Methods and apparatus are disclosed herein for encoding human readable text conveying a non-genetic message into nucleic acid sequences with a substantially reduced probability of biological impact and decoding such text from nucleic acid sequences. In one embodiment, each symbol of a symbol set of human readable symbols uniquely maps to a respective codon identifier. Mapping may ensure that each symbol will not map to a codon identifier that generates an amino acid residue which has a single-letter abbreviation that is the equivalent to the respective symbol. Synthetic nucleic acid sequences comprising such human readable text, and recombinant or synthetic cells comprising such sequences are provided, as well as methods of identifying cells, organisms, or samples containing such sequences.
Synthetic nucleic acid sequence 104 into a cell 114.

Extract nucleic acid or cells from living organism 116.

FIG. 1
Transcoder 200

Power Supply 202

Processor 204

Volatile memory 206

Non-volatile memory 208

I/O Module 216

Input 212

Output 214

Transcoder module 210

Symbol maps 220

Symbol frequency analyzer 230

FIG. 2
### Symbol Map 220

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Equivalent</th>
</tr>
</thead>
<tbody>
<tr>
<td>GGA = &quot;</td>
<td>GAA = *</td>
</tr>
<tr>
<td>AGG = I</td>
<td>AAA = {</td>
</tr>
<tr>
<td>AGC = &gt;</td>
<td>CAC = /</td>
</tr>
<tr>
<td>CCT = +</td>
<td>CCA = =</td>
</tr>
<tr>
<td>ACC = &amp;</td>
<td>ATG = *</td>
</tr>
<tr>
<td>TCT = 0</td>
<td>CTT = 1</td>
</tr>
<tr>
<td>GCC = 6</td>
<td>TAT = 7</td>
</tr>
<tr>
<td>TAG = A</td>
<td>AGT = B</td>
</tr>
<tr>
<td>TAC = G</td>
<td>TCA = H</td>
</tr>
<tr>
<td>CAA = M</td>
<td>TGC = N</td>
</tr>
<tr>
<td>GCT = S</td>
<td>TGA = T</td>
</tr>
<tr>
<td>CAT = Y</td>
<td>TGG = Z</td>
</tr>
<tr>
<td>TTC = #</td>
<td>CAG = :</td>
</tr>
<tr>
<td>CCG = (</td>
<td>GAC = )</td>
</tr>
<tr>
<td>CGA = .</td>
<td>GTG = ,</td>
</tr>
<tr>
<td>CGC = -</td>
<td>GAT = %</td>
</tr>
<tr>
<td>GGG = newline</td>
<td></td>
</tr>
</tbody>
</table>

Symbols 304

Codon identifiers 302

**FIG. 3**
Generate analysis of symbol frequency in reference language

Identify least frequently occurring symbol

Map least frequently occurring symbol to start codon

Receive next available codon identifier

Does codon identifier yield an amino acid with an abbreviation equal to unmapped symbol?

Any more unmapped symbols?

Map three most frequently occurring symbols to stop codons

Any more unmapped symbols?

Map unmapped symbol to codon identifier

FIG. 4
FIG. 5

Any more symbols in input sequence?  

Supportable input symbol?  

Generate codon identifier corresponding to input symbol  

Prepend all-6 reading frame stop codon containing sequence to encoded  

Append all-6 reading frame stop codon containing sequence to encoded  

Generate codon identifier message
START

Any more codon identifiers? 602

END

N

Y

Determine symbol corresponding to codon identifier 604

Output symbol 606

FIG. 6
START

Generate one or more codon identifiers for one or more symbols of a reference language

Identify or create a watermark containing one or more symbols of a reference language

Substitute the one or more symbols of a reference language with the one or more codon identifiers

Prepend all-6 reading frame stop codon containing sequence to encoded message

Append all-6 reading frame stop codon containing sequence to encoded message

Create synthetic nucleic acid sequence containing the encoded message

END

FIG. 7
Alt Full-Monty-Code

<table>
<thead>
<tr>
<th>1st Base in Full-Monty-Code</th>
<th>2nd Base in Full-Monty-Code</th>
<th>3rd Base in Full-Monty-Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>T</td>
<td>G</td>
</tr>
<tr>
<td>tab</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>T</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>T</td>
<td></td>
</tr>
<tr>
<td>space</td>
<td></td>
<td></td>
</tr>
<tr>
<td>G</td>
<td>SHIFT</td>
<td>T</td>
</tr>
<tr>
<td></td>
<td>CTRL</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>ALT</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>LOCK</td>
<td>new-line</td>
</tr>
</tbody>
</table>

FIG. 8A
### Ctrl Full-Monty-Code

#### 2nd Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>T</td>
</tr>
<tr>
<td>tab</td>
<td></td>
<td>A</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>C</td>
<td></td>
</tr>
</tbody>
</table>

#### 1st Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>space</td>
<td></td>
</tr>
<tr>
<td>shift</td>
<td>T</td>
</tr>
<tr>
<td>ctrl</td>
<td>C</td>
</tr>
<tr>
<td>alt</td>
<td>A</td>
</tr>
<tr>
<td>lock</td>
<td>new-line</td>
</tr>
</tbody>
</table>

#### 3rd Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
</tr>
<tr>
<td>C</td>
</tr>
<tr>
<td>A</td>
</tr>
</tbody>
</table>

**FIG. 8B**
### Default Full-Monty-Code

#### 2nd Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>c</td>
<td>0</td>
<td>7</td>
<td>;</td>
</tr>
<tr>
<td>tab</td>
<td>u</td>
<td>g</td>
<td>n</td>
</tr>
<tr>
<td>q</td>
<td>h</td>
<td>e</td>
<td>t</td>
</tr>
<tr>
<td>v</td>
<td>a</td>
<td>z</td>
<td>G</td>
</tr>
<tr>
<td>l</td>
<td>,</td>
<td>y</td>
<td>o</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>/</td>
<td>g</td>
</tr>
<tr>
<td>r</td>
<td>=</td>
<td>m</td>
<td>.</td>
</tr>
</tbody>
</table>

#### 3rd Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>b</td>
</tr>
<tr>
<td>S</td>
<td>&amp;</td>
<td>l</td>
<td></td>
</tr>
<tr>
<td>space</td>
<td>p</td>
<td>4</td>
<td>A</td>
</tr>
<tr>
<td>[</td>
<td>]</td>
<td>G</td>
<td></td>
</tr>
<tr>
<td>j</td>
<td>s</td>
<td>SHIFT</td>
<td>x</td>
</tr>
<tr>
<td>w</td>
<td>6</td>
<td>CTRL</td>
<td>f</td>
</tr>
<tr>
<td>9</td>
<td>k</td>
<td>ALT</td>
<td>A</td>
</tr>
<tr>
<td>,</td>
<td>5</td>
<td>LOCK</td>
<td>new-line</td>
</tr>
</tbody>
</table>

**FIG. 8C**
### Shift Full-Monty-Code

#### 2\textsuperscript{nd} Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>)</td>
<td>&amp;</td>
<td>:</td>
</tr>
<tr>
<td>tab</td>
<td>U</td>
<td>G</td>
<td>N</td>
</tr>
<tr>
<td>Q</td>
<td>H</td>
<td>E</td>
<td>T</td>
</tr>
<tr>
<td>V</td>
<td>A</td>
<td>Z</td>
<td>G</td>
</tr>
<tr>
<td>!</td>
<td>&quot;</td>
<td>Y</td>
<td>O</td>
</tr>
<tr>
<td></td>
<td></td>
<td>?</td>
<td>*</td>
</tr>
<tr>
<td>R</td>
<td>+</td>
<td>M</td>
<td>&gt;</td>
</tr>
<tr>
<td>G</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### 1\textsuperscript{st} Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>A</th>
<th>D</th>
<th>@</th>
<th>#</th>
<th>B</th>
<th>T</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>&amp;</td>
<td>L</td>
<td>C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>space</td>
<td>P</td>
<td></td>
<td>S</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>}</td>
<td></td>
<td>}</td>
<td>G</td>
<td></td>
</tr>
</tbody>
</table>

#### 3\textsuperscript{rd} Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>G</th>
<th>J</th>
<th>S</th>
<th>SHIFT</th>
<th>X</th>
<th>T</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>W</td>
<td>^</td>
<td>CTRL</td>
<td>P</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>(</td>
<td>K</td>
<td>ALT</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;</td>
<td>%</td>
<td>LOCK</td>
<td>new-line</td>
<td>G</td>
</tr>
</tbody>
</table>

**FIG. 8D**
## Monty Code

### 2nd Base in Full-Monty-Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>O</td>
<td>7</td>
<td>;</td>
</tr>
<tr>
<td>#</td>
<td>U</td>
<td>G</td>
<td>N</td>
</tr>
<tr>
<td>Q</td>
<td>H</td>
<td>E</td>
<td>T</td>
</tr>
<tr>
<td>V</td>
<td>@</td>
<td>A</td>
<td>Z</td>
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<tr>
<td>\</td>
<td>+</td>
<td>Y</td>
<td>O</td>
</tr>
<tr>
<td>/</td>
<td>-</td>
<td>/</td>
<td>8</td>
</tr>
<tr>
<td>R</td>
<td>=</td>
<td>M</td>
<td>A</td>
</tr>
<tr>
<td>1</td>
<td>)</td>
<td>:</td>
<td>&lt;</td>
</tr>
<tr>
<td>D</td>
<td>2</td>
<td>3</td>
<td>B</td>
</tr>
<tr>
<td>$</td>
<td>&amp;</td>
<td>\</td>
<td>&gt;</td>
</tr>
<tr>
<td>P</td>
<td>(</td>
<td>\</td>
<td>4</td>
</tr>
<tr>
<td>]</td>
<td>*</td>
<td>}</td>
<td>A</td>
</tr>
<tr>
<td>J</td>
<td>S</td>
<td>%</td>
<td>X</td>
</tr>
<tr>
<td>W</td>
<td>6</td>
<td></td>
<td>F</td>
</tr>
<tr>
<td>K</td>
<td>9</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>,</td>
<td>5</td>
<td>!</td>
<td>new-line</td>
</tr>
</tbody>
</table>

### 3rd Base in Monty Code

<table>
<thead>
<tr>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>O</td>
<td>7</td>
<td>;</td>
</tr>
<tr>
<td>#</td>
<td>U</td>
<td>G</td>
<td>N</td>
</tr>
<tr>
<td>Q</td>
<td>H</td>
<td>E</td>
<td>T</td>
</tr>
<tr>
<td>V</td>
<td>@</td>
<td>A</td>
<td>Z</td>
</tr>
<tr>
<td>\</td>
<td>+</td>
<td>Y</td>
<td>O</td>
</tr>
<tr>
<td>/</td>
<td>-</td>
<td>/</td>
<td>8</td>
</tr>
<tr>
<td>R</td>
<td>=</td>
<td>M</td>
<td>A</td>
</tr>
<tr>
<td>1</td>
<td>)</td>
<td>:</td>
<td>&lt;</td>
</tr>
<tr>
<td>D</td>
<td>2</td>
<td>3</td>
<td>B</td>
</tr>
<tr>
<td>$</td>
<td>&amp;</td>
<td>\</td>
<td>&gt;</td>
</tr>
<tr>
<td>P</td>
<td>(</td>
<td>\</td>
<td>4</td>
</tr>
<tr>
<td>]</td>
<td>*</td>
<td>}</td>
<td>A</td>
</tr>
<tr>
<td>J</td>
<td>S</td>
<td>%</td>
<td>X</td>
</tr>
<tr>
<td>W</td>
<td>6</td>
<td></td>
<td>F</td>
</tr>
<tr>
<td>K</td>
<td>9</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>,</td>
<td>5</td>
<td>!</td>
<td>new-line</td>
</tr>
</tbody>
</table>

**FIG. 8E**
ENCODING TEXT INTO NUCLEIC ACID SEQUENCES

CROSS-REFERENCES TO RELATED APPLICATIONS


FIELD OF THE INVENTION

[0002] The presently disclosed application relates generally to the field of molecular biology. More specifically, this application relates to synthetic nucleic acid sequences comprising non-genetic information.

REFERENCE TO SEQUENCE LISTING

[0003] This application contains references to amino acid sequences and/or nucleic acid sequences which have been submitted concurrently herewith as the sequence listing text file “SG1450_1.ST25.txt”, file size 09.02 KiloBytes (KB), created on Oct. 29, 2010. The aforementioned sequence listing is hereby incorporated by reference in its entirety pursuant to 37 C.F.R. §1.52(e)(5).

BACKGROUND OF THE INVENTION

[0004] Biological organisms comprise nucleic acid sequences that encode, among other things, genes that can be used to catalyze chemical reactions within the organism. Genes encompass a number of different regions, such as promoters, terminators, and perhaps most importantly, the open reading frame or coding region of the gene that contains the “text” for the protein of interest. Deoxynucleic acid sequences are transcribed to messenger RNA (mRNA) which is then translated into a protein sequence of interest. There are four bases used in deoxyribonucleic acid sequences, each of which can be used in three positions in a codon and, thus, there are in theory 64 possible codon permutations.

[0005] Because many cells have the ability to absorb and retain nucleic acid sequences, some have considered the prospect of using a biological organism as a memory source for storing human readable information. In order to achieve this end, various encoding schemes have been developed which attempt to map human readable symbols into nucleic acid sequences that can be stored within a living organism.

[0006] U.S. Pat. No. 6,312,911 discloses a steganographic method for creation of a secret code by producing a DNA molecule comprising a secret message DNA sequence flanked on each side by a primer sequence.

[0007] U.S. Pat. No. 7,056,724 discloses a method of storing data in Deinococcus radius by performing an evaluation of the genome of D. radius relative to one or more criteria for use as a storage medium, preparing a code based on the evaluation, encoding a DNA sequence in accordance to the code to represent the data, and incorporating the encoded DNA sequence into D. radius.

[0008] U.S. Published Application No. 20080124725 discloses a method of tagging a bacterium by exposing a portion of a CRISPR locus to at least one exogenous nucleic acid sequence to produce at least one tagged bacterium comprising a modified CRISPR locus.

[0009] U.S. Pat. No. 6,175,830 discloses a method for generating a searchable informational resource by assigning a categorical tag to each of a plurality of finite elements and arranging the results of a searchable step into a hierarchical structure according to information in categorical tags assigned to finite elements corresponding to searchable database records identified by searching.

[0010] U.S. Pat. No. 7,323,307 discloses a method for analyzing mRNA having one or more exons having an order defined by one or more signature sequences, by hybridizing labeled fragments to a nucleic acid array and determining the identity of sequence signatures and the order of one or more exons.

[0011] U.S. Pat. No. 6,607,878 discloses a composition of a mixture of different species of molecules, where at least some of the molecules are derived from a combinatorial synthesis process and some species are linked to a tag of linked information encoding elements, and where the physical property of the combination of elements identifies the species of molecule without determining the physical property of each element of the tag and, further, where the tags do not consist solely of nucleotides.

[0012] Clelland et al. (Nature, volume 399, pages 533-534 (1999)) discloses concealing secret messages hidden in DNA microdroms where the messages are flanked by PCR primer sequences.

[0013] Heider and Barnekow (BMC Molec. Biol., 9: 40 (2008) and Heider and Barnekow (BMC Bioinformatics, 8:176 (2007)) each disclose the application of watermarks based on DNA sequences using a binary code of the numbers 0 and 1.

[0014] Leier et al. (Biosystems, 57: 13-22 (2000)) disclose two different cryptographic techniques, each of which requires binary codes of the numbers 0 and 1.

[0015] Arita and Ohashi (Biotechnol. Prog. 20: 1605-1607 (2004)) disclose an encoding scheme in which the actual sequence of the gene that encodes the message is not independent of the genetic code of the organism; the correct decoding of an embedded sequence requires outside knowledge of the wild-type sequence of a carrier gene; and a minimum of 18 nucleotides is required to encode a single letter.

[0016] Unfortunately, conventional encoding schemes suffer from a one of two serious drawbacks, either they run the risk of causing a negative biological impact on a cell harboring nucleic acid sequences made using such encoding schemes, or they rely upon codon-redundancy in a carrier-gene of known function. Methods using a carrier-gene are characterized by extreme inefficiency of encoded information and are further limited by their requirement of encoding a message inside a carrier gene of known sequence and limited length. This imposes a limit on the length of the message that can be encoded that is further exacerbated by the inefficiency of the encoding scheme.

SUMMARY OF THE INVENTION

[0017] The present application is directed to generating an encoding scheme configured to translate human readable symbols into codon identifiers (i.e., discrete sequences of preferably three elements, where each element contains one of four selected nucleotide bases). In this manner, sequences of human readable symbols can be used to convey non-ge genetic messages (for example, text messages, trademarks, copyright notices, unique identifying information, etc.) by encoding the message into sequences of codon identifiers. These sequences of codon identifiers may then be used to generate synthetic nucleic acid sequences that are introduced
into a living cell or organism as free DNA or incorporated into other various types of cellular nucleic acid materials (e.g., plasmids, chromosomes, mitochondrial DNA, genomes, etc.). The resulting set of codons or codon identifiers effectively serves as a memory source for the encoded sequences of human readable symbols.

In another aspect, provided herein is a recombinant or synthetic organism containing a synthetic nucleic acid sequence as described herein.

In various embodiments, the recombinant or synthetic organism can be a prokaryotic cell, a eukaryotic cell, an archaeal cell or a virus. In certain preferred embodiments, the recombinant cell can be a bacterial cell, a yeast cell, a fungal cell, an algal cell, an animal cell, or a plant cell. In certain embodiments, the set of human readable symbols can be a watermark that allows the authentication or identification of the recombinant or synthetic cell or virus comprising the synthetic nucleic acid sequence containing the watermark, or of an organism comprising such a recombinant or synthetic cell or virus.

In another aspect, provided herein is a method of creating a recombinant or synthetic organism comprising a watermark that conveys a non-genetic message, comprising: generating a nucleic acid sequence comprising a sequence of codon identifiers selected based upon the text of the watermark such that a symbol mapping maps codon identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionately low frequency in the language of the watermark, and maps codon identifiers corresponding to stop codon(s) to human readable symbols that possess a disproportionately high frequency in the language of the watermark; synthesizing this nucleic acid sequence; and introducing this nucleic acid sequence into a recombinant or synthetic organism.

Alternatively, provided herein is a method of creating a recombinant or synthetic organism comprising a water mark that conveys a non-genetic message, comprising: generating a nucleic acid sequence comprising one or more codon identifiers from a set of human readable symbols of a reference language comprising said watermark, wherein a symbol mapping is configured to map a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and wherein the symbol mapping is further configured to map a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon; synthesizing this nucleic acid sequence; and introducing this nucleic acid sequence into a recombinant or synthetic organism.

In preferred embodiments, the symbol mapping does not map a three nucleotide codon identifier to a single letter representation of an amino acid residue normally assigned to that three nucleotide codon in the standard genetic code. In certain embodiments, the generating step is computer-assisted and comprises identifying the set of human readable symbols at a memory module and for each human readable symbol in the set, using a processor to read a symbol mapping for determining a codon identifier which maps to the respective human readable symbol.

A recombinant cell, a synthetic cell, a recombinant virus, a synthetic virus, or a recombinant or synthetic multicellular organism comprising such a non-genetic message can be used for any suitable purpose as is known in the art, for example, in connection with a recombinant plant or crop (e.g., corn, grapes, etc.); a modified animal (e.g., a genetically modified rodent, primate, poultry, large veterinary animal, etc.); a recombinant embryo; a genetically modified organism, cell, cell line or strain; a recombinant organism, cell, cell
line or strain; a synthetic organism, cell, cell line or strain; a recombinant virus or strain; a synthetic virus or strain; and the like.

[0029] In another aspect, provided herein is a method of determining the presence of a recombinant or synthetic organism, which may be a single cell, a multicellular organism or a virus, comprising a reference watermark that conveys a non-genetic message in a sample, said method comprising: sequencing nucleic acid material obtained from one or more organisms in said sample; transforming the nucleic acid sequence to a set of codon identifiers, wherein each codon identifier consists of three nucleotides of said sequence, and the transforming is performed in all three reading frames; determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determination is based at least in part upon a symbol mapping that map codons identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionately low frequency in the language of the watermark, and that maps codon identifiers corresponding to stop codon(s) to human readable symbols that possess a disproportionately high frequency in the language of the watermark; and comparing the human readable symbol sequence of all three reading frames to the reference watermark in said recombinant or synthetic organism, whereby the presence of the reference watermark in any reading frame of the nucleic acid material indicates the presence of the recombinant or synthetic organism in the sample.

[0030] Alternatively, provided herein is a method of determining the presence of a recombinant or synthetic organism, which may be a single cell, a multicellular organism or a virus, comprising a reference watermark that conveys a non-genetic message in a sample, said method comprising: sequencing nucleic acid material obtained from one or more organisms in said sample; transforming the nucleic acid sequence to a set of codon identifiers, wherein each codon identifier consists of three nucleotides of said sequence, and the transforming is performed in all three reading frames; determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determination is based at least in part upon a symbol mapping that is configured to map a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols and is further configured to map a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols; and comparing the human readable symbol sequence of all three reading frames to the reference watermark in said recombinant or synthetic organism, whereby the presence of the reference watermark in any reading frame of the nucleic acid material indicates the presence of the recombinant or synthetic organism in the sample.

[0031] A sample can be any sample that can contains a cell, multiple cells, a virus, or nucleic acid material from a cell, cells or virus, including without limitation, environmental samples, patient samples, veterinary samples, samples obtained from humans, animals, plants, viruses, bacteria, archaea, yeast, and any fractions or derivatives of any such samples. Samples can also be laboratory samples (e.g., for-profit and non-profit laboratories) and commercial samples.

[0032] In another aspect, provided herein is an apparatus for transforming a sequence of codon identifiers into a sequence of human readable symbols that conveys a non-genetic message, the apparatus comprising: a processor adapted to execute instructions; and a storage module, wherein the storage module comprises a data structure for mapping codon identifiers into human readable symbols, and a set of instructions which, when executed by the processor, generate a human readable symbol for each codon identifier read from a sequence of codon identifiers, wherein the human readable symbol generated is based at least in part upon the data structure; wherein the data structure is configured to map a start codon to a human readable symbol with a frequency of occurrence within a reference language that is less than a first predetermined threshold, and wherein the data structure is further configured to map a plurality of stop codons to human readable symbols with frequencies of occurrence within the reference language that are greater than a second predetermined threshold.

[0033] In another embodiment, the data structure maps a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols, and further maps a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols.

[0034] In one preferred embodiment, the data structure does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code. In another embodiment, the sequence of codon identifiers comprises at least one of an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence, and/or an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

[0035] In another aspect, provided herein is a computer-readable medium for use in a decoding machine, the computer-readable medium comprising instructions which, when executed by the decoding machine, perform a process comprising: identifying a sequence of codon identifiers in all three reading frames; and generating a human readable symbol for each codon identifier in the sequence; wherein the human readable symbol generated is based at least in part upon a mapping function configured to map a start codon to a human readable symbol that has a frequency of occurrence within a reference language that is smaller than every other human readable symbol of a first set of human readable symbols, and wherein the mapping function is further configured to map a stop codon to a human readable symbol that has a frequency of occurrence within the reference language that is larger than every other human readable symbol of the first set of human readable symbols.

[0036] Also provided herein is a computer-readable medium for use in a decoding machine, the computer-readable medium comprising instructions which, when executed by the decoding machine, perform a process comprising: identifying a sequence of codon identifiers in all three reading frames; and generating a human readable symbol for each codon identifier in the sequence of codon identifiers, wherein the human readable symbol generated is based upon a mapping function that maps codon identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionately low frequency in the language of the watermark, and that maps codon identifiers corresponding to stop codon(s) to human readable symbols that possess a disproportionately high frequency in the language of the watermark (which conveys a non-genetic message).

[0037] In another aspect, provided herein is a method of transforming a first signal adapted to indicate a sequence of
codon identifiers into a second signal adapted to indicate a sequence of human readable symbols that conveys a non-genetic message, the method comprising: receiving the first signal; determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determining is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a symbol sequence than one or more human readable symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the symbol sequence than one or more human readable symbols from the set of human readable symbols; and transforming the first signal into the second signal based upon the one or more determined human readable symbols.

[0038] Also provided herein is a method of transforming a first signal comprising a sequence of codon identifiers into a second signal to indicate a sequence of human readable symbols of a set of human readable symbols of a reference language that conveys a non-genetic message, the method comprising: identifying the first signal that indicates the sequence of codon identifiers; determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determining a human readable symbol is based at least in part upon a mapping function that maps a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols, and further maps a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols; and transforming the first signal into the second signal, wherein the second signal indicates the sequence of human readable symbols.

[0039] In another aspect, provided herein is an apparatus for converting a sequence of human readable symbols of a reference language that conveys a non-genetic message into a sequence of codon identifiers, the apparatus comprising: a processor configured to execute instructions; a memory module coupled to the processor and comprising instructions which, when executed by the processor, determine a codon identifier for each human readable symbol contained within the sequence of human readable symbols, wherein each codon identifier is determined upon reading a symbol map; and a data module coupled to the memory module, wherein the data module comprises the symbol map, wherein the symbol map is configured to map one or more start codons to respective human readable symbols that possess a disproportionally low frequency of occurrence in the reference language, and wherein the symbol map is further configured to map one or more stop codons to respective human readable symbols that possess a disproportionally high frequency in the reference language.

[0040] Also provided herein is an apparatus for converting a sequence of human readable symbols of a set of human readable symbols of a reference language that conveys a non-genetic message into a sequence of codon identifiers, the apparatus comprising: a processor that executes a sequence of instructions; a memory module coupled to the processor and comprising instructions for determining a codon identifier for each human readable symbol contained within the sequence of human readable symbols, wherein each codon identifier is determined upon reading a symbol map; and a data module coupled to the memory module, wherein the data module comprises the symbol map, the symbol map maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and the symbol map further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

[0041] In another aspect, provided herein is a computer-readable medium for use in an encoding machine, the computer-readable medium comprising instructions which, when executed by the encoding machine, perform a process comprising: receiving a sequence of human readable symbols that conveys a non-genetic message; and generating a codon identifier for each human readable symbol contained within the sequence, wherein the human readable symbol generated is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a reference language than one or more human readable symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the reference language than one or more human readable symbols from the set of human readable symbols.

[0042] Also provided herein is a computer-readable medium for use in an encoding machine, the computer-readable medium comprising instructions which, when executed by the encoding machine, perform a process comprising: generating a codon identifier for each human readable symbol in a set of human readable symbols that conveys a non-genetic message, wherein the human readable symbol generated is based upon a mapping function that maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and that further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

[0043] In another aspect, provided herein is a method of generating a sequence of codon identifiers from a sequence of human readable symbols that conveys a non-genetic message, the method comprising: receiving the sequence of human readable symbols at a memory module; loading a symbol map within the memory module, wherein the symbol map is configured to determine a codon identifier that maps to each human readable symbol within the sequence, wherein the symbol map is further configured to map a human readable symbol with a frequency of occurrence that is less than a first predetermined threshold within a reference language to a start codon, and wherein the symbol map is further configured to map a human readable symbol with a frequency of occurrence that is greater than a second predetermined threshold within the reference language to a stop codon; and outputting a sequence of codon identifiers corresponding to each human readable symbol within the sequence.

[0044] Also provided herein is a method of generating a sequence of codon identifiers from a sequence of human readable symbols of a set of human readable symbols of a
reference language that conveys a non-genetic message, the method comprising: identifying the sequence of human readable symbols at a memory module; and using a processor to read a symbol mapping for each human readable symbol in the sequence and determine a codon identifier which maps to the respective human readable symbol; wherein the symbol mapping maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and that further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

Various other aspects and embodiments will become more apparent with reference to the accompanying figures and detailed description provided below.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a functional sequence diagram illustrating an exemplary process of transcribing an input human readable symbol sequence and an encoded nucleic acid sequence.

FIG. 2 is a block diagram of an exemplary transcoder configured to encode an input human readable symbol sequence into a codon sequence with a low probability of biological impact.

FIG. 3 is a screen capture of an exemplary human readable symbol map which may be used to generate an encoded nucleic acid sequence with a low probability of biological impact.

FIG. 4 is a flow diagram of an exemplary method of creating a human readable symbol map which may be used to generate a nucleic acid sequence with a low probability of biological impact.

FIG. 5 is a flow diagram of an exemplary method of encoding an input human readable symbol sequence into a codon sequence with a low probability of biological impact.

FIG. 6 is a flow diagram of an exemplary method of decoding a nucleic acid sequence with a low probability of biological impact into a human readable symbol sequence.

FIG. 7 is a flow diagram of an exemplary method of encoding a watermark into a synthetic nucleic acid sequence with a low probability of biological impact.

FIGS. 8A-8E provide exemplary codon tables based on the English language. FIG. 8A depicts an exemplary code in the “Alt” format; FIG. 8B depicts an exemplary code in the “Ctrl” format; FIG. 8C depicts an exemplary code in the “Default” format; FIG. 8D depicts an exemplary code in the “Shift” format; and FIG. 8E depicts an exemplary code.

DETAILED DESCRIPTION

The present application provides a system for encoding basic text into a synthetic nucleic acid sequence of codon identifiers and, furthermore, for decoding text therefrom.

Previous attempts to create such a system have utilized standard amino acid encoding codon tables, which results in unwanted biological effects from the nucleic acid encoded text. The present system described herein is specifically designed to ensure that the encoded text does not correspond to the codons used by or otherwise biologically active in a host organism. One embodiment encodes all letters in the American English alphabet, as well as all 10 numerals, mathematical symbols, typographical characters and common punctuation marks. The codon usage schemes described herein are designed for use in a variety of host organisms, and can be specifically tailored for optimization in a particular host. In the following description, reference is made to the accompanying figures in which it is shown by way of illustration specific embodiments can be practiced. It is to be understood that other embodiments can be used and structural changes can be made without departing from the scope of the present application. Elements of the embodiments described herein can be combined to make additional embodiments not specifically described that are also within the scope of the invention. Headings within the application are solely for the convenience of the reader, and do not limit in any way the scope of the invention or its embodiments.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention relates. The following terms are defined for purposes of the invention as described herein.

As used herein, the terms “application”, “computer program”, “program”, and “software” include without limitation any sequence of human or machine recognizable steps that are adapted to be processed by a computer. Such may be rendered in any programming language or environment including, without limitation, C/C++, Fortran, COBOL, PASCAL, Perl, Prolog, Python, MATLAB, assembly language, scripting languages, markup languages (e.g., HTML, SGML, XML, VoXML), functional languages (e.g., API, Erlang, Haskell, Lisp, ML, F# and Scheme), as well as object-oriented environments such as the Common Object Request Broker Architecture (CORBA) and Java™ (including J2ME, JavaBeans, etc.).

As used herein, the term “display” includes any type of device or medium adapted to display information, including without limitation cathode ray tube displays (CRTs), liquid crystal displays (LCDs), thin film transistor displays (TFTs), digital light processor displays (DLPs), plasma displays, light emitting diodes (LEDs) or diode arrays, incandescent devices, and fluorescent devices. Display devices also include less dynamic devices such as printers, e-book devices, and other similar structures.

As used herein, the terms “local” and “remote” refer generally to devices, entities, or users that are serviced by separate sets of processes. These terms are intended to be relative, and bear no absolute reference or connection to the physical location of the executed processes of the served device, entities, or users.

As used herein, the term “memory” includes any type of integrated circuit or other storage device adapted for storing digital data including, without limitation, ROM, PROM, EEPROM, DRAM, SDRAM, DDR/2 SDRAM, EDO/FPM, RDRAM, FRAM, “flash” memory (e.g., NAND/NOR), and PSRAM.

As used herein, the term “module” refers to any type of software, firmware, hardware, or combination thereof that is designed to perform a desired function.

As used herein, the terms “processor,” “microprocessor,” and “digital processor” include all types of digital processing devices including, without limitation, digital signal processors (DSPs), reduced instruction set computers
(RISC), general-purpose (CISC) processors, microprocessors, gate arrays (e.g., FPGAs), programmable logic devices (PLDs), reconfigurable compute fabrics (RCFs), array processors, and application-specific integrated circuits (ASICs). Such processors may be contained on a single unitary IC die or distributed across multiple components.

As used herein, in the context of introducing nucleic acids into cells or organisms, the terms “introducing,” “transfection,” “transformation,” or “transduction,” refer to the introduction of one or more exogenous nucleic acid sequences or polynucleotides into a host cell or organism by using one or more physical or chemical methods as are known in the art. Many transfection techniques are known to those of ordinary skill in the art including, but not limited to, calcium phosphate DNA co-precipitation (see Methods in Molecular Biology, Vol. 7, Gene Transfer and Expression Protocols, Ed. E. J. Murray, Humana Press (1991)); DEAE-dextran; electroporation; cationic liposome-mediated transfection; tungsten particle-facilitated microparticle bombardment (Johnston: S. A., Nature 346: 776-777 (1990)); and streptomycin phosphate DNA co-precipitation (Brash D. E. et al., Molec. Cell. Biol. 7: 2031-2034 (1987)).

As used herein in the general context of transforming a sequence, set or signal (such as a sequence of nucleic acid residues or codon identifiers), the term “transform” refers simply to changing or converting a first sequence, set, or signal into a second sequence, set, or signal.

As used herein, “DNA isolation or extraction” refers to any procedure used to collect DNA from a sample for subsequent analysis. For example, there are three basic steps and one optional step in a DNA extraction: (i) breaking the cells open, typically referred to as cell disruption or cell lysis, to expose the DNA within (commonly achieved by physically grinding or sonicating the sample, or chemically treating the sample); (ii) removing membrane lipids by adding a detergent; (iii) removing proteins by adding a protease (optional); and (iv) precipitating the DNA with an alcohol (usually ice-cold ethanol or isopropanol). Since DNA is insoluble in these alcohols, it will aggregate together, resulting in a pellet upon centrifugation; this step also removes alcohol-soluble salt. Refinements of the technique include adding a chelating agent to sequester divalent cations such as Mg²⁺ and Ca²⁺; this stops DNase enzymes from degrading the DNA. Cellular and histone proteins bound to the DNA can be removed either by adding a protease or by having precipitated the proteins with sodium or ammonium acetate, or extracted them with a phenol-chloroform mixture prior to the DNA-precipitation. If desired, the DNA can be redissolved in a slightly alkaline buffer or in ultra-pure water.

As used herein, “RNA isolation or extraction” refers to any procedure used to collect RNA from a sample for subsequent analysis. Several methods can be used to isolate RNA from samples; the most common of these is guanidinium thioxyanate phenol-chloroform extraction.

As used herein, the term “DNA sequencing” refers to any sequencing method for determining the order of the nucleotide bases (adenine, guanine, cytosine, and thymine) in a molecule of DNA. Methods include, but are not limited to, Maxam-Gilbert sequencing, chain termination methods, dye-terminator sequencing, automated DNA-sequencing, in vitro cloning amplification, parallelized sequencing by synthesis, sequencing by ligation, microfluidic Sanger sequencing and sequencing by hybridization.

As used herein, “oligonucleotide synthesis” refers to the chemical synthesis of relatively short fragments of nucleic acids or codon identifiers with defined chemical structure (sequence). The technique is useful in because it provides a rapid and inexpensive access to custom-made oligonucleotides of a desired sequence. Whereas enzymes synthesize DNA and RNA in a 5’ to 3’ direction, chemical oligonucleotide synthesis is carried out in the opposite, 3’ to 5’ direction. Currently, the process is implemented as solid-phase synthesis using phosphoramidite method and A, C, G, T (2’-deoxy only), and U (ribo only) nucleoside phosphoramidites or 2’-deoxyxynucleoside phosphoramidites as building blocks. To obtain the desired oligonucleotide, the building blocks are sequentially coupled to the growing oligonucleotide chain in the order required by the sequence of the product.

As used herein, “nucleic acid synthesis” refers to the process of synthesizing an artificially designed sequence (e.g., a gene or a nucleic acid sequence which may contain a watermark) into a physical nucleic acid sequence.

The terms “cells”, “cell cultures”, “cell line”, “recombinant host cells”, “recipient cells” and “host cells” are often used interchangeably and will be clear from the context in which they are used. These terms include the primary subject cells and any progeny thereof, without regard to the number of transfers. It should be understood that not all progeny are exactly identical to the parental cell (due to deliberate or inadvertent mutations or differences in environment); however, such altered progeny are included in these terms, so long as the progeny retain the same functionality as that of the originally transformed cell. For example, though not limited to, such a characteristic might be the ability to include a non-genetic message, such as a watermark. The cell line can be any of those known in the art or described herein. A “clone” is a population of cells derived from a single cell or common ancestor by mitosis.

As used herein, the term “nucleotide” refers to a monomeric unit of a polynucleotide that consists of a heterocyclic base, a sugar, and one or more phosphate groups. The naturally occurring bases, (guanine, (G), adenine, (A), cytosine, (C), thymine, (T), and uracil (U)) are derivatives of purine or pyrimidine, though it should be understood that naturally and non-naturally occurring base analogs are also included. The naturally occurring sugar is the pentose (five-carbon sugar) deoxyribose (which forms DNA) or ribose (which forms RNA), though it should be understood that naturally and non-naturally occurring sugar analogs are also included. Nucleic acids are linked via phosphate bonds to form nucleic acids, or polynucleotides, though many other linkages are known in the art (such as, though not limited to phosphorothioates, boronophosphates and the like).

As used herein, the terms “nucleic acid”, “nucleotide” and “polynucleotide” refer to a polymeric form of nucleotides of any length, either ribonucleotides (RNA) or deoxyribonucleotides (DNA). These terms refer to the primary structure of the molecules and, thus, include double- and single-stranded DNA, and double- and single-stranded RNA. These terms include, as equivalents, natural or synthetic nucleic acids, analogs of either DNA or RNA made from nucleotide analogs and modified polynucleotides such as, though not limited to, methylated and/or capped polynucleotides. Nucleic acid sequences can be referred to as having a 5’ end and a 3’ end as is known in the art, which can be used as reference points for other sequences, for example,
as being the 5' (also upstream) or 3' (also downstream) to a codon identifier in a sequence.

[0074] As used herein, a “DNA molecule” refers to the polymeric form of deoxyribonucleotides (adenine, guanine, thymine, and cytosine) in its either single stranded form or a double-stranded helix. This term refers only to the primary and secondary structure of the molecule, and does not limit it to any particular tertiary forms. Thus, this term includes double-stranded DNA, found, inter alia, in linear DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes. In discussing the structure of particular double-stranded DNA molecules, sequences can be described herein according to the normal convention of giving only the sequence in the 5' to 3' direction along the non-transcribed strand of DNA (i.e., the strand having a sequence homologous to the mRNA). An “RNA molecule” refers to the polymeric form of ribonucleotides (adenine, guanine, uracil, and cytosine), which is typically, but not always, single-stranded.

[0075] Although embodiments may be described and illustrated in terms of deoxyribonucleic acid (DNA) sequences and corresponding nucleotide bases, it should be understood that the embodiments are not so limited, but are additionally applicable to other types of nucleic acids and nucleotide bases (including, for example, ribonucleic acid (RNA), such as messenger ribonucleic acid (mRNA)). Furthermore, although embodiments may be described and illustrated herein in terms of a single transcriptor configured to both encode and decode an input human readable symbol sequence, it should be understood that the associated encoding and decoding logic may be separated and/or distributed among multiple systems, devices, and/or computer networks.

[0076] As used herein, a nucleic acid “coding sequence” or “coding region” is a region of a nucleic acid sequence which can be transcribed and/or translated into a polypeptide when placed under the control of appropriate expression control sequences and in the presence of appropriate cellular machinery or enzymes. In other words, a coding sequence provides one type of genetic message to the cell containing the sequence. The boundaries of the coding sequence (the “open reading frame” or “ORF”) are determined by a start codon at the 5' terminus (encoding the amino terminus of a peptide or polypeptide) and a translation stop codon at the 3' terminus (encoding the carboxyl terminus of a peptide or polypeptide). For example, in most instances ATG and AUG denote sequences of DNA and RNA respectively that are the start codon or initiation codon encoding the amino acid methionine (Met) in eukaryotes and a modified Met (fMet) in prokaryotes, although alternate start codons, mainly UUG and UUG, may be used in prokaryotes. In the standard genetic code, there are three stop codons: UAG (in RNA)/TAG (in DNA) (“amber”), UAA/TAA (“ochre”), and UGA/TGA (“opal” or “umber”); although several variations to this most common set are known. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and synthetic DNA sequences. A polyadenylation signal and transcription termination sequence is, usually, located 3' to the coding sequence. As used herein, the term “non-coding sequence” or “non-coding region” refers to regions of a nucleic acid sequence that are not transcribed and/or translated into amino acids (e.g., un-translated regions, signal sequences, etc.).

[0077] As used herein term “reading frame” refers to one of the six possible reading frames, three in each direction (5' and 3'), of the nucleic acid molecule. The reading frame that is used determines which codons are used to encode amino acids within the coding sequence of a DNA molecule. When decoding sequences in methods and apparatus described herein, all three reading frames in the 5' direction are typically used to ensure detection of any non-genetic message encoded in a nucleic acid sequence. As used herein, an all-six or all-six reading frame stop codon containing sequence refers to a sequence that will mandatorily cause termination of transcription and translation proceeding in either the 5' or 3' direction, in any of the three respective reading frames (e.g., TTAAGTAGCTAA; SEQ ID NO: 1).

[0078] Using the exemplary sequence the three 5' reading frames would be:

[0079] TTA ACT AGC TAA (SEQ ID NO: 1), with the stop codon in the fourth triplet;
[0080] TAA CTA GCT AA- (SEQ ID NO: 2), with the stop codon in the first triplet; and
[0081] AAC TAG CTA AA-- (SEQ ID NO: 3), with the stop codon in the second triplet.

[0082] Using the same exemplary sequence the three 3' reading frames (anti-sense or complementary strand) would be:

[0083] TTA GCT AGT TAA (SEQ ID NO: 4), with the stop codon in the fourth triplet;
[0084] TAG CTA GCT AA-- (SEQ ID NO: 5), with the stop codon in the first triplet; and
[0085] ACG TAG TTA AA-- (SEQ ID NO: 6), with the stop codon in the second triplet.

[0086] As used herein, an “antisense” nucleic acid molecule comprises a nucleic acid sequence which is complementary to a “sense” nucleic acid encoding a protein, e.g., complementary to the coding strand of a double-stranded DNA molecule, complementary to an mRNA sequence or complementary to the coding strand of a gene. Accordingly, an antisense nucleic acid molecule can be hydrogen-bonded to a sense nucleic acid molecule.

[0087] As used herein, a “codon” refers to the three nucleotides which, when transcribed and translated, encode a single amino acid residue; or in the case of UUA, UGA or UAG encode a termination signal. As used herein, a “wobble position” refers to the third position of a codon. Codons of the standard genetic code encoding amino acids are well known in the art and are provided for convenience herein in Table 1.

TABLE 1

<table>
<thead>
<tr>
<th>Codon Table</th>
<th>Amino Acid</th>
<th>Amino Acid Abbr.</th>
<th>Codon</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>A</td>
<td>GCA</td>
<td></td>
</tr>
<tr>
<td>Cysteine</td>
<td>A</td>
<td>GCC</td>
<td></td>
</tr>
<tr>
<td>Aspartate</td>
<td>A</td>
<td>GCA</td>
<td></td>
</tr>
<tr>
<td>Glutamate</td>
<td>D</td>
<td>GAC</td>
<td></td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>F</td>
<td>UUC</td>
<td></td>
</tr>
<tr>
<td>Glycine</td>
<td>G</td>
<td>GGA</td>
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<tr>
<td></td>
<td>G</td>
<td>GGC</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G</td>
<td>GGG</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G</td>
<td>GUU</td>
<td></td>
</tr>
</tbody>
</table>

Nov. 3, 2011
from the Wisconsin Sequence Analysis Package, Version 8.1, Genetics Computer Group, Madison, Wis. Codon usage is also described in, for example, R. Nussinov, "Eukaryotic Dinucleotide Preference Rules and Their Implications for Degenerate Codon Usage," J. Mol. Biol. 149: 125-131 (1981). The codons which are most frequently used in highly expressed human genes are presumptively the optimal codons for expression in human host cells and, thus, form the bases for constructing a synthetic coding sequence. In alternate species, codon usage may vary (also known as codon bias), and sequences may be codon optimized to reflect such differences for use of sequences in different organisms. A useful source of information can be found on the internet at the www.URL.kazusa.or.jp/codon/ in a Codon Usage Database.

[0090] As used herein, a “codon identifier” refers to nucleotides that code for a single human readable symbol of a reference language, preferably a triplet or three nucleotides. A set or sequence of codon identifiers preferably do not correspond to the sequence of a naturally-occurring gene or other biologically active sequence. Rather, the one or more codon identifiers correspond to one or more letters, one or more numbers, one or more spaces, one or more punctuation marks (e.g., !, -, ;, :, ], {, }}, {, }, ?) and mathematical symbols ((), +, –, ×, ÷, S, %, etc.), one or more typographical characters (e.g., @, ©, ™, ®, §, etc.), one or more new lines, or a combination of any thereof and are made up of three nucleotides. Exemplary codon identifiers are provided in more detail below and in the accompanying figures. When combined into a synthetic nucleic acid sequence, a series of codon identifiers conveys a non-genetic message.

[0091] The following Table 2 illustrates exemplary human readable symbols recognizable in the English language. Any other characters or symbols could readily be incorporated as desired by the user.

<table>
<thead>
<tr>
<th>TABLE 2 Exemplary human readable symbols</th>
</tr>
</thead>
<tbody>
<tr>
<td>Punctuation</td>
</tr>
<tr>
<td>Math: symbols</td>
</tr>
<tr>
<td>Typographical characters</td>
</tr>
<tr>
<td>Examples</td>
</tr>
<tr>
<td>STOP STOP</td>
</tr>
</tbody>
</table>

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Abbreviation. It should be understood that the codons specified above are for RNA sequences. The corresponding codons for DNA have a T substituted for U. Each codon corresponds to an amino acid which can be abbreviated into a single letter of the alphabet. In preferred embodiments, three nucleotide codon identifiers do not correspond to these same single letters in the symbol mapping, such that any natural language information encoded as codon identifiers is extremely unlikely to correspond to a nucleic acid sequence with biological function. As such, the encoded sequence is unlikely to be lethal to a cell or organism comprising the sequence, or subject to genetic selection in a cellular context, or to correspond to a sequence that would come into existence naturally.

Optimal codon usage is indicated by codon usage frequencies for expressed genes, for example, as shown in the codon usage chart from the program "Human-High.cod"...
containing messages or watermarks can have lengths up to about 40 Kb, up to about 35 Kb, up to about 30 Kb, up to about 25 Kb, up to about 20 Kb, up to about 15 Kb, up to about 10 Kb, up to about 5 Kb, up to about 3 Kb, up to about 2 Kb, up to about 1 Kb, up to about 0.5 Kb, up to about 0.1 Kb, or any value therebetween. The length of the sequence generally does not exceed the length of a gene, genome, plasmid, or chromosome into which it is inserted. Insertion can be placement within a gene, genome, plasmid, or chromosome, or replacement of all or a portion thereof.

[0093] As used herein, a “reference language” refers to any language on the planet including, but not limited to, Afrikaans, Albanian, Arabic, Aranes (Occitan), Armenian, Basque, Cantonese Chinese, Catalan, Chipewyan, Cree, Croatian, Cyrryllic, Czech, Danish, Dutch, English, Faroese, Farsi, Finish, French, German, Galician, Gwich’in, Greek, Hebrew, Hindi, Hungarian, Korean, Icelandic, Inuinnaqtun, Inuktitut, Inuvialuktun, Italian, Japanese, Kalaallisut, Mandarin, Mazandaran, Norwegian, Persian, Polish, Portuguese, Punjabi, Romanian, Russian, Rusyn, Sami, Sanskrit, North and South Slavey, Slovone, Spanish, Swahili, Swedish, Tibetan, Tswana, Turkish, Ukrainian, Urdu, Uyghur, Uzbek, Venda, Vietnamese, Welsh, Xhosa, Yiddish, Zhuang, and Zulu.

[0094] As used herein, “isolated” (used interchangeably with “substantially pure”) in the context of an isolated biomolecule such as an isolated protein or nucleic acid, is a biomolecule removed from the context in which the biomolecule exists in nature. For example, an isolated protein or nucleic acid molecule is removed from the cell or organism with which it is associated in its natural state. An isolated biomolecule can be, in some instances, partially or substantially purified, for example, an isolated nucleic acid molecule can be a nucleic acid sequence that has been excised from the chromosome, genome, or episome that it is integrated into in nature.

Encoder

[0095] Provided herein are means for encoding a sequence of human readable symbols of a reference language that conveys a non-genetic message into one or more codon identifiers. Such means include, for example, an apparatus, systems, and a computer-readable medium for generating a sequence of codon identifiers from a reference language.

[0096] Provided herein is an apparatus for converting a sequence of human readable symbols of a reference language that conveys a non-genetic message into a sequence of codon identifiers, the apparatus comprising: a processor configured to execute instructions; a memory module coupled to the processor and comprising instructions which, when executed by the processor, determine a codon identifier for each human readable symbol contained within the sequence of human readable symbols, wherein each codon identifier is determined upon reading a symbol map; and a data module coupled to the memory module, wherein the data module comprises the symbol map, wherein the symbol map is configured to map one or more start codons to respective human readable symbols that possess a disproportionately low frequency of occurrence in the reference language, and wherein the symbol map is further configured to map one or more stop codons to respective human readable symbols that possess a disproportionately high frequency in the reference language.

[0097] The goal is to generate a nucleic acid sequence that is not genetically viable, and thus does not have a biological impact on a recombinant or synthetic cell, or on a recombinant or synthetic virus, comprising the sequence. To that end, the nucleic acid sequence should contain frequent occurrences of stop codons, and few occurrences of start codons. As an example, a start codon may be mapped to a character in the English language that is rarely used, such as *; such that a start codon would rarely be mapped into the synthetic nucleic acid sequence; the reverse complement of the start codon can be assigned to the rare "$"; and a stop codon may be mapped to a character in the English language that is most commonly used, such as the letter E, A or T, such that a stop codon would be frequently be mapped into the synthetic nucleic acid sequence. The reverse complement of two of the stop codons are the common characters “R” and “H”. These measures ensure that a watermark can be transcribed in either direction and any potential open reading frame will be short in the +0 and +0 reading frames. The code can be designed such that common two-character combinations such as “CH” ensure that the -1, -2, +1 and +2 reading frames do not tend to avoid stop codons. Common and uncommon characters can be distributed evenly across the chart to help guard against low-complexity sequences being added by patterns into a watermark text. A disproportionally low frequency of occurrence in the reference language typically refers to a symbol that has a frequency of distribution of less than one percent in the set of human readable symbols. A disproportionally high frequency of occurrence in the reference language typically refers to a symbol that has a frequency of distribution of more than five percent in the set of human readable symbols. For example, in the symbol map shown in FIG. 3, the character * would have disproportionally low frequency of occurrence in conventional text patterns in the English language, and the alphabetical characters E, A and T would have a disproportionally high frequency of occurrence in conventional text patterns in the English language.

[0098] Provided herein is an apparatus for converting a sequence of human readable symbols of a set of human readable symbols of a reference language that conveys a non-genetic message into a sequence of codon identifiers, the apparatus comprising: a processor that executes a sequence of instructions; a memory module coupled to the processor and comprising instructions for determining a codon identifier for each human readable symbol contained within the sequence of human readable symbols, wherein each codon identifier is determined upon reading a symbol map; and a data module coupled to the memory module, wherein the data module comprises the symbol map, the symbol map maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and the symbol map further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

[0099] Provided herein is a computer-readable medium for use in an encoding machine, the computer-readable medium comprising instructions which, when executed by the encoding machine, perform a process comprising: receiving a sequence of human readable symbols that conveys a non-genetic message; and generating a codon identifier for each human readable symbol contained within the sequence, wherein the human readable symbol generated is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a reference language than one or more human readable
symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the reference language than one or more human readable symbols from the set of human readable symbols.

[0100] Provided herein is a computer-readable medium for use in an encoding machine, the computer-readable medium comprising instructions which, when executed by the encoding machine, perform a process comprising: generating a codon identifier for each human readable symbol in a set of human readable symbols that conveys a non-genetic message, wherein the human readable symbol generated is based upon a mapping function that maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and that further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

[0101] Provided herein is a method of generating a sequence of codon identifiers from a sequence of human readable symbols, the method comprising: receiving the sequence of human readable symbols that conveys a non-genetic message at a memory module; loading a symbol map within the memory module, wherein the symbol map is configured to determine a codon identifier that maps to each human readable symbol within the sequence, wherein the symbol map is further configured to map a human readable symbol with a frequency of occurrence that is less than a first predetermined threshold within a reference language to a start codon, and wherein the symbol map is further configured to map a human readable symbol with a frequency of occurrence that is greater than a second predetermined threshold within the reference language to a stop codon; and outputting a sequence of codon identifiers corresponding to each human readable symbol within the sequence.

[0102] Provided herein is a method of generating a sequence of codon identifiers from a sequence of human readable symbols of a set of human readable symbols of a reference language that conveys a non-genetic message, the method comprising: identifying the sequence of human readable symbols at a memory module; and using a processor to read a symbol mapping for each human readable symbol in the sequence and determine a codon identifier which maps to the respective human readable symbol; wherein the symbol mapping maps a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and that further maps a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon.

Decoder

[0103] Provided herein are means for decoding a sequence of one or more codon identifiers into one or more human readable symbols of a reference language that conveys a non-genetic message. Such means include, for example, an apparatus, systems, and a computer-readable medium. When decoding a nucleic acid sequence comprising of one or more codon identifiers, it will initially be unknown from the source sequence which 5' reading frame may contain the non-genetic message or watermark, and therefore, all three 5' reading frames must be analyzed.

[0104] Provided herein is an apparatus for transforming a sequence of codon identifiers into a sequence of human readable symbols that conveys a non-genetic message, the apparatus comprising: a processor adapted to execute instructions; and a storage module, wherein the storage module comprises a data structure for mapping codon identifiers into human readable symbols, and a set of instructions which, when executed by the processor, generate a human readable symbol for each codon identifier read from a sequence of codon identifiers, wherein the human readable symbol generated is based at least in part upon the data structure; wherein the data structure is configured to map a start codon to a human readable symbol with a frequency of occurrence within a reference language that is less than a first predetermined threshold, and wherein the data structure is further configured to map a plurality of stop codons to human readable symbols with frequencies of occurrence within the reference language that are greater than a second predetermined threshold.

[0105] In another embodiment, the data structure maps a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols, and further maps a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols.

[0106] In one embodiment, the data structure does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code.

[0107] In another embodiment, the sequence of codon identifiers comprises at least one of an all-6 reading frame start codon containing sequence 5' to a first codon identifier in the sequence, and an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

[0108] Provided herein is a computer-readable medium for use in a decoding machine, the computer-readable medium comprising instructions which, when executed by the decoding machine, perform a process comprising: identifying a sequence of codon identifiers; and generating a human readable symbol for each codon identifier in the sequence; wherein the human readable symbol generated is based at least in part upon a mapping function configured to map a start codon to a human readable symbol that has a frequency of occurrence within a reference language that is smaller than every other human readable symbol of a first set of human readable symbols, and wherein the mapping function is further configured to map a stop codon to a human readable symbol that has a frequency of occurrence within the reference language that is larger than every other human readable symbol of the first set of human readable symbols.

[0109] Provided herein is a computer-readable medium for use in a decoding machine, the computer-readable medium comprising instructions which, when executed by the decoding machine, perform a process comprising: identifying a sequence of codon identifiers; and generating a human readable symbol for each codon identifier in the sequence of codon identifiers, wherein the human readable symbol generated is based upon a mapping function that maps codon identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionally low frequency in the language of the watermark, and that maps codon identifiers
corresponding to stop codon(s) to human readable symbols that possess a disproportionally high frequency in the language of the watermark.

[0110] In one embodiment, the mapping function does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code.

[0111] In another embodiment, the sequence of codon identifiers comprises at least one of an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence, and an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

[0112] Provided herein is a method of transforming a first signal adapted to indicate a sequence of codon identifiers into a second signal adapted to indicate a sequence of human readable symbols that conveys a non-genetic message, the method comprising: receiving the first signal; determining a human readable symbol for each codon identifier in the sequence, wherein said determining is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a human readable symbol sequence than one or more human readable symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the human readable symbol sequence than one or more symbols from the set of human readable symbols; and transforming the first signal into the second signal based upon the one or more determined human readable symbols.

[0113] Provided herein is a method of transforming a first signal comprising a sequence of codon identifiers into a second signal to indicate a sequence of human readable symbols of a set of human readable symbols of a reference language that conveys a non-genetic message, the method comprising: identifying the first signal that indicates the sequence of codon identifiers; determining a human readable symbol for each codon identifier in the sequence, wherein said determining a human readable symbol is based at least in part upon a mapping function that maps a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols, and further maps a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols; and transforming the first signal into the second signal, wherein the second signal indicates the sequence of human readable symbols.

[0114] In one embodiment, the mapping function does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code.

[0115] In another embodiment, the sequence of codon identifiers comprises at least one of an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence, and an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

Exemplary Embodiments of Encoding and Decoding

[0116] FIG. 1 is a functional sequence diagram illustrating an exemplary high-level process of translating an input symbol sequence into an encoded nucleic acid sequence that can be stored within any genetic material of an organism, such as that contained in a chromosome or genome in one or more cells of a sample, living organism and the like. The genetic material (e.g., DNA) of the cell/organism can be subsequently harvested or extracted using standard techniques which are well known in the art, so that the encoded nucleotide sequence can later be determined. The encoded nucleotide sequence may then be parsed and decoded in order to generate the original symbol sequence. Although the following description provides computer-assisted encoding and/or decoding processes, any of the methods described herein may be performed manually.

[0117] At block 102, a symbol sequence 104 is provided. The symbol sequence 104 may comprise any number of discrete representations or symbols, including alphabetic and non-standard symbols, ASCII or ANSI symbols, control symbols, and/or other types of metadata.

[0118] A wide variety of possible symbol sequences 104 may be utilized. For example, a symbol sequence 104 may include the names of people or organizations, trademarks and/or copyright notices, serial numbers, text messages, times and/or dates, tags and other indicators, classified information, data, digital computer instructions, graphics, video, information intended to be operated upon by a DNA based computer, etc. Myriad other types of content may also be contained within a symbol sequence 104 and are contemplated herein.

[0119] The symbol sequence 104 may also be provided in any number of ways. In some embodiments, for example, the symbol sequence 104 may be provided by a connected keyboard, touchpad, mouse, microphone, or other input peripheral. In other embodiments, the symbol sequence 104 may be read from one or more files or data streams. These files or data streams may be accessed on a local system (for example, on a local hard disk or other non-volatile memory source), a remote system (for example, on a networked system or server accessible over the Internet), or a removable media device (for example, a floppy disk, external hard drive, flash drive, smart card, or other serial bus device).

[0120] At block 106, an encoded sequence 108 of nucleotides is generated based upon the provided symbol sequence 104. A symbol map 220 (for example, as shown in FIG. 3) may be used to translate each symbol in the symbol sequence 106 into a sequence known as a codon identifier, preferably a tri-nucleotide. In some embodiments, the symbol map 220 may be stored locally (for example, within a lookup table, database, or other reference structure resident within a local memory module). In other embodiments, the symbol map 220 may be stored within the memory of one or more remote systems.

[0121] At block 110, a synthetic nucleic acid sequence 112 may then be created from the encoded sequence 108 specified. Conventional techniques in molecular biology and DNA synthesis may be used, for example, to create a synthetic nucleic acid sequence 112 which contains the same codon identifier ordering as the encoded sequence 108.

[0122] At block 114, the synthetic nucleic acid sequence 112 may then be introduced into a cell or living organism using standard techniques. The synthetic nucleic acid sequence 112 may be directly or indirectly introduced into the organism. Once the synthetic nucleic acid sequence 112 is introduced into the cell or organism, cells can then harbor the
synthetic nucleic acid sequence 112, effectively serving as a memory source for the encoded sequence 108.

[0123] In order to recover the symbol sequence 104 from the cell or organism (e.g., a virus or a multicellular organism), nucleic acid can be or cells can be extracted (as shown at block 116). Note that a variety of conventional extraction techniques may be used to extract the genetic material from a recombinant cell, a synthetic cell, or a recombinant or synthetic organism. The extracted cells 118 may then be analyzed at block 120 in order to recover the originally encoded sequence 108.

[0124] Once the encoded sequence 108 has been recovered, this sequence 108 may then be parsed and decoded at block 122. A symbol map 220 (for example, as that shown in FIG. 3) may be used to decode/translate each codon identifier 302 into a corresponding symbol 304 of the original symbol sequence 106. In this manner, all human readable symbols from the original symbol sequence 106 may be reproduced.

[0125] The symbol sequence 106 may then be output at block 124 in any number of ways. In some embodiments, for example, the symbol sequence 104 may be output directly on one or more output devices. Any device capable of writing or displaying data may be used for such purposes, including, for example, display devices (e.g., monitors), printers, projectors, televisions, speakers, networked devices (e.g., computers, digital cameras, personal data assistants, memory devices, etc.) and/or other output peripherals. In some embodiments, the symbol sequence 104 is configured to be written to one or more files that may be stored within a local memory source. In yet other embodiments, the symbol sequence 104 may be output by human writing on implements such as paper.

[0126] FIG. 2 is a block diagram of an exemplary transcoder 200 configured to encode an input symbol sequence into a codon sequence such that the codon sequence has substantially no biological impact on a host organism if introduced into the organism as a synthetic nucleic acid sequence (e.g., as free DNA). The transcoder 200 may also be configured to decode an input sequence of codons and thereby yield the originally input symbol sequence. Thus, the exemplary transcoder 200 depicted in FIG. 2 may be used to both encode a human readable symbol sequence into a codon sequence, and to decode a codon sequence into a human readable symbol sequence.

[0127] The power supply 202 provides a source of power to modules disposed within the transcoder 200. In some embodiments, power is supplied externally by one or more conductive wires, for example, by a power or serial bus cable. In other embodiments, a battery may be used as a source of power. In other embodiments, a human brain is used as a source of power.

[0128] One or more processors 204 are adapted to execute sequences of instructions by loading data from and storing data to a local memory module (for example, volatile memory 206, which may be implemented as any combination of static and/or dynamic RAM). Possible instructions may include, without limitation, instructions for data conversions, formatting operations, arithmetic operations, communication instructions, and/or storage and retrieval operations.

[0129] One or more I/O modules 216 may be used to interface a set of I/O peripherals with various programs, processes, or applications executing within the volatile memory 206 of the transcoder 200. In some embodiments, the I/O modules 216 may consist of one or more device drivers adapted to interface a set of hardware devices with an operating system associated with the transcoder 200. Note that the I/O modules 216 may be implemented as any combination of software, firmware, or hardware according to embodiments described herein.

[0130] A wide variety of input peripherals may be used to generate input 212 to the transcoder 200 according to embodiments disclosed herein. These input peripherals include, without limitation, keyboards, mice, trackballs, touch panels, microphones, controllers (e.g., joysticks), scanners, digital cameras, pencils, pens, markers, crayons, and communicative interfaces for networked devices (e.g., network or serial bus interfaces).

[0131] Similarly, a wide variety of output peripherals may be used to write and/or display output 214 according to various embodiments disclosed herein. These output peripherals include, without limitation, display devices (e.g., monitors), printers, projectors, televisions, speakers, local memory modules, pens, pens, markers, crayons, and networked devices (e.g., computers, digital cameras, personal data assistants, remote memory devices, high-speed serial bus devices, etc.).

[0132] A non-volatile memory module 208 may be used to persistently store data, instructions, process states, memory tables, and other information within the transcoder 200. The non-volatile memory module 208 may be implemented as any type or combination of memory adapted for persistent storage, including, without limitation, conventional hard disks, ROM (e.g., PROM, EPROM, EEPROM), flash memory, paper, etc. Note also that in some embodiments, all or a portion of the non-volatile memory module 208 may serve as virtual memory for the volatile memory module 206.

[0133] In some embodiments, the non-volatile memory 208 may include a symbol frequency analyzer 230 for determining how frequently certain symbols appear within one or more input symbol streams. For example, the symbol frequency analyzer 230 may be used to determine that the symbol “v” has a frequency of occurrence of approximately 1% within a specified symbol stream, while the symbol “e” has a frequency of occurrence of approximately 13% within the same stream. Note that an exemplary process of implementing the symbol frequency analyzer 230 has been described in more detail below (see FIG. 4 and accompanying text).

[0134] In some embodiments, the non-volatile memory 208 may include one or more symbol maps 220 which may be used to construct synthetic nucleic acid sequences with low probabilities of biological impact. An exemplary symbol map 220 has been provided with reference to FIG. 3. As shown by this figure, each symbol 304 from a domain of possible symbols (including letters, numbers, punctuation marks, symbols, and control symbols) uniquely maps to a single codon identifier 302. The symbol maps 220 may thus be used to translate a sequence of human readable symbols into a sequence of codons, or to translate a sequence of codons into a sequence of human readable symbols.

[0135] Note that while the symbol map 220 depicted in FIG. 3 illustrates a one-to-one mapping of sixty-four possible symbols 304 to sixty-four possible codons identifiers 302, the symbol map 220 depicted in FIG. 3 is merely exemplary in nature, and has been included herein so as to illustrate the broader principles of the application. It is to be understood that embodiments disclosed herein encompass a wide variety of possible mappings. Moreover, the domain of possible sym-
bols 304 and the range of possible codon identifiers 302 can also be smaller than or larger than sixty-four.

In some embodiments, for example, a reduced symbol domain may be utilized in order to further decrease the probability that a constructed synthetic nucleic acid sequence will have a detectable biologically impact an organism. This may be implemented, for example, by excluding from the symbol map 220 those permutations of nucleotides which could potentially be interpreted as a start codon by an organism’s internal biological processes (ATG, GTA, *AT, TG*, etc.).

In other embodiments, the symbol domain may be expanded in order to support a larger number of encodable symbols (for example, upper-case and lower-case symbols, non-standard symbols, etc.). This may be implemented, for example, by mapping each symbol 304 into a set of multiple codons instead of a single codon (for example, “A”=CAGCCG).

Returning now to FIG. 2, the non-volatile memory 208 may also include a transcoder module 210 for translating a sequence of symbols 304 into a sequence of codon identifiers 302 and/or for translating a sequence of codon identifiers 302 into a sequence of symbols. In some embodiments, the transcoder module 210 may utilize one or more symbol maps 220 as an input argument, value, or parameter. In other embodiments, the transcoder module 210 may contain internal logic supplying one or more encoding schemes (e.g., switch and/or case logic). Note that an exemplary process of encoding a sequence of symbols 304 into a sequence of codon identifiers 302 has been provided below with reference to FIG. 5, while an exemplary process of decoding a sequence of codon identifiers 302 into a sequence of symbols 304 has been provided below with reference to FIG. 6.

FIG. 4 is a flow diagram of an exemplary method of creating a symbol map which may be used to generate a nucleic acid sequence with a substantially reduced or a low probability of biological impact.

At block 402, an analysis of symbol frequency in a reference language or symbol stream is generated. This may be accomplished, for example, by parsing one or more input streams in order to determine the number of occurrences of a certain symbol relative to the total number of symbols analyzed. In some embodiments, a counter may be assigned to each unique symbol that is discovered within the one or more input streams. In other embodiments, counters may be assigned to only those symbols that are elements of an input symbol domain. Note that lower-case and upper-case equivalents may be treated as the same or separate symbols.

At block 404, the least frequently occurring symbol within the symbol domain may be determined. This may be accomplished, for example, by conventional sort routines (for example, bubble sort, insertion sort, selection sort, quick sort, etc.). In some embodiments, the least frequently occurring symbol is the asterisk symbol “*”. Note, however, that the least frequently occurring symbol may depend upon the one or more input streams analyzed and/or the symbol domain selected.

At block 406, the least frequently occurring symbol may be mapped to a specific permutation of nucleotides known as the start codon (i.e., ATG). A start codon is commonly used by an organism’s internal processes to indicate the beginning of a coding sequence. Mapping the least frequently occurring symbol to the start codon in this manner substantially reduces the number of start codons that will appear within the encoded sequence, thereby reducing the likelihood that an cell’s internal processes will interpret a portion of the encoded sequence as a genetic instruction.

At block 408, the three most frequently occurring symbols within the symbol domain may be determined. This may be implemented, for example, by reading the three most frequently occurring symbols within a symbol list that has been sorted by occurrence frequency (for example, by reading the sorted list generated at block 404). In some embodiments, the three most frequently occurring symbols are the letters “E”, “A”, and “I” (where the frequencies of upper-case and lower-case equivalents have been aggregated). As in the prior case, the three most frequently occurring symbols may depend upon the one or more input streams analyzed and/or the symbol domain selected.

At block 410, each of the three most frequently occurring symbols may then be mapped to a respective stop codon (i.e., TAA, TAG, and TGA). Mapping the most frequently occurring symbols to stop codons in this manner increases the likelihood that a stop instruction will appear within a given sequence of nucleotides, thereby substantially reducing the likelihood that an cell’s internal processes will interpret a portion of the encoded sequence as a genetic instruction.

Each unmapped symbol from the symbol domain may then be mapped to a codon identifier that has not yet been mapped to a human readable symbol. Optionally, logic may also be provided that is configured to prevent an unmapped symbol from mapping to a codon that generates an amino acid which has a single-letter abbreviation that is equivalent to the unmapped symbol. For example, if the next unmapped symbol were the letter “V”, the codons “GTT”, “GTC”, “GTA”, and “GTG” may be excluded from the range of possible candidates that could map to “V”, since each of these codons may ultimately yield the amino acid valine (commonly abbreviated as “V”). Exemplary logic for accomplishing this functionality is discussed below with reference to blocks 412-418.

At block 412, a decision may be made as to whether an unmapped human readable symbol presently exists. If there are no unmapped symbols remaining, the process can end. Otherwise, the next unmapped symbol may be retrieved and the process continued at block 414.

The next available codon identifier may then be determined at block 414. This may be implemented by retrieving the next codon identifier from a reference data structure (e.g., array, list, heap, stack, queue, etc.).

A decision may then be made at block 416 as to whether the next available codon identifier yields an amino acid with an abbreviation that is equivalent to the unmapped symbol. In some embodiments, both upper and lower case equivalents are considered in this decision. If the codon identifier does in fact yield an amino acid with an abbreviation that is equivalent to the unmapped symbol, a new codon identifier may be received at block 414, and the process may repeat until a suitable codon identifier is discovered. Otherwise, if the codon identifier does not yield an amino acid with an abbreviation that is equivalent to the unmapped symbol, the unmapped symbol may be mapped to the selected codon identifier at block 418, and the process repeated per block 412.

Note that in cases where each of the remaining unmapped codon identifiers yield an amino acid with an abbreviation that is equivalent to the unmapped symbol, conventional processes of backtracking may be utilized in order
to unmap and reassign previous symbols to alternate codon identifiers. In other embodiments, symbols with amino acid abbreviation equivalents may be mapped to codon identifiers before any other symbols are mapped, thereby obviating the need for backtracking logic.

At block 502, a decision is made as to whether an un-encoded symbol still exists within the input symbol sequence. If all human readable symbols from the input symbol sequence have been encoded, the process can end according to some embodiments. Alternatively, the process may further comprise inserting an all-6 reading frame stop codon at the beginning and/or end of the encoded sequence. This is shown at blocks 508 and 510, respectively. These all-6 reading frame stop codons may thus serve to designate the beginning and/or end of an encoded message, thereby allowing a message to be more easily detected among a large stream of successive nucleotides.

In some embodiments, the all-6 reading frame stop codons may be used to further decrease the probability that a cell’s internal processes or a virus will interpret a portion of the encoded sequence as a genetic instruction. In some embodiments, for example, one or more all-6 reading frame stop codons may be interleaved within the encoded message at periodic intervals, thereby ensuring that a stop codon occurs every “n” reading frames. A transcoder adapted to decode such a message may simply ignore these codons during the decoding process.

If an unencoded symbol still exists within the input sequence, at block 504, a decision may be made as to whether the input symbol is supportable (i.e., whether it exists within the domain of encodable symbols). In some embodiments, an error message may be generated when it is determined that a certain symbol cannot be encoded (e.g., as shown at block 512). In other embodiments, any non-encodable symbols may simply be ignored. In still other embodiments, a special symbol may be used to indicate that an unencodable symbol has been identified. This special symbol can thus serve as a replacement for every unencodable symbol encountered within the input symbol sequence.

At block 506, the codon identifier which corresponds to the input symbol can then be generated. According to some embodiments, the codon identifier which corresponds to the input symbol can be determined by consulting a symbol map that is stored within a local memory source (e.g., the symbol maps 220 depicted in FIG. 2 and FIG. 3). The process may then repeat at block 502 until all input symbols have finally been encoded.

FIG. 6 is a flow diagram of an exemplary method of decoding a nucleic acid sequence with a low probability of biological impact into a symbol sequence.

At block 602, a decision is made as to whether any additional codon identifiers exist within an input sequence of codon identifiers. If no codon identifiers exist, the process may then end. Otherwise, the process may continue per block 604. This process would be repeated for all three 5’ reading frames.

The symbol corresponding to the codon identifier may then be determined at block 604. According to some embodiments, the symbol which corresponds to the codon identifier may be determined by consulting a symbol map that is stored within a local memory source (e.g., the symbol maps 220 depicted in FIG. 2 and FIG. 3).

The determined symbol may then be output at block 606. In some embodiments, the symbol may be written to, or displayed upon, a connected output peripheral (for example, a display device, printer, television screen, paper, etc.). In other embodiments, the symbol may be written to a local memory source and stored within one or more files. In some embodiments, for example, as that shown in FIG. 6), symbols may be output as soon as they are determined. In other embodiments, the output may be generated after all symbols have been determined.

FIG. 7 is a flow diagram of an exemplary method of encoding a watermark into a synthetic nucleic acid sequence with a low probability of biological impact. Each symbol from a reference language can be assigned a codon identifier. A watermark containing a series of symbols from a reference language can be generated. Each symbol in the watermark can be substituted with a codon identifier and the resulting encoded sequence be appended with an all-6 reading frame stop codon containing sequence 508 and appended with an all-6 reading frame stop codon containing sequence 510 to create a synthetic nucleic acid sequence containing the encoded watermark message. Thus, within the context of a series of codon identifiers that are used to encode a message that exists between all-an all-6 reading frame stop codon containing sequence 5′ to a first codon identifier in the sequence and an all-6 reading frame stop codon containing sequence 3′ to the last codon identifier in the sequence, it would be possible to map symbols to codon identifiers which may represent a single letter abbreviation of an amino acid. In this context, it would be understood that the all-6 reading frame stop codon containing sequences would prevent an encoded message containing an amino acid abbreviation from being read as genetic material. Thus, a more static code that would typically not change from implementation to implementation can be created. Such a code can then become a standard such as a file format.

FIGS. 8A-B illustrate an exemplary embodiment contemplated for use with the system described above.

Synthetic Nucleic Acid Sequences

Messages can be input as synthetic nucleic acid fragments into a cell or virus and incorporated into a gene, a genome, a plasmid, or a chromosome, or any other genetic material in a cell. Incorporated nucleic acids are made up of codon identifiers that represent a series of human readable symbols from a human reference language. The sequence of codon identifiers creates a non-genetic message or watermark that can be used to identify or authenticate any cell or virus containing that message.

A synthetic nucleic acid sequence can further comprise an all-6 reading frame stop codon containing sequence 5′ (prior to) to a first codon identifier in the sequence, an all-6 reading frame stop codon containing sequence 3′ (subsequent to) to the last codon identifier in the sequence, or both.

Provided herein is a synthetic nucleic acid sequence, wherein said synthetic nucleic acid sequence comprises one or more codon identifiers corresponding to a set of human readable symbols of a reference language that conveys a non-genetic message, and further wherein said synthetic nucleic acid sequence is not genetically viable and does not have a biological impact upon a recombinant or synthetic cell,
or upon a recombinant or synthetic virus, comprising the synthetic nucleic acid sequence.

[0164] In one embodiment, a synthetic nucleic acid sequence cannot be biologically translated into a functional amino acid sequence by the recombinant or synthetic cell/virus.

[0165] The one or more codon identifiers do not correspond to sequence of a gene or other biologically active sequence. Rather, the one or more codon identifiers correspond to one or more letters, one or more numbers, one or more spaces, one or more punctuation marks (e.g., “,” “;” “{” “}” “(” “)” “:” “?” and “”), one or more mathematical symbols (e.g., “+”, “*”, “=”, “$”, “%”, etc.), one or more typographical characters (e.g., “@”, “©”, “™”, “®”, “€”, etc.), one or more new lines, or a combination of any thereof.

[0166] In one aspect, the set of human readable symbols comprises a watermark. Watermarks include, but are not limited to, a copyright notice, a trademark, a company identifier, a name, a phrase, a sentence, a quotation, genetic information, unique identifying information, data, digital computer instructions, texts, graphics, video, information intended to be operated upon by a DNA-based computer, or a combination of any thereof.

[0167] The synthetic nucleic acid sequence can further comprise an all-6 reading frame stop codon comprising sequence 5′ to a first codon identifier in the sequence, an all-6 reading frame stop codon comprising sequence 3′ to the last codon identifier in the sequence, or both.

[0168] One can empirically determine the size of a watermark on the size of one or more of the following: a gene, a genome, a plasmid, an artificial chromosome, a cell, or an organism. Message lengths can be up to about 40 Kb, up to about 35 Kb, up to about 30 Kb, up to about 25 Kb, up to about 20 Kb, up to about 15 Kb, up to about 10 Kb, up to about 5 Kb, up to about 3 Kb, up to about 2 Kb, up to about 1 Kb, up to about 0.5 Kb, up to about 0.1 Kb, or any value there between. In one embodiment, a message length can be up to about 5 Kb. In another embodiment, a message length can be up to about 2 Kb. A message length, typically, does not exceed the length of a gene, genome, plasmid, or chromosome into which it is to be incorporated or replace.

Recombinant and Synthetic Cells, Viruses, Organisms and Animals

[0169] One would understand that synthetic nucleic acid sequences conveying non-genetic messages can be used in any type of cell. In some instances, cells can be present in a population of cells (e.g., a cell culture, an embryo, a multicellular organism, a plant, an animal, etc.).

[0170] Provided herein is a recombinant or synthetic cell containing a synthetic nucleic acid sequence described herein. A recombinant or synthetic cell can be a prokaryotic cell, a eukaryotic cell, or an archaeal cell. Also provided herein is a recombinant or synthetic virus, multicellular organism, or animal containing a synthetic nucleic acid sequence described herein. A set of human readable symbols can be a watermark that allows the authentication or identification of said recombinant or synthetic cell, virus, organism or animal; or identification of an organism comprising recombinant or synthetic cells or viruses.

[0171] The recombinant or synthetic cells described herein are useful for tracking of cells or organisms for research and/or commercial use. Cells also include, but are not limited to cells and organisms in a research laboratory. Such cells and organisms may be distributed internally within a company or institute, or distributed externally as part of a collaboration or material transfer agreement. Other cells are described below with respect to samples which can be assessed for cells or organisms containing a watermark.

[0172] Provided herein is a recombinant animal that contains a synthetic nucleic acid sequence conveying a non-genetic message or watermark. Recombinant animals include, for example, transgenic rodents (e.g., mice, rats, ferrets, rabbits, etc.), horses (e.g., pure-bred, hybrid breed or thoroughbred), cows, bulls, dogs, cats, sheep, primates (e.g., gorillas, chimpanzees, monkeys, orangutans, etc.), fish (e.g., zebra fish or exotic fish), amphibians (e.g., frogs), insects, etc. Such watermarks can be used to identify, for example, the source of the animal, or identification of a specific genetic modification.

[0173] Provided herein is a recombinant embryo that contains a watermark; such watermarks can be used to track and identify embryos based on encoded information.

[0174] Also provided herein is a recombinant plant that contains a watermark; such watermarks can be used to track and identify plants based on encoded information.

[0175] A recombinant or synthetic cell can be a prokaryotic cell, a eukaryotic cell, or an archaeal cell.

[0176] A prokaryotic cell can be, for example, a bacterial cell that is Gram-positive or Gram-negative.

[0177] A eukaryotic cell can be, for example, a yeast cell, a fungal cell, an algal cell, an animal cell, or a plant cell.

[0178] Prokaryotic Cells

[0179] A prokaryotic cell can be, for example, a bacterial cell that is Gram-positive or Gram-negative, or may lack a cell wall. A synthetic nucleic acid sequence can be incorporated in a genome, a plasmid, or an artificial chromosome of any Gram-negative or Gram-positive bacterium.

[0180] Gram-negative bacterium include, but are not limited to Enterobacteriaceae spp. (e.g., E. coli, E. cloaca, E. intermedia, etc.), Hemophilus spp. (e.g., H. influenzae, etc.), Vibrionales spp. (e.g., V. cholera, etc.), Pseudomonadaceae spp. (e.g., Pseudomonas aeruginosa, Pseudomonas fluorescens, Pseudomonas putida, Pseudomonas stutzeri, etc.), Helicobacter spp. (e.g., H. pylori, etc.), Synechocystis spp., Acinetobacter baumannii, Acidovorax delfieldii, Aeromonas veroni, Aquaspirillum spp., Bordetella bronchiseptica, Flavobacterium odoratum, Cytophagaceae, Citrobacter braakii, Citrobacter freundii, Comamonas (Delfia) acidovorans, Burkholderia cepacia, Yersinia kristensenii, Stenotrophomonas spp., Serratia spp. (e.g., Serratia liquefaciens, Serratia marcescens, etc.), Salmonella spp. (e.g., Salmonella typhimurium, etc.), Ralstonia spp. (e.g., Ralstonia ettiophaga, Ralstonia pickettii, etc.), Proteus vulgaris, Providencia rettgeri, Pseudomonas spp., Pantoea ananatis, Paracoccus marucsi, Ochrobactrum anthropi, Morganella morganii, Neisseria spp. (e.g., Neisseria meningitides, etc.), Klebsiella spp. (e.g., Klebsiella oxytoca, Klebsiella pneumoniae, etc.) and Hydrogenophaga paloroni. One would understand that other genus and species of gram negative bacteria are included herein.

[0181] Gram-positive bacteria include, but are not limited to Streptococcus spp. (e.g., S. pneumoniae, S. sanguis, etc.), Enterococci spp. (e.g., E. faecalis, etc.), Bacteroides spp. and Clostridia spp. (e.g., C. sporogenes, etc.), Mycobacterium spp. (e.g., M. tuberculosis, M. avium, etc.), Corynebacterium spp. (e.g., C. renale, etc.), Peptostreptococcus spp., Listeria spp. (e.g., L. monocytogenes, etc.), Legionella spp.,
Alicyclobacillus acidocaldarius, Bacillus spp. (e.g., Bacillus licheniformis, Bacillus pumilus, Bacillus sphaericus, Bacillus subtilis, Bacillus thuringiensis, Bacillus cereus, Bacillus circulans, Bacillus diposorum, etc.), Brevibacillus chochinensis, Brevibacterium brevis, Deinococcus radiodurans, Staphylococcus spp. (e.g., Staphylococcus aureus, Staphylococcus capitis, Staphylococcus epidermidis, etc.), Rhodococcus equi, Propionibacterium acnes, Paenibacillus spp. (e.g., Paenibacillus glaucanolyticus, Paenibacillus polymyxa, etc.), Kocuria rosea, Microbacterium saperdae, Micrococcus species, Kocuria spp. (e.g., Kocuria kristinae, Kocuria rhizophila, etc.), and Geobacillus stercorarius. One would understand that other genus and species of gram positive bacteria are included herein.

[0182] Bacteria lacking a defined cell wall include, but are not limited to, Mycoplasma spp. (e.g., M. capricolum, M. gallisepticum, M. genitalium, M. hominis, M. kyupneumoniae, M. laboratorium, M. mycoides, M. ovipneumoniae, M. pneumoniae, etc.).

[0183] In some embodiments, photobacteria, including for example, green sulfur bacteria, purple sulfur bacteria, green nonsulfur bacteria, purple nonsulfur bacteria, or cyanobacteria may be used. Cyanobacterial species that can be used include, without limitation, Agmenellum, Anaabaena, Anaibaenopsis, Anacystis, Aphaniizonema, Anoplosira, Asterocarpa, Boreia, Calothrix, Chamaesiphon, Chlorogloeoopsis, Chroococcidiopsis, Chroococcus, Cricaulium, Cyanobacterium, Cyanobium, Cyanoaepria, Cyanothece, Cylindrospermopsis, Cylindromonas, Dactylolycoecopsis, Dermocarpa, Fischereilla, Freymaella, Geitleria, Geitleriina, Gloeobacter, Gloeocapsa, Gloeothecae, Halospirulina, Iyengariella, Leptolyngbya, Limnothrix, Lyngbya, Microcoleus, Microcystis, Microsorina, Nodularia, Nostoc, Nostocopsis, Oscillatoria, Phormidium, Plankothrix, Pleurocapsa, Prochlorococcus, Prochloron, Proclorothea, Pseudanabaena, Rivularia, Schizothrix, Scytonea, Spirulina, Stanieria, Starria, Stigonema, Symplocia, Synecococcus, Synecocystis, Tolypothrix, Trichodesmium, Tychothea, or Xenococcus species.

[0184] Each of the aforementioned prokaryotic cells and others known in the art are contemplated for use herein.

[0185] Archaia

[0186] Archaia are a group of single-celled microorganisms. They have no cell nucleus or any other organelles within their cells. Archaia include, but are not limited to cells of the phyla: Crenarchaeota, Euryarchaeota, Korarchaeota, Nanoarchaeota and Thaumarchaeota. Each of the aforementioned archaia and others known in the art are contemplated for use herein.

[0187] Viruses

[0188] Viruses are typically classified into the following groups: I: dsDNA viruses (e.g., Adenoviruses, Herpesviruses, Papovaviruses); II: ssDNA viruses (+)sense DNA (e.g., Parvoviruses); III: dsRNA viruses (e.g., Reoviruses); IV: (+)ssRNA viruses (+)sense RNA (e.g., Picornaviruses, Togaviruses); V: (--)ssRNA viruses (--)sense RNA (e.g., Orthomyxoviruses, Rhabdoviruses); VI: ssRNA-RT viruses (+)sense RNA with DNA intermediate in life-cycle (e.g., Retroviruses); and VII: dsDNA-RT viruses (e.g., Hepadnaviruses).

[0189] Each of the aforementioned viruses and others known in the art are contemplated for use herein.

[0189] Eukaryotic Cells

[0190] A eukaryotic cell contemplated herein can be any cell with a nucleus enclosed within a cell membrane, for example, a yeast cell, a fungal cell, an algal cell, an animal cell, or a plant cell.

[0192] Yeast

[0193] Yeast are unicellular microorganisms that belong to one of three classes: Ascomycetes, Basidiomycetes and Fungi Imperfecti. Pathogenic yeast strains and nonpathogenic yeast strains are considered herein.

[0194] Genera of yeast strains include, but are not limited to, Saccharomyces, Candida, Cryptococcus, Hansenula, Kluyveromyces, Pichia, Rhodotorula, Schizosaccharomyces and Yarrowia.

[0195] Non-limiting representative species of yeast strains include Saccharomyces cerevisiae, Saccharomyces carlsbergensis, Candida albicans, Candida kefyr, Candida tropicalis, Candida guillermondii, Candida parapsilosis, Cryptococcus laurentii, Cryptococcus neoformans, Cryptococcus humicola, Hansenula anomala, Hansenula polymorpha, Kluyveromyces fragilis, Kluyveromyces lactis, Kluyveromyces marxianus var. lactis, Pichia pastoris, Rhodotorula rubra, Rhodotorula glutinis, Schizosaccharomyces pombe and Yarrowia lipolytica. It is understood that a number of these species include a variety of subtypes, species, subtypes, etc. that are meant to be included within the aforementioned species.

[0196] Each of the aforementioned yeast genera and species and others known in the art are contemplated for use herein.

[0197] Algae

[0198] A synthetic nucleic acid sequence can be incorporated in a genome, a plasmid, or an artificial chromosome of any algal species.

[0199] Algae that can be used in the methods of the invention can be any algae, and can include microalgae, such as but not limited to, Achaeae, Amphipora, Amphora, Anistrosdesmus, Asterozemona, Boekelovia, Botryococcus, Breitococcus, Chaetoceros, Carteria, Chlamydomona, Chlorococcus, Chlororugium, Chlorella, Chroonomas, Chrysophyta, Cricophyta, Cryptocordium, Cryptomonas, Cyclotella, Dunaliella, Ellipsoïden, Elmiaria, Eremosphaera, Erodospora, Euglena, Fritschea, Fragilaria, Gloeothamnion, Haematococcus, Halocoleber, Hymenomonas, Isochrysis, Lepocinclis, Micrastrum, Monoraphidium, Nannochloris, Nannochloropsis, Navicula, Neochloris, Nephrochloris, Nostocariina, Nitzschia, Ochromonas, Oedogonium, Oocystis, Osteococcus, Pavlova, Parachlorocella, Pascheria, Phaeocytococcus, Phausis, Platymonas, Pleurochrysis, Pleurococcus, Prototheca, Pseudochlorella, Pyramimonas, Pyrophyto, Scenedesmus, Schizochytrium, Skeletonema, Spyrogyra, Stichococcus, Tetraselmis, Thraustochytrium, Thalassiosira, Virdiella, or Volvox species.

[0200] Each of the aforementioned algae and others known in the art are contemplated for use herein.

[0201] Plant Cells

[0202] Plant cells that can be used include those obtained from organisms such as trees, herbs, bushes, grasses, vines, ferns, and mosses. Diversity of living plant divisions includes non-vascular land plants or bryophytes, such as Marchantia (liverworts), Anthocerotophyta (hornworts), Bryophyta (mosses) and Hepaticophyta; and vascular plants or tracheophytes, such as Rhyniophyta, Zosterophyllyophyta, Lycophydiophyta (club mosses), Trimerophyta, Pteridophyta (ferns, whisk ferns & horsetails), Pterynophytophyta, and Seed plants or spermatophytes, such as Pteridophyta (ferns), Pinophyta (conifers),
Cycadophyta (cycads), Ginkgophyta (ginkgo), Gnetophyta (gnetophytes), and Magnoliophyta (flowering plants).

Each of the aforementioned plant cells and others known in the art are contemplated for use herein.

Animal Cells

Animal cells that can be used include, but are not limited to vertebrates, such as fish, amphibians, reptiles, birds and mammals (e.g., rodents, primates, sheep, horses, cows, pigs, dogs, cats, etc.); arthropods, such as insects (e.g., Drosophila melanogaster); and nematodes (e.g., Caenorhabditis elegans).

Each of the aforementioned animal cells and others known in the art are contemplated for use herein.

Fungi

Fungi that can be used include any of the phyla Microsporidia, Chytridiomycota, Blastocladiomycota, Neo-calciinastigomycteyta, Glomeromycota, Ascomycota, and Basidiomycota. Exemplary genera of fungi to be used in the compositions and methods described herein include, for example, Pullularia, Chaetomium, Aspergillus, Coniophora, Pseudocercosporella, Helminthosporium, Pyrenophora, Septoria, Helminthosporium, Fusarium, Rhizoctonia, Cercospora, Peronospora, Erysiphe, and Pestalozzia.

Species of fungi contemplated for use in the compositions and methods described herein include, for example, Pseudocercosporella herpotrichoides, Helminthosporium gramineum, Pyrenophora avenae, Septoria nodorum, Helminthosporium teres, Fusarium roseum, Fusarium nivale, Fusarium culmorum, Rhizoctonia cerealis, Pullularia pullulans, Chaetomium globosum, Coniophora puteana, Cercospora beticola, Peronospora tabacina, Erysiphe cichoracearum, Pyrenophora avenae, Whetzelinia soriotiorum, Monilia laxa, Mycosphaerella fijiensis, Marssonina panattoniana, Alternaria solani, Aspergillus niger, Cladosporium herbarum, Penicillium expansum, Phialophora cinereascens, Phoma betae, Phoma foveata, Phoma lingam, Verticillium dahliae, Ascochyta pisii, Guignardia bidwellii, Corticium rolfsii, Phomopsis viticola, Sclerotinia sclerotiorum, Sclerotinia minor, Phytophthora cinnamomi, Phytophthora cactorum, Phytophthora capsici, Phytophthora parasitica, Phytophthora megasperma, Phytophthora syringae, Coryneum cardinale, Septoria tritici, Botrytis cinerea, Fusarium oxysporum, Fusarium melonis, Rhizoctonia solani, and Helminthosporium gramineum.

Each of the aforementioned fungi and others known in the art are contemplated for use herein.

Methods of Creating a Recombinant or Synthetic Cell or Virus

Provided herein is a method of creating a recombinant or synthetic cell or virus comprising a watermark, comprising: (i) generating a nucleic acid sequence comprising one or more codon identifiers from a set of human readable symbols of a reference language comprising said watermark that conveys a non-genetic message, wherein a symbol mapping is configured to map a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and wherein the symbol mapping is further configured to map a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon; (ii) synthesizing said nucleic acid sequence; and (iii) introducing said nucleic acid sequence into a recombinant or synthetic cell or virus, thereby creating said recombinant or synthetic cell or virus comprising a watermark.

Alternatively, provided herein is a method of creating a recombinant or synthetic cell or virus comprising a watermark, comprising: (i) generating a nucleic acid sequence comprising one or more codon identifiers from a set of human readable symbols of a reference language comprising said watermark that conveys a non-genetic message, wherein a symbol mapping is configured to map a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols to a start codon, and wherein the symbol mapping is further configured to map a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols to a stop codon; (ii) synthesizing said nucleic acid sequence; and (iii) introducing said nucleic acid sequence into a recombinant or synthetic cell or virus, thereby creating said recombinant or synthetic cell or virus comprising a watermark.

In one embodiment, the symbol mapping does not map a three nucleotide codon identifier to a single letter representation of an amino acid residue normally assigned to that three nucleotide codon in the standard genetic code.

In another embodiment, the generating step (i) is computer-assisted and comprises identifying the set of human readable symbols at a memory module and for each human readable symbol in the set, using a processor to read a symbol mapping for determining a codon identifier which maps to the respective human readable symbol.

Methods of Use

Current technologies allow the generation of synthetic nucleic acid molecules and/or the ability to alter the nucleic acid sequences of existing nucleic acid molecules. With a careful coding scheme and arrangement, it is possible to encode important information as a synthetic nucleic acid sequence and store it in a living host safely and permanently. This technology can be used to identify origins of a host containing a watermark and protect research and development investments. It can also be used in environmental research to track generations of organisms and observe the ecological impacts of pollutants. Today, there are microorganisms that can survive under extreme conditions. As well, it is advantageous to consider multicellular organisms as hosts for stored information. These living organisms can provide as memory housing and protection for stored data or information. The present invention provides well for data storage in a living organism wherein at least one non-genetic message or watermark is encoded to represent information and incorporated into a living cell or organism.
One aspect provided herein is the storage of a non-genetic message in multicellular living organisms (e.g., rodents, primates, sheep, horses, cows, pigs, dogs, cats, etc.). This can be achieved by incorporating at least one nucleic acid sequence encoded to represent a non-genetic message into a germ cell; a precursor cell that gives rise to gametes that will then serve as specialized haploid cells (sperm or egg) in sexual reproduction, or stem cell; a relatively undifferentiated cell that will continue dividing indefinitely, throwing off (pro-ducing) daughter cells that will undergo terminal differentiation into particular cell types. The encoded nucleic acid sequence will then propagate into a multicellular living organism. This embodiment of the invention is a memory storage system that takes advantage of multicellular organisms (e.g., insect, rodent, etc.) and serves to propagate the encoded nucleic acid sequence in all daughter cells stemming from the original host stem cell.

Also provided herein is a memory storage system wherein a living organism comprises therein at least one nucleic acid sequence encoded to represent a non-genetic message. The stored non-genetic message resides in a living organism and remains there until recovery is desired. The non-genetic message is then retrieved and decoded so as to enable communication. Like a computer memory device that can store data and programs, the same or similar items can be contained in a nucleic acid memory system.

Non-genetic messages or watermarks described herein can be used to trace or monitor distribution of a particular cell or organism in situ, ex situ, in vitro, in vivo, or a combination thereof.

Provided herein is a method of determining the presence of a recombinant or synthetic organism comprising a reference watermark in an sample that conveys a non-genetic message, said method comprising: (i) sequencing nucleic acid material obtained from one or more organisms in said environmental sample; (ii) transforming the nucleic acid sequence obtained in step (i) to a set of codon identifiers, wherein each codon identifier of said set of codon identifiers consists of three nucleotides of said sequence in all three reading frames; (iii) determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determination is based at least in part upon a symbol mapping that is configured to map a start codon to a human readable symbol with a frequency of distribution of less than one percent in the set of human readable symbols and is further configured to map a stop codon to a human readable symbol with a frequency of distribution of more than five percent in the set of human readable symbols; and (iv) comparing the human readable symbol sequence of all three reading frames to the reference watermark in said recombinant or synthetic organism, whereby the presence of the reference watermark in any reading frame of the nucleic acid material obtained in step (i) indicates the presence of the recombinant or synthetic organism in the environmental sample.

A sample can be any sample that can contain a recombinant or synthetic cell such as, for example, an environmental sample, a sample deposited with a cell depository (e.g., American Type Culture Collection (ATCC) or another international depository, a laboratory sample, a food supplements, a commercial sample, recombinantly engineered crops and seeds, etc.

Environmental samples include, but are not limited to water samples, soil samples, crops, oil deposits or spills, coal deposits, mineral deposits, algae cells used for biofuel production, recombinantly engineered crops and seeds, crops that have come in contact with recombinantly engineered crops (e.g., corn, grapes, etc.) by virtue of reproduction, samples containing cells or organisms useful for improving soil quality, fungi used to enhance plant growth, etc.

Samples include, but are not limited to, cells and organisms in a research laboratory. Such samples may be distributed internally within a company, university or institute, or distributed externally as part of a collaboration or material transfer agreement.

Samples also include nutritional supplements such as, for example, algae stocks used in nutritional supplements, desiccated algae, bacteria for use in digestive supplements and/or yogurt cultures, animal food and animal supplements.

Samples also include bacterial and viral vaccines such as, for example, live vaccines, modified vaccines, inactivated vaccines, etc.

Also contemplated herein are samples containing organisms for making industrial enzymes. Such enzymes are useful, for example, in commercial detergents (e.g., dish soap, laundry detergent, etc.).

Samples also include recombinantly engineered animals (e.g., rodents, primates, sheep, horses, cows, bulls, pigs, dogs, cats, etc.). Provided herein is a method for genetic tagging of a non-human organism by introducing into the organism a nucleic acid molecule containing a watermark that can be decoded using the methods described herein.

Methods of Monitoring and Tracking

In one aspect, provided herein is a method of monitoring the source, ownership, or changes in a sample over time. The sample being monitored can be sample containing one or more of: a prokaryotic cell, a virus, an archaeal cell or a eukaryotic cell. Monitoring can determine whether the state of a sample has been changed over time. For example, a recombinantly engineered crop can be monitored to determine cells containing a modification are spread through the
environment via natural means or are transported illegally. Monitoring can be accomplished by any of the methods provided herein.

In another aspect, provided herein is a method of tracking a sample that is transported through either natural or artificial means.

Although embodiments of this application have been described with reference to the accompanying drawings, it is to be noted that various changes and modifications will become apparent to those skilled in the art. Such changes and modifications are to be understood as being included within the scope of embodiments as defined by the appended claims.

Terms and phrases used in this document, and variations thereof, unless otherwise expressly stated, should be construed as open ended as opposed to limiting. As examples of the foregoing, the term “including” should be read as mean “including, without limitation” or the like; the term “example” is used to provide exemplary instances of the item in discussion, not an exhaustive or limiting list thereof; and adjectives such as “conventional,” “traditional,” “normal,” “standard,” “known” and terms of similar meaning should not be construed as limiting the item described to a given time period or to an item available as of a given time, but instead should be read to encompass conventional, traditional, normal, or standard technologies that may be available or known now or at any time in the future. Likewise, a group of items linked with the conjunction “and” should not be read as requiring that each and every one of those items be present in the grouping, but rather should be read as “and/or” unless expressly stated otherwise. Similarly, a group of items linked with the conjunction “or” should not be read as requiring mutual exclusivity among that group, but rather should also be read as “and/or” unless expressly stated otherwise. Furthermore, although items, elements or components of the disclosure may be described or claimed in the singular, the plural is contemplated to be within the scope thereof unless limitation to the singular is explicitly stated. The presence of broadening words and phrases such as “one or more,” “at least,” “but not limited to” or other like phrases in some instances shall not be read to mean that the narrower case is intended or required in instances where such broadening phrases may be absent.

EXAMPLES

Elements of the present application are illustrated by the following examples, which should not be construed as limiting in any way.

Example 1

Encoding Methods

FIGS. 3 and 8 identify codon identifiers and the respective symbols encoded therefrom. By virtue of the design of the non-genetic message or watermark, the encoded text does not correspond to the sequences of a gene or other biologically active sequence when in the form of a nucleic acid in the cell or organism. The examples provided in the Figures encode all letters in the American English alphabet as well as all 10 numerals and common punctuation marks.

While the present Figures and Examples are described with respect to the English language, one would comprehend that the coding scheme can be adapted to any reference language as described above.

An encoded non-genetic message in the nucleic acid sequence is flanked by the sequence 5’-TTAACTAGCTAA-3’ (SEQ ID NO: 1) on both the 5’ and 3’ sides of the watermark since that sequence contains a stop codon in all 6 reading frames.

To encode a non-genetic message or watermark, one can substitute in a serial, one-to-one manner, a given symbol of human readable text using one or more of: the Roman alphabet, Arabic numerals, and certain common punctuation and formatting symbols for strings of three nucleotides. These substitutions are performed such that each successive codon identifier (three nucleotide sequence) is added to the 3’ end of the nucleic acid sequence.

For example, the encoding of the text “JCVI-Strain 012.3 All Rights Reserved, 2009” into a nucleic acid sequence watermark can be performed either by hand or by computer program as follows:

First the watermark begins with the DNA sequence 5’-TTAACTAGCTAA-3’ (SEQ ID NO: 1). Next, the first human readable symbol of the text is “J”. According to the exemplary table above, the letter “J” (either upper or lower case) corresponds to the 3-nucleotide string 5’-GTT-3’ in the code. Therefore the next three nucleotides of the watermark are 5’-GTT-3’, which are added to the 3’ end of the preceding watermark sequence. At this stage the still incomplete watermark’s sequence is 5’-TTAACTAGCTAAAGT-3’ (SEQ ID NO: 7).

The next human readable symbol of the text is “C” which corresponds to the 3-nucleotide string 5’-TTT-3’ in the code. Therefore the next three nucleotides of the watermark are 5’-TTT-3’. At this stage the still incomplete watermark’s sequence is 5’-TTAACTAGCTAAGTTTTTTTTT-3’ (SEQ ID NO: 8).

In this manner, one serially adds the appropriate three nucleotide strings that correspond to the human readable symbols of the text to the 3’ end of the growing watermark sequence. Human readable symbols that are not covered in the code are skipped. When all the human readable symbols of the text have been skipped or encoded into the watermark, the sequence 5’-TTAACTAGCTAA-3’ (SEQ ID NO: 1) is added to its 3’ end.

Thus, the completely encoded watermark sequence for the text “JCVI-Strain 012.3 All Rights Reserved, 2009” is:

5’-TTAACTAGCTAATTTTTTTTTGTGCGCCTTGACTATAGCTGTGCGATACCTCTCTAC TCGAAATATATAGAACAACATACACTACGT TACATGAAGCTATACATAGCTTAA CTTTGG TAAATGTTGATAACTTCCTGTGAGCAT AAACACTAGCTAA-3’ (SEQ ID NO: 9).

Decoding Methods

To decode a watermark one performs the same process as encoding as described in Example 1, but in reverse.

One substitutes in a serial one-to-one manner, each successive three nucleotides of the watermark for their respective human readable symbols of human readable text. These substitutions (performed either by hand or by a computer program) are made such that each human readable symbol is placed to the right of the preceding symbol as one substitutes along the watermark in a 5’ to 3’ direction. This is
process of substitution is performed after the sequence 5'-TTAACTAGCTAA-3' (SEQ ID NO: 1) is removed from both ends of the watermark.

[0245] For example, to decode the sequence 5'-TTAACCTAGCTAACTTCGGCCGGCTGTAATAGCTTGTCATGAACTGCTTCTTACGGTAAACTAGCTAA-3' (SEQ ID NO: 9), the first step removes the sequence 5'-TTAACCTAGCTAA-3' (SEQ ID NO: 1) from both ends of the watermark leaving the following watermark: 5'-GTTTCTTGTGTCGCCGGCTGTCATGAACTGGCTTTACGGTAAACTAGCTAA-3' (SEQ ID NO: 10).

[0246] Next, the first three nucleotides of the remaining watermark sequence are 5'-GTT-3' which corresponds to the code in the table "J". Thus, the first letter of the decoded text is "J". The next three nucleotides of the remaining watermark sequence are 5'-TTT-3' which corresponds to the code in the letter "C". Thus the decoded text now reads "JCC".

[0247] In this manner, one serially substitutes the appropriate human readable symbols that correspond to the three nucleotide strings of the watermark to the right side of the growing decoded text. In this exemplary case, the final decoded watermark reads: "JCVI-STRAIN 012.3 ALL RIGHTS RESERVED, 2009."

[0248] When one is unsure of the original reading frame to decode the sequence, this would be performed in all three reading frames: thus beginning the sequence with GTC as the first codon identifier (1), then with TTC as the first codon identifier (2), and then with TTT as the first codon identifier (3) to see if any of these yield a sequence of human readable symbols as follows:

\[
\begin{align*}
\text{GTT} & \text{ TTT} \text{ TGT} \text{ CGG} \text{ CCT} \ldots \ (1) = \text{JCVI-S} \\
\text{TTC} & \text{ TTT} \text{ TGC} \text{ TGC} \text{ CGG} \text{ CCT} \ldots \ (2) = \text{JCVI} \\
\text{TTC} & \text{ TTT} \text{ GCT} \text{ GCC} \text{ TTC} \ldots \ (3) = \text{JCVI}
\end{align*}
\]

wherein reading frame (1) would emerge as the recognizable sequence of human readable symbols, i.e., the watermark.

[0249] The decoded sequence is presented in uppercase letters because the code, in its basic form, does not distinguish between upper and lower case letters, causing the information of which letters were originally capitalized to be lost in the encoding process. However, punctuation marks such as commas, periods, hyphens, and spaces are all retained through the encoding and decoding procedures.

[0250] The above example demonstrates one of the useful features of the DNA watermarks: if the above watermark were encoded twice in the same genomic molecule at locations a convenient distance apart from one another (e.g., 2 kilobases) and on opposing strands, a single primer PCR reaction may be used as a diagnostic strain-specific test to identify the molecule in question. In this exemplary case, the primer would have a sequence that would place the end of the primer inside the portion of the watermark that encoded the strain number.

Example 3

Synthetic Cells Containing Watermarks

[0251] A 1.08 Mbp Mycoplasma mycoides genome was chemically synthesized, and assembled in yeast as a centro-meric plasmid; the genome was isolated as naked DNA and transplanted into Mycoplasma capricolum to create a new bacterial cell controlled only by the synthetic genome.

[0252] Described in International Patent Application PCT/US10/35490 is the design, synthesis and assembly of the 1,077,947-bp Mycoplasma mycoides JCVI-syn1 genome from 1.078 1-kb synthetic DNA cassettes. The assembly was facilitated by in vitro and in vivo assembly methods. Cassette were assembled in sets of ten were assembled by yeast recombination and propagated in a yeast/E. coli shuttle vector. The 10-kb assemblies were recombined in sets of ten to produce 100-kb assemblies. The resulting eleven 100-kb assemblies were recombined in a single final step into the complete genome. A yeast clone bearing the synthetic genome was selected and confirmed by multiplex PCR and restriction analysis.

[0253] The assembled synthetic genome was propagated in yeast as a centromeric plasmid and successfully transplanted into restriction-minus Mycoplasma capricolum cells. The new cells have the phenotypic properties expected for M. mycoides and the designed synthetic DNA sequence, including watermark sequences and other designed gene deletions and polymorphisms. This strain is referred to as M. mycoides JCVI-syn1; this was the second bacterial chromosome synthesized and the first over one million bp. It is a synthetic bacterial genome successfully transplanted into a recipient cell resulting in new cells that are controlled only by a synthetic chromosome. The new synthetic chromosome cells are capable of continuous self-replication. This study confirmed the ability to start with digitized genetic information, synthesize new DNA and transplant that synthetic DNA into cells replacing all of the existing genetic information and, as a result, created new cells controlled only by that synthetic designed DNA. The existing (endogenous) genetic information was lost and a result new cells were created which were controlled only by the designed synthetic chromosome.

[0254] Synthetic Donor Genome Design

[0255] Design of the M. mycoides JCVI-syn1 genome was based on the highly accurate finished genome sequences of two previously described laboratory strains of M. mycoides subspecies capri GM12 (Benders et al., Nucleic Acids Res. (2010); Lartigue et al., Science 325, 1693 (2009)). One was the genome donor used by Lartigue et al. [GenBank accession CP001621] (Lartigue et al., Science 317, 632 (2007)). The other was a strain created by transplantation of a genome that had been cloned and engineered in yeast, YCPMmyc 1.1- Δtopf[5]res, [GenBank accession CP001668] (Lartigue et al., Science 325, 1693 (2009)). Differences at 95 sites were identified between the M. mycoides genomic sequences. The sequence of the genome successfully transplanted from yeast (CP001668) was used as a design reference; all differences between previously synthesized cassettes that appeared to be of biological significance were corrected to match CP001668. Sequence differences between our synthetic cassettes and CP001668 that occurred at 19 sites appeared harmless, and so were not corrected. These provided 19 polymorphic differences between the synthetic genome (JCVI-syn1) and the natural genome that we have cloned in yeast and use as a standard for genome transplantation from yeast, YCPMmyc.1.1 (Lartigue et al., Science 325, 1693 (2009)).

[0256] Watermarks

[0257] To differentiate between a synthetic or non-naturally occurring genome and a natural one, four exemplary watermark sequences were designed; these sequences were added to a genome at places where insertion of an additional sequence, or where replacement of a sequence, would not interfere with viability.
Watermark-1, 321 unencoded characters, 1246 base pairs

Watermark-2 326 unencoded characters, 1081 base pairs
Following incubation on selective plates, approximately 100 colonies appeared.

Topological trapping and analysis was conducted. Yeast cultures (50 ml) were grown and processed as previously described. Yeast clones containing a completely assembled synthetic genome were screened by multiplex PCR with a primer set that produces 11 amplicons; one at each of the 11 assembly junctions. Primer pairs were designed to span each of the eleven 100-kb amplicons. Of 48 colonies screened, DNA extracted from one clone (sMmYCp235) produced all 11 amplicons. PCR of the WT positive control (YCPMmyc1.1) produced an indistinguishable set of 11 amplicons (data not shown).

To further demonstrate the complete assembly of a synthetic M. mycoides genome, intact DNA was isolated from yeast in agarose plugs and subjected to two restriction analyses; AscI and BssHII. Since these restriction sites are present in three of the four watermark sequences, this choice of digestion produces restriction patterns that are distinct from the natural M. mycoides genome. Natural (WT) and synthetic (235) M. mycoides genomes were isolated from yeast in agarose plugs. In addition, DNA was purified from the host strain alone. Agarose plugs were digested with AscI or BssHII and fragments were separated by clamped homogeneous electric field (CHEF) gel electrophoresis.

Transplantation of Synthetic Donor Genome into Recipient Cells

Additional agarose plugs used in the gel analysis above were also used in genome transplantation experiments. Intact synthetic M. mycoides genomes from the sMmYCp235 yeast clone were transplanted into restriction-minus M. capricolum recipient cells, as previously described (Lartigue et al., Science 325, 1693 (Sep. 25, 2009)). Results were scored by selecting for growth of blue colonies on SP4 medium containing tetracycline and X-gal at 37°C. Genomes isolated from this yeast clone produced 5-15 tetracycline-resistant blue colonies per agarose plug. This was comparable to the YCPMmyc1.1 control. Recovery of colonies in all transplantation experiments was observed when both M. capricolum recipient cells and on M. mycoides genome were present.

To rapidly distinguish the synthetic transplants from M. capricolum or natural M. mycoides, two analyses were performed. First, four primer pairs that were specific to each of the four watermarks were designed such that they produce four amplicons in a single multiplex PCR reaction.

Transplants containing a synthetic genome were screened by multiplex PCR with a primer set that produces 4 amplicons; one internal to each of the four watermarks. One transplant (syn1) originating from yeast clone sMmYCp235 was analyzed alongside a natural, non-synthetic genome (WT) transplanted out of yeast.

All four amplicons were produced by transplants generated from sMmYCp235, but not YCPMmyc1.1 (data not shown).

In summary, these results demonstrate that the synthetic genome was properly assembled and that transplantation to a natural host cell succeeded.

References


[0271] Second, the gel analysis with Ascl and BssHII, described above, was performed. Briefly, natural (WT) and synthetic (syn1) M. mycoides genomes were isolated from M. mycoides transplants in agarose plugs. Agarose plugs were digested with Ascl or BssHII and fragments were separated by CHEF gel electrophoresis. The restriction pattern obtained was consistent with a transplant produced from a synthetic M. mycoides genome (data not shown).

[0272] A single transplant originating from the sMnxYcp235 synthetic genome was sequenced. With the exception of the known polymorphisms that occurred during the synthesis process, and 8 new polymorphisms and an unexpected E. coli transposon insertion, the sequence matched the intended design. This strain is referred to as M. mycoides JCVI-syn1. Colonies (i.e., growing, dividing cells) were grown on SP4 agar containing Xgal to make the cells expressing beta-galactosidase blue. Thus, synthetic cells comprising four watermarks that did not biologically impact the viability of the cells had been produced.

[0273] While preferred embodiments have been shown and described herein, such embodiments are provided by way of example only. It should be understood that various alternatives and equivalents to the embodiments described herein can be employed.

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ORGANISM: Artificial sequence
FEATURE:
OTHER INFORMATION: Synthetic construct: 5 prime reading frame

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```
What is claimed is:

1. A synthetic nucleic acid sequence, wherein said synthetic nucleic acid sequence comprises one or more codon identifiers corresponding to a set of human readable symbols of a reference language, and further wherein said synthetic nucleic acid sequence is not genetically viable; does not have a biological impact upon a recombinant or synthetic organism comprising said synthetic nucleic acid sequence; and conveys a non-genetic message.

2. The synthetic nucleic acid sequence of claim 1, wherein said synthetic nucleic acid sequence cannot be biologically translated into a functional amino acid sequence by the recombinant or synthetic organism.

3. The synthetic nucleic acid sequence of claim 1, wherein said one or more codon identifiers:
   (i) do not correspond to sequence of a gene or other biologically active sequence;
   (ii) correspond to one or more letters, one or more numbers, one or more spaces, one or more punctuation marks, one or more mathematical symbols, one or more typographical characters, one or more new lines, or a combination of any thereof; or
   (iii) consists of three nucleotides.

4. The synthetic nucleic acid sequence of claim 1, wherein said set of human readable symbols comprises a watermark that is a copyright notice, a trademark, a company identifier,
a name, a phrase, a sentence, a quotation, genetic information, unique identifying information, data, or a combination of any thereof.

   5. The synthetic nucleic acid sequence of claim 1, further comprising an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence and/or an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

   6. A recombinant or synthetic organism comprising the synthetic nucleic acid sequence of claim 1.

   7. The recombinant or synthetic organism of claim 9, wherein said recombinant or synthetic organism is a prokaryotic cell, a eukaryotic cell, an archaeal cell or a virus.

   8. A method of creating a recombinant or synthetic organism comprising a watermarking that conveys a non-genetic message, said method comprising:

   (i) generating a nucleic acid sequence comprising a sequence of codon identifiers selected based upon the text of the watermark such that a symbol mapping maps codon identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionately low frequency in the language of the watermark, and maps codon identifiers corresponding to stop codon(s) to human readable symbols that possess a disproportionately high frequency in the language of the watermark;

   (ii) synthesizing said nucleic acid sequence; and

   (iii) introducing said nucleic acid sequence into a recombinant or synthetic organism, thereby creating said recombinant or synthetic organism comprising a watermark.

   9. The method of claim 8, wherein the symbol mapping does not map a three nucleotide codon identifier to a single letter representation of an amino acid residue normally assigned to that three nucleotide codon in the standard genetic code.

   10. The method of claim 8, wherein said generating step (i) is computer-assisted and comprises identifying the set of human readable symbols at a memory module and for each human readable symbol in the set, and using a processor to read a symbol mapping for determining a codon identifier which maps to the respective human readable symbol.

   11. A method of determining the presence of one or more recombinant or synthetic organisms comprising a reference watermark that conveys a non-genetic message in a sample, said method comprising:

   (i) sequencing nucleic acid material obtained from one or more organisms in said sample;

   (ii) transforming the nucleic acid sequence obtained in step (i) to a set of codon identifiers, wherein each codon identifier of said set of codon identifiers consists of three nucleotides, and said transforming is performed in all three reading frames;

   (iii) determining a human readable symbol for each codon identifier in the sequence in all three reading frames, wherein said determination is based at least in part upon a symbol mapping that maps codons identifiers corresponding to start codon(s) to human readable symbols that possess a disproportionately low frequency in the language of the watermark, and that maps codon identifiers corresponding to stop codon(s) to human readable symbols that possess a disproportionately high frequency in the language of the watermark; and

   (iv) comparing the human readable symbol sequence of all three reading frames to the reference watermark in said recombinant or synthetic organism, whereby the presence of the reference watermark in any reading frame of the nucleic acid material obtained in step (i) indicates the presence of the one or more recombinant or synthetic organism in the sample.

   12. An apparatus for transforming a sequence of codon identifiers into a sequence of human readable symbols that conveys a non-genetic message, the apparatus comprising:

   (i) a processor adapted to execute instructions; and

   (ii) a storage module, wherein the storage module comprises a data structure for mapping codon identifiers into human readable symbols, and a set of instructions which, when executed by the processor, generate a human readable symbol for each codon identifier read from a sequence of codon identifiers, wherein the human readable symbol generated is based at least in part upon the data structure;

   wherein the data structure is configured to map a start codon to a human readable symbol with a frequency of occurrence within a reference language that is less than a first predetermined threshold, and wherein the data structure is further configured to map a plurality of stop codons to human readable symbols with frequencies of occurrence within the reference language that are greater than a second predetermined threshold.

   13. The apparatus of claim 12, wherein the data structure does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code.

   14. The apparatus of claim 12, wherein the sequence of codon identifiers comprises an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence and/or an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

   15. A method of transforming a first signal adapted to indicate a sequence of codon identifiers into a second signal adapted to indicate a sequence of human readable symbols that conveys a non-genetic message, the method comprising:

   (i) receiving the first signal;

   (ii) determining a human readable symbol for each codon identifier in the sequence, wherein said determining is performed in all three reading frames and is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a human readable symbol sequence than one or more symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the human readable symbol sequence than one or more human readable symbols from the set of human readable symbols; and

   (iii) transforming the first signal into the second signal based upon the one or more determined human readable symbols.

   16. The method of claim 15, wherein the mapping function does not map a codon identifier to a single letter representation of an amino acid residue normally assigned to that codon identifier in the standard genetic code.
17. The method of claim 15, wherein the sequence of codon identifiers comprises an all-6 reading frame stop codon containing sequence 5' to a first codon identifier in the sequence and/or an all-6 reading frame stop codon containing sequence 3' to the last codon identifier in the sequence.

18. An apparatus for converting a sequence of human readable symbols of a reference language that conveys a non-genetic message into a sequence of codon identifiers, the apparatus comprising:
(i) a processor configured to execute instructions;
(ii) a memory module coupled to the processor and comprising instructions which, when executed by the processor, determine a codon identifier for each human readable symbol contained within the sequence of human readable symbols, wherein each codon identifier is determined upon reading a symbol map; and
(iii) a data module coupled to the memory module, wherein the data module comprises the symbol map, wherein the symbol map is configured to map one or more start codons to respective human readable symbols that possess a disproportionally low frequency of occurrence in the reference language, and wherein the symbol map is further configured to map one or more stop codons to respective human readable symbols that possess a disproportionally high frequency in the reference language.

19. A computer-readable medium for use in an encoding machine, the computer-readable medium comprising instructions which, when executed by the encoding machine, perform a process comprising:
(i) receiving a sequence of human readable symbols that conveys a non-genetic message; and
(ii) generating a codon identifier for each human readable symbol contained within the sequence, wherein the human readable symbol generated is based at least in part upon a mapping function configured to map a start codon to a first human readable symbol, wherein the first human readable symbol has a lower frequency of occurrence in a reference language than one or more human readable symbols from a set of human readable symbols containing the first human readable symbol, and wherein the mapping function is further configured to map a stop codon to a second human readable symbol, wherein the second human readable symbol is contained within the set of human readable symbols, and wherein the second human readable symbol has a higher frequency of occurrence in the reference language than one or more human readable symbols from the set of human readable symbols.

20. A method of generating a sequence of codon identifiers from a sequence of human readable symbols that conveys a non-genetic message, the method comprising:
(i) receiving the sequence of human readable symbols at a memory module;
(ii) loading a human readable symbol map within the memory module, wherein the human readable symbol map is configured to determine a codon identifier that maps to each human readable symbol within the sequence, wherein the human readable symbol map is further configured to map a human readable symbol with a frequency of occurrence that is less than a first predetermined threshold within a reference language to a start codon, and wherein the symbol map is further configured to map a human readable symbol with a frequency of occurrence that is greater than a second predetermined threshold within the reference language to a stop codon; and
(iii) outputting a sequence of codon identifiers corresponding to each human readable symbol within the sequence.