vivo or in vitro, synthesized, or otherwise modified using any available naturally mediated or laboratory-mediated technique. For example, in one embodiment, a shuffled recombinant nucleic acid made by recombining one or more target nucleic acid comprising a plurality of insertion sequences with one or more additional nucleic acid(s) is recombined with a third nucleic acid. The resulting secondary shuffled recombinant nucleic acid can be selected for a desired trait or property using any available selection method. In general, any recombinant nucleic acid can be selected for a desired trait or property.
Recombinant nucleic acids are also optionally expressed in a cell or in vitro, thereby producing a nucleic acid or protein. In one embodiment, the expressed protein can comprise intein and extein sequences. Typically, the intein (some times referred to as an "intervening protein sequence") is excised from an expressed protein sequences.

Concomitantly, the ligation of the flanking sequences (exteins) form a mature "extein protein" which is, optionally, active in one or more cell or in one or more in vitro reaction or system. Thus, expressed proteins can be proteolytically cleaved and ligated to produce an active protein, and/or to remove an intein from an expressed protein. This ligation reaction can occur in both cis- and trans-splicing reaction formats. Reactions occur in vitro or in vivo for cis or trans splicing inteins. For additional details regarding trans splicing of introns and inteins, see, Patten et al. “ENCRYPTION OF TRAITS USING SPLIT GENE SEQUENCES AND ENGINEERED GENETIC ELEMENTS” USSN 60/164,618 Filed November 10, 1999.

The presence of insertion sequences can be used to modulate recombination rates between regions of nucleic acids. For example, the cross over frequency between two points on a first and second target nucleic acids can typically be increased by placing insertion sequences between the two points. This is desirable, e.g., where low linkage rates between regions of nucleic acids to be recombined are desired, e.g., where one wishes to separately evolve different functional domains or elements of the nucleic acid.

Recombinant nucleic acids can be modified by removal of insertion sequences to improve expression or facilitate cloning of any final product. For example, where a nucleic acid encodes a plurality of intronic insertion sequences, the encoded mRNA can be reverse transcribed and the resulting cDNA cloned or otherwise manipulated. It should be noted that this process can result in a cDNA which does not hybridize to the recombinant nucleic acid comprising the introns. Indeed, the cDNA can be the result of several rounds of selection and recombination, resulting in a cDNA with a highly unique sequence which does not hybridize under e.g., stringent conditions, to any previously known sequence. Thus, sequence space which is inaccessible between two known nucleic acids is accessible by this procedure, resulting in recombinant products that could not otherwise be obtained.

The final product produced by any of the procedures herein can be a DNA (e.g., a genomic DNA, an artificial DNA, a cDNA, or the like), an RNA, an mRNA, a viral
RNA, a sn RNA, a tRNA, an rRNA, a gRNA, a protein, a proteolytically cleaved protein, a 
protein fragment, a spliced protein or any other molecule that can be encoded by a nucleic 
acid, including e.g., metabolic products and the like. As noted, target sequences can 
comprise homologous or non homologous nucleic acid subsequences which can be 
separated by homologous or non homologous insertion sequences. The target nucleic acids 
to be recombined can be homologous relative to each other, or comprise homologous and 
non-homologous sequences relative to each other. The nucleic acids can be present in 
vectors such as expression vectors, or can be free in solution. 

The nucleic acids to be recombined can be present in recombination 
mixtures. For example, one recombination mixture of the invention includes a first target 
nucleic acid comprising a plurality of insertion subsequences. Typically, the mixture also 
includes a second target nucleic acid having at least one region of sequence similarity to the 
first nucleic acid. The second target nucleic acid typically also includes a plurality of 
insertion subsequences. 

In one format, a recombination mixture resulting from fragmenting a first 
target nucleic acid comprising a plurality of insertion subsequences, and a second target 
nucleic acid comprising at least one region of sequence similarity to the first target nucleic 
acid is provided. For example, the first and second target nucleic acids can be fragmented 
with a DNase, or, e.g., cleaved chemically to produce nucleic acid fragments. Similarly, the 
first and second target nucleic acids can be “fragmented” by chemically synthesizing 
fragments of the first and second target nucleic acid. 

Recombinant nucleic acids produced by recombining the recombination 
mixtures of the invention are also provided. For example, the first or second nucleic acid 
can include one or more subsequence corresponding to one or more subsequence from one 
or more gene or cDNA such as a gene or cDNA encoding EPO, a gene or cDNA encoding 
an insulin protein, a gene or cDNA encoding a peptide hormone, a gene or cDNA encoding 
a cytokine, a gene or cDNA encoding an epidermal growth factor, a gene or cDNA 
encoding a fibroblast growth factor, a gene or cDNA encoding a hepatocyte growth factor, a 
gene or cDNA encoding insulin-like growth factor, a gene or cDNA encoding an interferon, 
a gene or cDNA encoding an interleukin, a gene or cDNA encoding a keratinocyte growth 
factor, a gene or cDNA encoding a leukemia inhibitory factor, a gene or cDNA encoding 
oncostatin M, a gene or cDNA encoding PD-ECSF, a gene or cDNA encoding PDGF, a
gene or cDNA encoding pleiotropin, a gene or cDNA encoding SCF, a gene or cDNA
encoding c-Kit ligand, a gene or cDNA encoding VEGF, a gene or cDNA encoding G-CSF,
a gene or cDNA encoding an oncogene, a gene or cDNA encoding a tumor suppressor, a
gene or cDNA encoding a steroid hormone receptor, a gene or cDNA encoding a plant
hormone, a gene or cDNA encoding a disease resistance gene, a gene or cDNA encoding an
herbicide resistance gene, a gene or cDNA encoding a bacterial gene, a gene or cDNA
encoding a monoxygenase, a gene or cDNA encoding a protease, a gene or cDNA
encoding a nuclease, a gene or cDNA encoding an RNase, and/or a gene or cDNA encoding
a lipase. Of course, many other nucleic acids/proteins can be made or modified by the
methods herein. The resulting recombinant nucleic acid can also comprise activities and
subsequences which correspond to these nucleic acids.

In one aspect, the invention provides methods of recombining a plurality of
sequence domains from a plurality of homologous or non-homologous nucleic acid
sequences. In the methods, a pre-mRNA comprising a plurality of sequence domains is
provided which correspond to a plurality of different parental nucleic acid sequences. The
pre-mRNA is alternatively spliced to produce a plurality of different mRNAs comprising a
plurality of different sets of sequence domains. Typically, the pre-mRNA has between
about 6 and about 20 exons or exteins, e.g., where the pre-mRNA has a plurality of mini
exons or exteins. Most typically, the plurality of different mRNAs are selected for a desired
trait or property. Optionally, the methods include cloning one or more of the plurality of
different mRNAs.

In this alternative splicing/recombination strategy, the methods typically
include recombining one or more of: the plurality of different mRNAs, the pre-mRNA, a
DNA encoding the mRNA, and a DNA encoding the pre-mRNA, with one or more
additional nucleic acid.

In one embodiment, the pre-mRNA is provided to a cell by transducing or
transfecting the cell with a vector comprising a DNA encoding the pre-mRNA. As
discussed throughout, in vitro formats are also available.

The present invention also provides methods of making a nucleic acid with a
desired splicing phenotype. In the methods, a plurality of homologous nucleic acids are
provided, each comprising a plurality of insertion nucleic acid sequences. The plurality of
homologous nucleic acids are recombined to produce a library of recombinant nucleic acids,
which are selected for production of a desired or selected mRNA or protein (or product thereof) when the selected recombinant nucleic acid is expressed in vitro or in a cell. As with any nucleic acid noted above, this selected nucleic acid is optionally recombined with an additional nucleic acid and the resulting secondary recombinant nucleic acid selected for production of a desired mRNA or protein (or product thereof).

The nucleic acids noted above which include insertion sequences will typically comprise as many as 10 insertion sequences and as many as 10 flanking sequences (e.g., exons or exteins) or more. Insertion nucleic acid sequences include those derived from bacterial introns, eukaryotic introns and archaeabacterial introns, as well as bacterial inteins, eukaryotic inteins and archaeabacterial inteins. The nucleic acids are recombined in vitro or vivo.

The present invention also provides apparatus, integrated systems and kits for practicing the methods herein, e.g., comprising use of the recombination mixtures herein, containers, instruction sets for practicing the methods herein, and the like.

**BRIEF DESCRIPTION OF THE FIGURES**

Fig. 1 is a schematic of a splicing procedure for creating non-hybridizing genes.

Fig. 2 is a schematic of an exemplar shuffling strategy for Cytokine G, an arbitrarily designated cytokine.

Fig. 3 is a schematic of a domain shuffling strategy.

Fig. 4 is a schematic of a trans-splicing library strategy.

Fig. 5 is a schematic of a combined trans-splicing/classical breeding approach to making a library, using roses as an example plant.

**DEFINITIONS**

The following definitions supplement those in the art.

An “insertion nucleic acid subsequence” is a nucleic acid subsequence found in a full-length nucleic acid sequence, which subsequence is derived from a nucleic acid sequence different from sequences immediately flanking the subsequence within the full-length nucleic acid sequence, or which subsequence is present in a different arrangement relative to the flanking subsequences from those found in nature. Examples include introns from genes e.g., which are heterologous to the sequences immediately flanking the
subsequence, inteins from genes e.g., which are heterologous to the sequences immediately flanking the subsequence, and the like. Examples of cis and trans splicing RNAs and proteins are well known in the art.

Nucleic acids and nucleic acid sequences are “homologous” when they are derived, naturally or artificially, from a common ancestor sequence. During natural evolution, this occurs when two or more descendant sequences diverge from a parent sequence over time, i.e., due to mutation and natural selection. Under artificial conditions, divergence occurs, e.g., in one of at least three different general ways. First, a given sequence can be artificially recombined with another sequence, as occurs, e.g., during cloning or during shuffling methods, e.g., to produce one or more descendant nucleic acids. Second, a nucleic acid can be synthesized de novo, by synthesizing a nucleic acid which varies in sequence from a selected parental nucleic acid sequence. Third, sequences can be modified e.g., in a computer by applying genetic operators which modify a given character string representing a nucleic acid.

When there is no explicit knowledge about the ancestry of two nucleic acids, homology is typically inferred by sequence comparison between two sequences. Where two nucleic acid sequences show sequence similarity it is inferred that the two nucleic acids share a common ancestor. The precise level of sequence similarity required to establish homology varies in the art depending on a variety of factors. For purposes of this disclosure, two nucleic acids are considered homologous where they share sufficient sequence identity to allow direct recombination to occur between the two nucleic acid molecules. Typically, nucleic acids utilize regions of close similarity spaced roughly the same distance apart to permit recombination to occur. The recombination can be in vitro or in vivo. Thus, two non-homologous sequences can be made “homologous” in the methods of the invention by placing homologous insertion sequences into the non-homologous sequences.

Nucleic acids “hybridize” when they associate, typically in solution, or with one of the nucleic acids fixed to a solid support. Nucleic acids hybridize due to a variety of well characterized physico-chemical forces, such as hydrogen bonding, solvent exclusion, base stacking and the like. An extensive guide to the hybridization of nucleic acids is found in Tijssen (1993) Laboratory Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Acid Probes part I chapter 2 “Overview of principles of
hybridization and the strategy of nucleic acid probe assays,” Elsevier, New York, as well as in Ausubel, supra.

Two nucleic acids “correspond” when they have the same sequence, or when one nucleic acid is a subsequence of the other, or when one sequence is derived, by natural or artificial manipulation from the other.

Two nucleic acids are “recombined” when sequences from each of the two nucleic acids are combined in a progeny nucleic acid. Two sequences are “directly” recombined when both of the nucleic acids are substrates for recombination. Two sequences are “indirectly recombined” when the sequences are recombined using an intermediate such as a sequence comprising a plurality of insertion sequences. For indirect recombination, no more than one of the sequences to be recombined is an actual substrate for recombination.

A collection of “fragmented nucleic acids” is a collection of nucleic acids derived by cleaving one or more parental nucleic acids (e.g., with a nuclease, or via chemical cleavage), or by synthesizing fragments of the one or more parent nucleic acids as oligonucleotides.

A “full-length protein” is a protein having substantially the same domains as a corresponding protein encoded by a natural gene. The protein can have modified sequences relative to the corresponding naturally encoded gene (e.g., due to recombination and selection), but is at least 95% as long as the naturally encoded gene (i.e., has at least 95% of the total number of amino acids as the corresponding naturally encoded gene.)

A “DNAse enzyme” is an enzyme which catalyzes cleavage of a DNA, in vitro or in vivo.

A “nucleic acid domain” is a nucleic acid region or subsequence. The domain can be conserved or not conserved between a plurality of homologous nucleic acids. Typically a domain is delineated by comparison between two or more sequences, i.e., a region of sequence diversity between sequences is a “sequence diversity domain,” while a region of similarity is a “sequence similarity domain.” Domain switching” refers to the ability to switch one nucleic acid region from one nucleic acid with a second domain from a second nucleic acid.

A region of “high sequence similarity” refers to a region that is 90% or more identical to a second selected region when aligned for maximal correspondence (e.g.,
manually or using the common program BLAST set to default parameters). A region of “low sequence similarity” is 40% or less identical to a second selected region when aligned for maximal correspondence (e.g., manually or using BLAST set with default parameters).

A “PCR amplicon” is a nucleic acid made using the polymerase chain reaction (PCR). A “PCR primer” is a nucleic acid which hybridizes to a template nucleic acid and permits chain elongation using a thermostable polymerase under appropriate reaction conditions.

A “library of oligonucleotides” is a set of oligonucleotides. The set can be pooled, or can be individually accessible.

**DETAILED DISCUSSION OF THE INVENTION**

The present invention provides a variety of new recombination formats and methods for controlling, modulating and tuning recombination rates. In the methods, a plurality of insertion sequences (e.g., about 2, 5, 10, 15, 20, or even more sequences) are integrated into a parental sequence of interest (e.g., in a physical molecule or in a corresponding computer character string). The integration of the insertion sequences alters the hybridization and recombination properties of the resulting molecule relative to the parental molecule and can also be used to influence expression, splicing and/or translation rates of encoded nucleic acids. In addition to integration of insertion sequences, other strategies which modify hybridization and which tune recombination rates can also be used in conjunction with the insertion strategies herein, such as codon modification of the parental nucleic acid to further tune and modify hybridization and/or, e.g., translation. Codon modification strategies are discussed in detail in “SHUFFLING OF CODON ALTERED GENES” by Phillip A. Patten and Willem P.C. Stemmer filed September 28, 1999, e.g., Application No: PCT/US99/22588 and USSN 09/407,800. This ability to make molecules with altered hybridization and recombination properties is useful for tuning recombination rates. In addition, many nucleic acids are proprietary and the ability to use non-proprietary hybridization altered nucleic acids is of considerable commercial value.

In a preferred embodiment, shuffled recombinant nucleic acids do not hybridize to the parental genes from which they were derived. This is desirable, e.g., where the original protein generates an immune response and a protein with similar functionality but a substantially different primary structure is desired, or, e.g., where the original nucleic acid is proprietary.
INSERTION SEQUENCES

As noted above, an insertion nucleic acid subsequence is a nucleic acid subsequence found in a full-length nucleic acid sequence. The subsequence is derived from a nucleic acid sequence different from sequences immediately flanking the subsequence within the full-length nucleic acid sequence, or which subsequence is present in a different arrangement relative to the flanking subsequences from those found in nature. Most typically, the insertion sequences are derived from nucleic acids which can either be removed from encoded mRNAs (e.g., where the insertion sequences are introns, or are spliced out of coding DNA by site-specific recombination), or which can be removed from encoded proteins (e.g., by protease-ligation strategies such as intein- extein protein splicing).

Intron/ exon strategies

Mitchell. Indeed, there is well over 20 years of literature on the topic of introns and RNA splicing; one of skill is presumed to be familiar with this available literature.

In one aspect of the invention, a modified gene sequence comprising insertion sequences is synthesized (by chemical synthesis, recombinantly, by PCR synthetic strategies, or a combination thereof, and/or, e.g., by “virtual” synthesis in a computer). In cases where hybridization to the parental gene is not desirable, the exons in the modified gene sequence are preferably about 24-50 bp in size (such small coding sequences in an intron-exon arrangement are typically referred to as “mini exons”). In nature, mini exons smaller than 50 bp are rare and usually occur as singular mini-exons surrounded by larger exons. Mini exons are also frequently skipped by cellular splicing machinery, for example in a tissue specific fashion, as a form of expression control, e.g., resulting in alternatively spliced products.

To ensure that mini exons in the described modified gene are spliced faithfully, regulatory elements supporting the inclusion of mini exons are included within the introns. These regulatory elements are Intron Splicing Enhancers (ISEs). For example, a 134 nt ISE found in the chicken cardiac troponin T (cTNT) gene ensures 100% retention of a heterologous 7 nt mini-exon in various cell lines (Carlo et al. (1996) RNA, 2:342-353). An example cTNT intronic splicing enhancer is:

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aaatccttcttctctgcectctgctgcagCAAAGGgaagtgcagctgcatgCCTCCCAACCA CACCTGTGCTGCATGACACCTGGGGCTGACCTGCAACAGAAGTGGGTGCTG AGGGAAAGACTGTCTCTGGGACTGTTGACTGTCAGACGCGGGGTTTGGTGACTCTC AGGATGCCAAAATGCCCA (SEQ ID NO.: 1).
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In another known example of ISEs, two copies of a 105 nt ISE from the c-src oncogene have been shown to stabilize the inclusion of a heterologous mini-exon in a heterologous cell line (Modefferi and Black (1997) Mol. Cell Biol. 17:6537-6545). Thus, in one class of embodiments, ISEs are included in some or all of the introns (placement and presence or lack of ISEs can also be used to modulate splicing variations, further increasing sequence diversity in resulting gene products). Variants of the modified gene are evolved, e.g., using DNA shuffling techniques. Resulting genes are typically transfected into cells and selected using appropriate functional assays.

The choice of ISE ensures faithful splicing of a mini exon array. The cTNT ISE is one preferred ISE. The cTNT ISE works in transiently transfected tissue culture cells
and in nuclear extracts. It is highly efficient, supporting up to 100% inclusion of a heterologous mini exon in a variety of cells (CHO, NIH 3T3, F9, S3, HEla, COS, M6). Furthermore, in the chicken cardiac troponin T (cTNT) gene, the cTNT ISE stabilizes the inclusion of a constitutively spliced mini exon and tissue specific factors do not appear to be required. In addition, a minimal cTNT ISE of 96 bp has been defined. This synthetic cTNT ISE contains 8 copies of a 7 nt sequence motif separated by spacing sub sequences. The synthetic cTNT ISE supports about 80% inclusion of a heterologous mini exon. When the complete 134 bp ISE is used, the overall intron size can be scaled down to about 170-180 nt. One valuable feature of the cTNT ISE is its position independence. The cTNT ISE supports the inclusion of a mini exon when inserted either upstream or downstream of a mini exon. By including cTNT ISEs in every intron in a non-hybridizing gene homologue, inclusion of each mini-exon is induced by both upstream and downstream ISEs. Thus, a synergistic effect supporting complete splicing of the mini exon array occurs.

In one embodiment, relevant features of chicken cardiac troponin T (cTNT)

intronic splicing enhancers include: 1) a 134 nt ISE contains 6 repeats of a 7nt sequence motif, that stabilizes inclusion of a constitutive 6 nt mini-exon in various cell lines, and 2) 8 repeats (96 nt synthetic ISE) which support 80% inclusion of a heterologous mini-exon (7nt).

In another embodiment, the c-src oncogene ISE is used. Relevant features include: 1) a 105 nt ISE, which stabilizes inclusion of a neuron-specific, 18 nt mini-exon, and 2) the c-src ISE is also active in non-neuronal cells: 2 105 nt ISEs induce 78% retention of a heterologous mini-exon in non-neuronal cells.

An exemplar splicing strategy is represented schematically in Fig. 1. As shown, an insertion-modified nucleic acid (e.g., which does not hybridize to a selected non

insertion-modified nucleic acid) is transfected into a cell. Following splicing, a spliced gene is produced. The splicing can be either nucleic acid splicing (as depicted) or intein splicing. Common exemplar steps in designing an RNA splicing system can include, e.g., the following: 1) building of splicing constructs; 2) transient transfection of constructs into, e.g., eukaryotic tissue culture cells; 3) isolation of RNA, RT-PCR splicing products; 4)

sequencing of splicing products; 5) putting splicing products into expression vectors and 6) testing chimeric gene products in functional assays.
An example of this strategy is schematically described in Fig. 2 for an example cytokine, arbitrarily designated “cytokine G.” Cytokines generally include, e.g., interleukins, interferons, chemokines, hematopoietic growth factors, tumor necrosis factors, transforming factors and the like. In general, these proteins are small molecular weight proteins that regulate maturation, activation, proliferation and/or differentiation of cells, e.g., of the immune system. Because, e.g., of the convenient size and relative commercial value of these proteins, cytokines represent a preferred target of the present invention. As shown, in step 1, maximum divergence from an initial cytokine type (e.g., the human form of cytokine G) is obtained by performing shuffling (e.g., in vitro, in vivo or in silico), e.g., by performing family shuffling. In step 2, introns are inserted into conserved regions and shuffling is performed to optimize splicing of the resulting nucleic acids. In step 3, translation is selected for, e.g., by FACS or other appropriate methods. In step 4, functional non-hybridizing nucleic acids (i.e., which do not hybridize to the initial cytokine G type) are isolated.

In the case of an exemplar 500 bp coding sequence, a non-hybridizing gene comprising 24 bp exons has a total size using the cTNT ISE intron arrangement discussed herein of about 3920 bp. Many genes of interest encode polypeptides in this approximate size range, including the genes for insulin proteins, peptide hormones, cytokines, epidermal growth factors, fibroblast growth factors, hepatocyte growth factors, insulin-like growth factors, interferons, interleukins, keratinocyte growth factors, leukemia inhibitory factors, oncostatin M, PD-ECSF, pleiotropin, SCF, c-kit ligand, VEGFs (VEGFE has a number of well-characterized splice variants with different activities), G-CSF, GM-CSF, many oncogene and tumor suppressor encoded peptides, plant hormones, disease resistance proteins, some bacterial proteins, some proteases, peptide ligands, angiogenesis inhibitors, C-X-C chemokines, C-C chemokines, cystein knot proteins (such as TGFβ, NGF, PDGFβ, etc.), Rnases and many others.

To facilitate assembly of the non-hybridizing gene, the introns can include suitable restriction sites or other features for sequence assembly. For example, in one embodiment, the gene is assembled from units consisting of a mini exon flanked by 2 intron halves.

A variety of variations can be performed to modify the efficiency of this system. For example, ISE sequences and trans acting splicing factors can be shuffled to
improve (or decrease, if desired) the efficiency of the mini-exon array. For this purpose, cTNT ISE sequences and genes encoding splicing factors from different species are isolates and subjected to shuffling. Structured introns can be used to guide splicing.

Similarly, the cell line can be selected for optimum splicing. Indeed, the cell line can be evolved, e.g., by whole genome shuffling as taught in Del Cardayre et al. 09/116,188 “EVOLUTION OF WHOLE CELLS AND ORGANISMS BY RECURSIVE SEQUENCE RECOMBINATION” filed 07-15-98 to optimize splicing for a particular ISE. Thus, either the ISE, or the cell line, or both, can be optimized for splicing prior to use in a particular gene.

Properly spliced sequences can be detected by detecting splicing products such as nucleic acids (e.g., by RT-PCR amplification), and proteins (e.g., by expressing gene products on the surface of cells and panning for particular epitopes, e.g., by ligand panning or FACS). For PCR amplification strategies, either those that rely on downstream analysis (e.g., gel electrophoresis, sequencing of products and the like) or those that rely on real time methods (e.g., FRET, Taqman, molecular beacons or the like), or both can be used. Phage display can also be used, as can polysome display.

Splice site consensus sequences extend for 1-2 nucleotides into exons at both the 5' and 3' ends. However, only the G residue at the 3' end of the exon is strongly conserved (about 79%). The other positions are more flexible (30-60%). Thus, by shuffling splice site sequences and using wobble positions at the 3' end of exons, introduction of efficient splice sites into the coding sequences is optimized.

Bacterial self splicing introns and trans splicing introns can also be used.

Protein Splicing
Analogous to RNA splicing approaches, the coding sequence of a gene can also be interrupted by integration of insertion sequences which encode one or more intein. Inteins, sometimes referred to as "intervening protein sequences", (Perler, et al., Nucleic Acids Research, 22:1125-1127 (1994)) or as “protein introns,” “protein spacers,” “protein inserts” or the like, are genetic elements that are eliminated from protein sequences to produce mature (typically functional) proteins. In other words, the proteins of intein interrupted genes are expressed as precursor proteins (and can be expressed as one or as more than one protein chain). Upon protein splicing, the inteins are removed to generate a functional protein.
Inteins were discovered, e.g., in yeast genes (Kane, et al., Science, 250:651-
VMA1 gene of Saccharomyces cerevisiae. Since then, many more cases of protein splicing
have been found (reviewed, e.g., in Clyman, ASM News, 61:344-347 (1995); Colston and
Davis, Mol. Microbiol., 12:359-363 (1994)). Inteins have been found in eukaryotes,
bacteria and archaeabacteria, i.e., in all kingdoms. Inteins are sometimes described as
protein sequences which are embedded, typically in frame, within a precursor protein, and
which are removed by protein splicing. During the protein splicing process, two terminal
polypeptide portions become ligated by a peptide bond, and form a “fusion protein” which
corresponds to one or more extein. The amino acids found at two hexapeptide motifs on
each end of a typical intein influence the splicing process. These regions, which are also
called intein splice sites, are somewhat conserved in inteins. The mechanism of protein
splicing involves several of these amino acids, typically including, e.g., the C-terminal
histidine, asparagine and cysteine/threonine/serine residues (Davis, et al., J. Bacteriol.,
(1992); Hodges, et al., Nucleic Acids Research, 20:6153-6157 (1992); Cooper, et al.,
EMBO Journal, 12:2575-2583 (1993)).

A “native” intein activity is one which occurs, in vitro or in vivo, as a result
of activity of proteins or other cellular factors participating in a splicing reaction. For
example, Splicing can be autocatalytic (a cis reaction) and, at least in this mode, typically
does not require host cell cofactors, as inteins can splice out of their precursor proteins in a
variety of in vivo and in vitro expression systems, including simple phosphate buffered
(1993); and reviewed in Colston and Davis, Mol. Microbiol., 12:359-363 (1994)). In
addition to such native or “spontaneous” intein splicing activity, “controllable” intein
splicing has also been proposed, in which artificial exposure to unusual levels of heat or
light, or artificial dephosphorylation or deglycosylation of proteins, or artificial treatment
with chemical reagents is used to control intein splicing reactions. See, e.g., Comb et al.
U.S. Pat. No. 5,834,247 and U.S. Pat. No. 5,496,714. Either native or controllable intein
splicing can be used in the context of the present invention. In addition to intein splicing,
chemical splicing of oligopeptides to form active proteins can also be performed in the
context of the present invention.

Additional details on protein splicing generally can be found at the Intein Databases web site (www.neb.com/neb/inteins/intein_intro.html); and in, e.g., Nucleic Acids Research 26(7):1741-1758.

Minimal functional inteins lacking a homing endonuclease have been defined. For example, a semi-synthetic intein 135 aa in length undergoes 50-90% intein removal in vitro (Lew et al. (1998) JBC 273:15887-15890). In the context of the present invention, inteins are optionally optimized by forced evolution (e.g., using shuffling) to achieve optimal protein splicing in any desired heterologous context. Selection of protein products is performed using an appropriate functional or physical assay for the protein.

In addition to cis-splicing of inteins, trans-splicing of inteins is also used in the present invention (as noted above, both cis and trans splicing of RNA can also be used to produce insertion modified nucleic acids). In one embodiment, proteins of interest are encoded by two or more separate nucleic acids which are expressed to produce two separate polypeptides. These two or more separate polypeptides are recombined to form the protein of interest. This is illustrated in Fig. 4. This figure shows screening of libraries encoded by split exteins that are trans-spliced. The gene family to be shuffled is divided into two (or more) exteins. Trans-splicable inteins are put between the exteins. Libraries of N- and C-terminal exteins fused to their respective inteins are constructed by standard methods. The protein fragments are expressed either in the same or in different cells. Trans splicing is promoted either in vivo or in vitro to yield a mature shuffled protein that is generated without the manipulation at any step of full length mature shuffled genes. Examples of
trans-intein splicing systems are available, such as the DnaE gene, encoded by dnaE-n and dnaE-c in the *Synchocystis* sp. PCC6803 genome. These and other systems can be shuffled to optimize splicing in general (including trans splicing), or in any specific system of interest.

In general, it is sometimes desirable to produce proteins which are not expressed in an active format. This is because, for example, some proteins are toxic to the cell in which the protein is expressed (e.g., RNases, DNases, toxins such as ricin, proteases, apoptosis inducing factors, microtubule proteins, etc.). In these situations, it is advantageous to express a protein in an inactive form that can be conditionally activated (e.g., by splicing of two inactive peptides to form an active protein, or by trans-splicing of RNAs to produce a coding nucleic acid). It is also possible to express different segments of the nucleic acid and/or protein in different systems (e.g., where expression is biased in one or more cell), e.g., to express different portions of the nucleic acids or proteins in different cells, or even in different in vitro expression systems. Thus, for example, one can express a portion of a protein in an *E. coli* and a portion in yeast. The separate portions are contacted and allowed to undergo trans-splicing. In some circumstances, it is convenient from a production standpoint to produce and store one portion of a protein or nucleic acid to be spliced and then to separately produce another portion at a different time. After the relevant portions are produced, they are combined, in vitro or in vivo, and the final product produced.

In addition, separate expression of protein fragments is useful for combinatorial purposes, i.e., the fragments can be separately evolved and the ability to recombine different fragments can increase the diversity in the final proteins which result from the splicing reactions. For example, the viability, activity and folding of proteins can be influenced by multiple regions of the protein, often simultaneously or in a cooperative fashion. By removing these cooperative effects until after the initial expression of protein fragments, these effects can be avoided or advantageously modulated. Thus, generally, separate large libraries of diversity can be generated and “recombined” by splicing the protein fragments encoded by the library members.

As an example, one of skill can transform mammalian cells with a library of “split gene” fragment members (i.e., encoding portions of a protein of interest) to generate large libraries of cells encoding a spliced proteins of interest. For example, the number of
cells that can be transfected in a typical DNA transfection experiment is generally a limiting feature in mammalian transfection/screening/cloning applications. For example, one might obtain only about 1000 stable transfectants following transfection of a population of cells with a library of $10^9$ different DNAs. However, using the present invention, one can sample much more than 1000 shuffled proteins by transfection, such that the typical transfectant receives 10 shuffled N-terminal and 10 shuffled C-terminal exteins. Each cell is then capable of making 100 unique trans-spliced mature proteins. This gives a large increase in the number and diversity of resulting recombined genes that can then be expressed and screened. It will be appreciated that the use of more than two fragments to generate the spliced protein of interest provides even larger numbers of resulting spliced proteins.

There is a large class of screening problems that are best performed in whole organisms, as there are not necessarily good in vitro or cellular assays for the activity of interest. For example, insect or herbicide resistance in plants or shuffled growth hormones in animals are advantageously screened in the plant or animal for which the activity is to be developed. Advantageously, an example of the “split” gene strategy embodied by the use of trans-splicing proteins is the screening of shuffled split gene libraries in whole organisms, e.g., for the development of these activities in the plant or animal of interest. In particular, using the split gene method, it is possible to access and screen diversity equal to (at least) the square of the number of transgenic organisms. For example, if 100 transgenic plants expressing 100 different shuffled N-terminal exteins are crossed to 100 transgenic plants expressing 100 different C-terminal exteins (e.g., by classical breeding) $10^4$ transgenic plants with unique spliced shuffled proteins will result upon splicing of the different N- and C-terminal portions of the proteins in the plants. This approach is illustrated schematically in Fig. 5 (only limited sets of breeding plants and progeny are represented for clarity of illustration) for sets of roses (any plant can be substituted for the roses which are depicted). For clarity of illustration, the complete sets are not shown. As depicted, e.g., 100 transgenic plants expressing 100 different shuffled N-terminal exteins are crossed to 100 transgenic plants expressing 100 different C-terminal exteins (e.g., by classical breeding), producing $10^4$ transgenic plants with unique spliced shuffled proteins in the F1 generation. See also, Patten et al. “ENCRYPTION OF TRAITS USING SPLIT GENE SEQUENCES AND ENGINEERED GENETIC ELEMENTS” USSN 60/164,618 Filed November 10, 1999.
One of skill will appreciate that, here again, fragmentation of the protein of interest into more than two extein fragments and the creation of libraries of organisms expressing each extein fragment (e.g., a C-terminal fragment, one or more middle fragment(s) and an N-terminal fragment) provides even larger numbers of proteins which can result from classical breeding of the different organism library members.

In addition to using intein-extein strategies to evolve proteins of interest, the present invention also provides methods and libraries for improving splicing activity. In particular, sequences encoding inteins and exteins can be shuffled (e.g., using any available shuffling method as set forth herein) and the splicing activity of the resulting shuffled sequences assessed, e.g., by assaying for a resulting spliced product (in this embodiment, splicing can be either cis or trans splicing). For example, the spliced product can encode a catalytic or other activity that can be measured using standard methods (e.g., in an assay which produces a detectable signal such as luminescence, fluorescence, or the like which is dependent on the presence of the spliced protein).

**Site Specific Recombination**
In site-specific recombination strategies, in vivo recombination is carried out to remove insertion sequences. In one aspect, enzymes carrying out site specific recombination are evolved by shuffling methods which select for increased performance and specificity. Selection of functional products is performed by detecting, e.g., mRNA or protein products, or by using an appropriate functional assay. Evolved variants of site-specific recombinases have acquired specificity for novel DNA recombination signals (e.g., altered loxP sites), so that the modified sites (e.g., lox P sites) encode a protein of interest.

**Alternative Splicing**
Alternative splicing of pre-mRNAs is used in cells to recombine functional gene segments encoded by separate exons. In the present invention, shuffling of protein domains or domains of non coding functional RNAs (viral RNAs, snRNAs, tRNAs, rRNAs, gRNAs, etc.) is performed based upon alternative splicing techniques. Synthetic genes containing multiple exons derived from related or unrelated (or distantly related) genes separated by introns are created. In one embodiment, alternative splicing is induced by transient transfection of the genes into, e.g., eukaryotic cells. Transiently transfected cells accomplish alternative splicing very efficiently and stable cell lines are not necessary. Although in vitro splicing of RNAs up to about 1 kb in size is possible in nuclear extracts,
in vitro splicing is relatively inefficient, especially when alternative splicing is desired. Interesting splicing products are assayed by functional assays or by detection of RNAs (e.g., by RT-PCR) or translated proteins. Splicing products are optionally recovered by RT-PCR, cloned and sequenced or further assayed. Splicing products of interest are, e.g., introduced into expression systems and gene products tested in functional assays.

The goal of domain recombination by alternative splicing is to create a large number of recombinant molecules without disrupting functional domains. The design of synthetic genes subjected to alternative splicing is optimized to achieve this goal.

The design of synthetic genes allows for modulation of patterns of splicing. Creating pre-mRNAs with e.g., about 6-20 exons provides many different exon combinations in spliced products. An even higher frequency of alternatively spliced RNAs occurs if mini exons are included. Exons smaller than 50 bp are inefficiently recognized (absent an ISE as discussed above) by the splicing machinery and as a consequence, are often skipped. Employing multiple mini exons induces exon skipping, creating additional splicing diversity.

An example of a synthetic gene undergoing multiple patterns of splicing is shown in Fig. 3. Exons are designated A, B and C and encode unrelated functional domains. B1 and B2 encode variants of the same functional domain, or for spacer regions. C1 and C2 encode variants of another functional domain. Exon 3 includes a polyadenylation site.

In addition to inducing exon skipping, selected exons can also be enriched in splicing products through the use of splicing enhancers. For example, purine rich sequence elements such as the sequence GAR, where R is a purine have been found within exons. These splicing enhancers activate the use of the upstream 3’ splice site, thus stabilizing the exon in which they are located. Purine rich splicing enhancers are found in genes from many higher eukaryotes and function in many tissues and cell lines. Incorporating splicing enhancers allows for anchoring of exons in a splicing product, preventing it from being skipped (which is especially desirable where combinations of the domain encoded by the exon with other domains is desirable). This biasing of splicing during recombination is a feature of the invention.

Flexible spacer regions between exons can also be used to increase the functionality of chimeric molecules.
MAKING INSERTION SEQUENCES AND GENES

As noted above, insertion sequences and nucleic acids comprising insertion sequences are made according to standard recombinant or synthetic methods, optionally in combination with shuffling or PCR synthetic methods. Generally, in silico, chemical synthetic, ligase and/or polymerase mediated approaches to synthesis are optionally performed to produce insertion modified nucleic acids.


Improved methods of cloning *in vitro* amplified nucleic acids are described in Wallace *et al.*, U.S. Pat. No. 5,426,039. Improved methods of amplifying large nucleic acids by PCR are summarized in Cheng *et al.* (1994) *Nature* 369: 684-685 and the references therein, in which PCR amplicons of up to 40kb are generated. One of skill will appreciate that essentially any RNA can be converted into a double stranded DNA suitable for restriction
digestion, PCR expansion and sequencing using reverse transcriptase and a polymerase. See, Ausubel, Sambrook and Berger, all supra.

In one preferred method, assembled sequences are checked for incorporation of insertion sequences. This can be done by cloning and sequencing the nucleic acids, and/or by restriction digestion, e.g., essentially as taught in Sambrook, Berger and Ausubel, above. The extent of PCR amplification can also be determined by incorporation of a label into one or more amplified elongated nucleic acid, or by hybridization to relevant probes, e.g., in a fluorogenic 5' nuclease assay, TaqMan, FRET, use of molecular beacons (or other real time PCR analysis methods), etc. In addition, sequences can be PCR amplified and sequenced, directly or indirectly. Thus, in addition to, e.g., Sambrook, Berger, Ausubel and Innis (id. and above), additional PCR sequencing PCR sequencing methodologies are also particularly useful. For example, direct sequencing of PCR generated amplicons by selectively incorporating boronated nuclease resistant nucleotides into the amplicons during PCR and digestion of the amplicons with a nuclease to produce sized template fragments has been performed (Porter et al. (1997) Nucleic Acids Research 25(8):1611-1617). In the methods, 4 PCR reactions on a template are performed, in each of which one of the nucleotide triphosphates in the PCR reaction mixture is partially substituted with a 2'deoxyribonucleoside 5'—[P-borano]—triphosphate. The boronated nucleotide is stochastically incorporated into PCR products at varying positions along the PCR amplicon in a nested set of PCR fragments of the template. An exonuclease which is blocked by incorporated boronated nucleotides is used to cleave the PCR amplicons. The cleaved amplicons are then separated by size using polyacrylamide gel electrophoresis, providing the sequence of the amplicon. An advantage of this method is that it uses fewer biochemical manipulations than performing standard Sanger-style sequencing of PCR amplicons. Again, real time PCR analysis (FRET, molecular beacons, etc.) provide a simple method for assessing sequences.

Codon Modification

In one aspect, the genes utilized in the methods herein have altered codon use as compared to the parental sequences from which the genes are derived. In particular, it is useful, e.g., to modify codon preference to optimize expression in a cell in which a recombinant product of an oligonucleotide shuffling procedure is to be assessed or otherwise selected. Conforming a recombinant nucleic acid to the codon bias of a particular cell in which selection is to take place typically results in maximization of expression of the
recombinant nucleic acid. Because the oligonucleotides used in the various strategies herein typically are made synthetically, selecting optimal codon preference is done simply by reference to well-known codon-bias tables.

In addition to selection of oligonucleotide sequences to optimize expression, codon preference can also be used to increase sequence similarity between distantly related nucleic acids which are to be recombined. By selecting which codons are used in particular positions it is possible to increase the similarity between the nucleic acids, which, in turn, increases the frequency of recombination between the nucleic acids. Additional details on codon modification procedures and their application to DNA shuffling are found in “SHUFFLING OF CODON ALTERED GENES” by Patten et al. filed September 28, 1999, USSN 09/407,800.

Expression in Vitro
Although often discussed herein in terms of cellular expression, the nucleic acids produced by the various procedures herein can be expressed in vivo or in vitro. Thus, in one embodiment of the invention, nucleic acids produced by the various diversity generation methods set forth herein (insertion of introns, inteins, or the like, optionally in combination with shuffling, mutation, or the like) are transcribed (i.e., where the nucleic acids are DNAs) into RNA, spliced if appropriate and translated into proteins (which can undergo splicing as discussed herein), which are, optionally, screened by any appropriate assay. Common in vitro transcription and/or translation reagents include reticulocyte lysates (e.g., rabbit reticulocyte lysates) wheat germ in vitro translation (IVT) mixtures, E. coli lysates, canine microsome systems, HeLa nuclear extracts, the “in vitro transcription component,” (see, e.g., Promega technical bulletin 123), SP6 polymerase, T3 polymerase, T7 RNA polymerase (e.g., Promega # TM045), the “coupled in vitro transcription/translation system” (Progen Single Tube Protein System 3) and many others. Many of translation systems are described, e.g., in Ausubel, supra. as well as in the references below, and many transcription/translation systems are commercially available.

Generally, cell-free transcription/translation systems can be employed to produce polypeptides from solid or liquid phase sets of DNAs or RNAs. Several transcription/translation systems are commercially available and can be adapted to the present invention by the appropriate addition of transcription and or translation reagents to insertion-modified nucleic acids. A general guide to in vitro transcription and translation
protocols is found in Tymms (1995) *In vitro Transcription and Translation Protocols: Methods in Molecular Biology* Volume 37, Garland Publishing, NY. Several in vitro transcription and translation systems are well known and described in Tymms (1995), *id.* For example, an untreated reticulocyte lysate is commonly isolated from rabbits after treatment of the rabbits with acetylphenylhydrazine as a cell-free in vitro translation system. Similarly, coupled transcription/translation systems often utilize an E. coli S30 extract. *See also*, the Ambion 1999 Product Catalogue from Ambion, Inc (Austin TX). A variety of commercially available in vitro transcription and translation reagents are commercially available, including the PROTEINscript-PRO™ kit (for coupled transcription/translation) the wheat germ IVT kit, the untreated reticulocyte lysate kit (each from Ambion, Inc (Austin TX)), the HeLa Nuclear Extract in vitro Transcription system, the TnT Quick coupled Transcription/translation systems (both from Promega, *see*, e.g., Technical bulletin No. 123 and Technical Manual No. 045), and the single tube protein system 3 from Progen. Each of these available systems (as well as many other available systems) have certain advantages which are detailed by the product manufacturer.


For example, an untreated rabbit reticulocyte lysate is suitable for initiation and translation assays where the prior removal of endogenous globin mRNA is not necessary. The untreated lysate translates exogenous mRNA, but also competes with endogenous mRNA for limiting translational machinery.

Similarly, The PROTEINscript-PRO™ kit from Ambion is designed for coupled in vitro transcription and translation using an E. coli S30 extract. In contrast to eukaryotic systems, where the transcription and translation processes are separated in time
and space, prokaryotic systems are coupled, as both processes occur simultaneously. During transcription, the nascent 5'-end of the mRNA becomes available for ribosome binding, allowing transcription and translation to proceed at the same time. This early binding of ribosomes to the mRNA maintains transcript stability and promotes efficient translation. Coupled transcription: translation using the PROTEINscript-PRO Kit is based on this E. coli model.

The Wheat Germ IVT™ Kit from Ambion, or other similar systems, is/are a convenient alternative, e.g., when the use of a rabbit reticulocyte lysate is not appropriate for in vitro protein synthesis. The Wheat Germ IVT™ Kit can be used, e.g., when the desired translation product comigrates with globin (approx. 12-15 kDa), when translating mRNAs coding for regulatory factors (such as transcription factors or DNA binding proteins) which may already be present at high levels in mammalian reticulocytes, but not plant extracts, or when an mRNA will not translate for unknown reasons and a second translation system is to be tested.

The TNT® Quick Coupled Transcription/Translation Systems (Promega) are single-tube, coupled transcription/translation reactions for eukaryotic in vitro translation. The TNT® Quick Coupled Transcription/Translation System combines RNA Polymerase, nucleotides, salts and Recombinant RNasin® Ribonuclease Inhibitor with the reticulocyte lysate to form a single TnT® Quick Master Mix. The TNT® Quick Coupled Transcription/Translation System is available in two configurations for transcription and translation of genes cloned downstream from either the T7 or SP6 RNA polymerase promoters. Included with the TNT® Quick System is a luciferase-encoding control plasmid and Luciferase Assay Reagent, which can be used in a non-radioactive assay for rapid (<30 seconds) detection of functionally active luciferase protein.

Many other systems are well known, well characterized and set forth in the references noted herein, as well as in other references known to one of skill. It will also be appreciated that one of skill can produce transcription/translation systems similar to those which are commercially available from available materials, e.g., as taught in the references noted above.

**DIVERSITY GENERATION STRATEGIES**

The present invention makes use of shuffling and other diversity generation methods to recombine, mutate, or otherwise create or diversify insertion modified nucleic
acids. Nucleic acids produced by the methods herein are optionally subjected to shuffling or other diversity generation methods (e.g., mutagenesis) to improve or modify the encoded function of the nucleic acids. For example, insertion sequences can be shuffled, e.g., by family shuffling techniques (e.g., family shuffling of homologous introns), mutated or otherwise diversified.


In addition, details and formats for DNA shuffling and other diversity generating protocols are found in a variety of PCT and foreign patent applications, including: Stemmer and Cramer, “DNA MUTAGENESIS BY RANDOM FRAGMENTATION AND REASSEMBLY” WO 95/22625; Stemmer and Lipschutz “END COMPLEMENTARY POLYMERASE CHAIN REACTION” WO 96/33207; Stemmer and Cramer “METHODS FOR GENERATING POLYNUCLEOTIDES HAVING DESIRED CHARACTERISTICS BY ITERATIVE SELECTION AND RECOMBINATION” WO 97/0078; Minshul and Stemmer, “METHODS AND COMPOSITIONS FOR CELLULAR AND METABOLIC ENGINEERING” WO 97/35966; Punnonen et al. “TARGETING OF GENETIC VACCINE VECTORS” WO 99/41402; Punnonen et al. “ANTIGEN LIBRARY IMMUNIZATION” WO 99/41383; Punnonen et al. “GENETIC VACCINE VECTOR ENGINEERING” WO 99/41369; Punnonen et al. OPTIMIZATION OF IMMUNOMODULATORY PROPERTIES OF GENETIC VACCINES WO 9941368;

Certain U.S. Applications provide additional details regarding DNA shuffling and related techniques, as well as other diversity generating methods, including “SHUFFLING OF CODON ALTERED GENES” by Patten et al. filed September 29, 1998, (USSN 60/102,362), January 29, 1999 (USSN 60/117,729), and September 28, 1999, USSN 09/407,800; “EVOLUTION OF WHOLE CELLS AND ORGANISMS BY RECURSIVE SEQUENCE RECOMBINATION”, by del Cardy et al. filed July 15, 1998 (USSN 09/166,188), and July 15, 1999 (USSN 09/354,922); “OLIGONUCLEOTIDE MEDIATED NUCLEIC ACID RECOMBINATION” by Cramer et al., filed February 5, 1999 (USSN 60/118,813) and filed June 24, 1999 (USSN 60/141,049) and filed September 28, 1999 (USSN 09/408,392); and filed January 18, 2000 (PCT/US00/01202) and “USE OF CODON-BASED OLIGONUCLEOTIDE SYNTHESIS FOR SYNTHETIC SHUFFLING” by Welch et al., filed September 28, 1999 (USSN 09/408,393); and “METHODS FOR MAKING CHARACTER STRINGS, POLYNUCLEOTIDES & POLYPEPTIDES HAVING DESIRED CHARACTERISTICS” by Selifonov and Stemmer, filed February 5, 1999 (USSN 60/118854) and filed October 12, 1999 (USSN 09/416,375) and USSN 09/484,850 and PCT/US00/01203 filed January 18, 2000.

As review of the foregoing publications, patents, published applications and U.S. patent applications reveals, recursive recombination of nucleic acids to provide new nucleic acids with desired properties can be carried out by a number of established methods and these procedures can be combined with any of a variety of other diversity generating methods.
In brief, at least 5 different general classes of recombination methods are applicable to the present invention and set forth in the references above. First, insertion-modified nucleic acids can be recombined in vitro by any of a variety of techniques discussed in the references above, including e.g., DNase digestion of nucleic acids to be recombined followed by ligation and/or PCR reassembly of the nucleic acids. Second, insertion modified nucleic acids can be recursively recombined in vivo, e.g., by allowing recombination to occur between nucleic acids in cells. Third, whole genome recombination methods can be used in which whole genomes of cells or other organisms are recombined, optionally including spiking of the genomic recombination mixtures with insertion modified nucleic acids. Fourth, synthetic recombination methods can be used, in which oligonucleotides corresponding to targets of interest (e.g., including one or more insertion-modified nucleic acid) are synthesized and reassembled in PCR and/or ligation reactions which include oligonucleotides which correspond to more than one parental nucleic acid (e.g., including one or more insertion-modified nucleic acid), thereby generating new recombined insertion modified nucleic acids. Oligonucleotides can be made by standard nucleotide addition methods, or can be made, e.g., by tri-nucleotide synthetic approaches. Fifth, in silico methods of recombination can be effected in which genetic algorithms are used in a computer to recombine sequence strings which correspond to nucleic acid homologues (or even non-homologous) insertion modified sequences. The resulting recombined sequence strings are optionally converted into nucleic acids by synthesis of nucleic acids which correspond to the recombined sequences, e.g., in concert with oligonucleotide synthesis/ gene reassembly techniques. Any of the preceding general recombination formats can be practiced in a reiterative fashion to generate a more diverse set of recombinant nucleic acids.

The above references provide these and other basic recombination formats as well as many modifications of these formats. Regardless of the format which is used, the nucleic acids of the invention can be recombined (with each other or with related (or even unrelated) nucleic acids to produce a diverse set of recombinant nucleic acids, including, e.g., sets of homologous or non-homologous insertion modified nucleic acids.

Following recombination, any nucleic acids which are produced can be selected for a desired activity. In the context of the present invention, this can include testing for and identifying any activity that can be detected, including in an automatable
format, by any of the assays in the art. A variety of related (or even unrelated) properties can be assayed for, using any available assay.


Any of these diversity generating methods can be combined, in any combination selected by the user, to produce nucleic acid diversity, which may be screened for using any available screening method.

**Example: Selection Strategies for Yeast**

Libraries of genes comprising insertion sequences can be recursively shuffled by recombination and selection to enrich for genes which splice efficiently. For example, a yeast two hybrid system can be used, by, for example, fusing reporter domains to the amino and carboxy terminus of the gene of interest. Stop codons can be incorporated into the introns so that all of the introns have to be removed by splicing to produce a full length protein that can function in the yeast two hybrid selection. Simply fusing a gene to a selectable reporter such as His3 in yeast allows for positive selection for efficient splicing. Similar approaches can be taken in other eukaryotes, or in bacteria or archaeabacteria.

**HYBRIDIZATION OF NUCLEIC ACIDS**

When there is no explicit knowledge about the ancestry of two nucleic acids, homology is typically inferred by sequence comparison between two sequences. Where two nucleic acid sequences show sequence similarity it is inferred that the two nucleic acids share a common ancestor. The precise level of sequence similarity required to establish homology varies in the art depending on a variety of factors. For purposes of this disclosure, two sequences are considered homologous where they share sufficient sequence identity to allow recombination to occur between two nucleic acid molecules. Typically,
nucleic acids require regions of close similarity spaced roughly the same distance apart to permit recombination to occur. Typically, regions of at least about 60% sequence identity or higher are optimal for recombination.

The terms "identical" or percent "identity," in the context of two or more nucleic acid or polypeptide sequences, refers to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same, when compared and aligned for maximum correspondence, as measured using one of the sequence comparison algorithms described below (or other algorithms available to persons of skill) or by visual inspection. Similarly, the phrase "substantially identical," in the context of two nucleic acids or polypeptides, refers to two or more sequences or subsequences that have at least about 40%, preferably 80%, most preferably 90-95% nucleotide or amino acid residue identity, when compared and aligned for maximum correspondence, as measured using one of the following sequence comparison algorithms, or by visual inspection. Such "substantially identical" sequences are typically considered to be homologous. Preferably, the "substantial identity" exists over a region of the sequences that is at least about 50 residues in length, more preferably over a region of at least about 100 residues, and most preferably the sequences are substantially identical over at least about 150 residues, or over the full length of the two sequences to be compared.

For sequence comparison and homology determination, typically one sequence acts as a reference sequence to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are input into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. The sequence comparison algorithm then calculates the percent sequence identity for the test sequence(s) relative to the reference sequence, based on the designated program parameters.

Group, 575 Science Dr., Madison, WI), or by visual inspection (see generally Ausubel et al., infra).

One example algorithm that is suitable for determining percent sequence identity and sequence similarity is the BLAST algorithm, which is described in Altschul et al., J. Mol. Biol. 215:403-410 (1990). Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., supra). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are then extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, a cutoff of 100, M=5, N=-4, and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength (W) of 3, an expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff (1989) Proc. Natl. Acad. Sci. USA 89:10915).

In addition to calculating percent sequence identity, the BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin & Altschul (1993) Proc. Nat’l. Acad. Sci. USA 90:5873-5787). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a
reference sequence if the smallest sum probability in a comparison of the test nucleic acid to
the reference nucleic acid is less than about 0.1, more preferably less than about 0.01, and
most preferably less than about 0.001.

Another indication that two nucleic acid sequences are substantially
identical/ homologous is that the two molecules hybridize to each other under stringent
conditions. The phrase “hybridizing specifically to,” refers to the binding, duplexing, or
hybridizing of a molecule only to a particular nucleotide sequence under stringent
conditions, including when that sequence is present in a complex mixture (e.g., total
cellular) DNA or RNA. “Bind(s) substantially” refers to complementary hybridization
between a probe nucleic acid and a target nucleic acid and embraces minor mismatches that
can be accommodated by reducing the stringency of the hybridization media to achieve the
desired hybridization of the target polynucleotide sequence. “Stringent hybridization
conditions” and “stringent hybridization wash conditions” in the context of nucleic acid
hybridization experiments such as Southern and northern hybridizations are sequence
dependent, and are different under different environmental parameters. Longer sequences
hybridize specifically at higher temperatures. An extensive guide to the hybridization of
nucleic acids is found in Tijssen (1993) Laboratory Techniques in Biochemistry and
Molecular Biology—Hybridization with Nucleic Acid Probes part I chapter 2 “Overview of
principles of hybridization and the strategy of nucleic acid probe assays,” Elsevier, New
York. Generally, highly stringent hybridization and wash conditions are selected to be about
5° C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic
strength and pH. Typically, under “stringent conditions” a probe will hybridize to its target
subsequence, but not to unrelated sequences.

The Tm is the temperature (under defined ionic strength and pH) at which
50% of the target sequence hybridizes to a perfectly matched probe. Very stringent
conditions are selected to be equal to the Tm for a particular probe. An example of stringent
hybridization conditions for hybridization of complementary nucleic acids which have more
than 100 complementary residues on a filter in a Southern or northern blot is 50%
formamide with 1 mg of heparin at 42°C, with the hybridization being carried out
overnight. An example of highly stringent wash conditions is 0.15M NaCl at 72°C for
about 15 minutes. An example of stringent wash conditions is a 0.2x SSC wash at 65°C for
15 minutes (see, Sambrook, infra., for a description of SSC buffer). Often, a high
stringency wash is preceded by a low stringency wash to remove background probe signal. An example medium stringency wash for a duplex of, e.g., more than 100 nucleotides, is 1x SSC at 45°C for 15 minutes. An example low stringency wash for a duplex of, e.g., more than 100 nucleotides, is 4-6x SSC at 40°C for 15 minutes. For short probes (e.g., about 10 to 50 nucleotides), stringent conditions typically involve salt concentrations of less than about 1.0 M Na ion, typically about 0.01 to 1.0 M Na ion concentration (or other salts) at pH 7.0 to 8.3, and the temperature is typically at least about 30°C. Stringent conditions can also be achieved with the addition of destabilizing agents such as formamide. In general, a signal to noise ratio of 2x (or higher) than that observed for an unrelated probe in the particular hybridization assay indicates detection of a specific hybridization. Similarly, if the signal to noise ratio is less than 25% as high as that observed for a perfectly matched probe under stringent conditions, the nucleic acids do not “hybridize under stringent conditions” as that term is used herein. This does not apply to highly stringent conditions, as the stringency can theoretically be increased until only a perfectly matched probe will hybridize.

In one example hybridization procedure, a target nucleic acid to be probed is blotted onto a filter by any conventional method. An unrelated nucleic acid such as a plasmid vector (assuming that the target nucleic acid has no homology with the target nucleic acid) is also blotted, in approximately equal amounts onto the filter. The filter is probed with a labeled probe complementary to the target nucleic acid. The experiment is repeated at gradually increasing stringency of hybridization and wash conditions until the hybridization of the labeled probe to the complementary target is 10-100X as high as to the unrelated plasmid vector nucleic acid. Once these conditions are determined as described above, a test nucleic acid is probed under the same conditions as the target. If signal from the labeled probe is 25% as high or higher than the signal from binding of the probe to the target, the test nucleic acid "hybridizes under stringent conditions" to the probe. If the signal is less than 25% as high, the test nucleic acid does not hybridize under stringent conditions to the probe.

**Post-Recombination Screening Techniques**
The precise screening method that is used in the various shuffling procedures herein is not a critical aspect of the invention. In general, one of skill can practice
appropriate screening (i.e., selection) methods, by reference to the activity to be selected for.

In any case, one or more recombination cycle(s) is/are optionally followed by at least one cycle of screening or selection for molecules having a desired property or characteristic. If a recombination cycle is performed *in vitro*, the products of recombination, *i.e.*, recombinant segments, are sometimes introduced into cells before the screening step. Recombinant segments can also be linked to an appropriate vector or other regulatory sequences before screening. Alternatively, products of recombination generated *in vitro* are sometimes packaged in viruses (e.g., bacteriophage) before screening. If recombination is performed *in vivo*, recombinant products can sometimes be screened in the cells in which recombination occurred. In other applications, recombinant segments are extracted from the cells, and optionally packaged as viruses, before screening.

The nature of screening or selection depends on what property or characteristic is to be acquired or the property or characteristic for which improvement is sought. It is not usually necessary to understand the molecular basis by which particular products of recombination (recombinant segments) have acquired new or improved properties or characteristics relative to the starting substrates. For example, a gene can have many component sequences, each having a different intended role (*e.g.*, coding sequence, regulatory sequences, targeting sequences, stability-conferring sequences, subunit sequences and sequences affecting integration). Each of these component sequences can be varied and recombined simultaneously. Screening/selection can then be performed, for example, for recombinant segments that have increased ability to confer activity upon a cell without the need to attribute such improvement to any of the individual component sequences of the vector.

Depending on the particular screening protocol used for a desired property, initial round(s) of screening can sometimes be performed using bacterial cells due to high transfection efficiencies and ease of culture. However, bacterial expression is often not practical or desired, and yeast, fungal or other eukaryotic systems are also used for library expression and screening. Similarly, other types of screening which are not amenable to screening in bacterial or simple eukaryotic library cells, are performed in cells selected for use in an environment close to that of their intended use. Final rounds of screening can be performed in the precise cell type of intended use.
If further improvement in a property is desired, at least one and usually a collection of recombinant segments (e.g., which include insertion modified sequences) surviving a first round of screening/selection are subject to a further round of recombination. These recombinant segments can be recombined with each other or with exogenous segments representing the original substrates or further variants thereof. Again, recombination can proceed in vitro or in vivo. If the previous screening step identifies desired recombinant segments as components of cells, the components can be subjected to further recombination in vivo, or can be subjected to further recombination in vitro, or can be isolated before performing a round of in vitro recombination. Conversely, if the previous screening step identifies desired recombinant segments in naked form or as components of viruses, these segments can be introduced into cells to perform a round of in vivo recombination. The second round of recombination, irrespective how performed, generates further recombinant segments which encompass additional diversity than is present in recombinant segments resulting from previous rounds.

The second round of recombination can be followed by a further round of screening/selection according to the principles discussed above for the first round. The stringency of screening/selection can be increased between rounds. Also, the nature of the screen and the property being screened for can vary between rounds if improvement in more than one property is desired or if acquiring more than one new property is desired.

Additional rounds of recombination and screening can then be performed until the recombinant segments have sufficiently evolved to acquire the desired new or improved property or function.

INTEGRATED ASSAYS AND INTEGRATED SYSTEM ELEMENTS

One aspect of the present invention is the alignment of nucleic acids using a computer and sequence alignment software. In addition, other integrated system components provide for high-throughput screening assays, as well as for the coupling of such assays to gene or nucleic acid selection, synthesis and recombination.

Of course, the relevant assay will depend on the application. Many assays for proteins, receptors, ligands and the like are known. Formats include binding to immobilized components, cell or organismal viability, production of reporter compositions, and the like.
In the high throughput assays of the invention, it is possible to screen up to several thousand different shuffled variants in a single day. In particular, each well of a microtiter plate can be used to run a separate assay, or, if concentration or incubation time effects are to be observed, every 5-10 wells can test a single variant. Thus, a single standard microtiter plate can assay about 100 (e.g., 96) reactions. If 1536 well plates are used, then a single plate can easily assay from about 100- about 1500 different reactions. It is possible to assay several different plates per day; assay screens for up to about 6,000-20,000 different assays (i.e., involving different nucleic acids, encoded proteins, concentrations, etc.) is possible using the integrated systems of the invention. In addition, “one pot” screening approaches can screen millions of cells or viruses for a desired property, with the cells or viruses being cloned by limiting dilution. Microfluidic approaches to reagent manipulation have also been developed, e.g., by Caliper Technologies (Mountain View, CA). Additional details regarding automated shuffling methods are found in Bass et al. “INTEGRATED SYSTEMS AND METHODS FOR DIVERSITY GENERATION AND SCREENING” USSN 60/175,551, filed January 11, 2000.

In one aspect, library members, e.g., cells, viral plaques, spores or the like, are separated on solid media to produce individual colonies (or plaques). Using an automated colony picker (e.g., the Q-bot, Genetix, U.K.), colonies or plaques are identified, picked, and up to 10,000 different mutants inoculated into 96 well microtiter dishes containing two 3 mm glass balls/well. The Q-bot does not pick an entire colony but rather inserts a pin through the center of the colony and exits with a small sampling of cells, (or mycelia) and spores (or viruses in plaque applications). The time the pin is in the colony, the number of dips to inoculate the culture medium, and the time the pin is in that medium each effect inoculum size, and each can be controlled and optimized. The uniform process of the Q-bot decreases human handling error and increases the rate of establishing cultures (roughly 10,000/4 hours). These cultures are then shaken in a temperature and humidity controlled incubator. The glass balls in the microtiter plates act to promote uniform aeration of cells and the dispersal of mycelial fragments similar to the blades of a fermenter. Clones from cultures of interest can be cloned by limiting dilution. As also described supra, plaques or cells constituting libraries can also be screened directly for production of proteins, either by detecting hybridization, protein activity, protein binding to antibodies, or the like.
A number of well known robotic systems have also been developed for solution phase chemistries useful in assay systems. These systems include automated workstations like the automated synthesis apparatus developed by Takeda Chemical Industries, LTD. (Osaka, Japan) and many robotic systems utilizing robotic arms (Zymate II, Zymark Corporation, Hopkinton, Mass.; Orca, Hewlett-Packard, Palo Alto, Calif.) which mimic the manual synthetic operations performed by a scientist. Any of the above devices are suitable for use with the present invention, e.g., for high-throughput screening of molecules assembled from the various oligonucleotide sets described herein. The nature and implementation of modifications to these devices (if any) so that they can operate as discussed herein with reference to the integrated system will be apparent to persons skilled in the relevant art.

High throughput screening systems are commercially available (see, e.g., Zymark Corp., Hopkinton, MA; Air Technical Industries, Mentor, OH; Beckman Instruments, Inc. Fullerton, CA; Precision Systems, Inc., Natick, MA, etc.). These systems typically automate entire procedures including all sample and reagent pipetting, liquid dispensing, timed incubations, and final readings of the microplate in detector(s) appropriate for the assay. These configurable systems provide high throughput and rapid start up as well as a high degree of flexibility and customization. The manufacturers of such systems provide detailed protocols the various high throughput. Thus, for example, Zymark Corp. provides technical bulletins describing screening systems for detecting the modulation of gene transcription, ligand binding, and the like.

Optical images viewed (and, optionally, recorded) by a camera or other recording device (e.g., a photodiode and data storage device) are optionally further processed in any of the embodiments herein, e.g., by digitizing the image and/or storing and analyzing the image on a computer. A variety of commercially available peripheral equipment and software is available for digitizing, storing and analyzing a digitized video or digitized optical image, e.g., using PC (Intel x86 or Pentium chip-compatible DOS™, OS2™ WINDOWS™, WINDOWS NT™ or WINDOWS95™ based machines), MACINTOSH™, or UNIX based (e.g., SUN™ work station) computers. One conventional system carries light from the assay device to a cooled charge-coupled device (CCD) camera, in common use in the art. A CCD camera includes an array of picture elements (pixels). The light from the specimen is imaged on the CCD. Particular pixels corresponding to
regions of the specimen (*e.g.*, individual hybridization sites on an array of biological polymers) are sampled to obtain light intensity readings for each position. Multiple pixels are processed in parallel to increase speed. The apparatus and methods of the invention are easily used for viewing any sample, *e.g.*, by fluorescent or dark field microscopic techniques.

Integrated systems for assay analysis in the present invention typically include a digital computer with high-throughput liquid control software, image analysis software, data interpretation software, a robotic liquid control armature for transferring solutions from a source to a destination operably linked to the digital computer, an input device (*e.g.*, a computer keyboard) for entering data to the digital computer to control high throughput liquid transfer by the robotic liquid control armature and, optionally, an image scanner for digitizing label signals from labeled assay components. The image scanner interfaces with the image analysis software to provide a measurement of probe label intensity.

Of course, these assay systems can also include integrated systems incorporating nucleic acid selection elements, such as a computer, database with nucleic acid sequences of interest, sequence alignment software, and oligonucleotide selection software. In addition, this software can include components for ordering the selected oligonucleotides, and/or directing synthesis of oligonucleotides by an operably linked oligonucleotide synthesis machine. Thus, the integrated system elements of the invention optionally include any of the above components to facilitate high throughput recombination and selection. It will be appreciated that these high-throughput recombination elements can be in systems separate from those for performing selection assays, or the two can be integrated.

**EXAMPLE: SHUFFLING OF A CYTOKINE**

Libraries of synthetic genes of a cytokine (designated Cytokine G) were generated that contain 3 introns at positions which are new as compared to any known natural form of the cytokine. The introns were derived from several human globin genes. In addition, splice site quasispecies were included to provide for multiple splice site sequences. The intron containing genes were cloned into a vector that allowed expression of the encoded proteins on the surface of transfected cells. The vector also encodes a C-terminal
epitope tag that allows the detection of expressed proteins by FACS or Western blot using an epitope-specific antibody.

The constructs were transfected into COS cells using protoplast fusion or superfect transfection. Upon splicing of the inserted introns in the transfected cells, translation of the spliced mRNA generated proteins presented on the cell surface that contained the epitope tag. Using FACS, expression of protein containing the epitope tag was detected in the libraries. Cells expressing protein were isolated by FACS sorting, and the transfected library inserts were isolated by PCR and recloned into the same expression vector. Upon retransfection of the PCR-rescued constructs, an enrichment of cells expressing epitope-tagged protein was observed. These observations suggest that library species which support splicing of the inserted introns were selected for.

 Modifications can be made to the method and materials as hereinbefore described without departing from the spirit or scope of the invention as claimed, and the invention can be put to a number of different uses, including:

The use of an integrated system to select and test insertion modified and/or shuffled nucleic acids for activity, including in an iterative process.

An assay, kit or system utilizing a use of any one of the selection strategies, materials, components, methods or substrates hereinbefore described. Kits will optionally additionally comprise instructions for performing methods or assays, packaging materials, one or more containers which contain assay, device or system components, or the like.

In an additional aspect, the present invention provides kits embodying the methods and apparatus herein. Kits of the invention optionally comprise one or more of the following: (1) a recombination or insertion component as described herein; (2) instructions for practicing the methods described herein, and/or for operating oligonucleotide synthesis or assembled gene selection procedures herein; (3) one or more assay component; (4) a container for holding nucleic acids, intron nucleic acids, intein sequences, enzymes, other nucleic acids, transgenic plants, animals, cells, or the like and, (5) packaging materials.

In a further aspect, the present invention provides for the use of any component or kit herein, for the practice of any method or assay herein, and/or for the use of any apparatus or kit to practice any assay or method herein.

While the foregoing invention has been described in some detail for purposes of clarity and understanding, it will be clear to one skilled in the art from a reading of this
disclosure that various changes in form and detail can be made without departing from the true scope of the invention. For example, all the techniques and materials described above can be used in various combinations. All publications and patent documents cited in this application are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication or patent document were so individually denoted.
WHAT IS CLAIMED IS:

1. A method of recombining a first and a second target nucleic acid, the method comprising:
   providing a first and a second target nucleic acid, wherein at least one of the first and second target nucleic acids comprise a plurality of homologous or non-homologous insertion nucleic acid sequences; and,
   recombining the first and second target nucleic acids, thereby providing a shuffled recombinant nucleic acid.

2. The method of claim 1, wherein the insertion nucleic acid sequences comprise at least one intein.

3. The method of claim 2, wherein the shuffled recombinant nucleic acid encodes a protein subsequence, which protein subsequence is spliced to a second protein subsequence to produce an active protein.

4. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in vitro.

5. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in vivo.

6. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in cis.

7. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in trans.

8. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in a spontaneous splicing reaction.

9. The method of claim 3, wherein the protein subsequence and the second protein subsequence are spliced in a controlled splicing reaction.
10. The method of claim 1, wherein the plurality of insertion nucleic acid sequences are present in both the first and second nucleic target acid.

11. The method of claim 1, wherein the plurality of insertion nucleic acid sequences at least partially comprise at least one nucleic acid subsequence which encodes one or more of: an intron, a self-splicing bacterial intron, a trans-splicing intron, an intein, an exon, an extein, a nucleic acid encoding a proteolyzed polypeptide sequence, a nucleic acid encoding a molecular tag, and a nucleic acid which is removed by site specific recombination.

12. The method of claim 1, wherein the first or second nucleic acid comprise at least about 10 mini exons or exteins.

13. The method of claim 1, the first or second target nucleic acid comprising at least about 10 insertion nucleic acid sequences.

14. The method of claim 1, wherein the insertion nucleic acid sequences modulate a recombination frequency between the first and second target nucleic acid.

15. The method of claim 1, wherein the insertion nucleic acid sequences modulate an expression level or expression pattern of the first target nucleic acid, the second target nucleic acid, or the shuffled recombinant nucleic acid in one or more cell.

16. The method of claim 1, wherein the insertion nucleic acid sequences modulate splicing of an RNA encoded by the first or second target nucleic acid.

17. The method of claim 16, wherein the insertion nucleic acid sequences comprise an intron.

18. The method of claim 17, wherein the intron is spliced in vitro or on vivo.

19. The method of claim 17, wherein the intron is spliced in cis or in trans.

20. The method of claim 1, wherein the insertion nucleic acid sequences are recombined with one or more parental nucleic acid to produce the first or second target nucleic acid.
21. The method of claim 20, wherein the parental nucleic acid corresponds to one or more of: a gene or cDNA encoding EPO, a gene or cDNA encoding an insulin protein, a gene or cDNA encoding a peptide hormone, a gene or cDNA encoding a cytokine, a gene or cDNA encoding an epidermal growth factor, a gene or cDNA encoding a fibroblast growth factor, a gene or cDNA encoding a hepatocyte growth factor, a gene or cDNA encoding insulin-like growth factor, a gene or cDNA encoding an interferon, a gene or cDNA encoding an interleukin, a gene or cDNA encoding a keratinocyte growth factor, a gene or cDNA encoding a leukemia inhibitory factor, a gene or cDNA encoding oncostatin M, a gene or cDNA encoding PD-ECSF, a gene or cDNA encoding PDGF, a gene or cDNA encoding pleiotropin, a gene or cDNA encoding SCF, a gene or cDNA encoding c-kit ligand, a gene or cDNA encoding VEGF, a gene or cDNA encoding G-CSF, a gene or cDNA encoding an oncogene, a gene or cDNA encoding a tumor suppressor, a gene or cDNA encoding a steroid hormone receptor, a gene or cDNA encoding a plant hormone, a gene or cDNA encoding a disease resistance gene, a gene or cDNA encoding an herbicide resistance gene, a gene or cDNA encoding a bacterial gene, a gene or cDNA encoding a monooxygenase, a gene or cDNA encoding a protease, a gene or cDNA encoding a nuclease, a gene or cDNA encoding a lipase, a gene or cDNA encoding a C-X-C chemokine, a gene or cDNA encoding a C-C chemokine, a gene or cDNA encoding an antibody V gene, a gene or cDNA encoding a cystein knot protein such as TGFβ, NGF, PDGFβ or the like, a gene or cDNA encoding a TNK family member, a gene or cDNA encoding CNTF, a gene or cDNA encoding 4F, a gene or cDNA encoding an RNase, an antibody, a peptide ligand, and an angiogenesis inhibitor.

22. The method of claim 1, the method further comprising: providing a first parental nucleic acid sequence and a second parental nucleic acid sequence, which first and second nucleic acid sequences are homologous or non-homologous; and, inserting a plurality of insertion nucleic acid sequences into one or more of the first and second parental nucleic acid sequences, thereby providing the first and the second target nucleic acids.

23. The method of claim 22, wherein the step of inserting the plurality of insertion nucleic acid sequences into one or more of the first and second parental nucleic acid
sequences is performed in a computer, wherein the first and second parental nucleic acid sequences are character strings in a computer data set.

24. The method of claim 22, wherein the step of inserting the plurality of insertion nucleic acid sequences into one or more of the first and second parental nucleic acid sequences is performed in a computer, wherein the first and second parental nucleic acid sequences are character strings in a computer data set, the method further comprising synthesizing the first and second target nucleic acids.

25. The method of claim 22, wherein the step of inserting the plurality of insertion nucleic acid sequences into one or more of the first and second parental nucleic acid sequences is performed by physically joining a plurality of subsequences of the first or second parental nucleic acid sequences to the plurality of insertion nucleic acid sequences.

26. The method of claim 22, wherein the first and second parental nucleic acid sequences hybridize under stringent conditions, and the first and second target nucleic acids do not hybridize under stringent conditions.

27. The method of claim 22, wherein the first and second parental nucleic acid sequences do not hybridize under stringent conditions, and wherein the first and second target nucleic acids hybridize under stringent conditions.

28. The method of claim 22, wherein the first and second nucleic target nucleic acid hybridize under stringent conditions, and wherein the first target nucleic acid does not hybridize under stringent conditions to the second parental nucleic acid, or wherein the second target nucleic acid does not hybridize under stringent conditions to the first parental nucleic acid.

29. The method of claim 22, wherein the first or second parental nucleic acid hybridizes to a third nucleic acid under stringent conditions, wherein the first and second target nucleic acids do not hybridize under stringent conditions to the third nucleic acid.

30. The method of claim 1, the method further comprising recombining the shuffled recombinant nucleic acid with a third nucleic acid, and, optionally, selecting the resulting secondary shuffled recombinant nucleic acid for a desired trait or property.
31. The method of claim 1, the method further comprising selecting the shuffled recombinant nucleic acid for a desired trait or property.

32. The method of claim 1, further comprising expressing the recombinant nucleic acid in a cell.

33. The method of claim 1, further comprising expressing the recombinant nucleic acid in a cell, thereby producing a protein, which protein is proteolytically cleaved to produce an active protein, or to remove an intein.

34. The method of claim 1, the first and second target nucleic acids each comprising a plurality of insertion nucleic acids, wherein, during recombination of the first and second target nucleic acids, the crossover frequency between the insertion sequences in the first and second nucleic acids is higher than the crossover frequency of non-insertion sequences in the first and second nucleic acids.

35. The method of claim 1, wherein the recombinant nucleic acid encodes a molecule which does not comprise or encode a translated insertion sequence.

36. The method of claim 35, wherein the molecule is selected from a DNA, an RNA, an mRNA, a viral RNA, an sn RNA, a tRNA, an rRNA, a gRNA, a protein, and a proteolytically cleaved protein.

37. The method of claim 1, wherein the recombinant nucleic acid encodes a protein with an activity selected from an insulin protein activity, a peptide hormone activity, a cytokine activity, an epidermal growth factor activity, a fibroblast growth factor activity, a hepatocyte growth factor activity, an insulin-like growth factor activity, an interferon activity, an interleukin activity, a keratinocyte growth factor activity, a leukemia inhibitory factor activity, an oncostatin M activity, a PD-ECSF activity, a pleiotropin activity, an SCF activity, a e-kit ligand activity, a VEGF activity, a G-CSF activity, a GM-CSF activity, a plant hormone activity, a disease resistance protein activity, a bacterial protein activity, a protease activity, a peptide ligand activity, an angiogenesis inhibitor activity, a C-X-C chemokine activity, a C-C chemokine activity, a cystein knot protein activity, and an EPO activity, wherein the recombinant nucleic acid does not hybridize under stringent conditions.
to a cDNA which encodes said activity, which cDNA is a copy of a naturally occurring mRNA.

38. The method of claim 1, wherein the first or second nucleic acid comprises at least one intronic splicing enhancer (ISE).

39. The method of claim 1, wherein the first or second nucleic acid comprises at least one chicken cardiac troponin T (cTNT) ISE.

40. The method of claim 1, wherein the first nucleic acid comprises two non-homologous subsequences and a plurality of insertion subsequences.

41. The method of claim 1, wherein the first or second nucleic acid or recombinant nucleic acid are present in an expression vector.

42. The method of claim 1, wherein the first and second target nucleic acids are derived from a first and second parental nucleic acid by integration of a plurality of insertion sequences into the first and second parental nucleic acid, wherein the first and second parental nucleic acid are less than 50% identical over the full length of the first and second parental nucleic acid, when the first and second nucleic acids are aligned for maximum identity.

43. The method of claim 1, wherein the first and second target nucleic acids are derived from a first and second parental nucleic acid by integration of a plurality of insertion sequences into the first and second parental nucleic acid, wherein the first and second parental nucleic acid are less than 25% identical over the full length of the first and second parental nucleic acid, when the first and second nucleic acids are aligned for maximum identity.

44. A recombination mixture comprising a first target nucleic acid comprising a plurality of insertion subsequences, the mixture further comprising a second target nucleic acid comprising at least one region of sequence similarity to the first nucleic acid.

45. The recombination mixture of claim 44, wherein the second target nucleic acid comprises a plurality of insertion subsequences.
46. A recombination mixture resulting from fragmenting a first target nucleic acid comprising a plurality of insertion subsequences, and a second target nucleic acid comprising at least one region of sequence similarity to the first target nucleic acid.

47. The recombination mixture of claim 46, wherein the first and second target nucleic acids are fragmented with a DNase.

48. The recombination mixture of claim 46, wherein the first and second target nucleic acids are fragmented by chemically synthesizing fragments of the first and second target nucleic acid.

49. A recombinant nucleic acid produced by recombining the recombination mixture of claim 46.

50. The recombination mixture of claim 46 wherein the first or second nucleic acid comprises one or more subsequence corresponding to one or more subsequence from one or more gene or cDNA selected from: a gene or cDNA encoding EPO, a gene or cDNA encoding an insulin protein, a gene or cDNA encoding a peptide hormone, a gene or cDNA encoding a cytokine, a gene or cDNA encoding an epidermal growth factor, a gene or cDNA encoding a fibroblast growth factor, a gene or cDNA encoding a hepatocyte growth factor, a gene or cDNA encoding insulin-like growth factor, a gene or cDNA encoding an interferon, a gene or cDNA encoding an interleukin, a gene or cDNA encoding a keratinocyte growth factor, a gene or cDNA encoding a leukemia inhibitory factor, a gene or cDNA encoding oncostatin M, a gene or cDNA encoding PD-ECSF, a gene or cDNA encoding PDGF, a gene or cDNA encoding pleiotropin, a gene or cDNA encoding SCF, a gene or cDNA encoding c-kit ligand, a gene or cDNA encoding VEGF, a gene or cDNA encoding G-CSF, a gene or cDNA encoding an oncogene, a gene or cDNA encoding a tumor suppressor, a gene or cDNA encoding a steroid hormone receptor, a gene or cDNA encoding a plant hormone, a gene or cDNA encoding a disease resistance gene, a gene or cDNA encoding an herbicide resistance gene, a gene or cDNA encoding a bacterial gene, a gene or cDNA encoding a monooxygenase, a gene or cDNA encoding a protease, a gene or cDNA encoding a nuclease, a gene or cDNA encoding an RNase, a gene or cDNA encoding a C-X-C chemokine, a gene or cDNA encoding a C-C chemokine, a gene or cDNA encoding an antibody V gene, a gene or cDNA encoding a cystein knot protein such as...
TGFβ, NGF, PDGFβ or the like, a gene or cDNA encoding a TNK or family member, a gene or cDNA encoding CNTF, a gene or cDNA encoding 4F and a gene or cDNA encoding a lipase.

51. The recombination mixture of claim 46 wherein the first or second target nucleic acids are present in one or more expression vector.

52. A method of recombining a plurality of sequence domains from a plurality of homologous or non-homologous nucleic acid sequences, the method comprising:
   providing a pre-mRNA comprising a plurality of sequence domains which correspond to a plurality of different parental nucleic acid sequences; and,
   alternatively splicing the pre-mRNA to produce a plurality of different mRNAs comprising a plurality of different sets of sequence domains.

53. The method of claim 52, the pre-mRNA comprising between about 6 and about 20 exons or exetins.

54. The method of claim 52, the pre-mRNA comprising a plurality of mini exons or exetins.

55. The method of claim 52, further comprising selecting one or more of the plurality of different mRNAs for a desired trait or property.

56. The method of claim 52, further comprising cloning one or more of the plurality of different mRNAs.

57. The method of claim 52, further comprising recombining one or more of: the plurality of different mRNAs, the pre-mRNA, a DNA encoding the mRNA, and a DNA encoding the pre-mRNA, with one or more additional nucleic acid.

58. The method of claim 52, wherein the pre-mRNA is provided in a cell by transducing the cell with a vector comprising a DNA encoding the pre-mRNA.

59. A method of making a nucleic acid with a desired splicing phenotype, the method comprising:
providing a plurality of homologous nucleic acids, each comprising a plurality of
insertion nucleic acid sequences;
recombining the plurality of homologous nucleic acids to produce a library of
recombinant nucleic acids, and
selecting the recombinant nucleic acids for a selected recombinant nucleic acid
which produces a desired mRNA or protein when the selected recombinant nucleic acid is
expressed in a cell.

60. The method of claim 59, the method further comprising recombining the desired
recombinant nucleic acid with an additional nucleic acid and selecting the resulting
secondary recombinant nucleic acid for production of a desired mRNA or protein.

61. The method of claim 59, wherein the plurality of homologous nucleic acids each
comprise at least about 10 exons or exteins.

62. The method of claim 59, wherein the insertion nucleic acid sequences are
selected from bacterial introns and eukaryotic introns.

63. The method of claim 59, wherein the plurality of homologous nucleic acids are
recombined in vivo.

64. The method of claim 59, wherein the plurality of homologous nucleic acids are
recombined in vitro.

65. A method of producing a selected protein, the method comprising:
providing a first shuffled nucleic acid subsequence encoding a first portion of the
selected protein and a second nucleic acid subsequence encoding a second portion of the
selected protein;
expressing the first and second subsequences to produce a first protein subsequence
and a second protein subsequence; and,
splicing the first and second protein subsequence to produce the selected protein.

66. The method of claim 65, comprising providing a third nucleic acid subsequence,
expressing the third nucleic acid subsequence to produce a third protein subsequence, and
splicing the third protein subsequence with the first or the second proteins subsequence, or both, to produce the selected protein.

67. The method of claim 66, wherein the second or third nucleic acid subsequence is a shuffled nucleic acid subsequence.

68. The method of claim 66, wherein the first, second and third nucleic acid subsequence are part of a single nucleic acid.

69. The method of claim 66, wherein the first, second and third nucleic acid subsequence are part of a more than one nucleic acid.

70. The method of claim 66, wherein one or more of the first, second or third nucleic acid subsequences comprise one or more homologous or non-homologous insertion nucleic acid sequences.

71. The method of claim 65, wherein the splicing is performed in vitro, in vivo, or both in vitro and in vivo.

72. The method of claim 65, wherein the splicing is performed by a spontaneous splicing reaction between two or more inteins or exteins.

73. The method of claim 65, wherein the splicing is performed by a controlled splicing reaction between two or more inteins or exteins.

74. The method of claim 65, wherein the splicing is performed by a cis splicing reaction between two or more inteins or exteins.

75. The method of claim 65, wherein the splicing is performed by a trans splicing reaction between two or more inteins or exteins.

76. A method of producing a selected RNA, the method comprising:

providing a first shuffled nucleic acid subsequence encoding a first portion of the selected RNA and a second nucleic acid subsequence encoding a second portion of the selected RNA; and,
splicing the first and second nucleic acid subsequences, or RNA copies thereof, to produce the selected RNA.

77. The method of claim 76, wherein the selected RNA encodes a protein.

78. The method of claim 76, wherein the selected RNA encodes a protein or protein subsequence comprising an intein.

79. The method of claim 76, wherein the selected RNA is expressed to produce a protein or a protein subsequence comprising an intein.

80. The method of claim 76, comprising providing a third nucleic acid subsequence, and splicing the third nucleic acid subsequence, or an RNA copy thereof, to produce the selected RNA.

81. The method of claim 80, wherein one or more of the first, second or third nucleic acid subsequences comprise one or more homologous or non-homologous insertion nucleic acid sequences.

82. The method of claim 80, wherein the second or third nucleic acid subsequence is a shuffled nucleic acid subsequence.

83. The method of claim 80, wherein the first, second and third nucleic acid subsequence are part of a single nucleic acid.

84. The method of claim 80, wherein the first, second and third nucleic acid subsequence are part of a more than one nucleic acid.

85. The method of claim 76, wherein the splicing is performed in vitro, in vivo, or both in vitro and in vivo.

86. The method of claim 76, wherein the splicing is performed by a trans splicing reaction between the nucleic acid subsequences.

87. The method of claim 76, wherein the splicing is performed by a cis splicing reaction between the nucleic acid subsequences.