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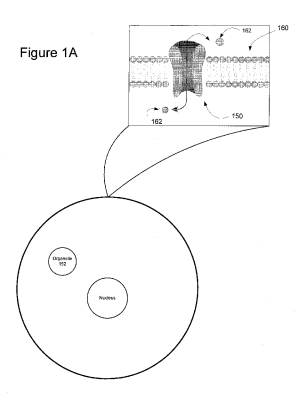
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[Continued on next page]

(54) Title: LIGHT-SENSITIVE ION-PASSING MOLECULES



(57) Abstract: The invention provides polynucleotides and methods for expressing light- activated proteins in animal cells and altering an action potential of the cells by optical stimulation. The invention also provides animal cells and non-human animals comprising cells expressing the light-activated proteins.

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LIGHT-SENSITIVE ION-PASSING MOLECULES

Related Patent Document

This patent document claims the benefit, under 35 U.S.C. § 119(e), of U.S. Provisional Patent Application Serial No. 61/314,969 filed on March 17, 2010, and entitled "Light-Sensitive Ion-Passing Molecules;" this patent document and the Appendices filed in the underlying provisional application are fully incorporated herein by reference..

Overview and Summary

Aspects of the present disclosure relate generally to systems and approaches for stimulating target cells, and more particularly to using optics to stimulate the target cells. The stimulation of various cells of the body has been used to produce a number of beneficial effects. One method of stimulation involves the use of electrodes to introduce an externally generated signal into cells. One problem faced by electrode-based brain stimulation techniques is the distributed nature of neurons responsible for a given mental process. Conversely, different types of neurons reside close to one another such that only certain cells in a given region of the brain are activated while performing a specific task. Alternatively stated, not only do heterogeneous nerve tracts move in parallel through tight spatial confines, but the cell bodies themselves may exist in mixed, sparsely embedded configurations. This distributed manner of processing seems to defy the best attempts to understand canonical order within the CNS, and makes neuromodulation a difficult therapeutic endeavor. This architecture of the brain poses a problem for electrode-based stimulation because electrodes are relatively indiscriminate with regards to the underlying physiology of the neurons that they stimulate. Instead, physical proximity of the electrode poles to the neuron is often the single largest determining factor as to which neurons will be stimulated. Accordingly, it is generally not feasible to absolutely restrict stimulation to a single class of neuron using electrodes.

Another issue with the use of electrodes for stimulation is that because electrode placement dictates which neurons will be stimulated, mechanical stability is frequently inadequate, and results in lead migration of the electrodes from the targeted area. Moreover, after a period of time within the body, electrode leads frequently become encapsulated with glial cells, raising the effective electrical resistance of the electrodes, and hence the electrical power delivery required to reach targeted cells. Compensatory

increases in voltage, frequency or pulse width, however, may spread the electrical current and increase the unintended stimulation of additional cells.

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Another method of stimulus uses photosensitive bio-molecular structures to stimulate target cells in response to light. For instance, light activated proteins can be used to control the flow of ions through cell membranes. By facilitating or inhibiting the flow of positive or negative ions through cell membranes, the cell can be briefly depolarized, depolarized and maintained in that state, or hyperpolarized. Neurons are an example of a type of cell that uses the electrical currents created by depolarization to generate communication signals (*i.e.*, nerve impulses). Other electrically excitable cells include skeletal muscle, cardiac muscle, and endocrine cells. Neurons use rapid depolarization to transmit signals throughout the body and for various purposes, such as motor control (*e.g.*, muscle contractions), sensory responses (*e.g.*, touch, hearing, and other senses) and computational functions (*e.g.*, brain functions). Thus, the control of the depolarization of cells can be beneficial for a number of different purposes, including (but not limited to) psychological therapy, muscle control and sensory functions.

Various aspects of the present invention are directed toward a blue-light sensing opsin capable of inhibiting neural activity. The opsin comes from cryptophytes *Guillardia theta* (*G. theta*). The opsin of interest is the third opsin isolated from *G. theta*, and is abbreviated GtR3. GtR3 is capable of mediating a hyperpolarizing current when illuminated with light. Characterization of the action spectra for GtR3 suggests that the absorption maxima are around 490nm, and GtR3 is not activated by yellow light.

Various aspects of the present invention are directed to a blue-light sensing channelrhodopsin capable of exciting neural activity. The channelrhodopsin is derived from *Dunaliella salina*. The channelrhodopsin of interest is abbreviated as DChR. DChR can be heterologously expressed in mammalian neurons and mediates a robust depolarizing current when illuminated with blue light. The action maxima for DChR are around 500nm.

Consistent with an embodiment of the present disclosure, an inhibitory current flow is created by engineering a protein derived from *Guillardia theta* that responds to light by producing an inhibitory current to dissuade depolarization of a neuron. The protein is delivered to a neuron of interest and the neuron is exposed to light.

Consistent with another embodiment of the present disclosure, a method of optical stimulation of a cell expressing a GtR3 proton pump comprises providing a sequence of stimuli to the cell, each stimulus increasing the probability of a depolarizing

event occurring in the cell. Light is provided to the cell to activate the expression of the GtR3 proton pump, thereby decreasing the probability of the depolarizing event occurring in the cell. In certain specific embodiments the light provided is in the blue light spectrum.

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Consistent with another embodiment of the present disclosure, a system for controlling an action potential of a neuron or other cell *in vivo* is disclosed. The system comprises a delivery device that introduces a protein responsive to blue light to the neuron or cell. The protein responsive to blue light produces an inhibitory current in response to blue light. The system includes a blue light source that generates light for stimulation of the blue light responsive protein and a control device that controls the generation of light by the light source.

Consistent with another embodiment of the present disclosure, a method for providing a light responsive protein for mammalian expression is disclosed. A light responsive protein is isolated from *G. theta*. The isolated protein has a C-terminus and an N-terminus. An endoplasmic reticulum (ER) export signal is added to the C-terminus of the isolated protein to create an enhanced light responsive protein. The enhanced protein is placed in an empty virus vector for delivery to a cell of interest. The virus vector with the enhanced protein is then delivered to the cell of interest.

Consistent with an embodiment of the present disclosure an animal cell is provided. The animal cell includes an integrated exogenous molecule which expresses a proton pump responsive to blue light. The exogenous molecule is derived from *G. theta*. In certain embodiments the animal cell is a neural cell. The animal cell may also be a muscle cell or a cell line, for example.

The above discussion of the present disclosure is not intended to describe each illustrated embodiment or every implementation of the present invention. The figures and detailed description that follow more particularly exemplify these embodiments.

Brief Description of the Drawings

The invention may be more completely understood in consideration of the detailed description of various embodiments of the invention that follows in connection with the accompanying drawings as follows:

- FIG. 1A shows an ion pump in an organelle membrane;
- FIG. 1B shows an ion pump in a cell membrane;

FIG. 2A shows cell populations expressing combinations of light responsive proteins;

- FIG. 2B shows a stimulus profile for use with certain embodiments in which two or more light responsive proteins are introduced into the same cell population;
- FIG. 3 shows a block diagram of a system for stimulating target cells, according to an example embodiment of the present invention;

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- FIG. 4 shows a block diagram of an implantable device for stimulating target cells, according to an example embodiment of the present invention;
- FIG. 5 shows a block diagram of an implantable device, according to an example embodiment of the present invention;
 - FIG. 6A shows a block diagram of an implantable device, according to an example embodiment of the present invention;
 - FIG. 6B shows a circuit diagram corresponding to the block diagram of FIG. 6A, according to an example embodiment of the present invention;
 - FIGs. 7A and FIG. 7B show a diagram of a mesh for containing photosensitive bio-molecules, according to an example embodiment of the present invention;
 - FIGs. 8A and FIG. 8B show a diagram of a viral matrix, according to an example embodiment of the present invention;
 - FIG. 9 shows a circuit diagram of a circuit that produces light in response to a magnetic field, according to an example embodiment of the present invention;
 - FIGs. 10A, 10B and 10C show a block diagram and circuits for the production of light in response to a RF signal, according to an example embodiment of the present invention;
- FIGs. 11A and FIG. 11B each show a diagram of a fiber-optic device, according to an example embodiment of the present invention;
 - FIGs. 12A, 12B, 12C and 12D depict various stages in the production of a photosensitive biological portion, according to an example embodiment of the present invention; and
- FIG. 13 shows an implantation device, according to an example embodiment of the present invention;
 - FIGs. 14A and FIG. 14B show a diagram for another implantation device, according to an example embodiment of the present invention;
 - FIG. 15 depicts an arrangement with multiple light sources, according to an example embodiment of the present invention;

FIG. 16 shows a system for controlling electrical properties of one or more cells in vivo, according to an example embodiment of the present invention;

- FIG. 17 shows a system for controlling electrical properties of one or more cells in vivo, according to an example embodiment of the present invention
- FIG. 18A shows a block diagram of a system for optical drug screening, according to an example embodiment of the present invention;

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- FIG. 18B shows a specific system diagram of a large-format, quasi-automated system for drug screening in accordance with the present methodology, according to an example embodiment of the present invention;
- FIG. 19 shows a system diagram of a small-format, fully automated drug screening system which operates in accordance with the invented methodology, according to an example embodiment of the present invention;
- FIG. 20A depicts the workings of an example of emitter/detector units, according to an example embodiment of the present invention;
- FIG. 20B depicts the workings of another embodiment of emitter/detector units, according to an example embodiment of the present invention;
- FIG. 21A and 21B depict an electronic circuit mechanism for activating the LED emitters used within the emitter/detector units, according to an example embodiment of the present invention;
- FIG. 22 shows a timeline for a sequence of events in the context of an example screening process, according to an example embodiment of the present invention;
- FIG. 23 illustrates an example of a layout of cell and drug samples within the wells of a well-plate, according to an example embodiment of the present invention; and
- FIG. 24 illustrates the context in which the disclosed invention may be employed within a larger system that facilitates high-throughput drug screening, according to an example embodiment of the present invention.
- FIG. 25D shows representative photocurrent traces showing in cells transduced with eNpHR3.0 and cells transduced with eNpHR2.0, and summary plots thereof;
- FIG. 25E shows representative hyperpolarization voltage traces showing in cells transduced with eNpHR3.0 and cells transduced with eNpHR2.0, and summary plots thereof;
- FIG. 26B shows a model of trans-synaptic gene activation by WGA-Cre-fusion. The schematic depicts two injection sites (one with WGA-Cre-fusion gene and another with Cre-dependent opsin virus) and long range projections; Cre can be *trans*-

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synaptically delivered from transduced cells to activate distant gene expression only in synaptically connected neurons that have received the Cre-dependent virus.

FIG. 26C shows construct design for the WGA-Cre and Cre-dependent AAV vectors optimized with mammalian codons;

FIG. 27A shows that six hundred thirty nanometer light evokes robust photocurrents in neurons transduced with eNpHR3.0 (representative voltage clamp trace at left). Summary plot comparing eNpHR2.0- and eNpHR3.0-expressing neurons (at right); eNpHR2.0, 42.7 ± 4.5 pA; eNpHR3.0, 239.4 ± 28.7 pA; unpaired t test p = 0.00004; n = 10).

FIG. 27B shows that six hundred thirty nanometer illumination evoked robust hyperpolarization (representative voltage clamp trace at left). Summary plot comparing eNpHR2.0- and eNpHR3.0-expressing neurons (right); 15.6 ± 3.2 mV for eNpHR2.0 and 43.3 ± 6.1 mV for eNpHR3.0; unpaired t test p = 0.00116; n = 10).

FIG. 27C shows a summary of outward photocurrents evoked by different wavelengths of red and far-red/infrared border illumination: 630 nm, 239.4 ± 28.7 pA (left, n = 10); 660 nm, 120.5 ± 16.7 pA (middle, n = 4); and 680 nm: 76.3 ± 9.1 pA (right, n = 4). Power density: 3.5 mW/mm² (630 nm) and 7 mW/mm² (660 nm, 680 nm).

FIG. 27D shows that illumination with red and far-red/infrared border light inhibited spiking induced by current injection in neurons expressing eNpHR3.0. Typical current-clamp traces show optical inhibition at 630 nm (top left), 660 nm (top right), and 680 nm (bottom). Power density: 3.5 mW/mm² (630 nm) and 7 mW/mm² (660 nm, 680 nm).

FIG. 27G shows that blue light (445 nm, 5 ms pulses) drove spiking at 20 Hz (left) and 10 Hz (right), while simultaneous application of yellow light (590 nm) inhibited spikes.

FIG. 27F shows activation spectrums for eNPAC, ChR2 (H134R), and eNpHR3.0 alone;

FIG. 28B shows five hundred sixty nanometer light induced outward photocurrents in eBR cells, and the corresponding sample trace in voltage clamp;

FIG. 28C shows five hundred sixty nanometer light induced hyperpolarizations in eBR cells, and the corresponding sample trace in current clamp;

FIG. 29A shows general subcellular targeting strategies for adapting microbial opsin genes to metazoan intact-systems biology;

FIG. 29B shows refinement of targeting at the tissue and subcellular levels;

FIG. 30A shows stability and recovery of potent photocurrents in cells expressing eNpHR3.0 when exposed to pairs of 10 second long yellow light pulses separated in time by: 2.5 seconds, 5 seconds, 10 seconds, and 20 seconds;

FIG. 30B shows a timecourse of eNpHR3.0 normalized photocurrents for long-term continuous light exposure;

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FIG. 30C shows stability of outward current of eNpHR3.0 over greater than 10 minutes; and

FIG. 31C shows sample current clamp and voltage clamp traces and summary data of GtR3 function under 472 nm light.

While the invention is amenable to various modifications and alternative forms, examples thereof have been shown by way of example in the drawings and will be described in detail. It should be understood, however, that the intention is not to limit the invention to the particular embodiments shown and/or described. On the contrary, the intention is to cover all modifications, equivalents, and alternatives falling within the spirit and scope of the invention.

Detailed Description

The present invention is believed to be useful for facilitating practical application of a variety of photosensitive bio-molecular structures, and the invention has been found to be particularly suited for use in arrangements and methods dealing with cellular membrane voltage control and stimulation. While the present invention is not necessarily limited to such applications, various aspects of the invention may be appreciated through a discussing of various examples using this context.

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As used herein, stimulation of a target cell is generally used to describe modification of properties of the cell. For instance, the stimulus of a target cell may result in a change in the properties of the cell membrane that can lead to the depolarization or polarization of the target cell. In a particular instance, the target cell is a neuron and the stimulus affects the transmission of impulses by facilitating or inhibiting the generation of impulses by the neuron.

Consistent with one example embodiment of the present invention, a light-responsive protein is engineered in a cell. The protein affects a flow of ions (anions, cations or protons) across the cell membrane in response to light. This change in ion flow creates a corresponding change in the electrical properties of the cells including, for example, the voltage and current flow across the cell membrane. In one instance, the protein functions *in vivo* using an endogenous cofactor to modify ion flow across the cell membrane. In another instance, the protein changes the voltage across the cell membrane so as to dissuade or encourage action potential firing in the cell. In yet another instance, the protein is capable of changing the electrical properties of the cell within several milliseconds of the light being introduced.

An inhibitory protein dissuades firing of the action potential by moving the potential of the cell away from the action potential trigger level for the cell. In many neurons, this means that the protein increases the negative voltage seen across the cell membrane. In a specific instance, the protein acts as a proton pump that actively transfers protons out of the cell. In this manner, the protein generates an inhibitory current across the cell membrane. More specifically, the protein responds to light by lowering the voltage across the cell, thereby decreasing the probability that an action potential or depolarization will occur.

Certain aspects of the present invention are based on the identification and development of a molecule/protein that functions as a proton pump. This proton pump is derived from the cryptophyte *Guillardia theta* (*G. theta*) and has been developed for

expression in target cells. In certain more specific embodiments the cell is a neuron. The engineered protein, GtR3, responds to blue light by producing an inhibitory current to dissuade depolarization of the cell. When expressed in neural cells the proton pump (GtR3) can be used to inhibit neural activity in response to blue light stimulation. The GtR3 pump responds to optical stimulus by creating an inhibitory current across the neural membrane. This current inhibits action potentials while the modified cell is exposed to (blue) light.

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The present disclosure also provides for the modification of light-activated proteins expressed in a cell by the addition of one or more amino acid sequence motifs which enhance transport to the plasma membranes of mammalian cells. Light-activated proteins derived from evolutionarily simpler organisms may not be expressed or tolerated by mammalian cells or may exhibit impaired subcellular localization when expressed at high levels in mammalian cells. Consequently, in some embodiments, the light-activated protein expressed in a cell is fused to one or more amino acid sequence motifs selected from the group consisting of a signal peptide, an endoplasmic reticulum (ER) export signal, a membrane trafficking signal, and an N-terminal golgi export signal. The one or more amino acid sequence motifs which enhance light-activated protein transport to the plasma membranes of mammalian cells can be fused to the N-terminus, the C-terminus, or to both the N- and C-terminal ends of the light-activated protein. Optionally, the light-activated protein and the one or more amino acid sequence motifs may be separated by a linker. Additional protein motifs which can enhance lightactivated protein transport to the plasma membrane of a cell are described in U.S. Patent Application No. 12/041,628 which is incorporated herein in its entirety.

The present disclosure additionally provides for light-activated proteins which contain amino acid substitutions, deletions, and insertions in the amino acid sequence of a native light-activated protein (such as, but not limited to, native GtR3, NpHR, DChR, and BR). Light-activated proteins include those in which one or more amino acid residues have undergone an amino acid substitution while retaining the ability to respond to light and the ability to control the polarization state of a plasma membrane. For example, light-activated proteins can be made by substituting one or more amino acids into the native or wild type amino acid sequence of the protein. In some embodiments, the invention includes proteins comprising altered amino acid sequences in comparison with a native amino acid sequence, wherein the altered light-activated protein retains the characteristic light-activated nature and/or the ability to regulate ion flow across plasma

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membranes of the precursor protein but may have altered properties in some specific aspects (for example, an increased or decreased sensitivity to light, an increased or decreased sensitivity to particular wavelengths of light, and/or an increased or decreased ability to regulate the polarization state of the plasma membrane of a mammalian cell, as compared to the native protein). Amino acid substitutions in a native protein sequence may be conservative or non-conservative and such substituted amino acid residues may or may not be one encoded by the genetic code. The standard twenty amino acid "alphabet" is divided into chemical families based on chemical properties of their side chains. These families include amino acids with basic side chains (e.g., lysine, arginine, histidine), acidic side chains (e.g., aspartic acid, glutamic acid), uncharged polar side chains (e.g., glycine, asparagine, glutamine, serine, threonine, tyrosine, cysteine), nonpolar side chains (e.g., alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan), beta-branched side chains (e.g., threonine, valine, isoleucine) and sidw chains having aromatic groups (e.g., tyrosine, phenylalanine, tryptophan, histidine). A "conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a chemically similar side chain (i.e., replacing an amino acid possessing a basic side chain with another amino acid with a basic side chain). A "non-conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a chemically different side chain (i.e., replacing an amino acid having a basic side chain with an amino acid having an aromatic side chain).

Certain aspects of the present invention are directed to an animal cell expressing the GtR3 molecule. In this manner, the animal cell includes an integrated exogenous molecule which expresses a proton pump responsive to blue light. In certain non-limiting embodiments the animal cell can be a neural cell, a muscle cell, a rod or cone cell or a cell line. In some embodiments, the animal cell is a mammalian cell.

Provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to blue light and is derived from *Guillardia theta*, wherein the protein is capable of mediating a hyperpolarizing current in the cell when the cell is illuminated with light. In some embodiments the light has a wavelength between about 450 and 495 nm. In some embodiments, the light has a wavelength about 490 nm. In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO: 1. In some embodiments, the light-activated protein comprises an amino acid sequence at

least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1. In some embodiments, the light-activated protein comprises substitutions, deletions, and/or insertions introduced into a native amino acid sequence to increase or decrease sensitivity to light, increase or decrease sensitivity to particular wavelengths of light, and/or increase or decrease the ability of the light-activated protein to regulate the polarization state of the plasma membrane of the cell. In some embodiments, the light-activated protein contains one or more conservative amino acid substitutions. In some embodiments, the light-activated protein contains one or more non-conservative amino acid substitutions. The light-activated protein comprising substitutions, deletions, and/or insertions introduced into the native amino acid sequence suitably retains the ability to hyperpolarize the plasma membrane of a neuronal cell in response to light.

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Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1 and an endoplasmic reticulum (ER) export signal. The ER export signal may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the ER export signal is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the ER export signal comprises the amino acid sequence FXYENE, where X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid sequence FCYENEV.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1 and a signal peptide (e.g., which enhances transport to the plasma membrane). The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core

amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the signal peptide comprises the amino acid sequence

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MDYGGALSAVGRELLFVTNPVVVNGSVLVPEDQCYCAGWIESRGTNG. In some embodiments, other signal peptides (such as signal peptides from other opsins) may be used.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1 and a trafficking signal (e.g., which enhances transport to the plasma membrane). The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the trafficking signal is derived from the amino acid sequence of the human inward rectifier potassium channel Kir2.1. In some embodiments, the trafficking signal comprises the amino acid sequence K S R I T S E G E Y I P L D Q I D I N V.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1 and two or more amino acid sequence motifs which enhance transport to the plasma membranes of mammalian cells selected from the group consisting of a signal peptide, an ER export signal, and a membrane trafficking signal. In some embodiments, the light activated protein comprises an N-terminal signal peptide and a C-terminal ER export signal. In some embodiments, the light activated protein comprises an N-terminal signal peptide and a C-terminal trafficking signal. In some embodiments, the light activated protein comprises an N-

terminal signal peptide, a C-terminal ER Export signal, and a C-terminal trafficking signal. In some embodiments, the light activated protein comprises a C-terminal ER Export signal and a C-terminal trafficking signal. In some embodiments, the C-terminal ER Export signal and the C-terminal trafficking signal are linked by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments the ER Export signal is more C-terminally located than the trafficking signal. In some embodiments the trafficking signal is more C-terminally located than the ER Export signal.

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Also provided herein are isolated polynucleotides encoding any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1. Also provided herein are expression vectors (such as a viral vector described herein) comprising a polynucleotide encoding any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 1. The polynucleotides may be used for expression of the light-activated protein in animal cells.

Provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to blue light and is derived from *Dunaliella salina*, wherein the protein is capable of mediating a depolarizing current in the cell when the cell is illuminated with light. In some embodiments the light has a wavelength between about 450 and 495 nm. In some embodiments, the light has a wavelength about 490 nm. In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 23-365, 21-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 1

365, 2-365, or 1-365 of SEQ ID NO: 2. In some embodiments, the light-activated protein comprises substitutions, deletions, and/or insertions introduced into a native amino acid sequence to increase or decrease sensitivity to light, increase or decrease sensitivity to particular wavelengths of light, and/or increase or decrease the ability of the light-activated protein to regulate the polarization state of the plasma membrane of the cell. In some embodiments, the light-activated protein contains one or more conservative amino acid substitutions. In some embodiments, the light-activated protein contains one or more non-conservative amino acid substitutions. The light-activated protein comprising substitutions, deletions, and/or insertions introduced into the native amino acid sequence suitably retains the ability to depolarize the plasma membrane of a neuronal cell in response to light.

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Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2 and an endoplasmic reticulum (ER) export signal. The ER export signal may be fused to the C-terminus of the core amino acid sequence or may be fused to the Nterminus of the core amino acid sequence. In some embodiments, the ER export signal is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the ER export signal comprises the amino acid sequence FXYENE, where X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid sequence VXXSL, where X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid sequence FCYENEV.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365,

10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2 and a signal peptide (e.g., which enhances transport to the plasma membrane). The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the signal peptide comprises the amino acid sequence

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MDYGGALSAVGRELLFVTNPVVVNGSVLVPEDQCYCAGWIESRGTNG. In some embodiments, other signal peptides (such as signal peptides from other opsins) may be used.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid 15 sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2 and a trafficking signal (e.g., which enhances transport to the plasma membrane). 20 The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, 25 for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the trafficking signal is derived from the amino acid sequence of the human inward rectifier potassium channel Kir2.1. In some embodiments, the trafficking signal comprises the amino acid sequence K S R I T S E G E Y I P L D Q I D I N V. 30

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-

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365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2 and two or more amino acid sequence motifs which enhance transport to the plasma membranes of mammalian cells selected from the group consisting of a signal peptide, an ER export signal, and a membrane trafficking signal. In some embodiments, the light activated protein comprises an N-terminal signal peptide and a C-terminal ER export signal. In some embodiments, the light activated protein comprises an N-terminal signal peptide and a C-terminal trafficking signal. In some embodiments, the light activated protein comprises an N-terminal signal peptide, a C-terminal ER Export signal, and a C-terminal trafficking signal. In some embodiments, the light activated protein comprises a C-terminal ER Export signal and a C-terminal trafficking signal. In some embodiments, the C-terminal ER Export signal and the C-terminal trafficking signal are linked by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments the ER Export signal is more C-terminally located than the trafficking signal. In some embodiments the trafficking signal is more C-terminally located than the ER Export signal.

Also provided herein are isolated polynucleotides encoding any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2. Also provided herein are expression vectors (such as a viral vector described herein) comprising a polynucleotide encoding any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in residues 25-365, 24-365, 23-365, 22-365, 21-365, 20-365, 19-365, 18-365, 17-365, 16-365, 15-365, 14-365, 13-365, 13-365, 12-365, 11-365, 10-365, 9-365, 8-365, 7-365, 6-365, 5-365, 4-365, 3-365, 2-365, or 1-365 of SEQ ID NO: 2.. The polynucleotides may be used for expression of the light-activated protein in animal cells.

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Provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to amber as well as red light and is derived from *Natronomonas pharaonis*, wherein the protein is capable of mediating a hyperpolarizing current in the cell when the cell is illuminated with light. In some embodiments the light has a wavelength between about 580 and 630 nm. In some embodiments, the light has a wavelength about 589 nm. In some embodiments, the light has a wavelength greater than about 630 nm (e.g. less than 740 nm). In some embodiments, the light has a wavelength around 630 nm. In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO: 3. In some embodiments, the light-activated protein comprises an amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3. In some embodiments, the light-activated protein comprises substitutions, deletions, and/or insertions introduced into a native amino acid sequence to increase or decrease sensitivity to light, increase or decrease sensitivity to particular wavelengths of light, and/or increase or decrease the ability of the light-activated protein to regulate the polarization state of the plasma membrane of the cell. In some embodiments, the lightactivated protein contains one or more conservative amino acid substitutions. In some embodiments, the light-activated protein contains one or more non-conservative amino acid substitutions. The light-activated protein comprising substitutions, deletions, and/or insertions introduced into the native amino acid sequence suitably retains the ability to hyperpolarize the plasma membrane of a neuronal cell in response to light.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3 and an endoplasmic reticulum (ER) export signal. The ER export signal may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the ER export signal is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the ER export signal comprises the amino acid sequence FXYENE, where

X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid sequence VXXSL, where X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid sequence FCYENEV.

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Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3 and a trafficking signal (e.g., which enhances transport to the plasma membrane). The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the trafficking signal is derived from the amino acid sequence of the human inward rectifier potassium channel Kir2.1. In some embodiments, the trafficking signal comprises the amino acid sequence K S R I T S E G E Y I P L D Q I D I N V.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3 and two or more amino acid sequence motifs which enhance transport to the plasma membranes of mammalian cells selected from the group consisting of an ER export signal and a membrane trafficking signal. In some embodiments, the light activated protein comprises a C-terminal ER Export signal and a C-terminal trafficking signal. In some embodiments, the C-terminal ER Export signal and the C-terminal trafficking signal are linked by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments the ER Export signal is more C-terminally located than the trafficking signal. In some embodiments the trafficking signal is more C-terminally located than the ER Export signal.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3, wherein the N-terminal signal peptide of SEQ ID NO:3 is deleted or substituted In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in residues 40-291, 39-291, 38-291, 37-291, 36-291, 35-291, 34-291, 33-291, 32-291, 31-291, 30-291, 29-291, 28-291, 27-291, 26-291, 25-291, 24-291, 23-291, 22-291, 21-291, 20-291, 19-291, 18-291, 17-291, 16-291, 15-291, 14-291, 13-291, 13-291, 12-291, 11-291, 10-291, 9-291, 8-291, 7-291, 6-291, 5-291, 4-291, 3-291, 2-291, or 1-291 of SEQ ID NO:3. In some embodiments, other signal peptides (such as signal peptides from other opsins) may be used. The light-activated protein may further comprise an ER transport signal and a membrane trafficking signal described herein.

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Also provided herein are polynucleotides encoding for any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 3, an ER export signal, and a membrane trafficking signal. The polynucleotides may be in an expression vector (such as a viral vector described herein). The polynucleotides may be used for expression of the light-activated protein in animal cells.

Provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to green light and is derived from *Halobacterium salinarum* wherein the protein is capable of mediating a hyperpolarizing current in the cell when the cell is illuminated with light. In some embodiments the light has a wavelength between about 520 and 570 nm. In some embodiments, the light has a wavelength about 560 nm. In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO: 4. In some embodiments, the light-activated protein comprises an amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 4. In some embodiments, the light-activated protein comprises substitutions, deletions, and/or insertions introduced into a native amino acid sequence to increase or decrease sensitivity to light, increase or decrease sensitivity to particular wavelengths of light, and/or increase or decrease the ability of the light-activated protein to regulate the

polarization state of the plasma membrane of the cell. In some embodiments, the light-activated protein contains one or more conservative amino acid substitutions. In some embodiments, the light-activated protein contains one or more non-conservative amino acid substitutions. The light-activated protein comprising substitutions, deletions, and/or insertions introduced into the native amino acid sequence suitably retains the ability to hyperpolarize the plasma membrane of a neuronal cell in response to light.

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Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 4 and an endoplasmic reticulum (ER) export signal. The ER export signal may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the ER export signal is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the ER export signal comprises the amino acid sequence FXYENE, where X can be any amino acid. In some embodiments, the ER export signal comprises the amino acid. In some embodiments, the ER export signal comprises the amino acid. In some embodiments, the ER export signal comprises the amino acid sequence FCYENEV.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 4 and a trafficking signal (e.g., which enhances transport to the plasma membrane). The signal peptide may be fused to the C-terminus of the core amino acid sequence or may be fused to the N-terminus of the core amino acid sequence. In some embodiments, the signal peptide is linked to the core amino acid sequence by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments, the trafficking signal is derived from the amino acid sequence of the human inward rectifier potassium channel Kir2.1. In some

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embodiments, the trafficking signal comprises the amino acid sequence K S R I T S E G E Y I P L D Q I D I N V.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 4 and two or more amino acid sequence motifs which enhance transport to the plasma membranes of mammalian cells selected from the group consisting of an ER export signal and a membrane trafficking signal. In some embodiments, the light activated protein comprises a C-terminal ER Export signal and a C-terminal trafficking signal. In some embodiments, the C-terminal ER Export signal and the C-terminal trafficking signal are linked by a linker. The linker can comprise any of 5, 10, 20, 30, 40, 50, 75, 100, 125, 150, 175, 200, 225, 250, 275, 300, 400, or 500 amino acids in length. The linker may further comprise a fluorescent protein, for example, but not limited to, a yellow fluorescent protein, a red fluorescent protein, a green fluorescent protein, or a cyan fluorescent protein. In some embodiments the ER Export signal is more C-terminally located than the trafficking signal. In some embodiments the trafficking signal is more C-terminally located than the ER Export signal.

Also provided herein is an animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the sequence shown in SEQ ID NO: 4, wherein the N-terminal signal peptide of SEQ ID NO:4 is deleted or substituted. In some embodiments, the light-activated protein comprises an amino acid sequence at least 95% identical to the sequence shown in residues 40-262, 39-262, 38-262, 37-262, 36-262, 35-262, 34-262, 33-262, 32-262, 31-262, 30-262, 29-262, 28-262, 27-262, 26-262, 25-262, 24-262, 23-262, 22-262, 21-262, 20-262, 19-262, 18-262, 17-262, 16-262, 15-262, 14-262, 13-262, 13-262, 12-262, 11-262, 10-262, 9-262, 8-262, 7-262, 6-262, 5-262, 4-262, 3-262, 2-262, or 1-262 of SEQ ID NO:4. In some embodiments, other signal peptides (such as signal peptides from other opsins) may be used. The light-activated protein may further comprise an ER transport signal and/or a membrane trafficking signal described herein.

Also provided herein are polynucleotides encoding for any of the proteins described herein, such as a light-activated protein comprising a core amino acid sequence at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to

the sequence shown in SEQ ID NO: 4, an ER export signal, and a membrane trafficking signal. The polynucleotides may be in an expression vector (such as a viral vector described herein). The polynucleotides may be used for expression of the light-activated protein in animal cells.

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In certain more particular embodiments, a method for providing a light responsive protein for mammalian expression is provided. A light responsive protein is isolated from *G. theta* (GtR3). GtR3 has a C-terminus and an N-terminus. A promoter is added to the C-terminus of GtR3. In certain embodiments the promoter is an endoplasmic reticulum (ER) export signal. For more specifics on optimizing GtR3 for expression in mammalian cells, *see* Appendix A as filed in the underlying provisional application and entitled, "Molecular and Cellular Approaches for Diversifying and Extending Optogenetics."

An excitatory protein encourages firing of the action potential by moving the potential of the cell toward the action potential trigger level for the cell. In many neurons, this means that the protein decreases the negative voltage seen across the cell membrane. In a specific instance, the protein acts an additional ion channel that transfers cations into the cell. In this manner, the protein generates an excitatory current across the cell membrane. More specifically, the protein responds to light by raising the voltage across the cell membrane, thereby increasing the probability that an action potential or depolarization will occur. In certain instances the voltage across the cell membrane can be increased to the action potential trigger level or higher, causing depolarization of the cell.

Certain aspects of the present invention are based on the identification and development of a channel (DChR) that is derived from *Dunaliella salina*.

Channelrhodopsin DChR mediates a robust depolarizing current when illuminated with blue light. The depolarizing current may cause a cell to become excited in response to exposure to blue light. In certain embodiments DChR is expressed as an exogenous protein in a cell of interest. For neural cells, DChR can function as an excitatory protein that uses an endogenous cofactor when delivered to a cell of interest.

The introduction of GtR3 and DChR to the field of optogenetics opens the door for a variety of applications. For instance, GtR3 and DChR can be used in combination with other optogenetic ion-passing molecules, such as ChR2 (developed from *Chlamydomonas reinhardtii*). GtR3 and DChR have different operating parameters, relative to other light-responsive molecules. This facilitates the development of a cell or

a cell population that has a precisely-tuned response to different wavelengths of light. The parameter that can be tuned include, but are not limited to, current density, hyperpolarization level, membrane conductance, stimulus frequency, resting potential, optical wavelength and/or optical intensity.

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In another example embodiment, a method for controlling action potential of a neuron involves the following steps: engineering a first light responsive protein in the neuron; producing, in response to light, an inhibitory current in the neuron and from the first light responsive protein; engineering a second light responsive protein in the neuron; and producing, in response to light, an excitation current in the neuron from the second light responsive protein. The first light responsive protein and the second light responsive protein may be responsive to different wavelengths of light. In certain instances, the use of DChR or GtR3 can facilitate specific current densities for this excitation or inhibitory current and/or optical wavelength at which such current is generated.

In another example embodiment, a method for controlling action potential of a neuron involves the following steps: engineering a first light responsive protein in the neuron; producing, in response to light, an inhibitory current in the neuron and from the first light responsive protein; engineering a second light responsive protein in the neuron; and producing, in response to light, a second inhibitory current from the second light responsive protein, the combination of the two currents more strongly inhibiting the neuron than a single light responsive protein. For instance, the first light responsive protein could be NpHR and the second light responsive protein could be GtR3.

Another method for controlling a voltage level across a cell membrane of a cell includes measuring the voltage level of the cell membrane. A light responsive protein is engineered and introducing the light responsive protein into the cell, where it is expressed. A light of a particular wavelength is provided, which produces a reaction in the cell by the light responsive protein. The response by the light responsive protein produces a current across the cell membrane. The current is responsive to the measured voltage level.

Another aspect of the present invention is directed to a system for controlling an action potential of a neuron *in vivo*. The system includes a delivery device, a light source, and a control device. The delivery device introduces a light responsive protein to the neuron, with the light responsive protein producing an inhibitory current. The light source generates light for stimulating the light responsive protein, and the control device

controls the generation of light by the light source. In other embodiments the light responsive protein introduced into the neuron produces an excitatory current.

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In more detailed embodiments, such a system is further adapted such that the delivery device introduces the light responsive protein by one of transfection, transduction and microinjection, and/or such that the light source introduces light to the neuron via one of an implantable light generator and fiber-optics. For additional detail regarding method of delivery and expression of genes, see, PCT Publication No. WO2010/019619 A1 (PCT/US2009/053474), entitled "Method and Composition for Controlling Gene Expression," which is fully incorporated herein by reference.

Another aspect of the present disclosure is directed to a method for treatment of a disorder. The method targets a group of neurons associated with the disorder; and in this group, the method includes engineering inhibitory proteins to respond to light by producing an inhibitory current to dissuade depolarization of the neurons, and exposing the neurons to light, thereby dissuading depolarization of the neurons.

Yet another aspect of the present disclosure is directed to a further method for treatment of a disorder. The method targets a group of neurons associated with the disorder; and in this group, the method includes engineering excitatory proteins to respond to light by producing an excitatory current to encourage depolarization of the neurons, and exposing the neurons to light, thereby encouraging depolarization of the neurons. In other aspects of the present invention both inhibitory and excitatory proteins, responsive to different wavelengths of light, are introduced into a targeted group of neurons. The neurons are exposed alternatively to different wavelengths of light to excite, and inhibit the depolarization of the neurons.

More detailed embodiments expand on such techniques. For instance, another aspect of the present invention co-expresses GtR3 and DChR with other known opsins, such as NpHR and VChR1 in the species (*e.g.*, a mouse and C. elegans). Also, opsins creating currents of opposite polarity and having different color sensitivity are integrated with calcium imaging in acute mammalian brain slices for bidirectional optical modulation and readout of neural activity. Likewise, the coupled opsins can be targeted to C. elegans muscle and cholinergic motoneurons to control locomotion bidirectionally. Together coupled opsins can be used as a complete and complementary optogenetic system for multimodal, high-speed, genetically-targeted, all-optical interrogation of living neural circuits. Further, more than one set of cells may be targeted with different

couplings of opsins, allowing for precise control of multiple sets of cells independently from each other.

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In addition to GtR3 and DChR, there are a number of channelrhodopsins, halorhodopsins, and microbial opsins that can be engineered to optically regulate ion flux or second messengers within cells. Various embodiments of the invention include codon-optimized, mutated, truncated, fusion proteins, targeted versions, or otherwise modified versions of such ion optical regulators. For example, GtR3 and DChR (e.g., Appendices B and C as filed in the underlying provisional application). The cited opsins are used as representative of a number of different embodiments. Discussions specifically identifying GtR3 and DChR are not meant to limit the invention to such specific examples of optical regulators unless specified. For further details regarding the above mentioned sequences, reference can be made to "Multimodal fast optical interrogation of neural circuitry" by Feng Zhang, et al, Nature (April 5, 2007) Vol. 446: 633-639, which is fully incorporated herein by reference.

Table 1 shows identified opsins for inhibition of cellular activity across the visible spectrum:

Table 1: Fast optogenetics: inibition across the visible spectrum				
Opsin Type	Biological Origin	Wavelength Sensitivity	Defined action	
NpHR	natronomonas pharaonis	680nm utility (with 3.0 series) 589 nm max	Inhibition (hyperpolarization)	
BR	halobacterium helobium	570nm max	Inhibition (hyperpolarization)	
AR	Acetabulaira acetabulum	518nm max	Inhibition (hyperpolarization)	
GtR3	Guillardia theta	472nm max	Inhibition (hyperpolarization)	

Table 2 shows identified opsins for excitation and modulation across the visible spectrum:

Table 2: Fast optogenetics: excitation and modulation across the visible spectrum				
Opsin Type	Biological Origin	Wavelength Sensitivity	Defined action	
VChR1	Volvox carteri	589nm utility	Excitation	
		535nm max	(depolarization)	
optoXRs	Bos taurus	505nm max	Modulation (Gs, Gq pathways)	
DChR	Dunaliella salina	500nm max	Excitation (depolarization)	
SFOs	Chlamydomonas reinhardtii	546nm deactivate 470nm activate	modulation (electrical up states)	
ChR2	Chlamydomonas reinhardtii	470nm max 380-405nm utility	Excitation (depolarization)	

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Opsins described in U.S. Patent Application Nos.12/988,567 and 12/996,753, U.S. Patent Application Publication Nos: 2007/0054319, 2010/0234273, 2007/0261127, 2007/0053996, 2010/0145418, 2009/0093403, 2008/0085265, 2010/0190229, 2009/0099038, and PCT Publication No. PCT/US09/64355 are incorporated herein by reference in their entirety.

In certain embodiments of the present invention, various combinations of the listed opsins, for example, are delivered to a cell population. A second combination can be delivered to a different cell population. Using the defined action and the wavelength sensitivity information for each opsin, combinations of opsins may be chosen to excite certain cell populations while inhibiting others, for example.

Consistent with one example embodiment of the present invention, target cells are stimulated using an implantable arrangement. The implantable arrangement includes a biological portion that facilitates the stimulation of the target cells in response to receipt of light. The implantable arrangement also includes a light generator for creating light to trigger the stimulus of the target cells. In certain embodiments the implantable arrangement includes a microcontroller and a feedback loop. The color of light created by the light generator is dependent on the voltage of the cell. The target cell can include multiple light sensitive proteins which respond to different wavelengths of light to inhibit or encourage depolarization of the target cell.

Consistent with another example embodiment of the present invention, a method is implemented for stimulating target cells *in vivo* using gene transfer vectors (for example, viruses) capable of inducing photosensitive ion channel growth (for example, DChR ion channels) or photosensitive ion pumps (for example, GtR3 proton pumps). The vectors are implanted in the body, along with the electronic components of the apparatus. A light producing device is implanted near the target cells. The target cells are stimulated in response to light generated by the light producing device. In certain more specific embodiments, gene transfer vectors are capable of inducing growth of more than one ion channel or ion pump.

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Consistent with a particular embodiment of the present invention, a protein is introduced to one or more target cells. When introduced into a cell, the protein changes the potential of the cell in response to light having a certain frequency. This may result in a change in resting potential that can be used to control (dissuade) action potential firing. In a specific example, the protein is GtR3 which acts as a membrane pump for transferring charge across the cell membrane in response to light. Membrane pumps are energy transducers which use electromagnetic or chemical bond energy for translocation of ions across the membrane. In other specific examples, the protein is a halorhodopsin which acts as the membrane pump. The halorhodopsin membrane pump moves specific ions across the membrane. For further information regarding halorhodopsin membrane pumps, reference can be made to "Halorhodopsin Is a Light-driven Chloride Pump" by Brigitte Schobert *et al*, The Journal of Biological Chemistry Vol. 257, No. 17. September 10, 1982, pp. 10306-10313, which is fully incorporated herein by reference.

In specific embodiments GtR3 and NpHR may be introduced into different target cell populations, respectively. The two inhibitors are responsive to different wavelengths of light allowing the two proteins to be used to inhibit different target cell populations independently. GtR3 and NpHR may also be introduced into the same cell population for gradient control over the inhibition of a cell population. For instance, GtR3 and NpHR molecules within the same cell can be activated relatively independent from one another. Thus, a first level of inhibition can be implemented by activating the molecule type (GtR3 or NpHR) that provides the lowest current levels for the cell conditions. The next level could be implemented by deactivating this first molecule type and activating the other molecule type. A third level can be implemented by activating both molecules simultaneously. Further gradient control can be achieved by varying the wavelength of light, or by varying the light intensity at or near the maxima of each protein, for example.

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The combination of multiple inhibitors can also lead to shunting inhibition, which combined with hyperpolarizing inhibition can lead to stronger hyperpolarization in the combination approach than from using a single inhibitor protein. In a cell membrane the reversal potential (also known as the Nernst potential) of an ion is the membrane potential at which there is no net (overall) flow of ions from one side of the membrane to the other. Each type of ion channel has a specific reversal potential. At the Nernst potential the outward and inward rates of ion movement are the same (the ion flux is in equilibrium for the ion type). A change of membrane potential on either side of the Nernst potential reverses the overall direction of ion flux. If the reversal potential of an ion channel is below the resting potential (or instant potential in the case where a light responsive protein ion channel been stimulated and changed the "resting" potential) of the cell membrane, the ion channel will contribute to inhibition through hyperpolarization of the cell. In contract, if the reversal potential for an ion channel is between the resting potential and the threshold for the generation of action potentials, the ion channel will have a shunting effect. The concentration gradient for any given ion (for example Cl-) is determined in part by the balance between the activity of the other ions (for example Na+ and K+). Shunting inhibition is termed "shunting" because the channel conductance short-circuits currents that are generated at adjacent excitatory channels. Accordingly, the addition of a light responsive protein channel with inhibitory characteristics can lower the membrane potential to a point where an existing ion channel reverses direction and "shunts" the current, creating an additional path which prevents the membrane potential from reaching the action potential.

Provided herein are populations of cells, tissues, and non-human animals comprising a cell expressing one or more of the light-activated proteins described herein on the cell membrane. Methods of using the cells, population of cells, tissues, and non-human animals are also provided.

Figure 1A shows a light responsive ion-passing molecule (a pump or channel) 150 in a cell membrane 160. In general, ion channels allow flow toward the channel's reversal potential (or equilibrium potential for the channel); the direction is based on whether the membrane potential is above or below the channel's reversal potential. Ion channels do not require additional energy to transport the ion. Ion pumps transport ions against the equilibrium flow and require introduction of some form of energy, such as ATP, to transport the ion across the membrane. Ions 162 may be anions or cations. The hydrogen ion is a proton (as well as a cation). The direction the ions flow along with the

ions polarity will determine whether the ion-passing molecule has a hyperpolarizing effect or a depolarizing effect on the cell. In certain embodiments, the ion-passing molecule 150 transports protons out of the cell, causing hyperpolarization of the cell and inhibition of the firing of the cell. The light responsive ion channel/pump 150 is not open/active unless light of the appropriate frequency is present.

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Figure 1B shows an alternative embodiment of the present invention where a light responsive ion-passing molecule 150 is expressed within membranes that are internal to a cell. Many of the internal components of the cell have membranes similar to the cell membrane that encloses the cell. The membrane of organelle 152 contains naturally occurring ion pump/channels before introduction of a light responsive protein. Depending on the promoters introduced along with the exogenous light responsive protein, the light responsive protein can express as an ion pump/channel in the membrane of variety of different organelles, in the cell membrane, or a combination of different organelles and/or the cell membrane. In certain embodiments the light responsive ion-passing molecule 150 is a proton pump. In other embodiments the pump can be a pump/channel for a specific cation or a specific anion. The ion pump/channel 150 is located in the membrane 156 of the organelle.

The transfer of protons across a membrane in which the ion-passing molecule is expressed can be used to change the membrane potential of either or both membrane 156 and membrane 160. The transport of protons out of the cytoplasm can cause a cell to become hyperpolarized, resulting in inhibition of the cell.

In certain instances, the pump/channel 150 can be used to change the pH of the organelle 152 and the surrounding cytoplasm 154. In addition, the pH of the organelle in which the protein is expressed can be changed and/or controlled. Because most enzymes within a cell are pH sensitive, the pH of each organelle critically determines the coordinated bio-chemical reactions occurring along the endocytic and secretory pathways. Aberrations of the normal organellar pH homeostasis can lead to significant functional changes. Depending on the level of expression of the light responsive protein, and the location (*i.e.* which organelle) various functions of the cell may be enhanced or retarded. At high enough levels of expression the cell may be killed. This can be desirable in unwanted (*i.e.* cancer) cells. For instance, cells can be locally transfected at a cancerous tumor location and then carefully directed light pulses can be used to shrink or eliminate the tumor.

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Figure 2A shows two examples of the use of light responsive proteins, each example including three cell populations, A, B, and C. As discussed briefly above, two or more light responsive proteins can be introduced into the same cell populations. The light responsive proteins introduced can cause the same response or different response when activated. The different proteins can also have different reaction times, or strengths of reactions. Two proteins of the same polarity can be activated at the same time causing an additive effect. Two depolarizing proteins, for example, may be alternatively stimulated. This can allow for an increase in frequency of firing. In example 1, cell populations A and B express the same inhibiting protein, but different excitatory proteins. Cell population C has an excitatory protein which reacts to light around the same wavelength as the inhibiting protein of cell populations A and B. The inhibitor of cell population C reacts to a wavelength similar to the excitatory protein of cell population A. This arrangement allows for a variety of combinations of cell population reactions to the light provided to stimulate the cell populations. As example 1 is set up, cell population B may be excited without exciting or inhibiting either of the other two populations. However, cell populations A and C react in almost an inverse way to the light provided.

Example 2 includes cell population A with three light responsive proteins and cell populations B and C with two light responsive proteins. Cell population A includes two excitatory proteins which react to different light wavelengths, and one inhibitory protein. Cell populations B and C each have one of the excitatory proteins of cell population A and an inhibitory protein which responses to wavelengths in the same spectrum as the excitatory protein present in the other cell population (B or C). This arrangement allows for the combination of cell populations A and B to be excited, or cell populations B and C to be excited, while the third cell population is inhibited. It allows for cell population A to be inhibited without affecting either cell population B or C, and for cell population A to be excited while both cell populations B and C are inhibited. The example combination of Figure 2A are not meant to be limiting, but to illustrate some of the many combinations of light responsive proteins in both a single cell population, and the combinations across cell populations.

Consistent with another example embodiment of the present invention, target cells are neurons located in the brain of a mammal. The target cells are genetically modified to express photosensitive bio-molecular arrangement, for example, DChR ion channels. Light can then be used to stimulate the neurons. Depending upon a number of

factors, such as the location within the brain and the frequency and length of stimulation, different objectives can be achieved. For instance, current techniques for deep brain stimulus (DBS) use electrodes to apply a current directly to the targeted area of the brain. The frequency of the electrical stimulus is sometimes referred to as either low-frequency DBS or high-frequency DBS. Studies have suggested that high-frequency DBS inhibits the generation of impulses from the stimulated cells, while low-frequency DBS facilitates the generation of impulses from the stimulated cells. The frequencies that produce the effects of high-frequency of low-frequency DBS have also been shown to vary depending upon the specific area of the brain being stimulated. According to one example of high-frequency DBS, the neurons are stimulated using electrodes supplying current pulses at frequencies around 100Hz or more. Such a frequency has been shown to be effective in certain applications, as discussed further herein.

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In various cell populations, similar to those illustrated in Fig. 2A, two or more light responsive proteins may be present. Some (or all) of the two or more light responsive proteins can direct the same action (*i.e.*, excitation or inhibition) within the cell population. For example cell population A of example 2 includes two excitatory light responsive proteins, DChR and VChR1. Fig. 2B shows a stimulus profile for use with certain embodiments in which two or more types of light responsive proteins, for example DChR and VChR1, may be introduced into the same cell population. Each ion-passing molecule has a respective stimulus frequency limit, which can be partially responsive to environmental factors like pH. This limit reflects the recovery time necessary for the molecule to transition between active and inactive states. The use of two types of light-responsive proteins, each responding to different optical wavelengths, can allow for each type of protein to be controlled separately. The respective ion channels can be activated alternatively, allowing for an increased frequency of stimulation of the brain.

For instance, example 3 shows that light pulses (solid vertical lines) of a first wavelength and provided at a frequency F. This wavelength is selected to activate a first type of molecule (triangles). A second set of light pulses (dotted vertical lines) of a second wavelength can also be provided at a frequency F. This second set of light pulses can be used to activate a second type of molecule (squares). The resulting activation frequency for both cells is thus twice the frequency of activation for each individual ion-passing molecule. Of course, the frequencies need not be identical between the two types of molecules and can be varied over time.

In another implementation, shown in example 4, a first type of ion-passing molecule can be activated for a time period followed by activation of a second type of ion-passing molecule. This can be particularly useful for allowing each of the ion-passing molecules to be inactive for the time period during which the other ion-passing molecule is being activated. In certain instances, this can facilitate recovery and sustained use of the ion-passing molecules. Moreover, the different current types, densities and ion-passing capabilities can be considered when deciding on the specific stimulation profile to create the desired response.

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A specific example of DBS is used for the treatment of Parkinson's disease. In this application, DBS is often applied to the globus pallidus interna, or the subthalamic nucleus within a patient's brain. By implanting a biological arrangement that modifies the cells to respond to light, a light flashing light can be used in place of electrodes. Thus, the targeted neuron cells and external electrical signal need not be directly applied to the targeted cells. Moreover, light can often travel from its point of origin farther than electricity, thereby increasing the effective area relative to the stimulation source and only those neurons that have been photosensitized are stimulated.

As with the electrode-based DBS methods, one embodiment of the present invention can be implemented using high-frequency DBS to inhibit neuron generated impulses. While high-frequency DBS has been accomplished at frequencies around 100Hz, high-frequency DBS using various embodiments of the present invention may not necessarily require the same frequency. For instance, it may be possible to reproduce the inhibiting effects of high-frequency DBS at lower frequencies (*e.g.*, 50Hz) when using light activated techniques. For example, activation of the GtR3 pump intrinsically favors hyperpolarization and resistance to action potential generation. Various frequencies can be used depending upon the particular application (*e.g.*, the targeted portion of the brain and the desired effect), and the stimulation modality being applied.

Consistent with another example embodiment of the present invention, gene transfer vectors inducing the expression of photosensitive bio-molecules are used to target a specific type of cell. For instance, viral-based proteins (*e.g.*, lentiviruses, adenoassociated viruses or retroviruses) can be created to target specific types of cells, based upon the proteins that they uniquely express. The targeted cells are then infected by the viral-based gene-transfer proteins, and begin to produce a new type of ion channel (for example DChR), thereby becoming photosensitive. This can be particularly useful for stimulating the targeted cells without stimulating other cells that are in proximity to the

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targeted cells. For example, neurons of disparate length, diameter, chronaxie, other membrane properties, electrical insulation, neurotransmitter output, and overall function, lie in close proximity to one another, and thus, can be inadvertently stimulated when using electrodes to provide the stimulation of the neurons. For further details on the generation of viral vectors and the *in vivo* modification and stimulation of neural cells, reference may be made to U.S. Patent Application No. 11/459,636 filed on July 24, 2006, "An optical neural interface: in vivo control of rodent motor cortex with integrated fiber optic and optogenetic technology" by Alexander M. Aravanis, et al, Journal Neural Engineering 4 (2007) S143-S156, "Neural substrates of awakening probed with optogenetic control of hypocretin neurons" by Antoine R. Adamantidis, et al, Nature, (November 15, 2007) Vol. 450: 420-424, "Targeting and Readout Strategies for Fast Optical Neural Control In Vitro and In Vivo" by Viviana Gradinaru, et al, The Journal of Neuroscience, (December 26, 2007) 27(52):14231–14238, "Multimodal fast optical interrogation of neural circuitry" by Feng Zhang, et al, Nature (April 5, 2007) Vol. 446: 633-639, "Circuit-breakers: optical technologies for probing neural signals and systems" by Feng Zhang, et al, Nature Reviews Neuroscience (August 2007) Vol. 8: 577-581, which are each fully incorporated herein by reference.

A specific embodiment of the present invention employs an implantable arrangement for *in vivo* use. A light-emitting diode, laser or similar light source is included for generating light (as shown, for example, light generator 104 in FIG. 3). A biological portion that modifies target cells to include light responsive molecules which facilitate stimulation of the target cells in response to light generated by the light source.

Another embodiment of the present invention employs an arrangement for stimulating target cells using a photosensitive protein that allows the target cells to be stimulated in response to light. A biological delivery device, such as those discussed in connection with biological portion 204 of FIG. 4, is used for implanting vectors that modify the target cells to include the photosensitive protein. An implantation component, such as that discussed in connection with biological portion 204, the mesh of FIG. 7 or viral matrix of FIG. 8, is used for implanting a light generating device near the target cells. A control device, such as that discussed in connection control circuit 208, is used for activating the light generating device to generate light to be received by the target cells, thereby stimulating the target cells in response to the generated light.

Returning now to the figures, FIG. 3 shows a block diagram of a system for stimulating target cells, according to an example embodiment of the present invention.

Block 102 represents a location internal to an organism (e.g., a mammal), as shown by the *in vivo* designation. Light generator 104 is an implantable device that generates light *in vivo*. The photosensitive biological portion 106 affects the target cells such that generated light strikes causes stimulation of the target. In one instance, the light generator 104 is a small electronic device on the order of a few millimeters in size. The small size is particularly useful for minimizing the intrusiveness of the device and associated implantation procedure. In another instance, the light generator 104 may include a fiber optic device that can be used to transmit light from an external source to the target cells.

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In one embodiment of the present invention, the target cells are modified to contain light-activated proton pump/channel proteins. A specific example of such protein is GtR3, which is a product based upon the cryptophytes *Guillardia theta*. Characterization of the action spectra for GtR3 suggests that the absorption maxima are around 490nm. Another specific example is DChR from *Dunaliella salina*. The action maxima for DChR are around 500nm.

These light sensitive proteins can be implanted using a number of different methods. Example methods include, but are not limited to, the use of various delivery devices, such as gelatin capsules, liquid injections and the like. Such methods also include the use of stereotactic surgery techniques such as frames or computerized surgical navigation systems to implant or otherwise access areas of the body. For further details on delivery of such proteins, reference may be made to U.S. Patent Application No. 11/459,636 filed on July 24, 2006 and entitled "Light-Activated Cation Channel and Uses Thereof", which is fully incorporated herein by reference.

FIG. 4 shows a block diagram of an implantable device for stimulating target cells, according to an example embodiment of the present invention. The figure includes control circuit 208, light source 206, biological portion 204 and target cells 202. Biological portion 204 affects the target cells 202 such that the target cells are stimulated in response to light

In one embodiment of the present invention, biological portion 204 may be composed of

In one embodiment of the present invention, biological portion 204 may be composed of target cells 202 that have been modified to be photosensitive. In another embodiment of the present invention, biological portion 204 may contain biological elements such as gene transfer vectors, which cause target cells 202 to become sensitive to light. An example of this is lentiviruses carrying the gene for DChR expression. In this manner, the stimulation of target cells 202 can be controlled by the implantable device. For

example, the control circuit 208 can be arranged to respond to an external signal by activating, or deactivating light source 206, or by charging the battery that powers light source 206. In one instance, the external signal is electromagnetic radiation that is received by control circuit 208. For example, radio frequency (RF) signals can be transmitted by an external RF transmitter and received by control circuit 208. In another example, a magnetic field can be used to activate and/or power the control circuit.

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Control circuit 208 can be implemented using varying degrees of complexity. In one instance, the circuit is a simple coil that when exposed to a magnetic field generates a current. The current is then used to power light source 206. Such an implementation can be particularly useful for limiting the size and complexity as well as increasing the longevity of the device. In another instance, control circuit 208 can include an RF antenna. Optionally, a battery or similar power source, such as a capacitive element, can be used by control circuit 208. While charged, the power source allows the circuitry to continue to operate without need for concurrent energy delivery from outside the body. This can be particularly useful for providing precise control over the light emitted by light source 206 and for increased intensity of the emitted light. In one embodiment of the present invention, light source 206 is implemented using a light-emitting-diode (LED). LEDs have been proven to be useful for low power applications and also to have a relatively fast response to electrical signals.

In another embodiment of the present invention, biological portion 204 includes a gelatin or similar substance that contains gene transfer vectors which genetically code the target cells for photosensitivity. In one instance, the vectors are released once implanted into the body. This can be accomplished, for example, by using a containment material that allows the vectors to be released into aqueous solution (*e.g.*, using dehydrated or water soluble materials such as gelatins). The release of the vectors results in the target cells being modified such that they are simulated in response to light from light source 206.

In another embodiment of the present invention, the biological portion 204 includes a synthetic mesh that contains the photosensitive cells. In one instance, the cells are neurons that have been modified to be photosensitive. The synthetic mesh can be constructed so as to allow the dendrites and axons to pass through the mess without allowing the entire neuron (*e.g.*, the cell body) to pass. One example of such a mesh has pores that are on the order of 3-7 microns in diameter and is made from polyethylene

terephthalate. In another example embodiment, the biological portion 204 includes an injection mechanism as discussed in further detail herein.

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FIG. 5 shows a block diagram of an implantable device, according to an example embodiment of the present invention. The implantable device of FIG. 3 is responsive to a field magnetic. More specifically, an inductor constructed from windings 302 and core 304 generates a current/voltage in response to a magnetic field. The current is passed to control circuit 310 through conductive path 306. In response, control circuit 310 activates light source 312 using conductive path 308. Light source 312 illuminates biological portion 314 in order to stimulate the target cells. In one instance, biological portion 314 includes a gelatin, synthetic mesh or injection mechanism as discussed in further detail herein.

In one embodiment of the present invention, the control portion can be a simple electrical connection, resistive element, or can be removed completely. In such an embodiment, the intensity, duration and frequency of light generated would be directly controlled by the current generated from a magnetic field. This can be particularly useful for creating inexpensive, long lasting and small devices. An example of such an embodiment is discussed further in connection with FIG. 6A and FIG. 6B.

In another embodiment of the present invention, the control portion can be implemented as a more complex circuit. For instance the control circuit may include and otherwise implement different rectifier circuits, batteries, pulse timings, comparator circuits and the like. In a particular example, the control circuit includes an integrated circuit (IC) produced using CMOS or other processes. Integrated circuit technology allows for the use of a large number of circuit elements in a very small area, and thus, a relatively complex control circuit can be implemented for some applications.

In a particular embodiment of the present invention, the inductor (302 and 304 of FIG. 5) is a surface mount inductor, such as a 100uH inductor part number CF1008-103K supplied by Gowanda Electronics Corp. The light generating portion is a blue LED, such as LEDs in 0603 or 0805 package sizes. A particular example is a blue surface mount LED having part number SML0805, available from LEDtronics, Inc (Torrance, CA). Connective paths 306 and 308 can be implemented using various electrical conductors, such as conductive epoxies, tapes, solder or other adhesive materials. LEDs emitting light in the amber spectrum (as applicable to NpHR channels) and other spectrums are available through commercial sources including this same manufacturer.

FIG. 6A shows a block diagram of an implantable device, according to an example embodiment of the present invention. FIG. 6A shows an inductor comprising coils 402 and core 404 connected to LED 408 using conductive paths shown by 406. FIG. 6B shows a circuit diagram corresponding to the block diagram of FIG. 6A. Inductor 412 is connected in parallel to LED 410. Thus, current and voltage generated by changing a magnetic field seen at inductor 412 causes LED 410 to produce light. The frequency and strength of the changing magnetic field can be varied to produce the desired amount and periodicity of light from LED 410.

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FIG. 7A and FIG. 7B show a diagram of a mesh for containing photosensitive bio-molecules, according to an example embodiment of the present invention. Mesh 502 is constructed having holes 504 of a size that allows illumination to pass but is small enough to prevent cells 506 to pass. This allows for cells 506 to be implanted while still receiving light from a light generator.

In one embodiment of the present invention, the cells 506 are stem cells that are modified to be photosensitive. The stem cells are allowed to mature as shown by FIG. 7B. In a particular instance, the stem cells mature into neurons having a cell body 512, axons/dendrites 508 and 510. The neurons are genetically modified to be photosensitive. Holes 504 are on the order of 3-7 microns in diameter. This size allows some axons and dendrites to pass through holes 504, while preventing the cell body 512 to pass.

FIG. 8A and FIG. 8B show a diagram of a viral matrix, according to an example embodiment of the present invention. The viral matrix includes structure 602, which contains viral vectors 604. In one instance, structure 602 includes a gel or fluid substance that contains viral vectors 604 until they are implanted in a mammal 606. Once viral vectors 604 are released, they infect target cells 608 in the vicinity of the implanted viral matrix as shown by FIG. 8B. Infected target cell 610 becomes photosensitive, and thus, light can be used to control the stimulation of target cell 610.

According to one embodiment of the present invention, structure 602 is a gelatin that has been impregnated, or otherwise sealed with viral vectors 604 contained within the gelatin. When structure 602 is implanted, the gelatin is hydrated and or dissolved, thereby releasing viral vectors 604. Standard commercially available gelatin mix may be used, in addition to compounds such as Matrigel by BD Biosciences division of Becton Dickenson and Company (Franklin Lakes, NJ)

FIG. 9 shows a circuit diagram of a circuit that produces light in response to a magnetic field, according to an example embodiment of the present invention. FIG. 9

includes an input circuit 720 and an output circuit 730. Inductor 704 generates current in response to magnetic field 702. Due to properties of magnetic fields, the current produced by inductor 704 is an alternating current (AC) signal. Full-wave bridge rectifier 706 rectifies the AC signal and along with an RC circuit generates a relatively stable voltage from the AC signal. This generated voltage is responsive to magnetic field 702 and output circuit 730 which generates light when the generated voltage is at a sufficient level. More specifically, power from battery 708 is used to drive LED 710 in response to magnetic field 702. This is particularly useful for applications where the magnetic field 702 seen by inductor 704 is less powerful (*e.g.*, due to the *in vivo* location of inductor 704).

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FIG. 10A shows a circuit diagram of a circuit that produces light in response to RF signal 801, according to an example embodiment of the present invention. Antenna 802 is used to receive RF transmission 801 and convert the signal to electricity. The received transmission is rectified by diode 803 and further filtered by capacitor 805. In a one instance, diode 803 can be implemented using a diode having a low forward bias and fast switching capabilities, such as a Schottky diode.

In a particular embodiment of the present invention, RF transmission 801 contains a power component for charging battery 815 and a signal component for controlling LED 825. Capacitor 805 can be selected to separate these components for use by the circuit. For instance, the power component may be a relatively low-frequency, large-amplitude signal, while the signal component is a relatively high-frequency, small-amplitude signal. Capacitor 805 can be selected to filter the power component of the signal to create a corresponding voltage. The remaining high-frequency component of the RF transmission is added to this voltage. The power component of the transmission can then be used to charge the battery 815, and the signal component of the transmission is used to enable LED 825. The light generated by LED 825 triggers stimulus of the target cells 827.

Figure 10B illustrates an alternative embodiment radio-frequency energy accumulator, which charges a battery, which in turn, powers a digital pulse generator, which powers a LED. An electromagnetic signal 850 is received by loop antenna 852 generating a corresponding electrical signal. The voltage generated from loop antenna 852 is limited by the reverse bias voltage of the diodes 855 and 856 and stored in capacitor 854. In a particular instance these diodes have a low reverse bias voltage that is relatively precise, such as a Zener diode. Electromagnetic signal 850 is rectified via

diode rectifier bridge 858 and filtered by voltage regulator 859 to produce a DC voltage. The DC can be used to charge power source 860.

Battery 860 is coupled to the input of Schmidt trigger 865 through capacitor 862. Feedback from the output of the Schmidt trigger is provided through resistor 864 relative to the charge on capacitor 863. Accordingly, the frequency of the square-wave output of Schmidt trigger 865 is determined by the values of the resistor-capacitor network including capacitor 863 and resistor 864. Resistor 864 and capacitor 863 may be fixed or variable. The output of Schmidt trigger 865 is fed through digital inverter 867 which powers LED 866. Light from LED 866 is transmitted to light-sensitive neurons 868 relative to the frequency of the square-wave output of Schmidt trigger 865.

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Figure 10C illustrates block diagram for an electromagnetic field (EMF) energy accumulator and pulsing approach in which the received EMF 897 (for example radiofrequency energy) includes not only energy for accumulation, but also an encoded signal regarding instructions to microcontroller 895. In step 885 (Energy plus Parameter Control Signal: Encoding and transmission), a control instruction signal is encoded to ride upon the energy component by methods known in the art, for example, by frequency modulation. Energy receiver block 890 uses a portion of the EMF signal to provide power to block 893. Control signal receiver block 891 uses a portion of the EMF signal to provide control instructions to microcontroller block 895.

The control instruction can be used to transmit information regarding the various parameters of the generated light, such as frequency, strength, duration, color, and the like. These instructions can be decoded and processed using a microcontroller or logic circuitry as shown by block 895. Block 895 can generate control signal(s) in response to the decoded instructions. Accordingly, the frequency (and other parameters) of the light generated by LED 896 rate need not be fixed for the given implanted device. Antenna 889 delivers input to the Energy Receiver 890 (providing power to voltage regulator and battery circuitry 893). Concurrently, antenna 889 delivers encoded data to Control Signal Receiver 891, which provides control input to microcontroller 895 that drives LED 896. Selected wavelength light 897 is then delivered to electrically excitable cell 898. The battery in the voltage regulator and battery circuitry 893 provides power to the microcontroller 895 and the Control Signal Receiver 891.

The circuit diagrams of FIG. 9 and FIG. 10A, 10B and 10C are merely illustrative of a few particular embodiments of the present invention, and various other implementations are envisioned. For example, particular embodiments implement a light

source that uses a blue LED; however, other colors and light sources can be implemented depending upon the particular application. In other particular embodiments the light source includes more than one color light source. The circuitry controls not only when light is provided, but which color, depending on whether the instructions require inhibition or excitation of the cell. For example, in a specific example the target cell express both an inhibitor (e.g., GtR3) and an exciter (e.g., VChR1), which respond to different wavelengths of light. The system can also control the level at which the cells are excited or inhibited. This can be done by including multiple light responsive proteins in the target cells that excite (or inhibit) and programming the instructions to provide more than one light wavelength at a single time.

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FIG. 11A and FIG. 11B each show a diagram of a fiber-optic device, according to an example embodiment of the present invention. The fiber-optic device includes a control portion 908, a light generator 906 and a fiber optic cable 902.

Fiber optic cable 902 can be positioned near a photosensitive biological portion, such as a viral matrix or synthetic mesh as discussed herein. This allows for control portion 908 and light generator 906 to be located at a distance from the target cells 910 (e.g., at a distance corresponding to the length of fiber-optic cable 902). This can be particularly useful for minimizing the size of the portion of the implanted device that is near the target cells, for example, where the target cells are located at or near a sensitive location within the brain. In some instances, the remote location of portions 908 and 906 also facilitates modifications of the device, including, but not limited to, replacement of various components (e.g., batteries), changes in stimulation frequency and length.

Control portion 908 can be configured to respond to an external signal, such as magnetic field or RF signals. Alternatively, control portion 908 can be configured to enable light generator 906 according to a programmed schedule or a combination of an external signal and a programmed response.

FIGs. 12A-12D depicts various stages in the production of a photosensitive biological portion, according to an example embodiment of the present invention. More specifically, FIG. 12A shows molding structure 1004 having several molds 1002. Molds 1002 are constructed to various sizes depending upon the particular application. In one such application, the molds are a few millimeters or less in diameter.

FIG. 12B shows the molds 1002 from FIG. 12A after applying a layer of gelatin or similar substance as shown by 1006 and 1008. Moreover, viral vectors (shown by 'v') are in the upper two molds. These viruses may be suspended within media 1012, which

may be a liquid or gelatinous media. Such liquids include normal saline, HEPES-buffered saline and other known viral substances and transfer media. Suitable gelatinous media includes Matrigel (BD Biosciences, San Jose CA). These viral vectors are designed transfer genes for light-sensitization to the membranes of targeted cells after implantation.

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FIG. 12C shows a side view of mold 1006. 1016 represents the molding structure that forms the shape of gelatin layer 1014. Gelatin layer 1014 traps viral vectors contained within media 1012. A top gelatin layer 1010 is applied to fully contain the viral vectors.

FIG. 12D shows the resulting viral vector capsule. The viral vectors 1018 are contained within area 1022 by casing 1020. Casing 1020 can be designed to dissolve or otherwise allow viral vectors 1018 to disseminate towards the target cells once implanted. In one instance, the capsule is constructed of a water soluble material, for example, gelatin, so that upon implantation the viral vectors are allowed to escape into the body. Water soluble capsule materials are well known in the pharmaceutical industry.

FIG.13 shows an implantation device, according to an example embodiment of the present invention. Biological portion 1102 and light generation device 1108 are implanted using the implantation device. For example, the shaft of the device 1114 is positioned near the target cells. Next, a user of the device presses on portion 1116 which causes portion 1112 to place biological portion 1102 and light generation device 1108 near the target cells. The implantation device can then be removed.

FIG. 14A and FIG. 14B show a diagram for another implantation device, according to an example embodiment of the present invention. Implantable light generating device 1204 is surrounded by, and permeated by fluid channels 1202. Fluid channels 1202 allow a solution 1210 containing bio-molecular material (*e.g.*, photosensitizing viral vectors) to be injected immediately proximal to light generating device 1204 and the target cells. The fluid channels can be located outside of device 1204 and/or within device 1204, as shown by 1212 and 1214 respectively. In this manner, the viral vectors can be injected in large quantities or over a period of time. For instance, cells infected by viral vectors can revert back to their pre-infection state after a period of time. Using the device of FIG. 12A, the viral vectors can be periodically reintroduced to the target cells. Alternatively, different viral vectors can be introduced through the fluid channels, allowing for targeting of different cells at the implantation

site. This can be particularly useful for staged treatment through stimulation of different types of cells.

A specific embodiment of the present invention relates to a method for genetically modifying neurons to express light-sensitive ion channel DChR. In this method, pulses of blue light causes DChR neurons to fire action potentials corresponding to each pulse. Depolarization and repolarization occur on a millisecond timescale making this method consistent with normal network neurophysiology.

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Specific targeted neurons are modified using viral vectors for gene transfer. For further details on the generation of viral vectors, reference can be made to Boyden *et al.* 2005, Zhang *et al.* 2006 entitled "Channelrhodopsin-2 and Optical Control of Excitable Cells," *Nature Methods* Vol. 3, No. 10, which is fully incorporated herein by reference. This transfection results in the introduction of a gene for a single protein, a cell membrane ion channel, known as "DChR". In nature, DChR resides on the cellular membrane of *Dunaliella salina*. Upon absorption of blue/green light (500nm), this ion channel briefly opens, allowing cation influx. When transfected into a mammalian nerve cell, affected nerves become photosensitive, producing light-triggered action potentials.

A neuronal-type specific feature which is also a robust promoter is inserted adjacent to the DChR code within the virus, and the line is propagated by calcium-phosphate cotransfection of 293FT cells. The supernatant is then centrifuged into viral pellets, which are placed within phosphate-buffered saline.

In a particular instance, application of a DChR is used for photostimulation. The amino-acid residues comprising DChR channelrhodopsin from *Dunaliella salina* can be used to impart fast photosensitivity upon mammalian nerve cells, by using a viral vector to insert the gene for DChR into targeted nerve cells which may subsequently express this gene. Upon illumination with approximately 500 nm blue/green light, ATR isomerizes and triggers a conformational change to open the channel pore. As DChR is a light-sensitive ion channel, it allows an inward current to be evoked.

In another instance, application GtR3 (derived from *Guillardia theta*) is used for photostimulation. This proton pump can be imparted upon mammalian nerve cells by using a viral vector to insert the gene for GtR3 into targeted nerve cells, which may subsequently express this gene. Upon illumination with approximately 472 nm blue light, active pumping of protons out of the neuronal cytoplasm results in hyperpolarization of the cell.

Since sensitivity to blue light via DChR or GtR3 is induced when a viral vector inserts the respective gene into a previously normal cell, the insertion may be genetically targeted to the products expressed by specific cellular subtypes. For example, it might be advantageous to cause only dopaminergic neurons, and not cholinergic neurons to react to blue light.

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As discussed above, certain embodiments of the present invention involves the use of an optically responsive ion pump or ion channel that is expressed in a cell. In a particular instance, the cell is either a neural cell or a stem cell. A specific embodiment involves *in vivo* animal cells expressing an ion pump. Certain aspects of the present invention are based on the identification and development of a proton pump, derived from *Guillardia Theta*, for example, for temporally-precise optical inhibition of neural activity. The pump allows both knockout of single action potentials within rapid spike trains and sustained blockade of spiking over many minutes.

According to an example embodiment of the present invention, an optically responsive ion pump and/or channel is expressed in one or more stem cells, progenitor cells, or progeny of stem or progenitor cells. Optical stimulation is used to activate expressed pumps/channels. The activation can be used to control the proton concentration in the cells. It may also be used to control the pH of particular subcellular organelles. This can be particularly useful for affecting the survival, proliferation, differentiation, de-differentiation, or lack of differentiation in the cells. Thus, optical stimulus is implemented to provide control over the (maturation) of stem or progenitor cells.

According to other example embodiments of the present invention, methods for generating an inhibitory neuron-current flow involve, in a neuron, engineering a protein that responds to light by producing an inhibitory current to dissuade depolarization of the neuron. In one such method, the protein is GtR3 is an exogenous molecule in the neuron, and in another method the protein is an inhibitory protein that uses an endogenous cofactor.

In another example embodiment, a method for controlling action potential of a neuron involves the following step: engineering a first light responsive protein in the neuron; producing, in response to light, an inhibitory current in the neuron and from the first light responsive protein; engineering a second light responsive protein in the neuron; and producing, in response to light, an excitation current in the neuron from the second light responsive protein. The combination of light responsive proteins may depend on the

light sensitivity of the protein, the reaction time of the protein, and defined action of the protein, for example. Multiple light responsive proteins of the same type (inhibitory vs. excitatory) may be used in the same cell population to allow for, for example, long term blockage of action potential or single action potential inhibition depending on the light wavelength used.

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In another method for controlling a voltage level across a cell membrane of a cell, the method comprises: engineering a first light responsive protein in the cell; measuring the voltage level across the cell membrane; and producing, in response to light of a first wavelength and using the first light responsive protein, a current across the cell membrane that is responsive to the measured voltage level.

Another aspect of the present invention is directed to a system for controlling an action potential of a neuron *in vivo*. The system includes a delivery device, a light source, and a control device. The delivery device introduces a light responsive protein to the neuron, with the light responsive protein producing an inhibitory current. The light source generates light for stimulating the light responsive protein, and the control device controls the generation of light by the light source. In a particular embodiment the light source is a blue light source and the light responsive protein is responsive to blue light.

In more detailed embodiments, such a system is further adapted such that the delivery device introduces the light responsive protein by one of transfection, transduction and microinjection, and/or such that the light source introduces light to the neuron via one of an implantable light generator and fiber-optics.

Another aspect of the present invention is directed to a method for treatment of a disorder. The method targets a group of neurons associated with the disorder; and in this group, the method includes engineering an inhibitory proteins that use an endogenous cofactor to respond to light by producing an inhibitory current to dissuade depolarization of the neurons, and exposing the neurons to light, thereby dissuading depolarization of the neurons.

In another aspect of the present invention, GtR3 is optimized for mammalian expression. A light responsive protein is isolated from *Guillardia theta*. The isolated protein has a C-terminus and an N-terminus. An endoplasmic reticulum (ER) export signal promoter is added to the C-terminus of the isolated protein, creating an enhanced light responsive protein. In various embodiments the promoter added to the isolated GtR3 varies depending on the desired location of expression.

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FIG. 15 depicts an arrangement with multiple light sources, according to an example embodiment of the present invention. Fig. 15 shows light sources 1602 and 1604 that illuminate proteins 1610 and 1614. The proteins 1610 and 1614 are engineered within cell 1612 to control current across the cell membrane in response to light from light sources 1602 and 1604, respectively. In one instance, the first protein 1610 functions to dissuade action potential firing, while the second protein 1614 functions to encourage action potential firing. Each of proteins 1610 and 1614 are responsive to light. In a particular instance, the first protein is responsive to light from light source 1602 having a wavelength A and the second protein is responsive to light from light source 1604 having a wavelength B. Thus, the light sources can be used to control each protein independently. This can be useful for both encouraging and dissuading action potentials in the cell. In another instance, having both types of proteins allows for both positive and negative control of the cell membrane voltage. Thus, the different light sources and proteins could be used to control the voltage or current level (*e.g.*, clamping) of the cell membrane.

One method of determining responsiveness involves quantifying the responsiveness in terms of the intensity of light required to produce a given response. In some instances, the first or second protein can still be responsive to the alternate wavelength of light although the responsiveness of the protein may be less than that of the primary wavelength. Accordingly, a protein of a first type may have some responsiveness to the wavelength corresponding to the other type of protein while still maintaining sufficient independence of operation. In one such instance, control of the cell can be implemented by shifting either the wavelength of light or the intensity of the light. For instance, the wavelength can be shifted between A and B to induce a corresponding increase or decrease of the membrane voltage potential. Alternatively, multiple proteins having similar actions, but different activation wavelength may be introduced into the same cell. The response may vary as the wavelength is shifted, in some places the combination of the two responses create a greater response than from an individual protein.

According to one embodiment of the present invention, pump 1614 can optionally be implemented for purposes other than dissuading action potential firing, such as controlling the voltage level of cell 1612. More specifically, a sensor can be used provide feedback to the light source 1602. For instance, this feedback could be a measurement of the voltage or current across the cell membrane. Thus, the light source

could be configured to maintain a constant current or voltage (e.g., clamp) across the cell. Moreover, the amount of responsiveness can be controlled by modifying one or more of the intensity and wavelength of the light.

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FIG. 16 shows a system for controlling electrical properties of one or more cells in vivo, according to an example embodiment of the present invention. Control/Interface unit 1702 enables/disables light source 1704 to illuminate target cells 1708. A delivery mechanism, such as fiber optic cable 1706, routes or otherwise directs the light to target cells 1708. Fiber optic cable 1706 may include a bundle of optical cables, each capable of carrying and directing light independently. Thus, fiber optic cable 1706 can be configured to deliver light having one or more wavelengths to multiple locations. Sensor 1710 can be implemented *e.g.*, as an optical device such as an optical scope or as a voltmeter, to provide feedback to control unit 1702. In a particular instance, the feedback includes optical imaging of the target cells or of other related cells. In another instance, the feedback could monitor the voltage response of the target cells, including the amount of action potential firing.

Fig. 17 shows a system for controlling electrical properties of one or more cells in vivo, according to an example embodiment of the present invention. Control/Interface unit 1802 enables/disables implantable light source 1804, which in turn illuminates target cells 1806. Light source 1804 is shown with two light source, inhibitory current light source 1808 and excitation current light source 1810. Light source 1808 produces light at a wavelength and intensity that an inhibitory protein is responsive to, while light source 1810 produces light at a wavelength and intensity that an excitation protein is responsive to. One skilled in the art would recognize that various configurations of light source 1810 are possible, including a single inhibitory light source or an array of light sources having one or more wavelengths. Control/Interface unit 1802 communicates with light source 1804 through any suitable communication mechanisms, such as wired communications or wireless communications using radio-frequency signals, magnetic signals and the like. As discussed above in connection with Fig. 16, sensor 1812 can optionally be implemented for providing feedback to control unit 1802.

Certain embodiments of the present invention can be useful in drug screening. The various light-sensitive proteins, serving to regulate membrane voltage using ion switches that, when activated (or deactivated) in response to light, function as channels or pumps and are referred to hereafter as light-responsive ion switches or light-activated membrane potential switches (LAMPS).

Consistent with one example embodiment of the present invention, a system screens for ion-channel and ion-pump affecting compounds. The system introduces one or more drug candidates that could either block or enhance the activity of ion-channels or ion-pumps to cells that were made optically responsive by the addition of a combination of the above mentioned proteins (ChR2, DChR and NpHR among others), for the purpose of screening the drug candidates. Light triggers optically responsive ion channels in the cells causing a change in the voltage seen across the cell membrane. The voltage change stimulates voltage-gated ion channels in the cells which will then cause a change in ion concentrations that can be read as optical outputs. These optical signals are detected and used to determine what effect, if any, the drug candidates have on the voltage-gated ion channels. In a more specific embodiment a protein expressing a proton pump is introduced into the cell.

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In one instance, the system allows for different drug candidates to be screened without necessitating extensive setup between screenings. For example, an assay may be performed using optics both to stimulate the optically responsive cells and to detect the effectiveness of the drug. The use of optics instead of manual contacts, *e.g.*, using a whole-cell patch clamp, can be particularly useful in increasing the throughput of the assay screening process. For instance, the time between screenings can be reduced by minimizing or eliminating physical manipulations otherwise necessary to stimulate or detect ion flow in the target cells. The cells can also be prepared prior to the screening process because the test equipment need only be optically coupled to the prepared cells. In another instance, throughput may be increased by screening a number of different drugs simultaneously using, for example, an array of photo detectors and a corresponding array of modified cells exposed to different drugs.

A cell line based approach is not limited to a particular ion channel. For example, cell lines can be created for voltage-gated sodium (*e.g.*, Na_v1.1 through Na_v1.9), potassium (*e.g.*, K_v such as hERG, TASK1, Shaker, or KvLQT1), or chloride conducting channels/pumps (*e.g.*, members of the CLC family of chloride channels). The methods of introducing such genes into the cell line are known in the art and may include, for example liposomal transfection, or viral gene transfer. For further information in this regard, reference may be made to one or more of the following references:

- <u>Warren Pear,</u> Transient Transfection Methods for Preparation of High-Titer Retroviral Supernatants, Supplement 68, Current Protocols in Molecular Biology, 9.11.1-9.11.18, John Wiley & Sons, Inc. (1996).
- R.E. Kingston, C.A. Chen, H. Okayama, and J.K. Rose, *Transfection of DNA into Eukarotic Cells*. Supplement 63, *Current Protocols in Molecular Biology*, 9.1.1-9.1.11, John Wiley & Sons, Inc. (1996).
- R. Mortensen, J.D. Chesnut, J.P. Hoeffler, and R.E. Kingston, Selection of Transfected Mammalian Cells, Supplement 62, Current Protocols in Molecular Biology, 9.5.1-09.5.19, John Wiley & Sons, Inc. (1997).
- H. Potter, Transfection by Electroporation, Supplement 62, Current Protocols in Molecular Biology, 9.3.1-9.3.6, John Wiley & Sons, Inc. (1996).
- T. Gulick, Transfection using DEAE-Dextran, Supplement 40, Current Protocols in Molecular Biology, 9.2.1-9.2.10, John Wiley & Sons, Inc. (1997).
- R.E. Kingston, C.A. Chen, H. Okayama, *Transfection and Expression of Cloned DNA*, Supplement 31, Current Protocols in Immunology (CPI), 10.13.1-10.13.9, John Wiley & Sons, Inc.

Each of the above references is incorporated by reference in its entirety.

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These and other transfer vectors may be generated using various genetic engineering techniques. For instance, the transfer vectors may be derived from a provirus clone of a retrovirus, such as an immunodeficiency virus (*e.g.*, HIV-1 or HIV-2, or SIV). For further details on the use of 293T cells and transfection thereof, reference can be made to U.S. Pat. No. 6,790,657 (entitled, *Lentivirus Vector System*, to Arya), which is fully incorporated herein by reference.

In one embodiment of the invention, optical stimulation of the modified cells may be altered to determine specific properties of an introduced drug candidate. For example, the intensity of the optical stimulus may be modified to change the corresponding level of depolarization. The level of desired depolarization can be tuned to further characterize the effectiveness of the drug under test. In another example, the optical stimulus may include rapid pulsing of the light. By correlating the temporal relationship between the optical stimulus and the resultant detected fluorescence, the drug may be further characterized in terms of a kinetic response. Thus, the drug may be characterized for a variety of different aspects including, but not limited to, the steady state effect on ion concentrations, a change in the level of depolarization necessary to trigger the voltage gated ion channels and the effect on repeated depolarization.

In one embodiment, the system allows for simple calibration of the optical stimulation and/or detection. The modified cells may be optically stimulated prior to introduction of the drug candidate. The ion channel responsiveness is detected and recorded. The recorded values may be used as a baseline for comparison to the ion channel responsiveness of the same modified cells after the introduction of the drug under test. The recorded values may also be used to modify the optical stimulus or the sensitivity of the optical detector. Such modifications may be applied to an individual test sample or an array of test samples. For such an array of test samples, each test sample may be individually calibrated by adjusting the corresponding optical stimulus. Similarly, each corresponding photo detector may be individually adjusted.

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FIG. 18A shows a basic block diagram of a system for screening for ion-channel affecting drugs, according to an embodiment of the invention. Optical control 1904 communicates with database 1902, optical source 1906 and optical detector 1909. Optical source 1906 provides optical stimulus to test sample 1908. Test sample 1908 includes the drug under test, cells with optically responsive ion channels, and a voltage/ion indicator. In one instance, the indicator fluoresces in response to light from optical source 1906. Optical control 1904 may also include a reconfigurable readout, so that as different LAMPS and different LEIAs are used, the same control system can be readily adapted to each paradigm. Optical detector 1909 produces a signal responsive to such florescence, and optical control 1904 receives the produced signal. The optical control 1904 stores data obtained from the signal in database 1902. The information stored may include factors such as the intensity, duration and wavelength of the detected light. In a particular instance, the stored data can be compared against baseline data, where the baseline data corresponds to data recorded prior to the introduction of the drug to the test sample 1908. In another instance, optical source 1906 may vary the intensity, duration or other parameters related to the control of optical source 1906. These and other parameters may be stored in database 1902.

It should be apparent that optical source 1906 may be implemented using a single light source, such as a light-emitting diode (LED), or using several light sources. Similarly, optical detector 1909 may use one or more detectors and database 1902 may be implemented using any number of suitable storage devices.

FIG. 18B shows a system diagram of a large-format, quasi-automated system for drug screening in accordance with a specific embodiment of the invention. Control device 1901 (e.g., a computer or control logic) controls various processes, and serves as

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the central point of system input/output functions. The environment may be maintained at an appropriate temperature, humidity, carbon dioxide level and ambient light level within the walls of the climate control chamber 1905, with the help of one or more sensors 1914 (e.g., thermostat, carbon dioxide sensor and humidity sensor), carbon dioxide and humidifier apparatus 1912, and heater 1910. Multi-well tray 1941 contains test wells 1940 for holding cultured cells, drugs, and other ingredients needed for each test. Tray 1941 rests upon X-Y-Z table 1925, the movement of which is carried out by table actuators 1920, under control of computer 1901. Xenon lamp 1955 emits highintensity white light 1956, which is passed through color filter 1960. In the case that DChR is used for stimulating the cells within wells 1940, color filter 1960 is blue, causing blue light 1961 to exit the filter, and strike dichroic mirror 1970. Blue light 1961 then passes upward, through microscope objective lens apparatus 1930, and through bottom of transparent tray 1941. In this fashion, the contents of wells 1940, with their transparent undersides, are illuminated. When a separate wavelength of light is required to stimulate a fluorescent light-emitting indicator of cellular activity, a filter of the appropriate specification may be substituted for the previous filter 160, causing light of the proper wavelength for this latter task to be piped toward well 1940. If the cells within well 1940 have been light-sensitized, and if the drug being tested in each of these wells does not suppress the process, a light-emitting indicator of cellular activity (LEIA), which has also been added to each well or expressed by the cells via genetic modification, will emit light in accordance with the voltage change caused by the effect of the light. This second wavelength of light, which may be much smaller in magnitude than the stimulation light, is collected by microscope turret 1935, and will also be passed through dichroic mirror 1975, onto the lens of (CCD) camera 1980.

Dichroic mirror 1970 allows for upward reflection of both the wavelength required to stimulate the optical gating of the membrane (e.g., blue-green for DChR), and the wavelength required by any LEIA used (e.g., ultraviolet for FURA-2). This dichroic mirror may be arranged to allow passage of the output spectrum of the LEIA (e.g., blue-green for FURA-2) with minimal reflection or absorption.

FIG. 19 is a system diagram of an automated-drug-screening system, according to an example embodiment of the invention. Emitter/detector units 2050 make up the emitter/detector array 2051. Emitter/detector array 2051 matches the number, size, and layout of the wells on tray 2040. Tray holding device 2025 permits tray swapping mechanism 2020 to rapidly move a new tray into position once testing of a given tray has

been completed. The entire process may be automated, and under the control of device 2001. Device 2001 can be implemented using a computer, control logic, programmable logic arrays, discreet logic and the like. The introduction of the drug candidates under test can also be automated using a machine that provides a reservoir for storing the drugs and a dispensing nozzle for injecting the drugs into the tray. In a manner similar to that shown by FIG. 18, the environment within the walls of the climate control chamber 2005 may be maintained at an appropriate temperature, humidity, carbon dioxide level and ambient light level, with the help of thermostat, carbon dioxide sensor and humidity sensor 2014, carbon dioxide and humidifier apparatus 2012, and heater 2010. The use of multiple stimulator/detector elements simultaneously and in parallel, can be particularly useful for augmenting the speed of the overall process. Low cost elements may be used to make multiple parallel detectors (*e.g.*, the components detailed below in description of FIG. 20A and 20B); the multiple parallel emitter/detector units may also be quite economically feasible.

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FIG. 20A depicts the workings of emitter/detector units, such as those shown in FIG. 19, according to an example embodiment of the invention. An LED stimulates light-sensitive ion channels of cells located within a well, and a photodiode detects the response of a LEIA. In this embodiment, device 2101 includes LED 2110, which produces light pulses 2111, at the proper wavelength, pulse frequency and intensity, so as to stimulate light-sensitive transgenic cells 2105 in culture within well 2106. Due to the presences of an LEIA (*e.g.*, a voltage-sensitive dye or a calcium dye), light 2116 is returned from cells 2105, and is detected by photodiode 2115. In the case that RH 1691 being used, red light is fluoresced and detected by photodiode 2115. In the absence of cellular depolarization, no fluorescence is detected by photodiode 2115. Other light detecting technologies may also be used instead of a photodiode including phototransistors, and CCD elements.

The combination of photostimulation with optical imaging techniques of LEIAs may be useful for a number of different reasons. For example, photostimulation may simplify the study of excitable cells by reducing the need to use mechanical electrodes for stimulation. Several commercially available LEIAs are suitable for photogrammetrically indicating the activation of electrically excitable cells. One such LEIA is calcium dye Fura-2, which may be stimulated with violet/ultraviolet light around 340nm, and whose fluorescent output is detectable as blue-green light around 535 nm. Another example is voltage sensitive dye RH 1691, which may be stimulated with green

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light at about 550nm, and whose fluorescent output is detectable as red light at about 70 nm. Another example is voltage sensitive dye di-4-ANEPPS, which is stimulated by blue light at about 560nm, and whose fluorescent output is detectable as red light at about 640nm.

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FIG. 20B depicts the workings of another embodiment of the emitter/detector units shown in the FIG. 19, in which multiple effects are tested within the context of a single well. For example, the cells 2155 in the wells 2156 may express both GtR3 and VChR1, and hence be sensitive to both the hyperpolarizing effects of blue light, and the depolarizing effects of amber light. Device 2151 includes LED 2160, which is used for the stimulation of the targeted proton pump (e.g., GtR3) of light-sensitive transgenic cells 2155. Additional LED 2175 may be used to stimulate a second targeted ion channel or pump (e.g., VChR1). Yet another LED 2180 may be used to stimulate a voltage sensitive dye (e.g., RH1691 or calcium dye, such as Fura-2). Each LED may be arranged to output specific wavelengths and intensities for stimulus of respective targeted compounds. In one instance, an LED may affect more than one target, depending upon the specific sensitivities of each compound used. Photodiode 2165 detects the fluorescence of a selected voltage dye, while photodiode 2170 is sensitive to the spectrum fluoresced by a selected calcium dye. The use of multiple LEDs for the same cell allows for the stimulation of LEIAs at different wavelengths. Multiple LEDs may also be used to detect different light wavelengths emitted by the LEIA.

FIG. 21A depicts an electronic circuit mechanism for activating the LED emitters used within the emitter/detector units, according to an example embodiment of the invention. Control device 2201 generates a "light on signal" 2202 to transistor base 2205. This light on signal 2202 will remain on for the duration of a light flash desired, or alternatively may turn on and off in order to produce rhythmic light flashes at a specified frequency. Light on signal 2202 permits (conventional) current to flow from power source 2210, through resister 2211, and through transistor collector 2207 and transistor emitter 2212, to ground 2213. Current is also thereby permitted to pass through resistor 2215, and into LED 2220. LED 2220 emits light 2221, which falls upon well 2225. In a particular instance, the transistor functions as transconductance amplifier of signal 2202. In this manner, light of the appropriate wavelength, intensity and frequency is delivered to cells within the well 2225, so as to cause them to stimulate the particular channel (e.g., DChR) or pump (e.g., GtR3), or other photoactive membrane structure being used to regulate the activity of electrically excitable cells. Various other

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circuits are also possible. For example, other circuits can be used in place of circuit 2206 to control LED 2220 including, but not limited to, replacing the transistor with an operational amplifier, a field-effect-transistor, a resistor divider network, transistor-transistor logic, push-pull driver circuits and switches.

FIG. 21B depicts an example electronic circuit mechanism for light detection by the emitter/detector units, according to one embodiment of the invention. Control device 2250 may (optionally, depending upon specific implementation) provide power to photodiode 2255. Photodiode 2255 receives fluoresced (emitted) light 2256 from the LEIA on the cells within well 2257. The received light results in an output signal. This output passes through resistor 2260, and is input to Schmitt triggered hex inverter 2270, which conditions the signal, providing a clean "high" or "low value" to be input to computer 2250.

Operation of the photodetector is shown in photovoltaic mode, but the element may also be used in the photoconductive mode of operation. Of course, many other light- detection devices and methods may also be used, including phototransistors, photothyristors, and charged-coupled device (CCD) elements, or arrays of elements.

Alternatively, the circuit of FIG. 21B can be used without Schmitt-triggered hex inverter 2270, permitting a continuum of signal intensities to be transmitted directly to an analog input to computer 2250 or to an analog-to-digital converter. Various other signal conditioning circuits are also possible.

FIG. 22 shows a sequence of steps using the embodiment shown in FIGs. 19, 20 and 21, in the context of projected high-throughput process time course 2300 and in accordance with one embodiment of the invention. In step 2305, light of the appropriate wavelength and intensity for the targeted ion channel is flashed-in this case for approximately three seconds. Concurrently, a LEIA stimulation flash 2310 may optionally be triggered, depending upon the specific voltage or calcium dye, etc. being used. This LEIA compound may have been previously added to the well, or may be (artificially) genetically imparted upon the cells such that the chemical is produced/expressed by the cells. In step 2315, the light signal produced by the LEIA is detected by the photodetector element (*e.g.*, photodiode). For example, RH1691, fluoresces red light at about 70 nm.

In step 2320, the signal resulting from the impingement of light onto the photodetector element is sent back to the computer. This may be a binary (*e.g.*, "high" versus "low" signal intensity), or may be graded to reflect a continuum of activation

levels. In the case that multiple photodetectors are used to determine energies at different wavelengths, the individual readings of these photodetectors may be logged in parallel or in sequence for appropriate interpretation in a later stage of the automated process. In step 2330, the system calls for the next tray to be placed by the automated system. The next tray is moved into position at step 2335 and the process may be repeated until all trays in a batch have been processed.

The amount of time allotted for light delivery may vary, and depends on factors

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including the level of light-gated proton or ion channel/pump expression, and the density and characteristics of other proton/ionic channel characteristics of that cell population. The amount of time allotted for light receipt may vary, and depends upon factors including the degree of accuracy required for the screening session. The amount of time allotted for well-plate (tray) changing may vary, and depends upon factors including the mechanical speed of the automated apparatus. If fast neurons are used as the cells being tested, the cellular stimulation and LEIA detection process may be accomplished in milliseconds.

The process above may be repeated under varying conditions. For example, a given set of cells may be tested with no drug present, and subsequently with one or more drugs present. The response of electrically-excitable cells under those conditions may be thereby documented, compared and studied. If the invention is implemented with at least one emitter/detector for each well on a tray and at least two concurrently operating devices, continuous operation may be maintained for extended periods of time.

FIG. 23 illustrates an example of a layout of cell and drug samples within the wells of a well-plate which is suitable for use within an embodiment of the invention. In this figure, well-plate 2401 (also referred to herein as a "tray" contains wells 2405 (examples), which are organized into columns 2425, labeled with numbers 1-12 and rows 2420, labeled with letters A-H. More specifically, an example column and row are defined by 2410 and 2415 respectively.

As an example of a functional layout of contents introduced into these wells, rows A-H of a single plate might be used for the testing of two different drugs. To represent a baseline condition, column 1 might contain optically gated cells, an endogenous or exogenous LEIA, but no drug. Columns 2-6 might be used for five different concentrations of Drug X, one concentration level per column. Likewise, columns 7-11 might be use for five different concentrations of Drug Y, one

concentration per column. Column 12, while fully usable, is left unused in this particular example.

Variables in the various wells might include the type of cell being tested, the type of ion channel being tested for, the type of drug placed in the cell, the concentration of the drug placed in the well, the specific LEIA used, and the optical gating stimulation parameters (*e.g.*, wavelength, intensity, frequency, duration) applied to the cells in that well.

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FIG. 24 illustrates the context in which the disclosed invention may be employed within a larger system which facilitates high-throughput drug screening. Well-plate 2506 contains wells 2505. These are carried forward by conveyer 2520, which may be a device such as a conveyor belt, robotic transporter or other delivery mechanism. Pipettes 2510 are held in array by robotic member 2515, and serve to inject the proper number of cultured cells and media into wells 2505. Subsequently, well-plate 2506 is moved down conveyer 2520, where robotic member 2525, analogous to robotic member 2515 and also containing pipettes, injects the proper amount of a LEIA into wells 2505. Conveyer 2520 then brings well-plate 2505 into screening chamber 2530. An emitter/detector apparatus, such as those described in connection with FIG. 17, FIG. 18A, FIG. 18B, FIG. 19A, and FIG. 19B, is located within chamber 2530. Additionally, portions of the processes described in FIG. 20A and FIG. 20B may occur within this chamber. Subsequently, well-plates 2535 is moved out of screening chamber 2530 by conveyor 2540, and discarded at 2545. In an alternative embodiment, one or more robotic devices may move pipettes 2510, screening chamber 2530, etc. to the locations of well-plate 2506, rather than vice-versa.

Consistent with the above discussion, example screening methods could include the collection of multiple data points without having to switch samples. Because control over the samples is reversible in the same sample preparation by simply turning the activating light on and off with fast shutters, the same samples can be reused. Further, a range of patterns of stimulation can be provided to the same cell sample so that testing can be performed for the effect of drugs without concern with regards to differences across different sample preparations. By modulating the level of excitation (*e.g.*, by ramping the level from no light to a high or maximum intensity), the effect of the drug across a range of membrane potentials can be tested. This permits for the identification of drugs that are efficacious during hyperpolarized, natural, or depolarized membrane potentials.

The cell lines described herein may be a particularly useful for detailed characterization of drug candidates in a high-throughput manner. Optical control is relatively fast, thereby allowing for the testing the drug's activity under more physiological forms of activation. For example, different frequencies of depolarization and/or hyperpolarization may be used to determine how a drug interacts with the channel under physiological forms of neural activity. In some instances, the process may be accomplished without the application of expensive chemical dyes to the cell lines.

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In conjunction with the various properties discussed herein, the use of various embodiments of the invention may be particularly useful for improving screening throughput by eliminating the need for cumbersome mechanical manipulation and liquid handling. Various embodiments may also be useful for repeatable the screening assay using the same samples, reducing screening cost by eliminating the need for chemically-based fluorescence reports, producing high temporal precision and low signal artifact (due to the optical nature of the voltage manipulation), modulating the level of depolarization by attenuating the light intensity used for stimulation, and ascertaining the kinetics of the drug's modulation on the ion channel through the use of pulsed light patterns.

The existence of multiple independently controllable excitation proteins and inhibition proteins opens the door for a variety of applications including, but not limited to, applications for treatment of a variety of disorders and the use of a plurality of lightresponsive proteins that can be selected so as to respond to a plurality of respective optical wavelengths. Although not always expressly stated, inhibition can be used in combination with, in addition to, or in place of excitation in the applications. The family of single-component proteins has been shown to respond to multiple wavelengths and intensities of light. Aspects of the invention allow for further mutations and/or searches for sequences that allow for additional optical wavelengths and/or individually controllable protein channels. Variations on the optical stimulus (e.g., a wavelength, intensity or duration profile) can also be used. For instance, stimulation profiles may exploit overlaps in the excitation wavelengths of two different ion channel proteins to allow excitation of both proteins at the same time. In one such instance, the proteins may have different levels of responsibility. Thus, in a neural application, one set of ion channels may produce spiking at a different success percentage relative to a second set of ion channels. Similarly, the overlaps in inhibition wavelengths of two different ion channels (or pumps) allows for inhibition of both proteins at the same time.

Alternatively, multiple light sources may be used allowing for stimulations of the light responsive proteins in the combination desired, while leaving other proteins unstimulated.

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Many human applications of the present invention require governmental approval prior to their use. For instance, human use of gene therapy may require such approval. However, similar gene therapies in neurons (non-proliferative cells that are non-susceptible to neoplasms) are proceeding rapidly, with active, FDA-approved clinical trials already underway involving viral gene delivery to human brains. This is likely to facilitate the use of various embodiments of the present invention for a large variety of applications. The following is a non-exhaustive list of a few examples of such applications and embodiments.

Addiction is associated with a variety of brain functions, including reward and expectation. Additionally, the driving cause of addiction may vary between individuals. According to one embodiment, addiction, for example nicotine addiction, may be treated with optogenetic stabilization of small areas on the insula. Optionally, functional brain imaging, for example cued-state PET or fMRI, may be used to locate a hyper metabolic focus in order to determine a precise target spot for the intervention on the insula surface.

Optogenetic excitation of the nucleus accumbens and septum may provide reward and pleasure to a patient without need for resorting to use of substances, and hence may hold a key to addiction treatment. Conversely, optogenetic stabilization of the nucleus accumbens and septum may be used to decrease drug craving in the context of addiction. In an alternative embodiment, optogenetic stabilization of hyper metabolic activity observed at the genu of the anterior cingulate (BA32) can be used to decrease drug craving. Optogenetic stabilization of cells within the arcuate nucleus of the medial hypothalamus which contain peptide products of pro-opiomelanocortin (POMC) and cocaine-and-amphetamine-regulating transcript (CART) can also be used to decrease drug addiction behavior. For further information in this regard, reference may be made to: Naqvi NH, Rudrauf D, Damasio H, Bechara A. "Damage to the insula disrupts addiction to cigarette smoking." Science. 2007 Jan 26; 315(5811):531-534, which is fully incorporated herein by reference.

Optogenetic stimulation of neuroendocrine neurons of the hypothalamic periventricular nucleus that secrete somatostatin can be used to inhibit secretion of growth hormone from the anterior pituitary, for example in acromegaly. Optogenetic stabilization of neuroendocrine neurons that secrete somatostatin or growth hormone can

be used to increase growth and physical development. Among the changes that accompany "normal" aging, is a sharp decline in serum growth hormone levels after the 4th and 5th decades. Consequently, physical deterioration associated with aging may be lessened through optogenetic stabilization of the periventricular nucleus.

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Optogenetic stabilization of the ventromedial nucleus of the hypothalamus, particularly the pro-opiomelanocortin (POMC) and cocaine-and-amphetamine-regulating transcript (CART) of the arcuate nucleus, can be used to increase appetite, and thereby treat anorexia nervosa. Alternatively, optogenetic stimulation of the lateral nuclei of the hypothalamus can be used to increase appetite and eating behaviors.

Optogenetic excitation in the cholinergic cells of affected areas including the temporal lobe, the NBM (Nucleus basalis of Meynert) and the posterior cingulate gyrus (BA 31) provides stimulation, and hence neurotrophic drive to deteriorating areas. Because the affected areas are widespread within the brain, an analogous treatment with implanted electrodes may be less feasible than an opto-genetic approach.

Anxiety disorders are typically associated with increased activity in the left temporal and frontal cortex and amygdala, which trends toward normal as anxiety resolves. Accordingly, the affected left temporal and frontal regions and amygdala may be treated with optogenetic stabilization, so as to dampen activity in these regions.

In normal physiology, photosensitive neural cells of the retina, which depolarize in response to the light that they receive, create a visual map of the received light pattern. Optogenetic ion channels can be used to mimic this process in many parts of the body, and the eyes are no exception. In the case of visual impairment or blindness due to damaged retina, a functionally new retina can be grown, which uses natural ambient light rather than flashing light patterns from an implanted device. The artificial retina grown may be placed in the location of the original retina (where it can take advantage of the optic nerve serving as a conduit back to the visual cortex). Alternatively, the artificial retina may be placed in another location, such as the forehead, provided that a conduit for the depolarization signals are transmitted to cortical tissue capable of deciphering the encoded information from the optogenetic sensor matrix. Cortical blindness could also be treated by simulating visual pathways downstream of the visual cortex. The stimulation would be based on visual data produced up stream of the visual cortex or by an artificial light sensor.

Treatment of tachycardia may be accomplished with optogenetic stimulation to parasympathetic nervous system fibers including CN X or Vagus Nerve. This causes a

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decrease in the SA node rate, thereby decreasing the heart rate and force of contraction. Similarly, optogenetic stabilization of sympathetic nervous system fibers within spinal nerves T1 through T4, serves to slow the heart. For the treatment of pathological bradycardia, optogenetic stabilization of the Vagus nerve, or optogenetic stimulation of sympathetic fibers in T1 through T4 will serve to increase heart rate. Cardiac disrhythmias resulting from aberrant electrical foci that outpace the sinoatrial node may be suppressed by treating the aberrant electrical focus with moderate optogenetic stabilization. This decreases the intrinsic rate of firing within the treated tissue, and permits the sinoatrial node to regain its role in pacing the heart's electrical system. In a similar way, any type of cardiac arrhythmia could be treated. Degeneration of cardiac tissue that occurs in cardiomyopathy or congestive heart failure could also be treated using this invention; the remaining tissue could be excited using various embodiments of the invention.

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Optogenetic excitation stimulation of brain regions including the frontal lobe, parietal lobes and hippocampi, may increase processing speed, improve memory, and stimulate growth and interconnection of neurons, including spurring development of neural progenitor cells. As an example, one such application of the present invention is directed to optogenetic excitation stimulation of targeted neurons in the thalamus for the purpose of bringing a patient out of a near-vegetative (barely-conscious) state. Growth of light-gated ion channels or pumps in the membrane of targeted thalamus neurons is affected. These modified neurons are then stimulated (e.g., via optics which may also gain access by the same passageway) by directing a flash of light thereupon so as to modulate the function of the targeted neurons and/or surrounding cells. For further information regarding appropriate modulation techniques (via electrode-based treatment) or further information regarding the associated brain regions for such patients, reference may be made to: Schiff ND, Giacino JT, Kalmar K, Victor JD, Baker K, Gerber M, Fritz B, Eisenberg B, O'Connor JO, Kobylarz EJ, Farris S, Machado A, McCagg C, Plum F, Fins JJ, Rezai AR "Behavioral improvements with thalamic stimulation after severe traumatic brain injury," Nature, Vol. 448, Aug 2, 2007, pp. 600-604.

In an alternative embodiment, optogenetic excitation may be used to treat weakened cardiac muscle in conditions such as congestive heart failure. Electrical assistance to failing heart muscle of CHF is generally not practical, due to the thin-stretched, fragile state of the cardiac wall, and the difficulty in providing an evenly distributed electrical coupling between an electrodes and muscle. For this reason,

preferred methods to date for increasing cardiac contractility have involved either pharmacological methods such as Beta agonists, and mechanical approaches such as ventricular assist devices. In this embodiment of the present invention, optogenetic excitation is delivered to weakened heart muscle via light emitting elements on the inner surface of a jacket surround the heart or otherwise against the affected heart wall. Light may be diffused by means well known in the art, to smoothly cover large areas of muscle, prompting contraction with each light pulse.

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Optogenetic stabilization in the subgenual portion of the cingulate gyrus (Cg25), yellow light may be applied with an implanted device. The goal would be to treat depression by suppressing target activity in manner analogous to what is taught by Mayberg HS *et al.*, "Deep Brain Stimulation for Treatment-Resistant Depression," Neuron, Vol. 45, 651–660, March 3, 2005, pp. 651-660, which is fully incorporated herein by reference. In an alternative embodiment, an optogenetic excitation stimulation method is to increase activity in that region in a manner analogous to what is taught by Schlaepfer *et al.*, "Deep Brain stimulation to Reward Circuitry Alleviates Anhedonia in Refractory Major Depression," Neuropsychopharmacology 2007, pp. 1-10, which is fully incorporated herein by reference.

In yet another embodiment, the left dorsolateral prefrontal cortex (LDPFC) is targeted with an optogenetic excitation stimulation method. Pacing the LDLPFC at 5-20 Hz serves to increase the basal metabolic level of this structure which, via connecting circuitry, serves to decrease activity in Cg 25, improving depression in the process. Suppression of the right dorsolateral prefrontal cortex (RDLPFC) is also an effective depression treatment strategy. This may be accomplished by optogenetic stabilization on the RDLPFC, or suppression may also be accomplished by using optogenetic excitation stimulation, and pulsing at a slow rate (e.g., 1 Hz or less) improving depression in the process. Vagus nerve stimulation (VNS) may be improved using an optogenetic approach. Use of optogenetic excitation may be used in order to stimulate only the vagus afferents to the brain, such as the nodose ganglion and the jugular ganglion. Efferents from the brain would not receive stimulation by this approach, thus eliminating some of the side-effects of VNS including discomfort in the throat, a cough, difficulty swallowing and a hoarse voice. In an alternative embodiment, the hippocampus may be optogenetically excited, leading to increased dendritic and axonal sprouting, and overall growth of the hippocampus. Other brain regions implicated in depression that could be treated using this invention include the amygdala, accumbens, orbitofrontal and

orbitomedial cortex, hippocampus, olfactory cortex, and dopaminergic, serotonergic, and noradrenergic projections. Optogenetic approaches could be used to control spread of activity through structures like the hippocampus to control depressive symptoms.

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So long as there are viable alpha and beta cell populations in the pancreatic islets of Langerhans, the islets can be targeted for the treatment of diabetes. For example, when serum glucose is high (as determined manually or by closed loop glucose detection system), optogenetic excitation may be used to cause insulin release from the beta cells of the islets of Langerhans in the pancreas, while optogenetic stabilization is used to prevent glucagon release from the alpha cells of the islets of Langerhans in the pancreas. Conversely, when blood sugars are too low (as determined manually or by closed loop glucose detection system), optogenetic stabilization may be used to stop beta cell secretion of insulin, and optogenetic excitation may be used to increase alpha-cell secretion of glucagon.

For treatment of epilepsy, quenching or blocking epileptogenic activity is amenable to optogenetic approaches. Most epilepsy patients have a stereotyped pattern of activity spread resulting from an epileptogenic focus. Optogenetic stabilization could be used to suppress the abnormal activity before it spreads or truncated it early in its course. Alternatively, activation of excitatory tissue via optogenetic excitation stimulation could be delivered in a series of deliberately asynchronous patterns to disrupt the emerging seizure activity. Another alternative involves the activation of optogenetic excitation stimulation in GABAergic neurons to provide a similar result. Thalamic relays may be targeted with optogenetic stabilization triggered when an abnormal EEG pattern is detected.

Another embodiment involves the treatment of gastrointestinal disorders. The digestive system has its own, semi-autonomous nervous system containing sensory neurons, motor neurons and interneurons. These neurons control movement of the GI tract, as well as trigger specific cells in the gut to release acid, digestive enzymes, and hormones including gastrin, cholecystokinin and secretin. Syndromes that include inadequate secretion of any of these cellular products may be treated with optogenetic stimulation of the producing cell types, or neurons that prompt their activity. Conversely, optogenetic stabilization may be used to treat syndromes in which excessive endocrine and exocrine products are being created. Disorders of lowered intestinal motility, ranging from constipation (particularly in patients with spinal cord injury) to megacolan may be treated with optogenetic excitation of motor neurons in the intestines.

Disorders of intestinal hypermotility, including some forms of irritable bowel syndrome may be treated with optogenetic stabilization of neurons that control motility.

Neurogenic gastric outlet obstructions may be treated with optogenetic stabilization of neurons and musculature in the pyloris. An alternative approach to hypomobility syndromes would be to provide optogenetic excitation to stretch-sensitive neurons in the walls of the gut, increasing the signal that the gut is full and in need of emptying.

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In this same paradigm, an approach to hypermobility syndromes of the gut would be to provide optogenetic stabilization to stretch receptor neurons in the lower GI, thus providing a "false cue" that the gut was empty, and not in need of emptying. In the case of frank fecal incontinence, gaining improved control of the internal and external sphincters may be preferred to slowing the motility of the entire tract. During periods of time during which a patient needs to hold feces in, optogenetic excitation of the internal anal sphincter will provide for retention. Providing optogenetic stimulation to the external sphincter may be used to provide additional continence. When the patient is required to defecate, the internal anal sphincter, and then external anal sphincter should be relaxed, either by pausing the optogenetic stimulation, or by adding optogenetic stabilization.

Conductive hearing loss may be treated by the use of optical cochlear implants. Once the cochlea has been prepared for optogenetic stimulation, a cochlear implant that flashes light may be used. Sensorineural hearing loss may be treated through optical stimulation of downstream targets in the auditory pathway.

Another embodiment of the present invention is directed toward the treatment of blood pressure disorders, such as hypertension. Baroreceptors and chemoreceptors in regions such as the aorta (aortic bodies and paraaortic bodies) and the carotid arteries ("carotic bodies") participate in the regulation of blood pressure and respiration by sending afferents via the vagus nerve (CN X), and other pathways to the medulla and pons, particularly the solitary tract and nucleus. Optogenetic excitation of the carotid bodies, aortic bodies, paraortic bodies, may be used to send a false message of "hypertension" to the solitary nucleus and tract, causing it to report that blood pressure should be decreased. Optogenetic excitation or stabilization directly to appropriate parts of the brainstem may also be used to lower blood pressure. The opposite modality causes the optogenetic approach to serve as a pressor, raising blood pressure. A similar effect may also be achieved via optogenetic excitation of the Vagus nerve, or by optogenetic stabilization of sympathetic fibers within spinal nerves T1-T4. In an

alternative embodiment, hypertension may be treated with optogenetic stabilization of the heart, resulting in decreased cardiac output and lowered blood pressure. According to another embodiment, optogenetic stabilization of aldosterone-producing cells within the adrenal cortex may be used to decrease blood pressure. In yet another alternative embodiment, hypertension may be treated by optogenetic stabilization of vascular smooth muscle. Activating light may be passed transcutaneously to the peripheral vascular bed.

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Another example embodiment is directed toward the treatment of hypothalamicpituitary-adrenal axis disorders. In the treatment of hypothyroidism, optogenetic excitation of parvocellular neuroendocrine, neurons in the paraventricular and anterior hypothalamic nuclei can be used to increase secretion of thyrotropin-releasing hormone (TRH). TRH, in turn, stimulates anterior pituitary to secrete TSH. Conversely, hyperthyroidism may be treated with optogenetic stabilization of the provocellular neuroendocrine neurons. For the treatment of adrenal insufficiency, or of Addison's disease, optogenetic excitation of parvocellular neuroendocrine neurons in the supraoptic nucleus and paraventricular nuclei may be used to increase the secretion of vasopressin, which, with the help of corticotropin-releasing hormone (CRH), stimulate anterior pituitary to secrete ACTH. Cushing syndrome, frequently caused by excessive ACTH secretion, may be treated with optogenetic stabilization of the parvocellular neuroendocrine neurons of supraoptic nucleus via the same physiological chain of effects described above. Neuroendocrine neurons of the arcuate nucleus produce dopamine, which inhibits secretion of prolactin from the anterior pituitary. Hyperprolactinemia can therefore be treated via optogenetic excitation, while hypoprolactinemia can be treated with optogenetic stabilization of the neuroendocrine cells of the arcuate nucleus.

In the treatment of hyperautonomic states, for example anxiety disorders, optogenetic stabilization of the adrenal medulla may be used to reduce norepinephrine output. Similarly, optogenetic stimulation of the adrenal medulla may be used in persons with need for adrenaline surges, for example those with severe asthma, or disorders that manifest as chronic sleepiness.

Optogenetic stimulation of the adrenal cortex will cause release of chemicals including cortisol, testosterone, and aldosterone. Unlike the adrenal medualla, the adrenal cortex receives its instructions from neuroendocrine hormones secreted from the pituitary and hypothalamus, the lungs, and the kidneys. Regardless, the adrenal cortex is amenable to optogenetic stimulation. Optogenetic stimulation of the cortisol-producing

cells of the adrenal cortex may be used to treat Addison's disease. Optogenetic stabilization of cortisol-producing cells of the adrenal cortex may be used to treat Cushing's disease. Optogenetic stimulation of testosterone-producing cells may be used to treat disorders of sexual interest in women: Optogenetic stabilization of those same cells may be used to decrease facial hair in women. Optogenetic stabilization of aldosterone-producing cells within the adrenal cortex may be used to decrease blood pressure. Optogenetic excitation of aldosterone-producing cells within the adrenal cortex may be used to increase blood pressure.

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Optogenetic excitation stimulation of specific affected brain regions may be used to increase processing speed, and stimulate growth and interconnection of neurons, including spurring the maturation of neural progenitor cells. Such uses can be particularly useful for treatment of mental retardation.

According to another embodiment of the present invention, various muscle diseases and injuries can be treated. Palsies related to muscle damage, peripheral nerve damage and to dystrophic diseases can be treated with optogenetic excitation to cause contraction, and optogenetic stabilization to cause relaxation. This latter relaxation via optogenetic stabilization approach can also be used to prevent muscle wasting, maintain tone, and permit coordinated movement as opposing muscle groups are contracted. Likewise, frank spasticity can be treated via optogenetic stabilization.

In areas as diverse as peripheral nerve truncation, stroke, traumatic brain injury and spinal cord injury, there is a need to foster the growth of new neurons, and assist with their integration into a functional network with other neurons and with their target tissue. Re-growth of new neuronal tracts may be encouraged via optogenetic excitation, which serves to signal stem cells to sprout axons and dendrites, and to integrate themselves with the network. Use of an optogenetic technique (as opposed to electrodes) prevents receipt of signals by intact tissue, and serves to ensure that new target tissue grows by virtue of a communication set up with the developing neurons, and not with an artificial signal like current emanating from an electrode.

Obesity can be treated with optogenetic excitation to the ventromedial nucleus of the hypothalamus, particularly the pro-opiomelanocortin (POMC) and cocaine-and-amphetamine-regulating transcript (CART) of the arcuate nucleus. In an alternative embodiment, obesity can be treated via optogenetic stabilization of the lateral nuclei of the hypothalamus. In another embodiment, optogenetic stimulation to leptin-producing

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cells or to cells with leptin receptors within the hypothalamus may be used to decrease appetite and hence treat obesity.

Destructive lesions to the anterior capsule and analogous DBS to that region are established means of treating severe, intractable obsessive-compulsive disorder 48 (OCD48). Such approaches may be emulated using optogenetic stabilization to the anterior limb of the internal capsule, or to regions such as BA32 and Cg24 which show metabolic decrease as OCD remits.

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Chronic pain can be treated using another embodiment of the present invention. Electrical stimulation methods include local peripheral nerve stimulation, local cranial nerve stimulation and "sub threshold" motor cortex stimulation. Reasonable autogenic approaches include optogenetic stabilization at local painful sites. Attention to promoter selection would ensure that other sensory and motor fibers would be unaffected. Selective optogenetic excitation of interneurons at the primary motor cortex also may provide effective pain relief. Also, optogenetic stabilization at the sensory thalamus, (particularly medial thalamic nuclei), periventricular grey matter, and ventral raphe nuclei, may be used to produce pain relief. In an alternative embodiment, optogenetic stabilization of parvalbumin-expressing cells targeting as targeting strategy, may be used to treat pain by decreasing Substance P production. The release of endogenous opiods may be accomplished by using optogenetic excitation to increase activity in the nucleus accumbens. In an alternative embodiment, when POMC neurons of the arcuate nucleus of the medial hypothalamus are optogenetically excited, beta endorphin are increased, providing viable treatment approaches for depression and for chronic pain.

Certain personality disorders, including the borderline and antisocial types, demonstrate focal deficits in brain disorders including "hypofrontality." Direct or indirect optogenetic excitation of these regions is anticipated to produce improvement of symptoms. Abnormal bursts of activity in the amygdala are also known to precipitate sudden, unprompted flights into rage: a symptom of borderline personality disorder, as well as other conditions, which can benefit from optogenetic stabilization of the amygdala. Optogenetic approaches could improve communication and synchronization between different parts of the brain, including amygdala, striatum, and frontal cortex, which could help in reducing impulsiveness and improving insight.

The amygdalocentric model of post-traumatic-stress disorder (PTSD) proposes that it is associated with hyperarousal of the amygdala and insufficient top-down control

by the medial prefrontal cortex and the hippocampus. Accordingly, PTSD may be treated with optogenetic stabilization of the amygdale or hippocampus.

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Schizophrenia is characterized by abnormalities including auditory hallucinations. These might be treated by suppression of the auditory cortex using optogenetic stabilization. Hypofrontality associated with schizophrenia might be treated with optogenetic excitation in the affected frontal regions. Optogenetic approaches could improve communication and synchronization between different parts of the brain which could help in reducing misattribution of self-generated stimuli as foreign.

Optogenetic stabilization of cells within the arcuate nucleus of the medial hypothalamus, which contain peptide products of pro-opiomelanocortin (POMC) and cocaine-and-amphetamine-regulating transcript (CART), can be used to reduce compulsive sexual behavior. Optogenetic excitation of cells within the arcuate nucleus of the medial hypothalamus which contain peptide products of pro-opiomelanocortin (POMC) and cocaine-and-amphetamine-regulating transcript (CART) may be used to increase sexual interest in the treatment of cases of disorders of sexual desire. In the treatment of disorders of hypoactive sexual desire testosterone production by the testes and the adrenal glands can be increased through optogenetic excitation of the pituitary gland. Optogenetic excitation of the nucleus accumbens can be used for the treatment of anorgasmia.

The suprachiasmatic nucleus secretes melatonin, which serves to regulate sleep/wake cycles. Optogenetic excitation to the suprachiasmic nucleus can be used to increase melatonin production, inducing sleep, and thereby treating insomnia. Orexin (hypocretin) neurons strongly excite numerous brain nuclei in order to promote wakefulness. Optogenetic excitation of orexin-producing cell populations can be used to treat narcolepsy, and chronic daytime sleepiness.

Optogenetic stimulation of the supraoptic nucleus may be used to induce secretion of oxytocin, can be used to promote parturition during childbirth, and can be used to treat disorders of social attachment.

Like muscular palsies, the motor functions that have been de-afferented by a spinal cord injury may be treated with optogenetic excitation to cause contraction, and optogenetic stabilization to cause relaxation. This latter relaxation via optogenetic stabilization approach may also be used to prevent muscle wasting, maintain tone, and permit coordinated movement as opposing muscle groups are contracted. Likewise, frank spasticity may be treated via optogenetic stabilization. Re-growth of new spinal

neuronal tracts may be encouraged via optogenetic excitation, which serves to signal stem cells to sprout axons and dendrites, and to integrate themselves with the network.

Stroke deficits include personality change, motor deficits, sensory deficits, cognitive loss, and emotional instability. One strategy for the treatment of stroke deficits is to provide optogenetic stimulation to brain and body structures that have been deafferented from excitatory connections. Similarly, optogenetic stabilization capabilities can be imparted on brain and body structures that have been deafferented from inhibitory connections.

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Research indicates that the underlying pathobiology in Tourette's syndrome is a phasic dysfunction of dopamine transmission in cortical and subcortical regions, the thalamus, basal ganglia and frontal cortex. In order to provide therapy, affected areas are preferably first identified using techniques including functional brain imaging and magnetoencephalography (MEG). Whether specifically identified or not, optogenetic stabilization of candidate tracts may be used to suppress motor tics. Post-implantation empirical testing of device parameters reveals which sites of optogenetic stabilization, and which are unnecessary to continue.

In order to treat disorders of urinary or fecal incontinence optogenetic stabilization can be used to the sphincters, for example via optogenetic stabilization of the bladder detrussor smooth muscle or innervations of that muscle. When micturation is necessary, these optogenetic processes are turned off, or alternatively can be reversed, with optogenetic stabilization to the (external) urinary sphincter, and optogenetic excitation of the bladder detrussor muscle or its innervations. When a bladder has been deafferentated, for example, when the sacral dorsal roots are cut or destroyed by diseases of the dorsal roots such as tabes dorsalis in humans, all reflex contractions of the bladder are abolished, and the bladder becomes distended. Optogenetic excitation of the muscle directly can be used to restore tone to the detrussor, prevent kidney damage, and to assist with the micturition process. As the bladder becomes "decentralized" and hypersensitive to movement, and hence prone to incontinence, optogenetic stabilization to the bladder muscle can be used to minimize this reactivity of the organ.

In order to selectively excite/inhibit a given population of neurons, for example those involved in the disease state of an illness, several strategies can be used to target the optogenetic proteins/molecules to specific populations.

For various embodiments of the present invention, genetic targeting may be used to express various optogenetic proteins or molecules. Such targeting involves the

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targeted expression of the optogenetic proteins/molecules via genetic control elements such as promoters (*e.g.*, Parvalbumin, Somatostatin, Cholecystokinin, GFAP), enhancers/silencers (*e.g.*, Cytomaglovirus Immediate Early Enhancer), and other transcriptional or translational regulatory elements (*e.g.*, Woodchuck Hepatitis Virus Post-transcriptional Regulatory Element). Permutations of the promoter+enhancer+regulatory element combination can be used to restrict the expression of optogenetic probes to genetically-defined populations.

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Various embodiments of the present invention may be implemented using spatial/anatomical targeting. Such targeting takes advantage of the fact that projection patterns of neurons, virus or other reagents carrying genetic information (DNA plasmids, fragments, etc), can be focally delivered to an area where a given population of neurons project to. The genetic material will then be transported back to the bodies of the neurons to mediate expression of the optogenetic probes. Alternatively, if it is desired to label cells in a focal region, viruses or genetic material may be focally delivered to the interested region to mediate localized expression.

Various gene delivery systems are useful in implementing one or more embodiments of the present invention. One such delivery system is Adeno-Associated Virus (AAV). AAV can be used to deliver a promoter+optogenetic probe cassette to a specific region of interest. The choice of promoter will drive expression in a specific population of neurons. For example, using the CaMKIIa promoter will drive excitatory neuron specific expression of optogenetic probes. AAV will mediate long-term expression of the optogenetic probe for at least one year or more. To achieve more specificity, AAV may be pseudotyped with specific serotypes 1, 2, 3, 4, 5, 6, 7, and 8, with each having different trophism for different cell types. For instance, serotype 2 and 5 is known to have good neuron-specific trophism.

Another gene delivery mechanism is the use of a retrovirus. HIV or other lentivirus-based retroviral vectors may be used to deliver a promoter+optogenetic probe cassette to a specific region of interest. Retroviruses may also be pseudo-typed with the Rabies virus envelope glycoprotein to achieve retrograde transport for labeling cells based on their axonal projection patterns. Retroviruses integrate into the host cell's genome, therefore are capable of mediating permanent expression of the optogenetic probes. Non-lentivirus based retroviral vectors can be used to selectively label dividing cells.

Gutless Adenovirus and Herpes Simplex Virus (HSV) are two DNA-based viruses that can be used to deliver promoter+optogenetic probe cassette into specific regions of the brain as well. HSV and Adenovirus have much larger packaging capacities and therefore can accommodate much larger promoter elements and can also be used to deliver multiple optogenetic probes or other therapeutic genes along with optogenetic probes.

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Focal Electroporation can also be used to transiently transfect neurons. DNA plasmids or fragments can be focally delivered into a specific region of the brain. By applying mild electrical current, surrounding local cells will receive the DNA material and expression of the optogenetic probes.

In another instance, lipofection can be used by mixing genetic material with lipid reagents and then subsequently injected into the brain to mediate transfection of the local cells.

Various embodiments involve the use of various control elements. In addition to genetic control elements, other control elements (particularly promoters and enhancers whose activities are sensitive to chemical, magnetic stimulation or infrared radiation) can be used to mediate temporally-controlled expression of the optogenetic probes. For example, a promoter whose transcriptional activity is subject to infrared radiation allows one to use focused radiation to fine tune the expression of optogenetic probes in a focal region at only the desired time.

Parkinson's Disease can be treated by expressing optogenetic stabilization in the glutamatergic neurons in either the subthalamic nucleus (STN) or the globus pallidus interna (GPi) using an excitatory-specific promoter such as CaMKIIα, and apply optogenetic stabilization. Unlike electrical modulation in which all cell-types are affected, only glutamatergic STN neurons would be suppressed.

Aspects of the present invention are directed towards testing a model of a neural circuit or disease. The model can define output response of the circuit as a function of input signals. The output response can be assessed using a number of different measurable characteristics. For instance, characteristics can include an electrical response of downstream neurons and/or behavioral response of a patient. To test the model, optogenetic probes are expressed at an input position for the model. The optogenetic probes are stimulated and the output characteristics are monitored and compared to an output predicted by the model.

In certain implementations, the use of optogenetic probes allows for fine tuning of models defined using electrical probes. As electrical probes provide only limited ability to direct the stimulus and thus are not well suited for stimulus of certain areas without also directly stimulating nearby areas. Optogenetic probes disclosed herein provide a mechanism for more precise selection of the stimulus location. For instance, the stimulus from the optogenetic probes can be directed to very specific types of circuits/cells, such as afferent fibers. The following description provides an example implementation consistent with such an embodiment and is meant to show the feasibility and wide-ranging applicability for aspects of present invention.

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According to one embodiment of the present invention, the invention may be used in animal models of DBS, for example in Parkinsonian rats, to identify the target cell types responsible for therapeutic effects (an area of intense debate and immense clinical importance). This knowledge alone may lead to the development of improved pharmacological and surgical strategies for treating human disease.

One such application involves long-term potentiation (LTP) and/or long-term depression (LTD) between two neural groups. By targeting the expression of VChR1 and ChR2 to different neural populations and stimulating each with a different frequency of light, LTP or LTD can be accomplished between the two groups. Each group can be individually controlled using the respective wavelength of light. This can be particularly useful for applications in which the spatial arrangement of the two groups presents issues with individual control using the same wavelength of light. Thus, the light delivery device(s) are less susceptible to exciting the wrong neural group and can be less reliant upon precise spatial location of the optical stimulus.

The delivery of the proteins to cells in vivo can be accomplished using a number of different deliver devices, methods and systems. On such delivery device is an implantable device that delivers a nucleotide sequence for modifying cells *in vivo*, such as a viral-vector. The implantable device can also include a light delivery mechanism. The light delivery can be accomplished using, for example, light-emitting diodes (LEDs), fiber optics and/or Lasers.

Another embodiment of the present invention involves the use of VChR1 in affecting stem cell fate including survival/death, differentiation and replication. The modulation of electrical properties has been shown to control stem cell fate. Various techniques can be used to provide stimulus patterns that modify stem cell fate. A specific example is consistent with the techniques explained in Deisseroth, K. *et al.*

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"Excitation-neurogenesis coupling in adult neural stem/progenitor cells," Neuron 42, pp. 535-552 (2004), which is fully incorporated herein by reference.

Another embodiment of the present invention is directed to the use of DChR and/or GtR3 to assess the efficacy of treatments. This can include, but is not limited to, drug screening, treatment regimens or modeling of treatments/disorders. In a specific embodiment, DChR is used as the primary optically responsive protein in such assessments. In alternate embodiments, DChR is used with other types of optically responsive proteins (*e.g.*, VCHR1, GtR3, ChR2 and/or NpHR) that respond to different wavelengths.

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Also provided herein are methods of identifying transsynaptic connection between neuronal cells in an animal or a tissue, comprising: a) administering a first viral vector encoding a Cre recombinase fused to a transcellular tracer protein to neuronal cells in region A of the animal or tissue; b) administering a second viral vector encoding a light-activated protein to neuronal cells in region B of the animal or tissue, wherein the expression of the light-activated protein depends on the presence of the Cre recombinase; and c) identifying neuronal cells expressing the light-activated protein in region B, wherein the expression of the light-activated protein in the neuronal cells indicating that these cells are in transsynaptic connection with the cells in region A.

Also provided herein are methods of generating optical control of targeted neuronal cells in an animal or tissue, comprising: a) administering a first viral vector expressing a Cre recombinase fused to a transcellular tracer protein to region A of the animal or tissue; b) administering a second viral vector encoding a light-activated protein to region B of the animal or tissue, wherein the expression of the light-activated protein depends on the presence of the Cre recombinase, and wherein the neuronal cells in region A and in region B are in transsynaptic connected; and c) controlling action potential of a neuronal cell in region B with light that activates the protein.

Also provided herein are methods of controlling action potential of a neuron in an animal, comprising activating a light-activated protein in the neuron with light to generate action potential change, wherein expression of the light-activated protein in the neuron is generated by a) administering a first viral vector expressing a Cre recombinase fused to a transcellular tracer protein to region A of the animal, b) administering a second viral vector encoding a light-activated protein to region B of the animal which contains the neuron, wherein the expression of the light-activated protein depends on the presence

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of the Cre recombinase, wherein the neurons in region A and region B are in transsynaptic connection.

In some embodiments, the viral vector is a viral vector selected from the group consisting of AAV, HSV, and lentivirus. In some embodiments, the transcellular tracer protein is wheat germ agglutinin (WGA) or tetanus toxin-fragment C (TTC).

It is to be understood that one, some, or all of the properties of the various embodiments described herein may be combined to form other embodiments of the present invention. These and other aspects of the invention will become apparent to one of skill in the art.

It is understood that aspect and variations of the invention described herein include "consisting" and/or "consisting essentially of" aspects and variations.

As used herein and in the appended claims, the singular forms "a," "an," and "the" include plural reference unless the context clearly indicates otherwise. For example, reference to an "animal cell" is a reference to from one to many cells.

An "isolated" polynucleotide is one which has been identified and separated and/or recovered from a component of its natural environment.

Reference to "about" a value or parameter herein includes (and describes) variations that are directed to that value or parameter per se. For example, description referring to "about X" includes description of "X". The term "about" has its normal meaning of approximately. In some embodiments, "about" means $\pm 10\%$ or $\pm 5\%$.

Table 3: Sequence for GtR3

>GtR3

Table 3: Sequence for GtR3

Translation Map

GtR3

- 1 ATGGACTACGGAGGAGCACTGTCTGCTGTGGGCCGTGAATTACTCTTTGTGACCAATCCA
 1 M D Y G G A L S A V G R E L L F V T N P
- ${\bf 61~GTCGTTGTAAATGGGAGCGTCCTGGTGCCGGAGGATCAATGCTACTGCGCCGGTTGGATT}$
- 21 V V V N G S V L V P E D Q C Y C A G W I
- 121 GAAAGCAGAGGCACGAATGGGGCCTCATCCTTCGGCAAGGCCCTACTGGAGTTTGTCTTC
- 41 ESRGTNGASSFGKALLEFVF
- 181 ATCGTCTTCGCGTGTATCACATTACTGTTGGGAATTAACGCTGCGAAATCAAAGGCTGCA
- 61 I V F A C I T L L L G I N A A K S K A A
- 241 TCTAGGGTGCTGTTTCCCGCTACTTTCGTCACTGGAATCGCAAGTATCGCATATTTTTCC
- 81 S R V L F P A T F V T G I A S I A Y F S
- 301 ATGGCAAGCGGCGGGGGGGGTGATTGCCCCTGACTGTCGGCAGCTCTTTGTGGCCCGC
- 101 M A S G G G W V I A P D C R Q L F V A R
- 361 TATCTGGACTGGCTCATTACTACACCACTTCTACTCATAGATTTGGGTCTGGTTGCAGGG
- 121 Y L D W L I T T P L L L I D L G L V A G
- 421 GTCAGTCGGTGGGATATAATGGCCCTCTGCCTGTCTGATGTCCTGATGATTGCTACGGGT
- 141 V S R W D I M A L C L S D V L M I A T G
- 161 A F G S L T V G N V K W V W W F F G M C
- 541 TGGTTTCTTCACATAATCTTCGCGCTTGGGAAAAGTTGGGCAGAAGCAGCCAAGGCCAAG
- 181 W F L H I I F A L G K S W A E A A K A K
- 601 GGCGGCGACTCTGCTTCTGTGTACTCCAAAATCGCCGGCATCACCGTGATTACATGGTTC
- 201 G G D S A S V Y S K I A G I T V I T W F
- 661 TGTTATCCCGTGGTATGGGTCTTCGCTGAGGGCTTCGGAAACTTTTCCGTAACCTTCGAA
- 221 CYPVVWVFAEGFGNFSVTFE

174(c), 183(c), 678(c)

- 721 GTTCTCATCTATGGAGTGTTGGATGTTATTTCAAAGGCCGTTTTTGGCCTTATACTGATG
- 241 V L I Y G V L D V I S K A V F G L I L M
- 781 TCAGGGGCCGCCACCGGATACGAGTCCATT
- 261 S G A A T G Y E S I

Restriction Sites

Bbsl

Name	Seq.	Locatio	ns
Aatl	AGGCCT	-	none
Accl	GTMKA	C	none
AfIII	CTTAAG		none
Agel	ACCGGT	-	none
AlwI	GGATC	92	
AlwNI	CAGNNI	NCTG	none
Apal	GGGCCC		none
ApaLl	GTGCAC	:	none
Ascl	GGCGCC	GCC	none
Asel	ATTAAT	none	
Aval	CYCGRG		none
Avall	GGWCC	none	
Avrll	CCTAGG		none
BamHl	GGATCC		none

GAAGAC

Table :	3: Sequence for	or GtR3	
Bbvl	GCAGC 341, 5	85, 219(c), 234(c)
3cll	TGATCA	none	
Bgll	GCCNNNNNG	GC .	none
Bglll	AGATCT	none	
BlpI	GCTNAGC	none	
Bsal	GGTCTC	none	
BsmAl	GTCTC none		
BsmBl	CGTCTCnone		
BstEll	GGTNACC	none	
BstXI	CCANNNNNNT	GG	none
Clal	ATCGAT	none	
Dralll	CACNNNGTG	none	
Eagl	CGGCCG	none	
Earl	CTCTTC none		
EcoRI	GAATTC	none	
EcoRV	GATATC	none	
Fokl	GGATG 740, 14		
Fsel	GGCCGGCC	none	
	AAGCTT	none	
Kasl	GGCGCC	none	
Kpnl	GGTACC	none	
Miul	ACGCGT	none	
Narl	GGCGCC	none	
Ncol	CCATGG	298	
Ndel	CATATG	none	
Nhel	GCTAGC	none	
Notl	GCGGCCGC	none	
Nsil	ATGCAT		
	TTAATTAA	none	
Paci Deil	ACATGT	none	
Pcil Dmol		none	
Pmel	GTTTAAAC	none	
Pstl	CTGCAG	none	
Pvul	CACCTC	none	
Pvull	CAGCTG	none	
Sacl	GAGCTC	none	
SacII	CCGCGG	none	
Sali	GTCGAC	none	
Sapl	GCTCTTC	none	
Sfil	GGCCNNNNNG		none
SgrAl	CRCCGGYG	none	
Smal	CCCGGG	none	
Spel	ACTAGT	none	
SphI	GCATGC	none	
Sspl	AATATTnone		
Stul	AGGCCT	none	
Swal	ATTTAAAT	none	
Tlil	CTCGAG	none	
Xbal	TCTAGA	none	
Xhol	CTCGAG	none	
Xmal	cccggg	none	

Table 3: Sequence for GtR3								
Xmnl	GAANNNNTTC	697						
Availl	atgcat none							
Afel	AGCGCT	none						
Avrll	CCTAGG	none .						
BspEl	TCCGGA	none						
BsrGl	TGTACA	none						

Table 4: Sequence for DChR

>DChR

GGATTTTTACTACCTGGCTGGTTCAGATAAGGAATGGGTCGTTGGCCCTAGCGACCAGTGTTACTGCCG TTCGCCATTTGTATCGTACAACTGGTTTATTTCATGTTTGCCGCTTGGAAGGCAACGGTCGGATGGGAG GAAGTCTACGTGAACATCATTGAGCTGGTGCACATTGCCCTGGTGATTTGGGTCGAGTTCGATAAACCC GCCATGCTCTACCTTAACGACGGTCAGATGGTTCCATGGTTGCGCTATAGTGCATGGCTCCTTTCCTGCC CAGTCATCCTAATTCACCTGAGCAACTTAACAGGGGCTAAAGGGGGACTATAGTAAGAGAACCATGGGG CTTTTGGTCTCTGACATCGGAACCATAGTGTTTGGTACAAGCGCCGCACTCGCTCCGCCAAACCATGTC AAAGTCATCTTATTTACAATTGGGTTGCTGTATGGACTCTTCACTTTTTTCACGGCAGCGAAGGTATATA TTGAGGCCTACCACACCGTTCCAAAAGGCCAATGTAGAAACCTCGTGAGGGCTATGGCCTGGACTTATT TCGTAAGTTGGGCGATGTTCCCCATCCTGTTTATCCTGGGAAGAGAGGGTTTTGGCCATATTACATATTT TGGCTCATCCGTCGGACACTTCATACTGGAGATATTTTCAAAAAATCTGTGGAGTCTACTGGGCCACGG ATTACGGTATCGCATAAGGCAGCATATCATCATTCATGGCAATTTGACAAAGAAGAATAAGATTAATAT CGCAGGGGACAACGTCGAAGTGGAAGAGTACGTGGATTCTAACGACAAGGACAGCGACGTT

>DChR

GGATTTTTACTACCTGGCTGGTTCAGATAAGGAATGGGTCGTTGGCCCTAGCGACCAGTGTTACTGCCG AGCATGGTCTAAATCACACGGCACCGACAGAGAGGGGCGAGGCGGCTGTGGTGTGGGCGTACATCGTA TTCGCCATTTGTATCGTACAACTGGTTTATTTCATGTTTGCCGCTTGGAAGGCAACGGTCGGATGGGAG GAAGTCTACGTGAACATCATTGAGCTGGTGCACATTGCCCTGGTGATTTGGGTCGAGTTCGATAAACCC GCCATGCTCTACCTTAACGACGGTCAGATGGTTCCATGGTTGCGCTATAGTGCATGGCTCCTTTCCTGCC CAGTCATCCTAATTCACCTGAGCAACTTAACAGGGGCTAAAGGGGGACTATAGTAAGAGAACCATGGGG CTTTTGGTCTCTGACATCGGAACCATAGTGTTTGGTACAAGCGCCGCACTCGCTCCGCCAAACCATGTC AAAGTCATCTTATTTACAATTGGGTTGCTGTATGGACTCTTCACTTTTTTCACGGCAGCGAAGGTATATA TTGAGGCCTACCACACCGTTCCAAAAGGCCAATGTAGAAACCTCGTGAGGGCTATGGCCTGGACTTATT TCGTAAGTTGGGCGATGTTCCCCATCCTGTTTATCCTGGGAAGAGAGGGTTTTGGCCATATTACATATTT TGGCTCATCCATCGGACACTTCATACTGGAGATATTTTCAAAAAATCTGTGGAGTCTACTGGGCCACGG ATTACGGTATCGCATAAGGCAGCATATCATCATTCATGGCAATTTGACAAAGAAGAATAAGATTAATAT CGCAGGGGACAACGTCGAAGTGGAAGAGTACGTGGATTCTAACGACAAGGACAGCGACGTT

Table 4: Sequence for DChR

Translation Map

DChR

- 1 M R R R E S Q L A Y L C L F V L I A G W
- 21 A P R L T E S A P D L A E R R P P S E R
- 121 AACACCCCTTACGCCAATATTAAAAAGGTGCCCAATATAACTGAACCCAACGCCAATGTG
- 41 N T P Y A N I K K V P N I T E P N A N V
- 61 Q L D G W A L Y Q D F Y Y L A G S D K E
- 241 TGGGTCGTTGGCCCTAGCGACCAGTGTTACTGCCGAGCATGGTCTAAATCACACGGCACC
- 81 W V V G P S D Q C Y C R A W S K S H G T
- 301 GACAGAGAGGGCGAGGCGGCTGTGGTGTGGGCGTACATCGTATTCGCCATTTGTATCGTA
- 101 D R E G E A A V V W A Y I V F A I C I V
- 361 CAACTGGTTTATTTCATGTTTGCCGCTTGGAAGGCAACGGTCGGATGGGAGGAAGTCTAC
- 121 Q L V Y F M F A A W K A T V G W E E V Y
- 421 GTGAACATCATTGAGCTGGTGCACATTGCCCTGGTGATTTGGGTCGAGTTCGATAAACCC
- 141 V N I I E L V H I A L V I W V E F D K P
- 481 GCCATGCTCTACCTTAACGACGGTCAGATGGTTCCATGGTTGCGCTATAGTGCATGGCTC
- 161 A M L Y L N D G Q M V P W L R Y S A W L
- 541 CTTTCCTGCCCAGTCATCCTAATTCACCTGAGCAACTTAACAGGGGCTAAAGGGGGACTAT
- 181 L S C P V I L I H L S N L T G L K G D Y
- 601 AGTAAGAGAACCATGGGGCTTTTGGTCTCTGACATCGGAACCATAGTGTTTGGTACAAGC
- 201 S K R T M G L L V S D I G T I V F G T S
- 661 GCCGCACTCGCCCAAACCATGTCAAAGTCATCTTATTTACAATTGGGTTGCTGTAT
- 221 A A L A P P N H V K V I L F T I G L L Y
- 721 GGACTCTTCACTTTTTTCACGGCAGCGAAGGTATATATTGAGGCCTACCACACCGTTCCA
- 241 G L F T F F T A A K V Y I E A Y H T V P
- 781 AAAGGCCAATGTAGAAACCTCGTGAGGGCTATGGCCTGGACTTATTTCGTAAGTTGGGCG
- 261 K G Q C R N L V R A M A W T Y F V S W A
- 841 ATGTTCCCCATCCTGTTTATCCTGGGAAGAGAGGGTTTTGGCCATATTACATATTTTGGC
- 281 M F P I L F I L G R E G F G H I T Y F G
- 901 TCATCCATCGGACACTTCATACTGGAGATATTTTCAAAAAATCTGTGGAGTCTACTGGGC
- 301 S S I G H F I L E I F S K N L W S L L G
- 961 CACGGATTACGGTATCGCATAAGGCAGCATATCATCATCATGGCAATTTGACAAAGAAG
- 321 H G L R Y R I R Q H I I I H G N L T K K
- 1021 AATAAGATTAATATCGCAGGGGACAACGTCGAAGTGGAAGAGACGTGGATTCTAACGAC
- 341 N K I N I A G D N V E V E E Y V D S N D
- 1081 AAGGACAGCGACGTT
- 361 K D S D V

Restriction Sites

Name Seq. Locations

Aatl AGGCCT

760

GTMKAC Accl

414, 949

AfIII CTTAAG none

Agel ACCGGT

none

GGATC none Alwi

AlwNI CAGNNNCTG none

Table 4	4: Sequence for	or DChR
Apal	GGGCCC	58
ApaLl	GTGCAC	438
Ascl	GGCGCGCC	none
Asel	ATTAAT1026	
Aval	CYCGRG	none
Avall	GGWCCnone	
Avrll	CCTAGG	none
	GGATCC	none
Bbsl	GAAGAC	none
Bbvl	GCAGC 741, 9	
Bcll	TGATCA	none
	GCCNNNNNG	
Bgll Bglll	AGATCT	
_	GCTNAGC	none
BlpI Bool		none
Bsal	GGTCTC 14 62	623
	GTCTC 14, 62	4
	CGTCTCnone	****
BstEll	GGTNACC	none
BstXI	CCANNNNNNT	
Clal	ATCGAT	none
Dralll	CACNNNGTG	none
Eagl	CGGCCG	none
Earl	CTCTTC 723, 80	65(c), 1056(c)
EcoRI	GAATTC	none
EcoRV	GATATC	none
Fokl	GGATG 402, 55	54(c), 848(c), 901(c)
Fsel	GGCCGGCC	none
HindIII	AAGCTT	none
Kasl	GGCGCC	none
Kpnl	GGTACC	none
Mlul	ACGCGT	none
Varl	GGCGCC	none
Ncol	CCATGG	513, 610
Ndel	CATATG	none
Nhel	GCTAGC	none
Notl	GCGGCCGC	none
Vsil	ATGCAT	none
Pacl	TTAATTAA	none
Pcil	ACATGT	none
Pmel	GTTTAAAC	none
Pstl	CTGCAG	none
Pvul	CGATCG	none
Pvull	CAGCTG	none
Sacl	GAGCTC	none
Sacil	CCGCGG	none
Sall	GTCGAC	none
Sapl	GCTCTTC	none
Sfil Sahi	GGCCNNNNNG	
SgrAl	CRCCGGYG	
		none
Smal	CCCGGG	none

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Table 4: Sequence for DChR						
Spel	ACTAGT	none				
Sphl	GCATGC	none				
Sspl	AATATT135					
Stul	AGGCCT	760				
Swal	ATTTAAAT	none				
Tlil	CTCGAG	none				
Xbal	TCTAGA	none				
Xhol	CTCGAG	none				
Xmal	CCCGGG	none				
Xmnl	GAANNNNTTC	none				
Avalll	atgcat none					
Afel	AGCGCT	none				
Avrll	CCTAGG	none				
BspEI	TCCGGA	none				
BsrGl	TGTACA	none				

Provided herein is an animal cell including, but not limited to, neural cells, cell lines and muscle cells, the animal cell comprising: an integrated exogenous molecule which expresses a proton pump responsive to blue light, the exogenous molecule derived from Guillardia theta.

Also provided herein is a method comprising: modifying a light responsive protein derived from Guillardia theta to add an endoplasmic reticulum (ER) export signal at the C-Terminus of the light responsive protein.

Also provided herein is a system for controlling an action potential of a neuron in vivo, the system comprising: a delivery device that introduces a protein to the neuron, the protein being responsive to blue light, wherein the produces an inhibitory current a blue light source that generates light for stimulation of the blue light responsive protein; and a control device that controls the generation of light by the light source.

Also provided herein are systems, methods, arrangements, or kits directed toward: optical stimulation of a cell expressing a GtR3 proton pump.

Also provided herein are systems, methods, arrangements, or kits directed toward control of subcellular processes using GtR3 and/or DChR.

Also provided herein are systems, methods, arrangements, or kits directed toward the use of GtR3 and DChR in different cell populations of a common cellular network.

Also provided herein are systems, methods, arrangements, or kits directed toward DChR.

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Also provided herein is a protein consistent with any of the sequences described herein (e.g., sequences shown in Table 1).

Also provided herein are systems, methods, arrangements, or kits directed toward combinations of DChR or GtR3 with other light-responsive opsin types, which can be based on reaction profiles thereof.

Also provided herein are systems, methods, arrangements, or kits directed toward combinations of three or more light-responsive opsin types.

Also provided herein are methods for providing tiered-levels of activity using multiple opsin types in same cell but reactive to different light.

Also provided herein are methods for providing tiered-levels of activity using multiple opsin types in same cell to provide increase frequency of channel function through alternating stimulation of each opsin type.

Also provided herein are systems, methods, arrangements, or kits directed toward muscle control.

Also provided herein are systems, methods, arrangements, or kits directed toward combination control of several different cell groups, each group having a different combination of opsins so that each groups be controlled independently of at least one other group.

Also provided herein is a system comprising a feedback loop for control of a cell (or cell population) with the cell(s) having multiple light-responsive proteins types expressed therein.

Also provided herein are therapeutic applications including, but not limited to, treatment of Parkinson's disease.

Also provided herein are systems, methods, arrangements, or kits directed toward implantation in retinal cells/retraining brain to respond to intensity of multiple wavelengths.

Also provided herein are systems, methods, arrangements, or kits directed toward drug testing.

Also provided herein are systems, methods, arrangements, or kits directed toward the use of GtR3 and/or DChR with transgenic animals.

Also provided herein are systems, methods, arrangements, or kits directed toward control of pH levels in cells including, but not limited to, control over pH levels in an organelle, which can be implemented, without limitation, to encourage or inhibit functions thereof or to kill or maim the cell.

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Examples

According to certain embodiments, subcellular and transcellular trafficking strategies now permit (1) optical regulation at the far-red/infrared border and extension of optogenetic control across the entire visible spectrum; (2) increased potency of optical inhibition without increased light power requirement (nanoampere-scale chloride-mediated photocurrents that maintain the light sensitivity and reversible, step-like kinetic stability of earlier tools); and (3) generalizable strategies for targeting cells based not only on genetic identity, but also on morphology and tissue topology, to allow versatile targeting when promoters are not known or in genetically intractable organisms. These results illustrate the use of cell-biological principles to enable expansion of the versatile fast optogenetic technologies suitable for intact-systems biology and behavior.

Specific aspects of the present disclosure are directed to applications of molecular trafficking strategies, to derive a panel of tools that both quantitatively and qualitatively enhance the power of optogenetics and open distinct avenues of investigation. In particular, tools are developed that allow targeting of cells solely by virtue of their topological relationships within tissue and that extend the reach of optical control to the infrared border, with effector function enhanced beyond the other known tools and covering the entire visible spectrum.

According to the present disclosure, examination of eNpHR2.0-expressing hippocampal neurons revealed the absence of globular ER accumulations with persistent intracellular labeling and poor membrane localization, suggesting that additional modifications subsequent to the ER export step are important. Examination of primary-sequence differences between two forms of an inward rectifier potassium channel with differential membrane localization (Kir2.1 and Kir2.4) revealed differences not only in C-terminal ER export motifs but also in N-terminal Golgi export signals and in C-terminal trafficking signals (Hofherr et al., 2005). Surprisingly, provision of the Golgi export signal did not significantly affect surface expression, but that addition of the trafficking signal from Kir2.1 either between eNpHR and the EYFP fusion, or at the C terminus of the fusion protein, dramatically reduced intracellular labeling and increased apparent surface membrane expression and also improved labeling of cellular processes. Indeed, high-resolution confocal imaging revealed marked localization in processes, with identifiable labeled membranes spanning intracellular regions apparently devoid of the

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opsin-EYFP fusion protein, in a pattern never previously observed with NpHR or its derivatives.

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We examined photocurrents, using whole-cell patch clamp recordings to quantify bona fide functional plasma membrane localization of halorhodopsin pump molecules. Photocurrents were indeed profoundly increased (to a level ~20-fold larger than the initially described NpHR currents; mean \pm standard error of the mean [SEM]. photocurrent 747.2 ± 93.9 pA in lentivirally transduced hippocampal pyramidal neurons under the human synapsin I promoter; n = 10). FIG. 25D shows representative traces in the left portion of the figure, and summary plots in the right portion of the figure. The representative traces and summary plots of FIG. 25D shows the average photocurrent levels in cells expressing eNPHR3.0 (747.2 \pm 93.9 pA), shown in black, and eNpHR2.0, shown in gray, (214.1 \pm 24.7 pA; unpaired t test p, 0.0005; n = 10). Membrane input resistance was similar for all neurons patched (eNpHR: $193.1 \pm 36.6 \text{ M}\Omega$; eNpHR3.0: $151.6 \pm 28.5 \text{ M}\Omega$; unpaired t test p = 0.37). At action spectrum peak described below, nanoampere- scale mean outward currents were readily observed with 3.5 mW/mm² yellow light, an order of magnitude lower intensity than required by proton pumps to attain this level of photocurrent (maintaining low light intensities becomes an important issue only for in vivo experiments, wherein safe control of significant tissue volumes is paramount) (Aravanis et al., 2007; Adamantidis et al., 2007; Chow et al., 2010). FIG. 25E shows representative voltage traces, in the left portion of the figure, and summary plots, in the right portion, of eNpHR3.0 (black) and eNpHR2.0 (gray). In virally transduced neurons, light-induced hyperpolarizations by >100 mV were routinely achievable, at the same modest light power levels, as shown in FIG. 25E (mean hyperpolarization in cells expressing eNpHR3.0: 101.0 ± 24.7 mV, n = 10; and eNpHr2.0: 57.2 ± 6.8 mV, unpaired t test p, 0.0005, n = 10). Membrane potential changes of this new magnitude represent a functionally distinct advance in optogenetic inhibition, and we accordingly designate this third-generation NpHR as eNpHR3.0 (the Natronomonas halorhodopsin was named NpHR in 2005 [Sato et al., 2005], and the first trafficking-enhanced version developed by Gradinaru et al. [2008] is now referred to as eNpHR2.0). As expected from prior work (Zhang et al., 2007a) showing that NpHR photocurrents were step-like and exhibited little inactivation over more than 10 min of continuous illumination (indeed, NpHR was selected for this reason, as described in Zhang et al. [2007a]), the eNpHR3.0 photocurrents were also step-like, resistant to

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inactivation, and highly stable over multiple light pulses and long (behaviorally relevant) timescales (Zhang et al., 2007a).

FIG. 30A shows stability and recovery for eNpHR3.0 over a short time scale. The representative traces in the left portion of FIG. 30A show photocurrents in cells expressing eNpHR3.0 when exposed to pairs of 10 second long yellow light pulses separated in time by, from the top to bottom of FIG. 30A: 2.5 seconds, 5 seconds, 10 seconds, and 20 seconds. The upper right portion of FIG. 30A shows the summary plot for pulses 20 second apart displaying normalized average photocurrent levels in cells expressing eNpHR3.0 (P1 = first pulse peak, 1.00, S1 = first pulse steady state, 0.74 \pm 0.01; P2 = second pulse peak, 0.86 \pm 0.02; n = 11). The lower right portion of FIG. 30A shows the summary plot for pulses 20 seconds apart displaying approximately 50% peak recovery (P2-S1)/(P1-S1). After 20 seconds, the peak recovers to (45.2 \pm 6.6)%. FIG. 30B shows the timecourse of NpHR3.0 normalized photocurrents for long-term continuous light exposure (n = 11; various plotted are mean \pm SEM). FIG. 30C shows the outward current of eNpHR3.0 stability over 10 minutes (light deliver of 593 nm is indicated by the solid bar; output power density: 2.5 mW/mm²).

To address whether the robust improved expression is preserved in the mammalian brain in vivo, we injected lentiviral vectors delivering the novel opsin gene under control of the CaMKIIa promoter to the CA1 region of the hippocampal formation in adult mice and examined distribution of the expressed EYFP fusion. As in cultured cells, strong expression was observed not only in dendrites but also in axons in vivo with both eNpHR3.0 and eNpHR3.1 (a shorter version of eNpHR3.0 with equivalent functionality but the N-terminal signal peptide removed). A major in vivo opportunity for systems neurobiology is controlling not just a projection from region A to region B, but a cell type itself that has (among its connections) a projection from A to B. This fundamentally distinct result requires multiplexing of optical control with other targeting methods. Such control would be of great value in systems neurobiology; for example, cortical excitatory pyramidal neurons form a genetically and anatomically defined class of cell, but within this class are cells that each project to multiple different areas of the brain (e.g., thalamus, spinal cord, striatum, and other cortical areas) and therefore have fundamentally distinct roles (Lein et al., 2007; Yoshimura et al., 2005). It is unlikely that genetic tools will advance far enough to separate all of these different cell classes, pointing to the need to inhibit or excite cells defined by connection topology (FIG. 26B). One way to achieve this goal is to capitalize on transcellular trafficking: to introduce into WO 2011/116238 PCT/US2011/028893

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the local cell-body location a Cre-dependent virus conditionally expressing the microbial opsin gene of choice (e.g., Tsai et al., 2009), and rather than additionally employing a Cre-drive mouse line, to instead introduce into a distant target structure (chosen to define the cells of interest by anatomical connectivity) a virus expressing Cre recombinase fused to a transcellular tracer protein, e.g., wheat germ agglutinin (WGA) (FIG. 26B) or tetanus toxin- fragment C (TTC) (Kissa et al., 2002; Maskos et al., 2002; Perreault et al., 2006; Sano et al., 2007; Sugita and Shiba, 2005). Cre recombinase in the fusion protein would be transported by presumed endosomal trafficking mechanisms along with the tracer to the local cell-body location if anatomically connected and activate opsin expression in the subset of local cells defined by this connectivity (FIG. 26B) (Gradinaru et al., 2007, 2009; Petreanu et al., 2007, 2009). Note that this approach does not require any specific promoter fragment or genetic definition of target cells (a clear advantage for use in less-genetically tractable species such as rats and primates); but, if needed, such additional genetic refinements can be readily added (for example, both the WGA-Creand the Cre-dependent opsin could be delivered under control of cell type-specific promoters where available), creating a versatile means for addressing cells defined at the intersection of connectivity, location, and genetics.

This concept was first validated in the rat by devising a strategy to selectively introduce eNpHR3.0 into those primary motor cortex (M1) microcircuits that are involved in cortico-cortical connections with primary sensory cortex (S1) (Colechio and Alloway, 2009). To do this, we injected the previously described Cre-dependent AAV, now conditionally expressing eNpHR3.0 into motor cortex, and injected a novel WGA-Cre-expressing AAV (AAV2-EF1α-mCherry-IRES-WGA-Cre) remotely into primary somatosensory cortex. Robust eNpHR3.0-EYFP expression was indeed observed in a distributed subset of the motor cortex neurons at 5 weeks after injection, despite the remoteness of the Cre recombinase AAV injection; in control animals without Cre recombinase, no expression is observed from these Cre-dependent AAVs (Tsai et al., 2009; Sohal et al., 2009). FIG. 26C shows the construct design for the WGA-Cre and Cre dependent AAV vectors, wherein the WGA and Cre genes are both optimized with mammalian codons. Consistent with the anticipated mode of trans-synaptic or transcellular transport of Cre, no mCherry-positive cell bodies were observed in motor cortex, and no EYFP-positive cell bodies were observed in S1 sensory cortex. Cre can be trans-synaptically delivered from transduced cells to activate distant gene expression only in synaptically connected neurons that have received the Cre-dependent virus, and

not in others. The expected EYFP-eNpHR3.0 axon terminals arising from M1 were present in S1. Simultaneous optrode stimulation/ recording (Gradinaru et al., 2007) was conducted to validate functionality of eNpHR3.0 under the WGA system; indeed, robust inhibition was readily observed in M1, as expected from the intense fluorescence of the XFP-opsin fusion protein. These data indicate that neurons involved in cortico-cortical connections can indeed be addressed and targeted not simply as a projection, but as a cell type defined by connectivity.

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To independently validate this targeting technology in a distinct circuit and with a different opsin, we next targeted hippocampal formation dentate gyrus neurons involved in interhemispheric projections. Within the dentate hilus, the only known monosynaptic contralateral projection arises from the hilar mossy cells, which terminate on granule cells of the contralateral dentate, in dendrites of the molecular layer (Freund and Buzsáki, 1996; Ratzliff et al., 2004). The WGA-Cre AAV was unilaterally injected into one dentate gyrus, while the Cre-dependent AAV was injected into the contralateral dentate gyrus of the same animal. Strikingly, opsin expression was observed only in hilar cells of the contralateral side. Indeed, in this case and at this time point, accumulation of Cre was retrograde and monosynaptic to the contralateral hilar cells, as no EYFP labeling was observed in the contralateral granule cell layer; moreover, pointing to lack of direct transduction of axon terminals with this AAV serotype, no mCherry was observed in the contralateral dentate. The only EYFP-expressing circuit elements in the ipsilateral dentate, affording precise opportunities for optical control, were axonal fibers observed to terminate in the molecular layer of the granule cell layer, precisely as expected for fibers arising from the contralateral dentate hilus. Indeed, in vivo optrode recordings confirmed the functionality of WGA/Creactivated ChR2 in driving lighttriggered spikes both at the opsin-expressing cell bodies and in neurons downstream to the axonal projections of ChR2-expressing cells, in the contralateral hemisphere; in line with previous optogenetic studies (Zhang et al., 2007a, 2007b), 470 nm light pulses at 30 Hz (5 ms pulse width) delivered through the optical fiber reliably drove neuronal firing in vivo.

The utility of these targeting strategies for engineered opsins within intact tissue raised the question of whether additional advantages might accrue with regard to volume of tissue modulatable in vivo. While membrane trafficking modifications alone will not shift the action spectrum, the capability to control neurons in the far red is a long-sought goal of optogenetics, as this will allow use of light that penetrates much more deeply into

scattering biological tissues (as with the far-red utility recently demonstrated for fluorescent proteins) (Shu et al., 2009), and therefore will allow recruitment of larger volumes (Aravanis et al., 2007; Adamantidis et al., 2007; Gradinaru et al., 2009). The massive photocurrents observed for eNpHR3.0 (~20x those initially reported for NpHR, which itself is capable of blocking spiking in response to 589 nm amber light), suggested optogenetic control with far-red light might be achieved. We therefore explored optical control in the far red with the trafficking-enhanced eNpHR3.0.

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Even in response to true-red (630 nm) light of only 3.5mW/mm², we observed potent ~250 pA outward photocurrents in virally transduced cells—still more than 6-fold larger than the first observed NpHR currents with yellow light (FIG. 27A), and maintaining the characteristic step-like, stable kinetics typical of NpHR (eNpHR3.0expressing and eNpHR2.0 expressing neurons: eNpHR3.0: 239.4 \pm 28.7 pA; eNpHR2.0: $42.7 \pm 4.5 \text{ pA}$; unpaired t test p = 0.00004; n = 10) (Zhang et al., 2007a). Moreover, we found that these photocurrents evoked by red light could be used to trigger large (>40 mV) hyperpolarizations in hippocampal pyramidal neurons (FIG. 27B) (eNpHR3.0expressing and eNpHR2.0 expressing neurons: eNpHR3.0: 43.3 ± 6.1 mV; eNpHR2.0: 15.6 ± 3.2 mV; unpaired t test p = 0.00116; n = 10). We therefore explored even further red-shifted light. We continued to observe robust photocurrents in the deep red with 660 nm light and at the red/infrared border with 680 nm light (FIG. 27C). At 680 nm, the photocurrents (~75 pA) were still larger than peak (yellow light) eNpHR2.0 currents previously reported at 7 mW/mm². Importantly, at all of the red and far-red wavelengths tested, eNpHR3.0 photocurrents readily blocked action potentials induced by current injection (FIG. 27D) with 7 mW/mm² or less, validating the extension of optogenetic control channels to far-red light. The outward photocurrents evoked by different wavelengths of red and far-red/infrared border illumination are: 239.4 ± 28.7 pA (n = 10) at 630 nm; 120.5 ± 16.7 pA (n = 4) at 660 nm; and 76.3 ± 8.1 pA (n = 4) at 680 nm.

One important feature of NpHR is its spectral compatibility with ChR2: the two opsins have largely separable action spectra and operate with similar light power density requirements, allowing bidirectional control of optical activity (Zhang et al., 2007a) in vitro or in vivo despite a small region of spectral overlap. To test whether eNpHR3.0 had become too potent, given the spectral overlap, to use in combination with ChR2 in the same cell, we created a bicistronic vector containing eNpHR3.0; analogous 2A-based combination vectors (Ryan and Drew, 1994) have been employed with earlier tools, and channelrhodopsin currents with this method have ranged from 150 to 240 pA and

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halorhodopsin currents from 11 to 40 pA (Tang et al., 2009, Han et al., 2009b). We transfected the eNpHR3.0-2A-ChR2 construct (abbreviated eNPAC) into hippocampal pyramidal neurons. With these experiments, trafficking of both opsin gene products to cellular processes was observed. To verify that independent excitation and inhibition was still possible despite the increased currents from eNpHR3.0, we mapped out the steadystate photocurrent action spectrum in detail for eNPAC and for ChR2(H134R) (Gradinaru et al., 2007) and eNpHR3.0 alone. FIG. 27F, shows the activation spectrums for eNPAC, the left portion of FIG. 27F, and for ChR2 (H124R) and eNpHR3.0, on the right portion of FIG. 27F. In the right portion of FIG. 27F, the activation spectrum of ChR2 is shown in dark gray, and the activation spectrum of eNpHR3.0 is shown in light gray. Maximal eNPAC steady-state excitatory and inhibitory currents were both approximately 60% of that observed when each opsin was expressed individually, yielding maximal photocurrents of >550 pA in each direction (FIGs. 27F-G); the modestly overlapping action spectra may provide a feature, in that potent shunting inhibition combined with hyperpolarizing inhibition is likely possible with this combination approach. More specifically, maximum eNPAC steady-state excitation was $567 \pm 49 \text{ pA}$ at 427 nm (n = 9), 62% of the value for ChR2(H134R) alone (916 ± 185 pA, n = 5). Similarly, maximum eNPAC inhibition was 679 ± 109 pA at 590 nm (n = 9), 61% of the value for eNpHR3.0 alone (1110 \pm 333 pA; n = 4). Output power density for peak current values was 3.5-5 mW/mm² (3.5 mW/mm² at 590 nm). Validation in vivo will require demonstration that the specific P2A method (or other linker approach) is functional in a particular circuit or cell type (yet to be determined); however, these data in cultured hippocampal neurons demonstrate that potent bidirectional photocurrents of >500 pA each can be achieved within a single cell, without incapacitating interference of the trafficking-enhanced opsin.

The known wide action spectrum of the microbial opsins poses challenges with regard to achieving multiple independent channels of control; interestingly, eNpHR3.0 becomes not only a potent far-red optical control tool, but also the most potent known blue light-driven opsin-based inhibitor (>400 pA at 472 nm). Indeed, the membrane trafficking strategies delineated here may form a generalizable strategy for adapting diverse microbial opsins with unique properties for optogenetic control purposes. In a final series of experiments, we explored whether these and other enhanced membrane trafficking principles could enable the addition of genetically and functionally distinct components to the optogenetic toolbox.

While a very large number of microbial opsins genes exist in nature, we and others have thus far found none that outperform eNpHR3.0 (as described herein) with regard to photocurrent size, light requirements, or kinetics (Zhang et al., 2007a; Han and Boyden, 2007; Chow et al., 2010). It is important to continue to expand the optogenetic toolbox, but we have found that most microbial opsins traffic poorly in mammalian cells. However, application of the trafficking principles for microbial opsin engineering outlined here may enable optogenetics to continue the genomics progress over the past few years (Zhang et al., 2008), capitalizing on the immense natural diversity of microbial opsins (Zhang et al., 2008; Chow et al., 2010). We sought to test the adaptability of the membrane trafficking principle with the best-characterized microbial opsin, bacteriorhodopsin (BR) (Stoeckenius and Bogomolni, 1982), from *H. salinarum*, a green light-activated regulator of transmembrane ion conductance (Marti et al., 1991).

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We found that expressed in unmodified form, prominent intracellular accumulations were observed, similar to those seen when the Natronomonas halorhodopsin is expressed at high levels, and no photocurrents were observed. However, addition of the trafficking signal (TS, as employed for eNpHR3.0) between BR and EYFP substantially improved membrane and process localization, with smaller persistent ER-like accumulations that were eliminated with further C-terminal addition of the ER export signal FCYENEV. The resulting construct (eBR, doubly engineered for optimal membrane trafficking) was well tolerated in cultured neurons, with marked membrane localization and process targeting. Validation of functional plasma membrane targeting revealed that eBR could typically deliver ~50 pA of outward photocurrent and ~10 mV hyperpolarizations that sufficed to block spiking in hippocampal pyramidal neurons when exposed to the optimal wavelength light of 560 nm, thereby providing another channel for optogenetic control and illustrating the potential generalizeability of the microbial opsin membrane trafficking approach. More specifically, as seen in FIG. 28B, five hundred sixty nanometer light induced outward photocurrents of 46.4 ± 7.2 pA in eBR cells (mean \pm SEM is plotted, n = 12). The membrane input resistance was similar for all neurons patched (131.6 \pm 19.5 m Ω). The light power density at sample was 7 mW/mm². As seen in FIG. 28C, the light induced hyperpolarizations were $10.8 \pm$ 1.0 mV (mean \pm SEM plotted, n = 12).

We also continued genomic strategies similar to those that allowed our identification of the red-shifted excitatory opsin VChR1 from *Volvox carteri* (Zhang et al., 2008); indeed, a number of microbes have been reported to display light sensitivities

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from violet to near infrared. We accordingly continued our broad genomic mining approach in environmental sequencing databases, plant/microbial expressed sequence tag (EST) libraries, and whole-genome shotgun (WGS) sequencing repositories to search for new rhodopsins with channel or pump properties and novel light sensitivities (Zhang et al., 2008). Using the primary amino acid sequences for ChRs, HRs, and BRs as the template sequence, we continued the search among evolutionarily distant species (Zhang et al., 2008; Chow et al., 2010). Among other candidate sequences from diverse hosts (Cryptomonas, Guillardia, Mesostigma, Dunaliella, Gloeobacter, etc.), one of these from Guillardia theta was different from the previously reported GtR1 and GtR2 (Sineshchekov et al., 2005) and showed high amino acid homology to ChR2. We designated this new protein as G. theta rhodopsin-3 (GtR3), optimized the codon bias of GtR3 for mammalian expression, and delivered the GtR3-EYFP fusion gene to hippocampal pyramidal neurons. In an emerging theme, GtR3 showed intracellular accumulations and no photocurrents. Provision of the TS signal between GtR3 and EYFP only mildly reduced accumulations, but, together with addition of the ER export signal FCYENEV to the C terminus, accumulations were abolished and increased surface and process localization observed.

The resulting modified GtR3 hyperpolarizes hippocampal neurons in response to 472 nm blue light, albeit with smaller currents than eBR, and could inhibit spiking as well. We also achieved blue inhibition of spiking with an opsin (AR) from Acetabularia acetabulum (Tsunoda et al., 2006; Chow et al., 2010) engineered for improved trafficking; AR generates little current alone but was initially aggregate free and required only addition of the TS signal between AR and EYFP for functional membrane localization and spike inhibition. Sample current clamp and voltage clamp traces and summary data show GtR3 function under 472 nm light (18.5 mW/mm²) are shown in the left and right portions, respectively, of FIG. 31C. A light induced outward photocurrent summary is shown in the left bar graph of FIG. 31C, and the corresponding hyperpolarization summary for a blue light peak is shown in the right bar graph. Corresponding photocurrents and hyperpolarization were: 0.5 ± 0.4 pA and 0.12 ± 0.09 mV for yellow light (589 nm; 7.5 mW/mm²); 20.0 ± 6.7 pA and 5.6 ± 1.2 mV for blue light (472 nm; 18.5 mW/mm²); 1.7 ± 0.9 pA and 0.6 ± 0.3 mV for purple light (406 nm; 3 mW/mm^2) (mean \pm SEM plotted; n = 10; input resistance was similar for all neurons: $113.5 \pm 24.2 \text{ m}\Omega$).

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While these and other published microbial opsin-derived inhibitors are not yet as potent as eNpHR3.0 (and for this reason we continue with eNpHR3.0 for optogenetic applications), the improved functionality achieved here by membrane trafficking modifications point to the potential versatility of this approach in unlocking the full potential of ecological diversity, and to the individualized strategies that will be indicated for different microbial opsin genes. FIG. 29A shows general subcellular targeting strategies for adapting microbial opsin genes to metazoan intact-systems biology. FIG. 29B shows refinement of targeting at the tissue and subcellular levels (subcellular opsin targeting methods have been previously described, see Gradinaru et al. [2007] and Lewis et al., [2009]; tissue/transcellular opsin targeting methods are described herein).

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Optogenetic approaches previously have found substantial utility in the control of biological processes and behavior in freely moving mammals (Adamantidis et al., 2007; Airan et al., 2009; Gradinaru et al., 2009; Petreanu et al., 2007, 2009; Sohal et al., 2009; Tsai et al., 2009) and other animals (Douglass et al., 2008; Hwang et al., 2007; Lerchner et al., 2007; Zhang et al., 2007a), with the high temporal precision that is important for intact-systems biology. We have found that engineering specific membrane-trafficking capabilities for microbial proteins is an important step in generating diverse optogenetic technologies for intact-systems biology. Not all trafficking strategies will be suitable for all microbial opsins, with different motifs required for opsins that encounter trafficking difficulty at different stages; therefore, careful subcellular analysis with rational selection of proper modifications together constitute a directed and principled strategy that may be applicable to all opsins of nonmammalian origin, thereby enabling systematic generation of novel optogenetic tools from genomic resources.

We have previously observed that inhibition with NpHR and NpHR2.0, while useful for many applications (Gradinaru et al., 2009; Sohal et al., 2009; Tønnesen et al., 2009; Arrenberg et al., 2009), can in some cases be overcome by very strong excitation (Sohal et al., 2009). Hyperpolarizations by greater than 100 mV with eNpHR3.0 provide a substantial step forward in the potency of optical inhibition. The inhibition now provided with eNpHR3.0, more than 20-fold stronger than the initial NpHR, remains tunable with light intensity or duty-cycle adjustments, as with any of the optogenetic tools. At the action spectrum peak, nanoampere- scale mean outward currents readily resulted with only 3.5 mW/mm² yellow light (10-fold less light power than required for approaching similar currents with previously described proton pumps). At the same time,

eNpHR3.0 preserved the step-like kinetics, fast recovery, and resistance to inactivation over long timescales of NpHR (Zhang et al., 2007a).

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eNpHR3.0 is particularly well suited for in vivo applications, as the most redshifted and potent optogenetic inhibitor to date, but further strategies to enhance potency of this and other tools will no doubt emerge, and membrane trafficking work as described here may enable even more potent inhibitors in the future. When employing eNpHR3.0, inhibition can be readily dialed down if needed by using weaker promoters, shorter expression times, or reduced light levels, while maintaining access to new wavelengths ~100 nm red-shifted from previous reports (680 nm versus 589 nm) to enable operating at the infrared border with deeper-penetrating and safer photons. Of course, not only the opsin gene and light source selected, but also the strategy for circuit element targeting, may determine effectiveness; for example, optogenetic work on Parkinsonian models (Gradinaru et al., 2009) showed that therapeutically effective deep brain stimulation (DBS) in the subthalamic nucleus (STN) is likely initiated by action on afferent axons (which in turn then modulate both downstream and upstream networks). While direct fiber-optic-based inhibition of local cell bodies in the STN did not show behavioral effects comparable to those observed with direct modulation of afferent axons, these results do not mean that inhibition of the STN is not important (indeed, optogenetic axonal modulation in the STN results in inhibition of STN spiking, as noted by Gradinaru et al. [2009]). Rather, these results informed the long-standing clinical significant questions surrounding the mechanism and target of DBS by showing that axonal modulation constitutes the likely therapeutic mechanism and a highly efficient means for a point source such as a DBS electrode (or optical fiber) to control a structure or network in diseased neural circuitry (Gradinaru et al., 2009). In addition to these circuit element targeting considerations, the light intensity and wavelength, promoter choice, virus titer, virus tropism, time of opsin expression, target cell biophysics, and local patterns of endogenous activity and modulation will all affect optogenetic inhibition efficacy and should be carefully considered for each experimental system (Cardin et al., 2010).

To enable control of cell types on the basis of connectivity properties, we leveraged transcellular delivery of a Cre recombinase. First, in rat, we selectively targeted those M1 neurons that are involved in cortico-cortical connections with S1, and second, we targeted hippocampal formation dentate gyrus neurons involved in interhemispheric projections; in both cases, cells were targeted defined only by

connectivity, without use of cell type-specific promoter fragments or transgenic animals. In each system, this approach must be validated for directionality (antero- or retrograde) and extent of Cre transport, which may depend on cell-specific endosomal dynamics and experimental timepoint; this strategy may also work only with viruses rather than mouse transgenesis, allowing Cre access to episomal DNA. This approach is best served by vectors that do not directly transduce axon terminals, which may not be the case for all AAV serotypes (as we and others have in some circuits observed [Paterna et al., 2004]). Any direct transduction of axon terminals may be detected or ruled out with appropriate XFP markers, and direct transduction of axon terminals in some cases may be desirable and can be achieved with HSVs, some AAV serotypes, and pseudotyped lentiviruses; however, such an approach (unlike the present strategy) does not allow selection of the cell type postsynaptic to the transduced terminals and is not as efficient with certain AAVs that are the vector of choice for many applications due to high titer, tissue penetration, safety, and tolerability.

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The cell-process targeting enabled by membrane trafficking modification allows for control of cells that are defined topologically—that is, by the essential character of their connections within tissue. At present, it is not guaranteed that transport is synaptic or monosynaptic (as in the hippocampal circuit experiments, such properties will need to be validated in each system); therefore, the term "topological targeting" rather than synaptic targeting is here used to underscore the deformation independence of the fundamental character of the connection—an axonal connection can take any path from A to B, and as long as the connection is present, the topological targeting strategy remains valid. This property is important in genetically less-tractable organisms, but also of substantial value even in animals such as mice, where genetic targeting tools are in many cases inadequate. Of course, genetic targeting strategies may be multiplexed with topological targeting; for example, expression from the Cre-dependent vector and the Cre-fusion vector may each be governed by specific genetic targeting sequences if available. Moreover, the availability of multiple channels of optical control opens the door to combinatorial topological targeting strategies.

Like ChR2, NpHR, and VChR1, we note that most microbial opsins can benefit from substantial protein engineering to achieve new kinds of functionality. Indeed, we and others have previously demonstrated molecular strategies for eliciting from microbial opsins increased light sensitivity (Berndt et al., 2009), increased photocurrent magnitude (Gradinaru et al., 2007, 2008; Zhao et al., 2008), faster kinetics (Gunaydin et

al., 2010; Lin et al., 2009), and bistable switching behavior (Berndt et al., 2009). Other possibilities such as shifted action spectrum (Zhang et al., 2008; Gunaydin et al., 2010), increased two-photon responsivity, and altered ion permeability (e.g., for Ca2+) may also be achieved in the future.

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While ion conductance-regulating opsins have been the most versatile for ready translation (employing a common electrical language), biochemical control with light in defined cell types is also possible (but with a different set of approaches, given that microbial signal transduction employs principles distinct from metazoan signaling). Indeed, optogenetic control of well-defined biochemical signaling pathways was recently achieved both in cultured cells and in freely moving mammals, using the optoXR method for optical control of specified G protein-coupled receptor signaling (Airan et al., 2009). A photoactivatable adenylyl cyclase has been studied from Euglena, although with high dark activity that limits in vivo application (Schröder-Lang et al., 2007), and subsequent work on light-sensitive PAS or LOV domains (Levskaya et al., 2009; Wu et al., 2009) may open up new ways to control protein-protein association if these approaches can be made to operate in living animals.

We have found that in the nervous system, optogenetic tools can be applied to probe the neural circuit underpinnings of information transmission, oscillations, locomotion, awakening, and reward, as well as to probe the operation of neural circuits important in a number of brain diseases including Parkinson's disease and epilepsy 20 (Adamantidis et al., 2007; Airan et al., 2009; Cardin et al., 2009; Gradinaru et al., 2009; Sohal et al., 2009; Tønnesen et al., 2009; Tsai et al., 2009). Moreover, results thus far point to substantial versatility of the optogenetic approach across animal species (Adamantidis et al., 2007; Airan et al., 2009; Aravanis et al., 2007; Arenkiel et al., 2007; Bi et al., 2006; Boyden et al., 2005; Chow et al., 2010; Douglass et al., 2008; Gradinaru 25 et al., 2008, 2009; Hägglund et al., 2010; Han et al., 2009a; Huber et al., 2008; Hwang et al., 2007; Ishizuka et al., 2006; Li et al., 2005; Nagel et al., 2003, 2005; Petreanu et al., 2007, 2009; Tsai et al., 2009; Wang et al., 2007; Zhang et al., 2006, 2007a; Zhang and Oertner, 2007; Zhao et al., 2008). Together with fiberoptic (Adamantidis et al., 2007; Aravanis et al., 2007) and integrated fiberoptic-electrode "optrode" assemblies 30 (Gradinaru et al., 2007), even cells located deep within large, dense organs can be readily accessed and interrogated in freely moving mammals. The additional resources defined here arise from the application of molecular, cellular, and genomic strategies to enable expansion of the capabilities of optical control, and, as this toolbox rapidly grows,

optogenetics may come to play an increasingly potent and versatile role in intact-systems biology for the fast control of defined cells within functioning tissues.

Experimental Procedures

Constructs

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All NpHR variants were produced by PCR amplification of the NpHR-EYFP construct previously published (Zhang et al., 2007b). All opsins described here have been optimized for mammalian expression by changing each gene's codon usage to conform to human codon usage distribution. Updated maps and vectors are available and freely distributed from the Deisseroth laboratory (affiliated with Stanford University, Stanford, California) and described at 2010 Scientific American, "Method of the Year," December 2010 (http://www.optogenetics.org/), the contents of which are hereby incorporated by reference in their entirety.

15	The amino acid sequence of GtR3 without the signal peptide sequence																		
	A	S	S	F	G	K	A	L	L	E	F	V	FΙ	V	F	Α	C	I	Т
	L	L	L	G	I	N	A	А	K	S	K	A	A S	R	V	L	F	P	A
	\mathbf{T}	F	V	T	G	I	A	S	I	А	Y	\mathbf{F}	S M	A	S	G	G	G	W
	V	I	А	P	D	С	R	Q	L	F	V	A	R Y	L	D	M	L	I	T
20	\mathbf{T}	Р	L	L	L	I	D	L	G	L	V	Α	G V	S	R	M	D	I	M
	А	L	С	L	S	D	V	L	M	I	A	Т	G A	F	G	S	L	Т	V
	G	N	V	K	W	V	W	W	F	F	G	М	C W	F	L	Н	I	I	F
	A	L	G	K	S	M	A	E	Α	A	K	A	K G	G	D	S	A	S	V
	Y	S	K	I	А	G	I	Т	V	I	Т	W	F C	Y	Р	V	V	M	V
25	F	A	Ε	G	F	G	N	F	S	V	\mathbf{T}	F	E V	L	I	Y	G	V	L
	D	V	Ι	S	K	A	V	F	G	L	I	L	M S	G	A	A	T	G	Y
	E S I (SEQ ID NO:1)																		

The amino acid sequence of GtR3 with the signal peptide sequence from ChR2 L L F V T Ν Ρ R E V 30 M D Y G G Α L S Α G Ι С Y C Α G W Ρ D Q G S V \mathbf{L} V Ε V V Ν V F Α \mathbf{L} \mathbf{L} E F V S S F G K Τ G Α Ε S R G N K S K Α I Ν Α Α A \mathbb{C} T \mathbf{L} L L G F Α Ι Ι V F V \mathbf{T} G Ι Α S Ι Α Y F S Α T F Ρ V L S R

Α Ρ С R L F V Α R Μ Α S G G G W V Ι D O L G V Α G L Ι Τ Τ Ρ L L Ι D \mathbf{L} Y D W L L G Ι Α L С L S D V \mathbf{L} Μ Ι Α Τ V S R W D Μ F G S \mathbf{L} Τ V G Ν V K W V W W F F G Μ C Α K K F G K S Α Ε Α Α Α 5 W F \mathbf{L} Η Ι Ι Α \mathbf{L} W S Ι T VΙ T W F G G D S Α S V Υ K Ι Α G F G Ν F S V Τ F Ε C Υ Ρ V V W V F Α Ε G Ι V G V \mathbf{L} D V Ι S K Α V F G \mathbf{L} \mathbf{L} Μ \mathbf{L} Ι Y S Ι (SEQ ID NO:5) S G Α Α Τ G Y Ε

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The amino acid sequence of DChR

С L F \mathbf{L} Ι Α W Μ R R R Ε S \mathbf{L} Α Y L V G S L Τ Ε S Α Ρ D L Α \mathbf{E} R R Ρ Ρ Ε R Α Ρ R Ι Α Ν V Ν Τ Ρ Y Α Ν Ι K K V Ρ Ν Τ Ε Ρ Ν F Y Y L A G S D K Ε Υ D 15 G W Α \mathbf{L} Q Q \mathbf{L} D Τ С С Α W S K S Η G W V V G Ρ S D Q Y R V V W Α Y Ι V F Α I Ι V Ε G Ε Α Α D R F K Α Τ V G W \mathbf{E} \mathbf{E} V Y V Y F Α Α W Q L Μ V Ι V Ι Α \mathbf{L} V Ι WV Ε F D K Ρ Ν Ι Ε \mathbb{L} Η Y N D G Q Μ V Ρ W \mathbf{L} R Υ S Α W \mathbf{L} 20 Α Μ \mathbf{L} \mathbf{L} S С Ρ V Ι \mathbf{L} Ι Η \mathbf{L} S Ν \mathbf{L} Τ G \mathbf{L} K G D Y L S Ι G Τ Ι V \mathbf{F} G Τ S S Τ \mathbf{L} V D K R G \mathbf{L} Μ Τ Α Α \mathbf{L} Α Ρ Ρ Ν Η V K V Ι \mathbf{L} F Ι G \mathbb{L} \mathbf{L} Y Ρ F Τ K V Y Ι Ε Α Y Η Τ V G \mathbf{L} F Τ F Α Α F S Α С V R Α Α W Τ Y V W 25 K G Q R Ν \mathbf{L} Μ F G Η Ι Τ Y F G F Ρ Ι F Ι \mathbf{L} G R Ε G Μ L Ε Ι F S K S L G S S Ι G Η F Ι L Ν L W \mathbf{L} Ι Ι G L T K K Η G L R Υ R Ι R Q Η Ι Η Ν Ε S Ν V Ε V Ε Y V D Ν D Ι Α G D Ν K Ι Ν S V (SEQ ID NO:2) 30 K D D

The amino acid sequence of NpHR

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

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The amino acid sequence of BR

M L E L L P T A V E G V S Q A Q I T G R P E W I W L A L G T
A L M G L G T L Y F L V K G M G V S D P D A K K F Y A I T T
L V P A I A F T M Y L S M L L G Y G L T M V P F G G E Q N P
I Y W A R Y A D W L F T T P L L L L D L A L L V D A D Q G T
I L A L V G A D G I M I G T G L V G A L T K V Y S Y R F V W
W A I S T A A M L Y I L Y V L F F G F T S K A E S M R P
E V A S T F K V L R N V T V V L W S A Y P V V W L I G S E G
A G I V P L N I E T L L F M V L D V S A K V G F G L I L R
S R A I F G E A E A P E P S A G D G A A A T S D (SEQ ID
NO:4)

Hippocampal Cultures

Primary cultured hippocampal neurons were prepared from P0 Spague-Dawley rat pups. The CA1 and CA3 regions were isolated, digested with 0.4 mg/ml papain (Worthington, Lakewood, NJ), and plated onto glass coverslips precoated with 1:30 Matrigel (Beckton Dickinson Labware, Bedford, MA) at a density of 65,000/cm². Hippocampal cultures grown on coverslips were transfected or transduced at 4 days in vitro (DIV) with titer-matched viruses for all constructs (final dilution 10⁴ infectious units (i.u.)/ml in neuronal growth medium). Whole-cell patch clamp recordings were performed as previously described (Zhang et al., 2007b). Primary hippocampal cultures were infected at 4 DIV with titer-matched virus (final dilution 104 i.u./ml in neuronal growth medium). At 14 DIV, cultures were fixed for 30 min with ice-cold 4% paraformaldehyde and then permeabilized for 30 min with 0.4% saponin in 2%normal donkey serum (NDS). Primary antibody incubations were performed overnight at 4°C;

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Cy3-conjugated secondary antibodies (Jackson Laboratories, West Grove, PA) were applied in 2% NDS for 1 hr at room temperature. Images were obtained on a Leica confocal microscope with a 63x/1.4 NA oil objective.

Stereotactic Injection into the Rodent Brain and Optrode Recordings Adult mice and Long-Evans rats were housed according to the approved protocols at Stanford. All surgeries were performed under aseptic conditions. The animals were anesthetized with intraperitoneal injections of a ketamine (80 mg/kg)/xylazine (15-20 mg/kg) cocktail (Sigma). The virus was delivered via a 10 μl syringe and a thin 34 gauge metal needle; the injection volume and flow rate (1 µl at 0.1 μl/min) were controlled with an injection pump from World Precision Instruments (Sarasota, FL). For validation of opsin functionality, simultaneous optical stimulation and electrical recording in living rodents was conducted as described previously (Gradinaru et al., 2007) with an optrode composed of an extracellular tungsten electrode (1 M Ω , ~125 μ m) attached to an optical fiber (~200 μ m) with the tip of the electrode deeper (~0.4 mm) than the tip of the fiber to ensure illumination of the recorded neurons. The optical fiber was coupled to a 473 nm (for ChR2) or 560 nm (for eNpHR3.0) laser diode (10 mW fiber output) from CrystaLaser. Optrode recordings were conducted in rodents anesthetized with 1.5% isoflurane, and the optrode was placed through small craniotomies created above target regions. pClamp 10 and a Digidata 1322A board were used to both collect data and generate light pulses through the fiber. The recorded signal was band-pass filtered at 300 Hz low/5 kHz high (1800 Microelectrode AC Amplifier).

Tissue Slice Preparation

For preparation of brain slices, mice or rats were sacrificed 4 to 5 weeks after viral injection. Rodents were perfused with 20 ml ice-cold PBS, followed by 20 ml 4% paraformaldehyde. The brains were then fixed overnight in 4% paraformaldehyde and transferred to 30% sucrose solution for 2 days. Brains were frozen and coronal slices (40 µm) were prepared with a Leica SM2000R cryostat and preserved in 4°C in cryoprotectant (25% glycerol and 30% ethylene glycol in PBS). Slices (DAPI stain 1:50,000) were mounted with PVA-DABCO on microscope slides, and single confocal optical sections (e.g., through dorsal CA1 region, ~1–2.5mm posterior to bregma or the dorsal subiculum, 2.7–3 mm posterior to bregma) were acquired using a 10x air and 40x/1.4 NA oil objectives on a Leica confocal microscope.

Extended Experimental Procedures

Opsin Sources

All opsins described here have been optimized for mammalian expression by changing each gene's codon usage to conform to human codon usage distribution (http://www.kazusa.or.jp/codon/cgi-bin/showcodon.cgi?species=9606). The GenBank accession code for the original AR, BR, and GtR3 sequences are DQ074124, M11720, and EG722553.

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DNA Constructs

All NpHR variants were produced by PCR amplification of the NpHR-EYFP construct previously published (Zhang et al., 2007b) and cloned in-frame into the AgeI and EcoRI restriction sites of a lentivirus carrying the CaMKIIa or Synapsin-1 promoters according to standard molecular biology protocols. A similar strategy was used for BR and AR. GtR3 was identified through genomic searches. All opsins described here have been optimized for mammalian expression by changing each gene's codon usage to conform to human codon usage distribution (http://www.kazusa.or.jp/codon/cgibin/showcodon.cgi?species=9606), and the optimized sequence was custom synthesized (DNA2.0, Inc., Menlo Park, CA). The GenBank accession codes for the original AR, BR, and GtR3 sequences are DQ074124, M11720, and EG722553. pAAV-EF1amCherry-IRES-WGA-Cre vector was constructed using standard molecular biology protocols. Codons for the WGA and Cre genes were optimized for expression in mammalian cells. The genes were synthesized by DNA2.0 (Menlo Park, CA). Cre was fused in-frame to the C-term of WGA, which in turn was fused to IRES. The mCherry-IRES-WGA-Cre bicistronic expression cassette was designed using the EMCV IRES sequence. The pAAV-EF1a plasmid backbone is the same as described previously (Sohal et al., 2009; Tsai et al., 2009). pAAV-hSyn-eNpHR3.0-EFYP-P2A-ChR2H134RmCherry was constructed with a 120-mer primer

(5'caagttetgetaegagaacgaggtgggeteeggagecacgaacttetetetgttaaagcaageagg agacgtggaagaaaacceeggteecatggactatggeggegetttgtetgeeg 3') that contained the p2A region with the ER export sequence at the 5' end and 20 bases of the start of hChR2 at the 3' end. First, the ChR2(H134R)-mCherry fragment was amplified using the 120-mer as the forward primer and 5'-atategaatteteattacttgtacagetegt-3' as the reverse primer.

Second, this amplified product was used as a reverse primer along with the forward

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primer 5'-ccggatcccgggtaccggtaggccaccatgacagagaccctgcct—3' to fuse eNpHR 3.0-EYFP to ChR2 (H134R)-mCherry with the p2A region interposed. The 3.4 Kb fragment was then purified and cloned into the BamHI and EcoRI sites of the pAAV-hSyn vector. All constructs were fully sequenced for accuracy of cloning; updated maps are available online at http://www.optogenetics.org, the contents of which are hereby incorporated by reference in their entirety.

Lentivirus Preparation and Titering

Lentiviruses for cultured neuron infection and for in vivo injection were produced as previously described (Zhang et al., 2007b). Viral titering was performed in HEK293 cells that were grown in 24-well plates and inoculated with 5-fold serial dilutions in the presence of polybrene (8 μ g/ml). After 4 days, cultures were resuspended in PBS and sorted for EYFP fluorescence on a FACScan flow cytometer (collecting 20,000 events per sample) followed by analysis using FlowJo software (Ashland, OR). The titer of the virus was determined as follows: [(% of infected cells) x (total number of cells in well) x (dilution factor)]/ (volume of inoculum added to cells) = infectious units/ml. The titer of viruses for culture infection was 10^5 i.u./ml. The titer of concentrated virus for in vivo injection was 10^{10} i.u./ml.

Hippocampal Cultures

Primary cultured hippocampal neurons were prepared from P0 Spague-Dawley rat pups. The CA1 and CA3 regions were isolated, digested with 0.4 mg/mL papain (Worthington, Lakewood, NJ), and plated onto glass coverslips precoated with 1:30 Matrigel (Beckton Dickinson Labware, Bedford, MA) at a density of 65,000/cm². Cultures were maintained in a 5%CO₂ humid incubator with Neurobasal-A medium (Invitrogen Carlsbad, CA) containing 1.25% FBS (Hyclone, Logan, UT), 4% B-27 supplement (GIBCO, Grand Island, NY), 2 mM Glutamax (GIBCO), and FUDR (2 mg/ml, Sigma, St. Louis, MO).

Calcium Phosphate Transfection

6-10~div hippocampal neurons were grown at 65,000 cells/well in a 24-well plate. DNA/CaC CI₂ mix for each well: 1.5-3 µg DNA (QIAGEN endotoxin-free preparation) + 1.875 µl 2M CaC CI₂ (final Ca²⁺ concentration 250 mM) in 15 µl total H₂0. To DNA/CaCI₂ was added 15 µl of 2X HEPES-buffered saline (pH 7.05), and the final volume was mixed well by pipetting. After 20 min at RT, the 30 µl DNA/CaCI₂/HBS

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mixture was dropped into each well (from which the growth medium had been temporarily removed and replaced with 400 μ l warm MEM) and transfection allowed to proceed at 37C for 45-60 min. Each well was then washed with 3 X 1mL warm MEM and the growth medium replaced. Opsin expression was generally observed within 20-24 hr.

Electrophysiology

Hippocampal cultures grown on coverslips were transduced at 4 div with titermatched viruses for all constructs (final dilution 10⁴ i.u./ml in neuronal growth medium) and allowed to express for one week. Whole-cell patch clamp recordings were performed as previously described (intracellular solution: 129 mM K-gluconate, 10 mM HEPES, 10 mM KCI, 4 mM MgATP, 0.3 mM Na₃GTP, titrated to pH 7.2; extracellular Tyrode: 125 mM NaCl, 2 mM KCl, 3 mM CaCl2, 1 mM MgCl₂, 30 mM glucose, and 25 mM HEPES, titrated to pH 7.3). For voltage clamp recordings cells were held at -70mV. Light in the visible range was delivered from a 300W DG-4 lamp (Sutter Instruments, Novato, CA) through filters of different wavelength selectivity (Semrock, Rochester, NY) and a Leica 40X/0.8NA water objective. Filters, except for power spectra, (given here as wavelength in nm / bandwidth in nm / output power in mW/mm²) were: 406 / 15 / 3; 472 / 30 / 18.5; 560 / 14 / 7; 589 / 15 / 7.5; 593 / 40 / 15.5; 630 / 20 / 3.5. Far-red and near-infrared light delivery: light (7 mW/ mm2) for 660nm inhibition was delivered using a light emitting diode and a 40x/.8 NA water objective. Light (7mW/mm²) for 680nm inhibition was delivered using the X-Cite 120W halogen light source through a 680 ± 13 nm filter and a 40x/0.8 NA water objective. Light delivery for eNPAC, ChR2(H134R), and eNpHR3.0 power spectra was delivered from a 300W DG-4 lamp fitted with a Lambda 10-3 filter wheel (Sutter Instruments) with a 10-position wheel for 25-mm filters of different wavelengths and a 40X/0.8NA water objective. Filters (given here as wavelength in nm / bandwidth in nm / output power in mW/mm²) were: 387 / 10 / 3.5; 406 / 15 / 3.5; 427 / 10 / 4.0; 445 / 20 / 4.0; 470 / 22 / 4.0; 494 / 20 / 4.5; 520 / 15 / 4.5; 542 / 27 / 5.0; 560 / 20 / 5.0; 590 / 20 / 3.5; 630 / 20 / 3.5. For Figures 1, 3A–3D, and 4 and Figure S2 (see, Table 5 below), confocal images and whole-cell patch clamp data are from cultured hippocampal neurons either transfected (confocal data) or transduced (patch data) with lentiviral NpHR, BR, GtR3 and AR-based constructs, and allowed to express for one week. Expression was driven by the human Synapsin I promoter and visualized by fusion to EYFP.

Table 5 shows additional trafficking-enhanced tools for blue inhibition, and a new opsin sequence: G. *theta* rhodopsin-3 or GtR3.

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Table 5

ChR2 GtR3	MDYGGALSAVGRELLFVTNPVVVNGSVLVPEDQCYCAGWIESRGTNGAQTASNVLQWLAAEGAKLDVHGCKTVDMASSFGKALLEFVF	60 60
ChR2 GtR3	GFSILLLMFYAYQTWKSTCGWEEIYVCAIEMVKVILEFFFEFKNPSMLYLATGHRVQWLR IVFACITLLLGINAAKSKAASRVLFPATFVTGIASIAYFSMASGGGWVIAPDCRQLFVAR	120 120
ChR2	YAEWLLTCPVILIHLSNLTGLSNDYSRRTMGLLVSDIGTIVWGATSAMATGYVKVIFFCL	180
GtR3		177
0.021.0	TM4	
ChR2	GLCYGANTFFHAAKAYIEGYHTVPKGRCRQVVTGMAWLFFVSWGMFPILFILGPEGFGVL	240
GtR3	GMCWFLHIIFALGKSWAEAAKAK-GGDSASVYSKIAGITVITWFCYPVVWVFA-EGFGNF	235
CLIC	TM6	
ChR2	SVYGSTVGHTIIDLMSKNCWGLLGHYLRVLIHEHILIHGDIRKTTKLNIGGTEIEVETLV	300
GtR3	SVTFEVLIYGVLDVISKAVFGLILMSGAATGYESI	270
	TM7	
ChR2	EDEARAGAVP 310	
GtR3		

New opsin sequence: *G. theta* rhodopsin-3 or GtR3. The EST sequence included all seven transmembrane helices; the 5' amino acid sequence was provided from ChR2 (transmembrane motifs: bars; conserved residues: highlighted; truncation site for peptide: *; signal peptide provided from ChR2: gray.

Immunohistochemistry

Primary hippocampal cultures were infected at 4 div with titer matched virus (final dilution 10⁴ i.u./ml in neuronal growth medium). At 14 div cultures were fixed for 30 min with ice-cold 4% paraformaldehyde and then permeabilized for 30 min with 0.4% saponin in 2% normal donkey serum (NDS). Primary antibody incubations were performed overnight at 4°C using a monoclonal marker of endoplasmic reticulum recognizing endogenous ER-resident proteins containing the KDEL retention signal (KDEL 1:200, Abcam, Cambridge, MA). Cy3-conjugated secondary antibodies (Jackson Laboratories, West Grove, PA) were applied in 2% NDS for 1 hr at room temperature. Images were obtained on a Leica confocal microscope using a 63X/1.4NA oil objective.

Stereotactic Injection into the Rodent Brain

Adult mice and Long-Evans rats were housed according to the approved protocols at Stanford. All surgeries were performed under aseptic conditions. The

animals were anesthetized with intraperitoneal injections of a ketamine (80 mg/kg)/xylazine (15-20 mg/kg) cocktail (Sigma). The head was placed in a stereotactic apparatus (Kopf Instruments, Tujunga, CA; Olympus stereomicroscope). Ophthalmic ointment was applied to prevent eye drying. A midline scalp incision was made and a small craniotomy was performed using a drill mounted on the stereotactic apparatus 5 (Fine Science Tools, Foster City, CA). The virus was delivered using a 10 µl syringe and a thin 34 gauge metal needle; the injection volume and flow rate (1 µl at 0.1 CI₂l/min) was controlled with an injection pump from World Precision Instruments (Sarasota, FL). After injection the needle was left in place for 5 additional minutes and then slowly withdrawn. The skin was glued back with Vetbond tissue adhesive. The animal was kept 10 on a heating pad until it recovered from anesthesia. Buprenorphine (0.03 mg/kg) was given subcutaneously following the surgical procedure to minimize discomfort. For the experiment in Figure 2A, to cover a large area in dorsal CA1, 1µl of concentrated lentivirus (10¹⁰ i.u./ml) carrying eNpHR3.1 (a shorter form of eNpHR3.0 with the Nterminal signal peptide, the first 17 amino acids of original NpHR, removed) under the 15 CaMKIIa promoter was microinjected into 2 sites in each hippocampus (site one: anteroposterior -1.5 mm from bregma; lateral, \pm 1 mm; ventral, 1.5 mm; site two: AP, -2.5 mm from bregma; lateral, $\pm 2 \text{ mm}$; ventral, 1.5 mm). For Figures 2D and 2E, two different adeno-associated viruses (AAVs) (virus titer 2x10¹² g.c./mL), were stereotactically injected during the same surgery with an injection speed of 0.15ul/min. 20 High-titer (2x10¹² g.c./mL) AAV was produced by the UNC VectorCore. For Figure 2D, double-floxed cre-dependent AAV5 carrying eNpHR3.0-EYFP (AAV5-Ef1a-DIOeNpHR3.0-EYFP) was injected into M1, and AAV2-Ef1α-mCherry-IRESWGA- Cre was injected into S1 of adult Long-Evans rats. 1 μ l of virus was delivered at five different sites defined by the following coordinates: M1 injection I: AP, +1 mm from 25 bregma; lateral, 1.5 mm; ventral, 2 mm; M1 injection II: AP, +2 mm; lateral, 1.5 mm; ventral, 2 mm; S1 injection I: AP, -0.3 mm; lateral, 3.4 mm; ventral, 2 mm; S1 injection II: AP, -1.3 mm; lateral, 3 mm; ventral, 2 mm; S1 injection III: AP, -2.12 mm; lateral, 3 mm; ventral, 2 mm. For Figure 2E, 1 µl of virus was injected bilaterally into the dentate gyrus (DG) of adult BL6 mice. AAV8-EF1a-DIO-ChR2-EYFP was injected in the right 30 DG and of AAV2-EF1a-mCherry-IRES-WGA-Cre was injected in the left DG with the following coordinates: AP, -2.1 from bregma; lateral, \pm 1.05 mm; ventral, 2.1 mm.

In Vivo Optrode Recordings

To validate opsin functionality in the WGA-Cre system simultaneous optical stimulation and electrical recording in living rodents was conducted as described previously (Gradinaru et al., 2007) using an optrode composed of an extracellular tungsten electrode (1M Ω , ~125 µm) attached to an optical fiber (~200 µm) with the tip of the electrode deeper (~0.4 mm) than the tip of the fiber to ensure illumination of the recorded neurons. The optical fiber was coupled to a 473 nm (for ChR2) or 560 nm (for eNpHR3.0) laser diode (10 mW fiber output) from CrystaLaser. Optrode recordings were conducted in rodents anesthetized with 1.5% isoflurane and the optrode was placed through small craniotomies created above target regions. pClamp 10 and a Digidata 1322A board were used to both collect data and generate light pulses through the fiber. The recorded signal was band pass filtered at 300Hz low/5 kHz high (1800 Microelectrode AC Amplifier). For precise placement of the fiber/electrode pair, stereotactic instrumentation was used.

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Tissue Slice Preparation

For preparation of brain slices, mice or rats were sacrificed 4 to 5 weeks after viral injection. Rodents were perfused with 20ml of ice-cold PBS, followed by 20 ml of 4% paraformaldehyde. The brains were then fixed overnight in4% paraformaldehyde, and transferred to 30% sucrose solution for 2 days. Brains were frozen and coronal slices (40 µm) were prepared using a Leica SM2000R cryostat, and preserved in 4°C in cryoprotectant (25% glycerol, 30% ethylene glycol, in PBS). Slices (DAPI stain 1:50,000) were mounted with PVA-DABCO on microscope slides, and single confocal optical sections (e.g., through dorsal CA1 region, ~1-2.5mm posterior to bregma or the dorsal subiculum, 2.7-3mm posterior to bregma) were acquired using a 10X air and 40X/1.4NA oil objectives on a Leica confocal microscope.

For further details and discussion of the above-noted embodiments, reference can be made to "Molecular and Cellular Approaches for Diversifying and Extending Optogenetics" by Viviana Gradinaru *et al.*, Cell 141, 154–165, April 2, 2010, which is fully incorporated herein by reference.

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All references, publications, and patent applications disclosed herein are hereby incorporated by reference in their entirety.

The various embodiments described above are provided by way of illustration only and should not be construed to limit the invention. Based on the above discussion and illustrations, those skilled in the art will readily recognize that various modifications and changes may be made to the present invention without strictly following the exemplary embodiments and applications illustrated and described herein. For instance, such changes may include the use of digital logic or microprocessors to control the emitted light. Such modifications and changes do not depart from the true spirit and scope of the present invention, which is set forth in the following claims. As discussed above, specific applications and background details relative to the present invention are discussed above, in the description below and throughout the references cited herein. The embodiments in the Appendices may be implemented in connection with one or more of the above-described embodiments and implementations, as well as with those shown in the figures and described below. Reference may be made to the Appendices (A, B and C) which were filed in the underlying provisional application and incorporated herein by reference.

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WHAT IS CLAIMED IS:

- An animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to blue light and is derived from *Guillardia theta*, wherein the protein is capable of mediating a hyperpolarizing current in the cell when the cell is illuminated with light.
 - 2. The animal cell of claim 1, wherein the protein comprises a core amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO:1.
- 3. The animal cell of claim 2, wherein the protein comprises the core amino acid sequence and an endoplasmic reticulum (ER) export signal at the C-terminus of the core amino acid sequence.
 - 4. The animal cell of claim 3, wherein the core amino acid sequence is linked to the ER export signal through a linker.
- The animal cell of claim 3, wherein the ER export signal comprises the aminoacid sequence FCEYENEV.
 - 6. The animal cell of claim 3, wherein the protein comprises the amino acid sequence shown in SEQ ID NO:5.
 - 7. The animal cell of claim 1, wherein the animal cell is a neuronal cell, a muscle cell, or a stem cell.
- 20 8. The animal cell of claim 1, further comprising a second light-activated protein expressed on the cell membrane.
 - 9. The animal cell of claim 8, wherein the second light-activated protein is a protein capable of mediating a depolarizing current in the cell when the cell is illuminated with light.
- The animal cell of claim 9, wherein the second light-activated protein is selected from the group consisting of a VChR1, a DChR, a SFO, and a ChR2.
 - 11. A population of cells comprising the cell of any one of claims 1-10.

- 12. An isolated polynucleotide comprising a nucleotide sequence encoding a light-activated protein comprising a core amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO:1.
- 13. An expression vector comprising a nucleotide sequence encoding a lightactivated protein comprising a core amino acid sequence at least 95% identical to
 the sequence shown in SEQ ID NO:1.
 - 14. The expression vector of claim 13, wherein the expression vector is a viral vector.
 - 15. The expression vector of claim 14, wherein the viral vector is an AAV vector, a retroviral vector, an adenoviral vector, a HSV vector, or a lentiviral vector.
- 10 16. A method of using the cell of any one of claims 1-7, comprising activating the light-activated protein with light.
 - 17. A method of using the cell of any one of claims 8-10, comprising activating the light-activated proteins at their respective wavelengths.
 - 18. A non-human animal comprising a cell of any one of claims 1-10.
- 19. An animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein is responsive to blue light and is derived from *Dunaliella salina*, wherein the protein is capable of mediating a depolarizing current in the cell when the cell is illuminated with light.
- The animal cell of claim 19, wherein the protein comprises a core amino acid sequence at least 95% identical to the sequence shown in residues 25-365 of SEQ ID NO:2.
 - 21. The animal cell of claim 20, wherein the protein comprises the core amino acid sequence and an endoplasmic reticulum (ER) export signal at the C-terminus of the core amino acid sequence.
- 25 22. The animal cell of claim 21, wherein the core amino acid sequence is linked to the ER export signal through a linker.
 - 23. The animal cell of claim 22, wherein the ER export signal comprises the amino acid sequence FCEYENEV.

- 24. The animal cell of claim 19, wherein the animal cell is a neuronal cell, a muscle cell, or a stem cell.
- 25. The animal cell of claim 19, further comprising a second light-activated protein expressed on the cell membrane.
- The animal cell of claim 25, wherein the second light-activated protein is a protein capable of mediating a hyperpolarizing current in the cell when the cell is illuminated with light.
 - 27. The animal cell of claim 25, wherein the second light-activated protein is selected from the group consisting of a NpHR, a BR, an AR, and a GtR3.
- 10 28. A population of cells comprising the cell of any one of claims 19-27.
 - 29. An isolated polynucleotide comprising a nucleotide sequence encoding a light-activated protein comprising a core amino acid sequence at least 95% identical to the sequence shown in residues 25-365 of SEQ ID NO:2.
- 30. An expression vector comprising a nucleotide sequence encoding a lightactivated protein comprising a core amino acid sequence at least 95% identical to
 the sequence shown in residues 25-365 of SEQ ID NO:2.
 - 31. The expression vector of claim 29, wherein the expression vector is a viral vector.
 - 32. The expression vector of claim 30, wherein the viral vector is an AAV vector, a retroviral vector, an adenoviral vector, a HSV vector, or a lentiviral vector.
- 20 33. A method of using the cell of any one of claims 19-24, comprising activating the light-activated protein with light.
 - 34. A method of using the cell of any one of claims 24-27, comprising activating the light-activated proteins at their respective wavelengths.
 - 35. A non-human animal comprising a cell of any one of claims 19-27.
- 25 36. An animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least

- 95% identical to the sequence shown in SEQ ID NO:3, an ER export signal, and a membrane trafficking signal.
- 37. The animal cell of claim 36, wherein the light-activated protein comprises, from the N-terminus to the C-terminus, the core amino acid sequence, the ER export signal, and the membrane trafficking signal.

- 38. The animal cell of claim 36, wherein the light-activated protein comprises, from the N-terminus to the C-terminus, the core amino acid sequence, the membrane trafficking signal, and the ER export signal.
- The animal cell of claim 36, wherein the ER export signal comprises the aminoacid sequence FCEYENEV.
 - 40. The animal cell of claim 36, wherein the membrane trafficking signal comprises the amino acid sequence KSRITSEGEYIPLDQIDINV.
 - 41. The animal cell of claim 36, wherein the animal cell is a neuronal cell, a muscle cell, or a stem cell.
- 15 42. The animal cell of claim 36, further comprising a second light-activated protein expressed on the cell membrane.
 - 43. The animal cell of claim 42, wherein the second light-activated protein is a protein capable of mediating a depolarizing current in the cell when the cell is illuminated with light.
- The animal cell of claim 42, wherein the second light-activated protein is selected from the group consisting of a VChR1, a DChR, a SFO, and a ChR2.
 - 45. A population of cells comprising the cell of any one of claims 36-44.
- 46. A polynucleotide for use in preparation of the cell of any one of claims 36-44 or the population of cells of claim 45, comprising a nucleotide sequence encoding a light-activated protein comprising a core amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO:3, an ER export signal, and a membrane trafficking signal.

- 47. The polynucleotide of claim 46, wherein the polynucleotide is in an expression vector.
- 48. The polynucleotide of claim 47, wherein the expression vector is a viral vector.
- 49. The polynucleotide of claim 48, wherein the viral vector is an AAV vector, a retroviral vector, an adenoviral vector, a HSV vector, or a lentiviral vector.

- 50. A method of using the cell of any one of claims 36-41, comprising activating the light-activated protein with light having a wavelength between about 580 and about 630 nm.
- 51. A method of using the cell of any one of claims 42-44, comprising activating the light-activated proteins at their respective wavelengths.
 - 52. A non-human animal comprising a cell of any one of claims 36-44.
 - 53. An animal cell comprising a light-activated protein expressed on the cell membrane, wherein the protein comprises a core amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO:4, an ER export signal, and a membrane trafficking signal.
 - 54. The animal cell of claim 53, wherein the light-activated protein comprises, from the N-terminus to the C-terminus, the core amino acid sequence, the ER export signal, and the membrane trafficking signal.
- 55. The animal cell of claim 53, wherein the light-activated protein comprises, from the N-terminus to the C-terminus, the core amino acid sequence, the membrane trafficking signal, and the ER export signal.
 - 56. The animal cell of claim 53, wherein the ER export signal comprises the amino acid sequence FCEYENEV.
- 57. The animal cell of claim 53, wherein the membrane trafficking signal comprises the amino acid sequence KSRITSEGEYIPLDQIDINV.
 - 58. The animal cell of claim 53, wherein the animal cell is a neuronal cell, a muscle cell, or a stem cell.

- 59. The animal cell of claim 53, further comprising a second light-activated protein expressed on the cell membrane.
- 60. The animal cell of claim 59, wherein the second light-activated protein is a protein capable of mediating a depolarizing current in the cell when the cell is illuminated with light.
- 61. The animal cell of claim 59, wherein the second light-activated protein is selected from the group consisting of a VChR1, a DChR, a SFO, and a ChR2.
- 62. A population of cells comprising the cell of any one of claims 53-61.

- 63. A polynucleotide for use in preparation of the cell of any one of claims 53-61 or the population of cells of claim 62, comprising a nucleotide sequence encoding a light-activated protein comprising a core amino acid sequence at least 95% identical to the sequence shown in SEQ ID NO:4, an ER export signal, and a membrane trafficking signal.
- 64. The polynucleotide of claim 63, wherein the polynucleotide is in an expression vector.
 - 65. The polynucleotide of claim 64, wherein the expression vector is a viral vector.
 - 66. The polynucleotide of claim 65, wherein the viral vector is an AAV vector, a retroviral vector, an adenoviral vector, a HSV vector, or a lentiviral vector.
- 67. A method of using the cell of any one of claims 53-58, comprising activating the light-activated protein with light.
 - 68. A method of using the cell of any one of claims 59-61, comprising activating the light-activated proteins at their respective wavelengths.
 - 69. A non-human animal comprising a cell of any one of claims 59-61.
- 70. A method of identifying transsynaptic connection between neuronal cells in an animal or a tissue, comprising: a) administering a first viral vector encoding a Cre recombinase fused to a transcellular tracer protein to neuronal cells in region A of the animal or tissue; b) administering a second viral vector encoding a light-activated protein to neuronal cells in region B of the animal or tissue, wherein the

expression of the light-activated protein depends on the presence of the Cre recombinase; and c) identifying neuronal cells expressing the light-activated protein in region B, wherein the expression of the light-activated protein in the neuronal cells indicating that these cells are in transsynaptic connection with the cells in region A.

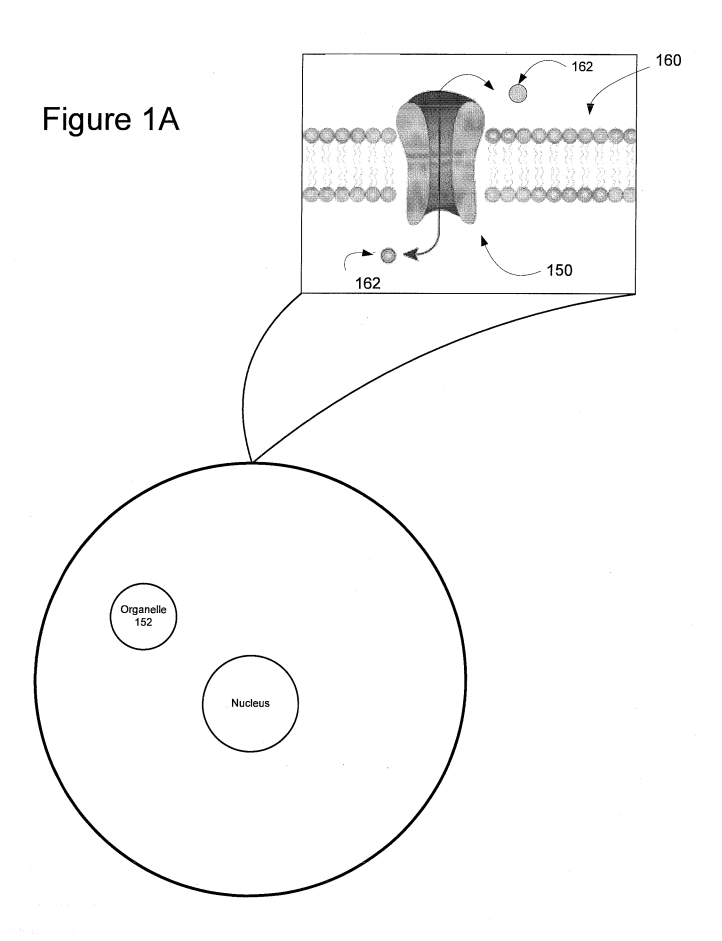
- 71. A method of generating optical control of targeted neuronal cells in an animal or tissue, comprising: a) administering a first viral vector expressing a Cre recombinase fused to a transcellular tracer protein to region A of the animal or tissue; b) administering a second viral vector encoding a light-activated protein to region B of the animal or tissue, wherein the expression of the light-activated protein depends on the presence of the Cre recombinase, and wherein the neuronal cells in region A and in region B are in transsynaptic connected; and c) controlling action potential of a neuronal cell in region B with light that activates the protein.
- A method of controlling action potential of a neuron in an animal, comprising activating a light-activated protein in the neuron with light to generate action potential change, wherein expression of the light-activated protein in the neuron is generated by a) administering a first viral vector expressing a Cre recombinase fused to a transcellular tracer protein to region A of the animal, b) administering a second viral vector encoding a light-activated protein to region B of the animal which contains the neuron, wherein the expression of the light-activated protein depends on the presence of the Cre recombinase, wherein the neurons in region A and region B are in transsynaptic connection.
- 73. The method of any one of claims 70-72, wherein the viral vector is a viral vector selected from the group consisting of AAV, HSV, and lentivirus.
 - 74. The method of any one of claims 70-72, wherein the transcellular tracer protein is wheat germ agglutinin (WGA) or tetanus toxin-fragment C (TTC).
- 75. An animal cell comprising an integrated exogenous molecule which expresses a proton pump responsive to blue light, the exogenous molecule is derived from *Guillardia theta*.

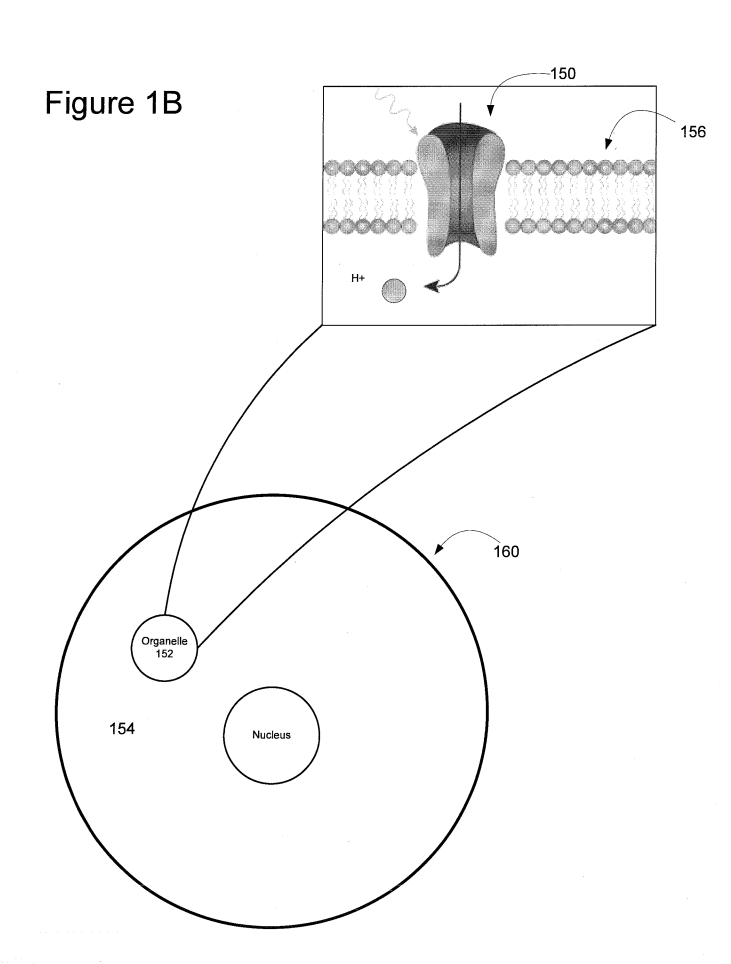
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76. The animal cell of claim 75, wherein the animal cell is a neural cell, a muscle cell, or a cell line.

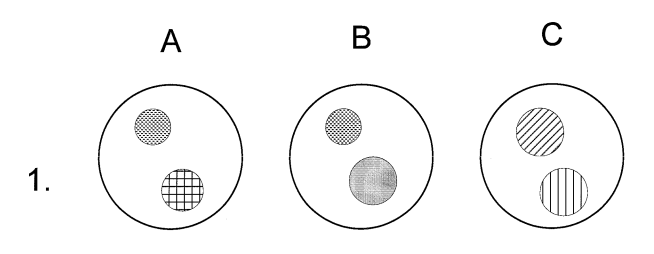
77. The animal cell of claim 75, wherein the exogenous molecule contains GtR3 amino acid sequence shown in Table 3.

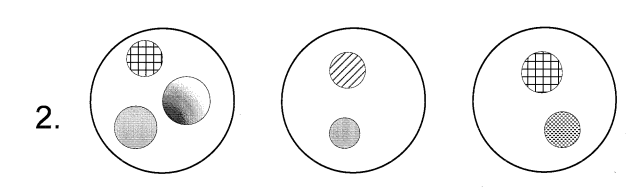
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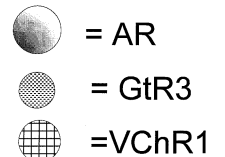


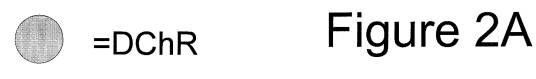
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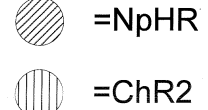


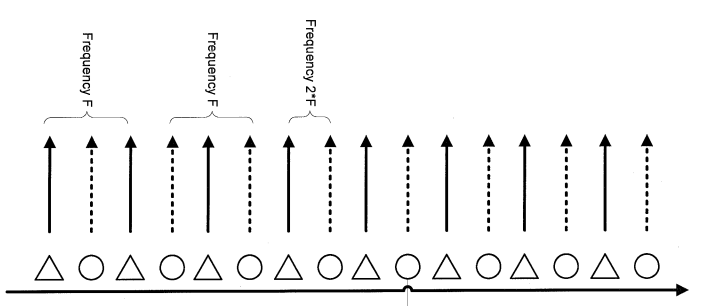




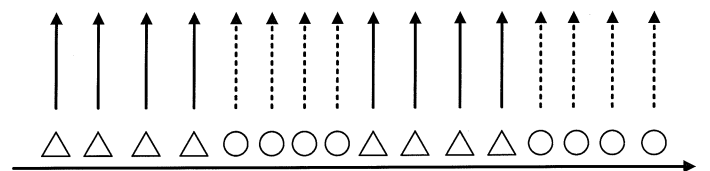








Example 3



Example 4

Figure 2B

FIG. 3

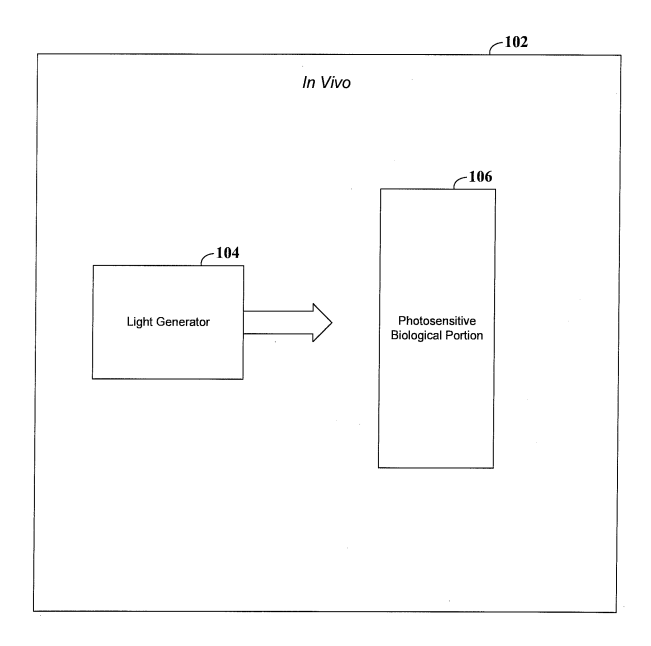
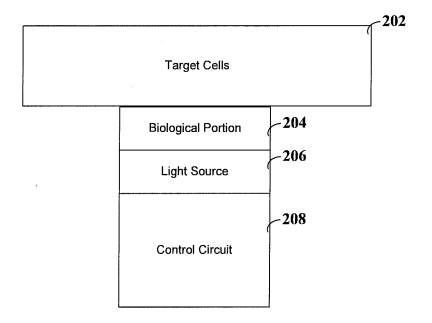


FIG. 4



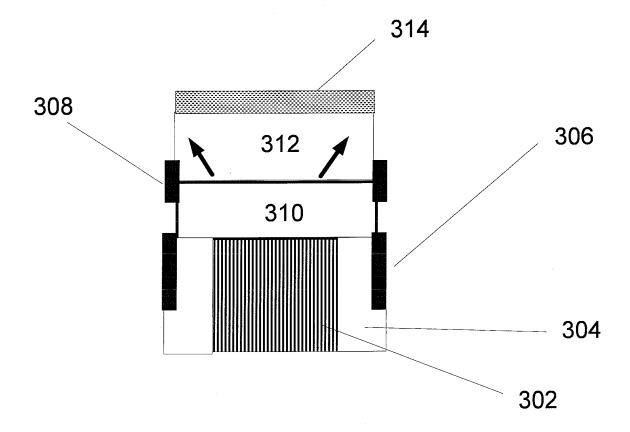
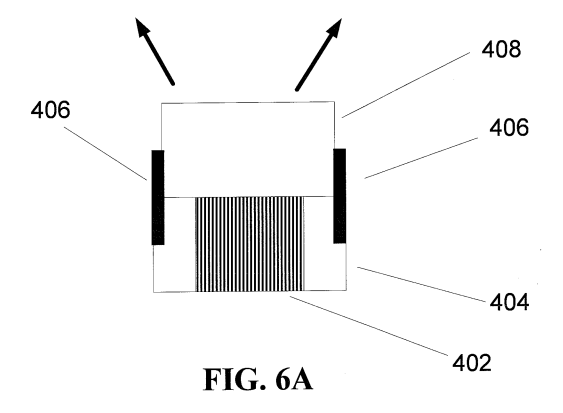
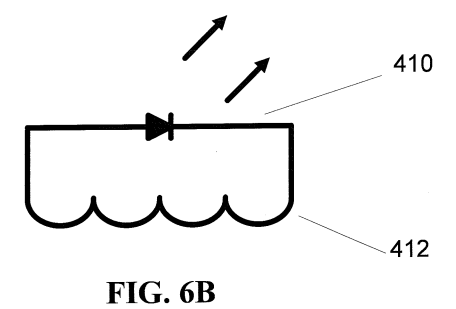
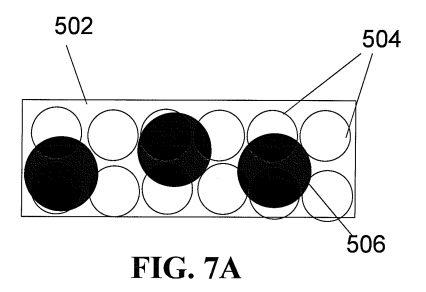


FIG. 5







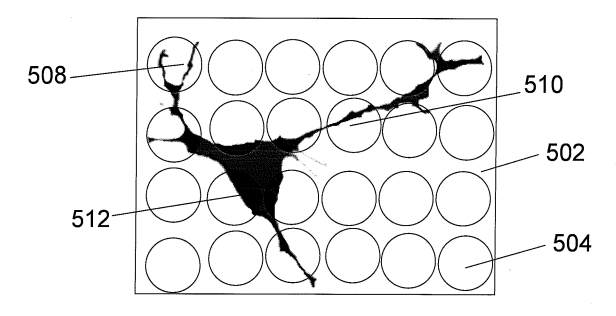


FIG. 7B

FIG. 8A

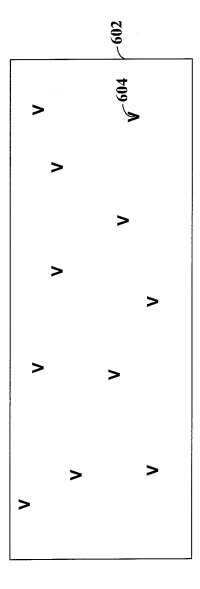
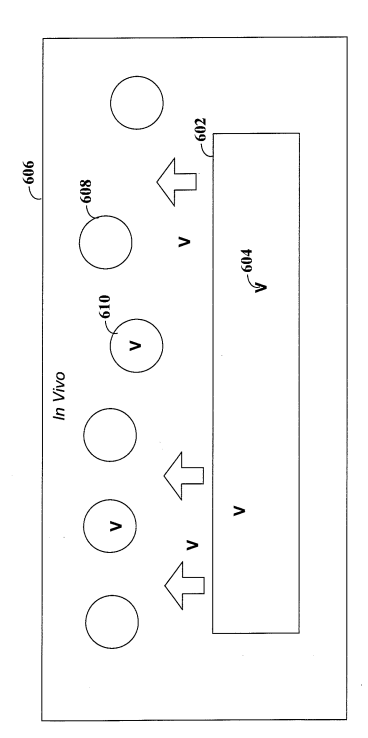
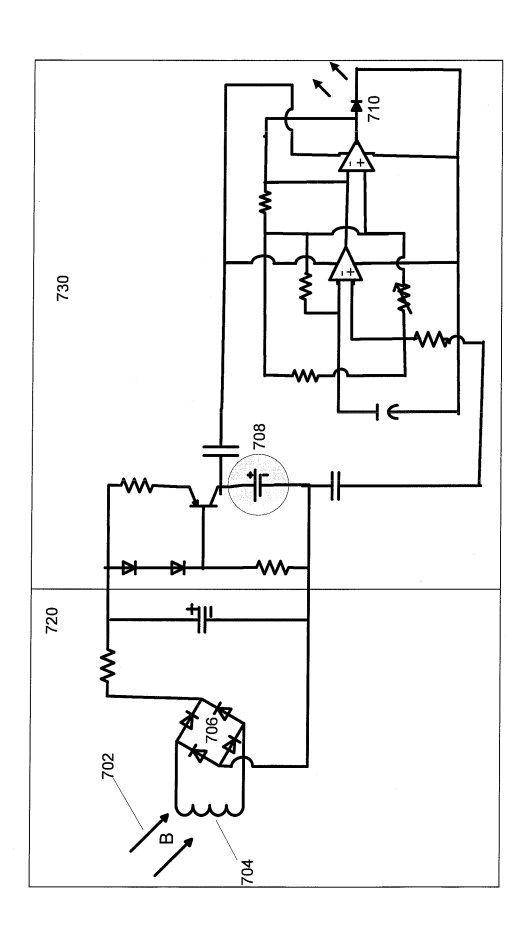
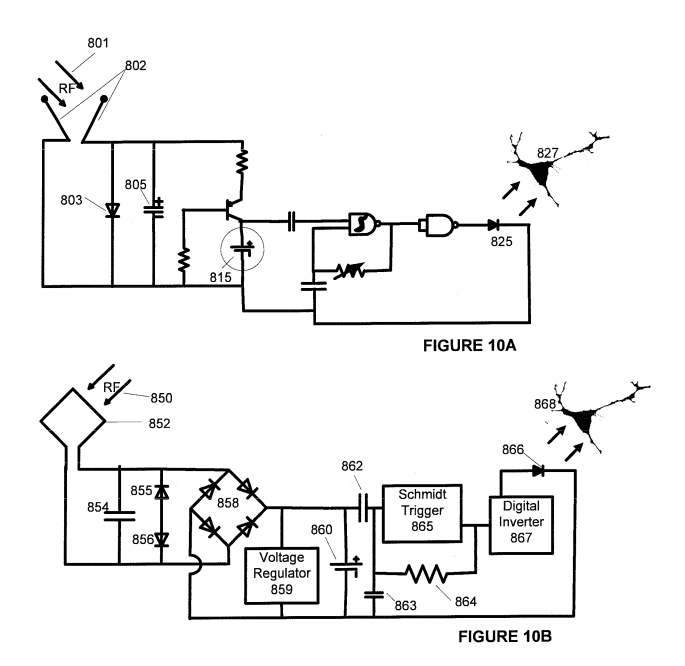


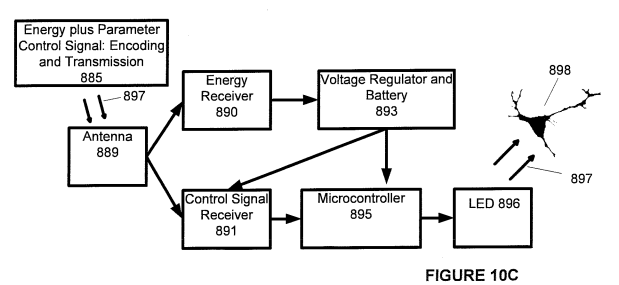
FIG. 8B











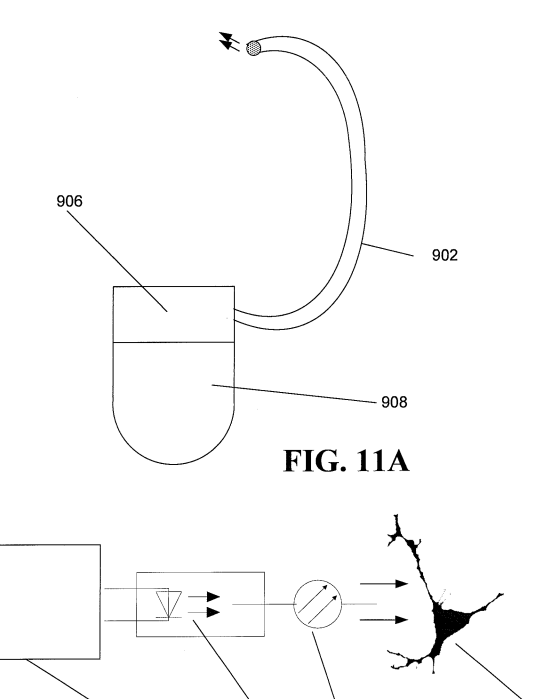


FIG. 11B

902

906

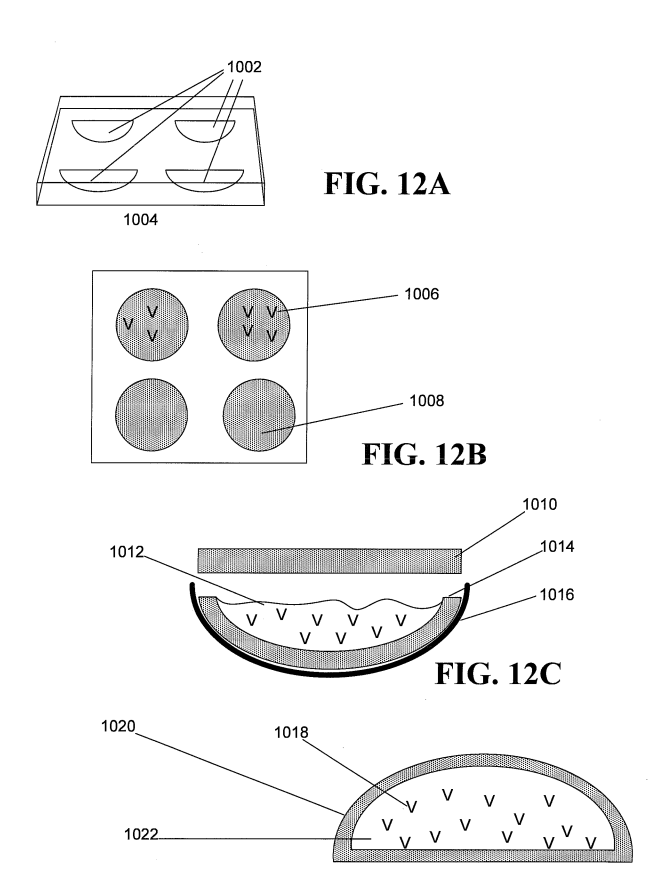
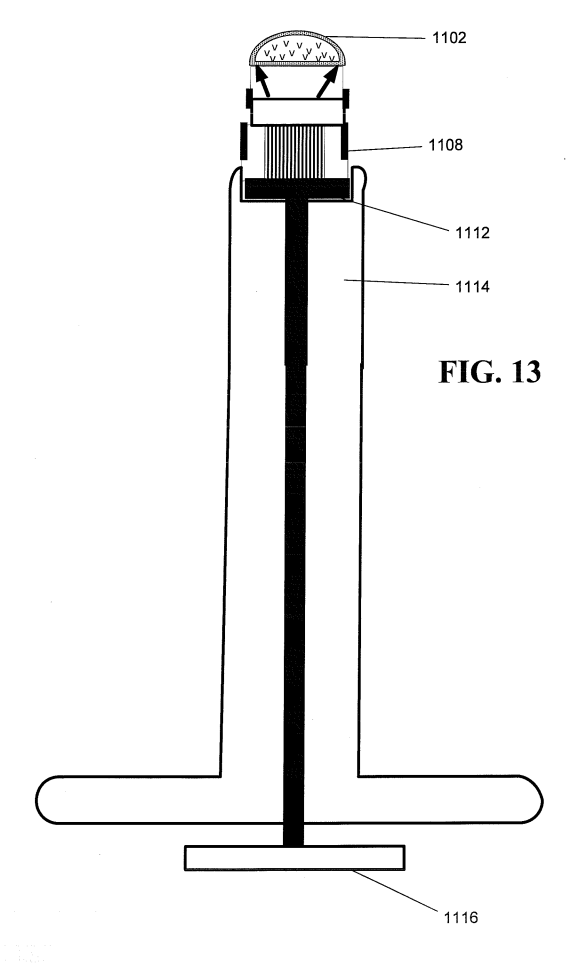
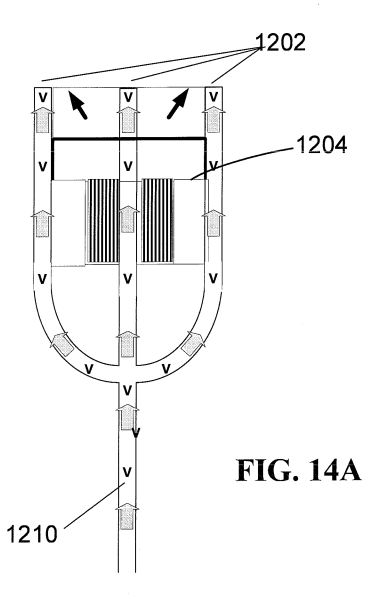
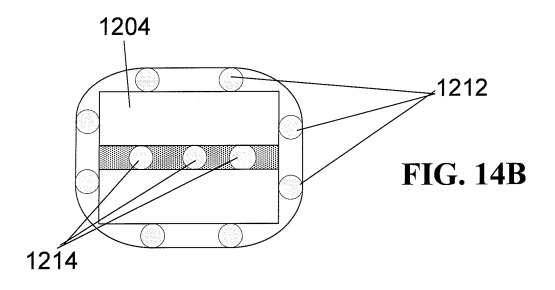
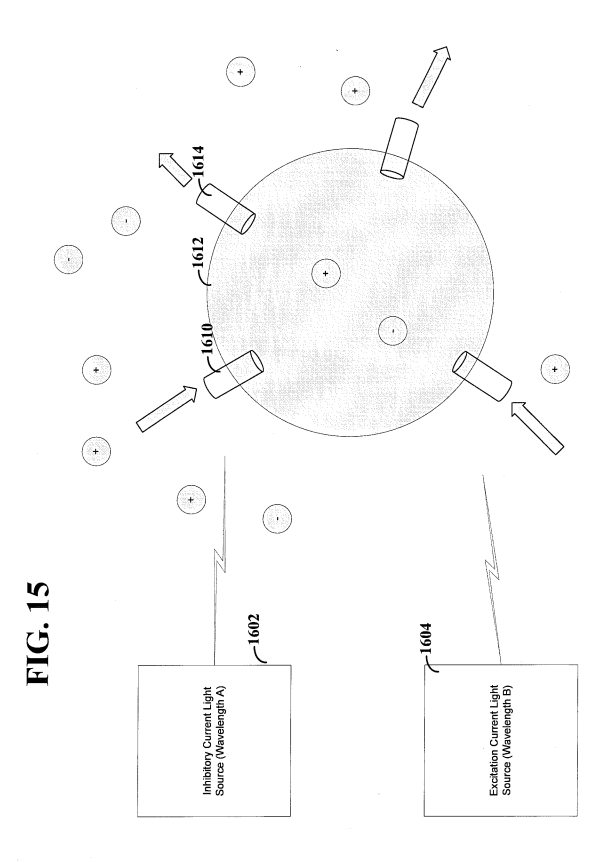


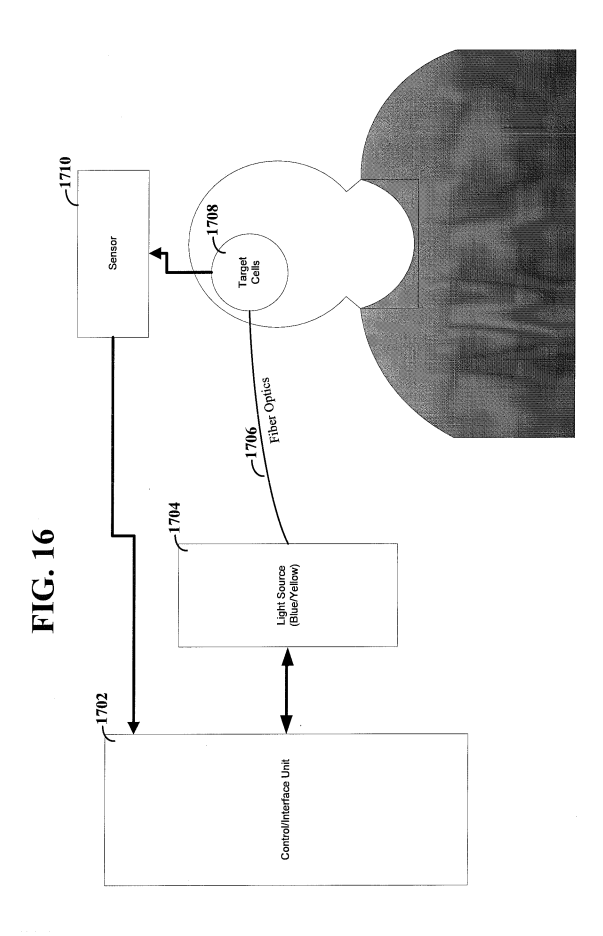
FIG. 12D

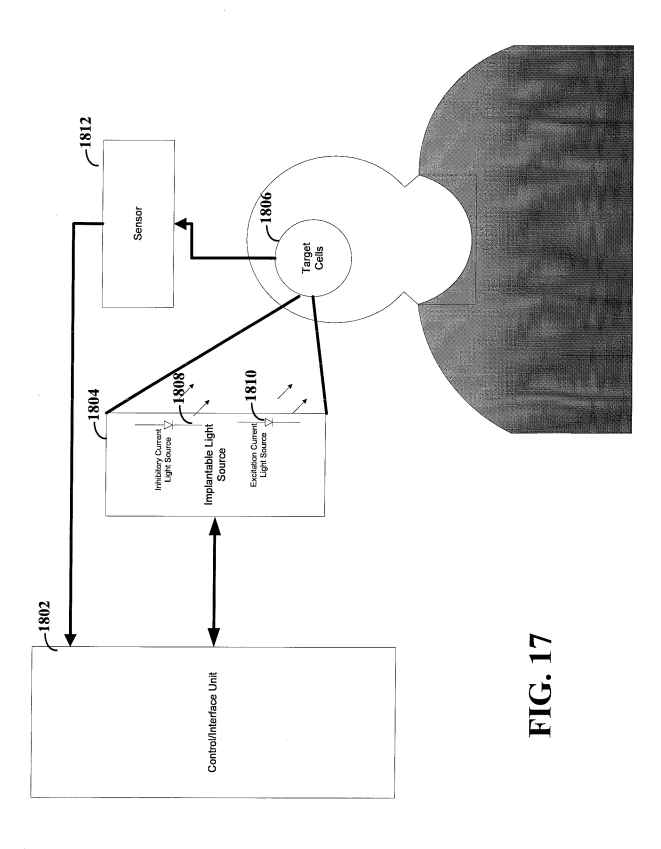


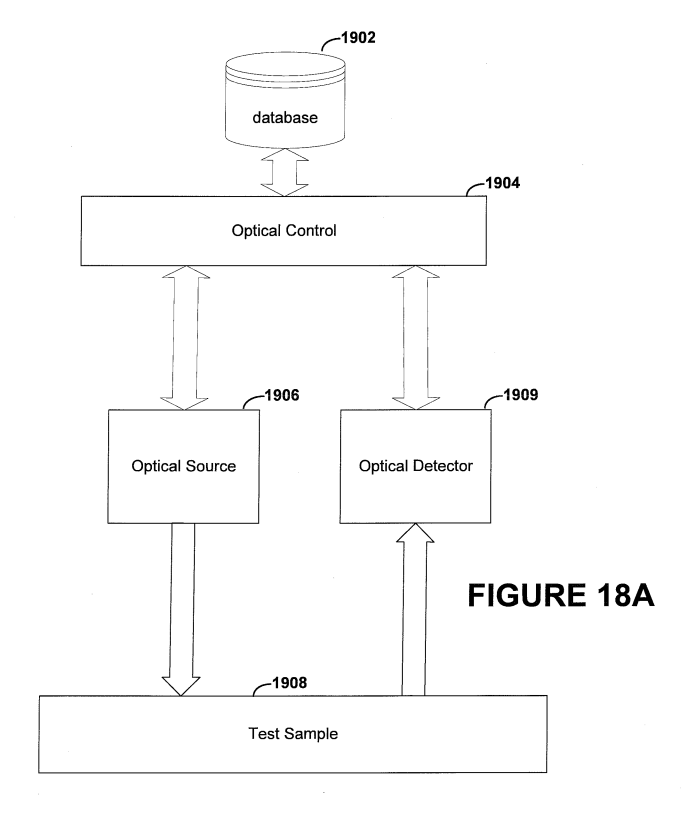














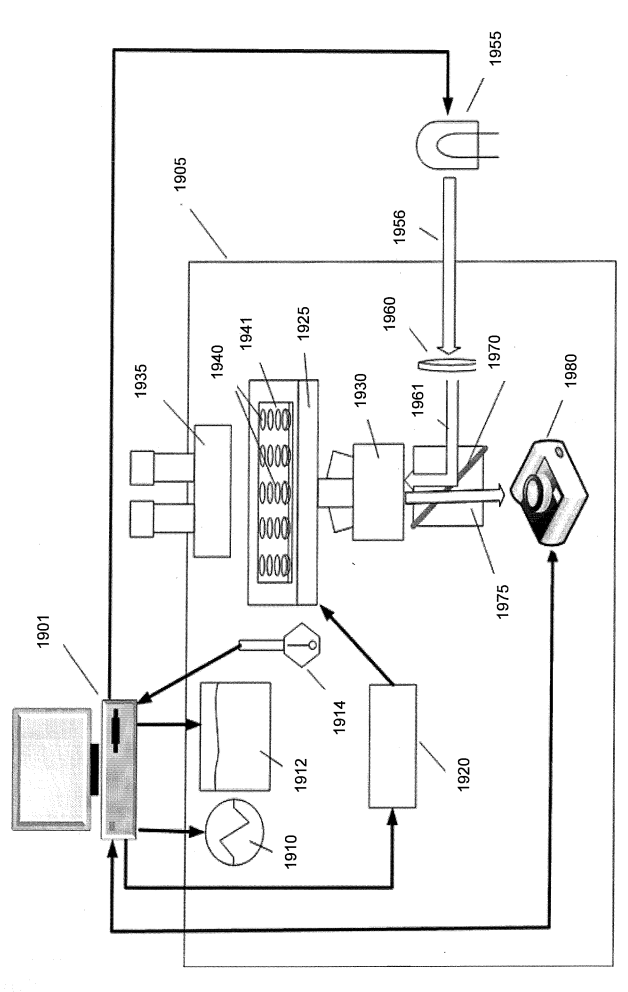
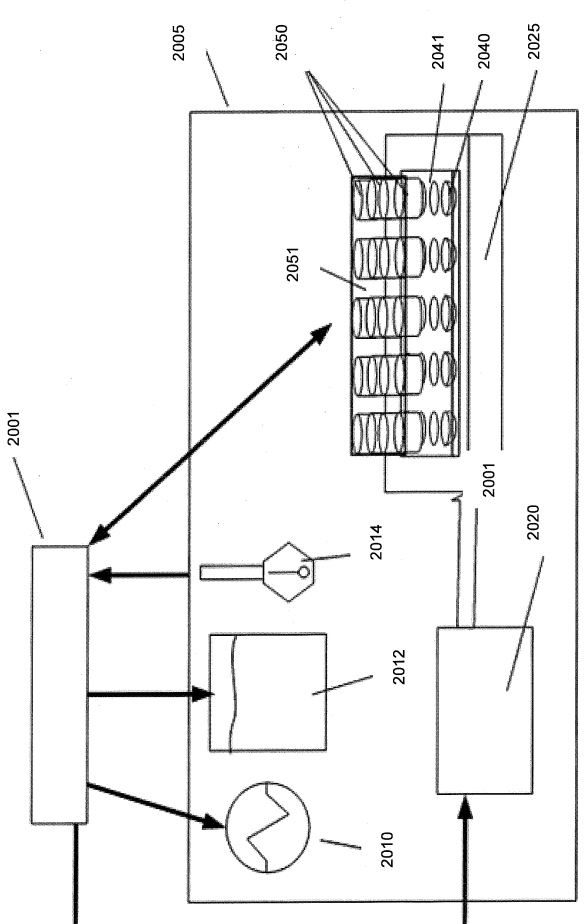




Figure 19



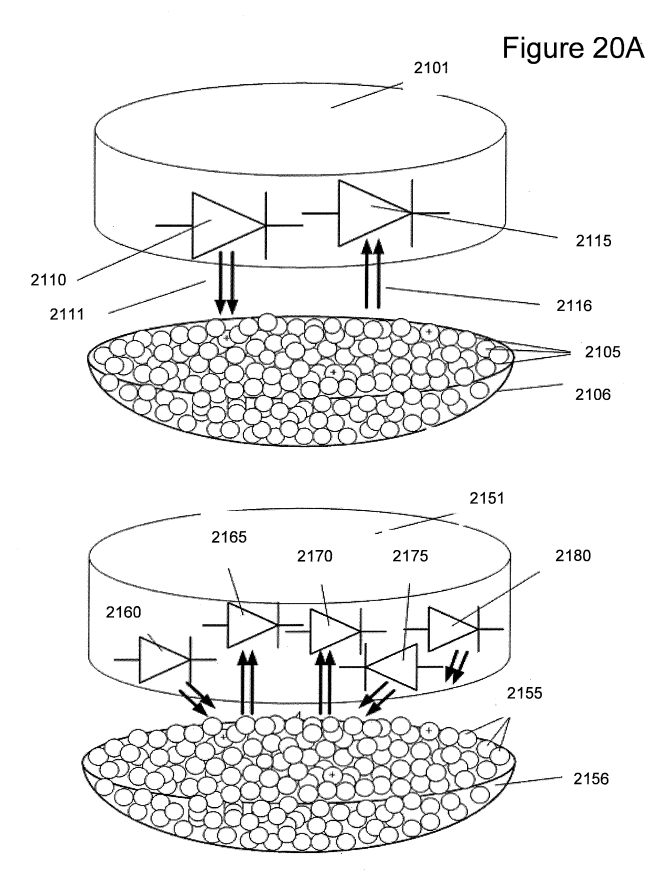


Figure 20B

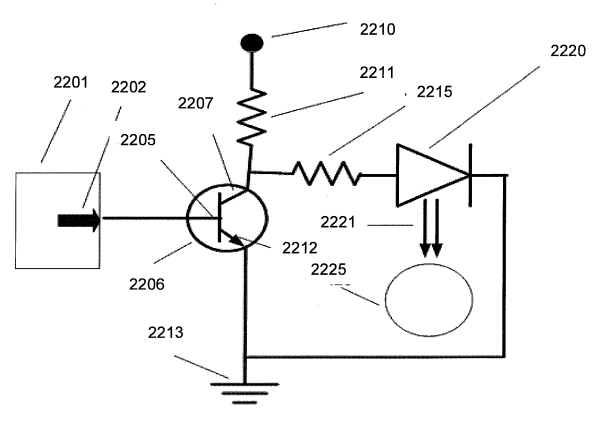


Figure 21A

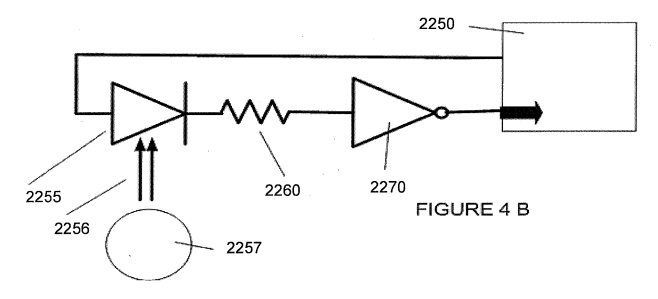


Figure 21B

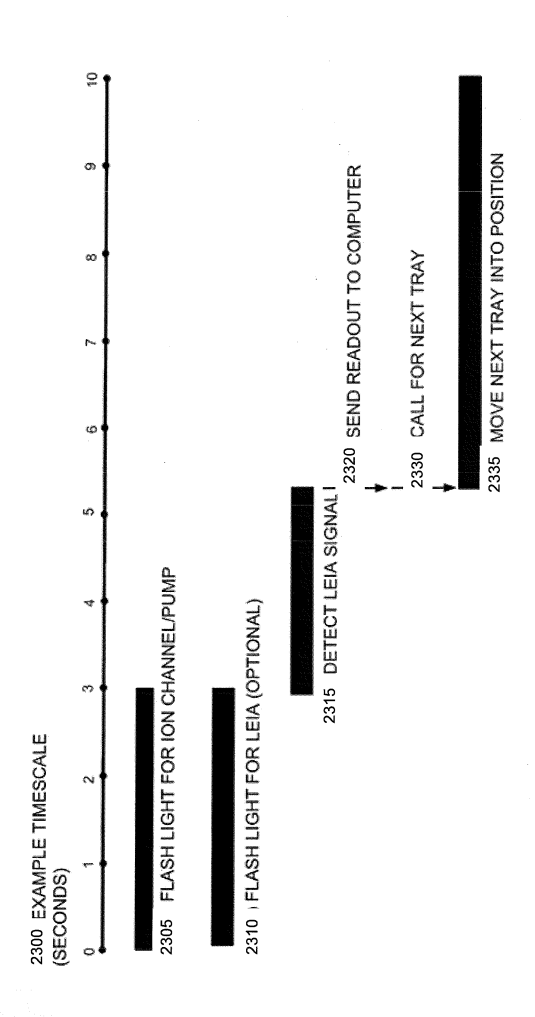
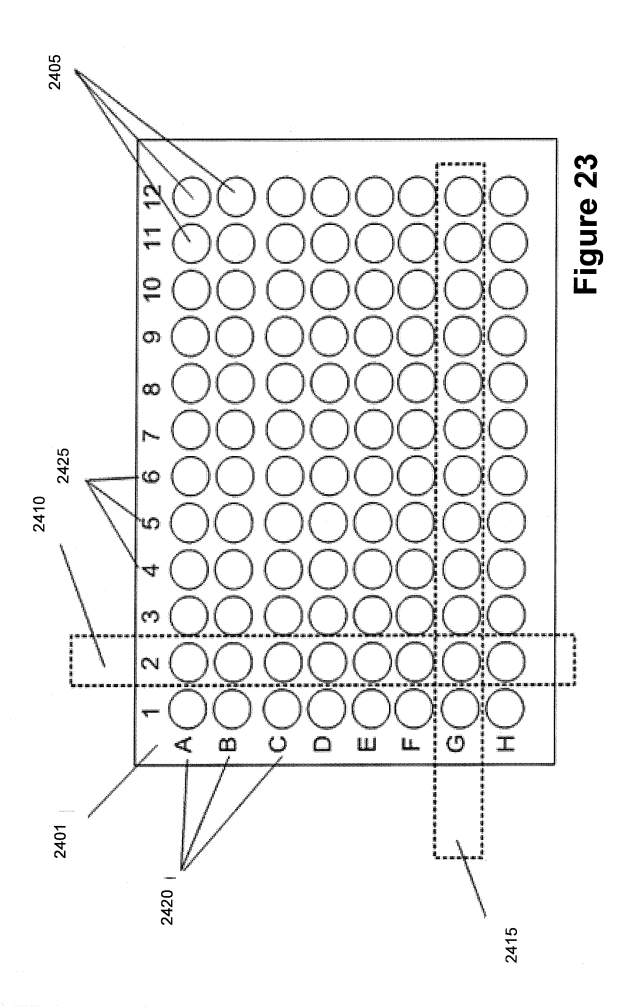
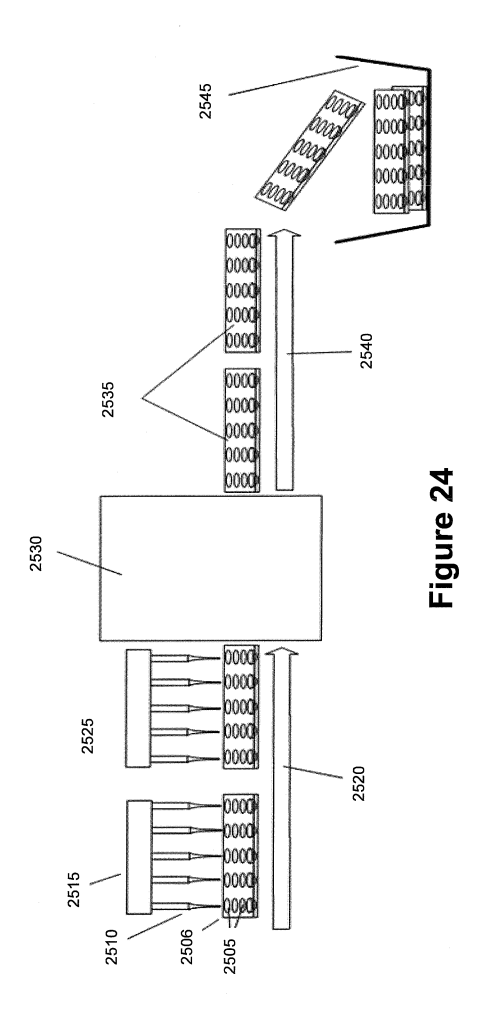


Figure 22





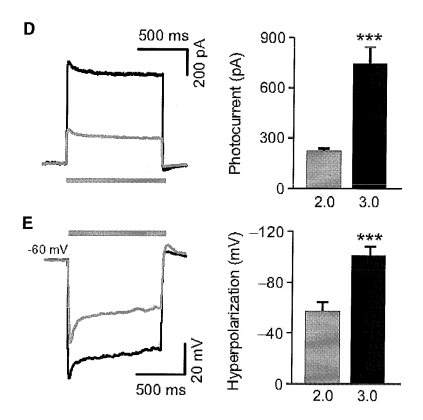


FIG. 25

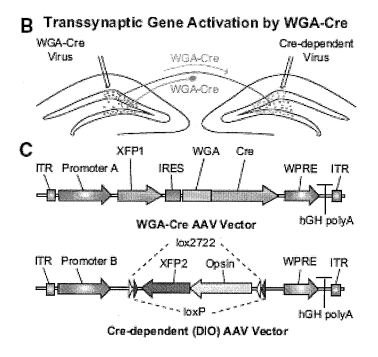


FIG. 26

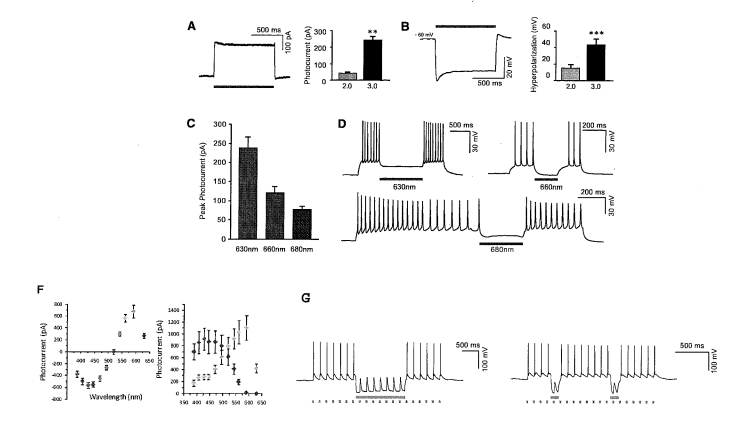


FIG. 27

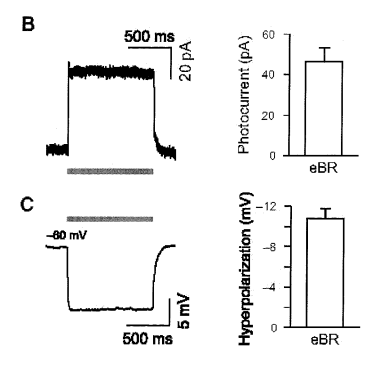


FIG. 28

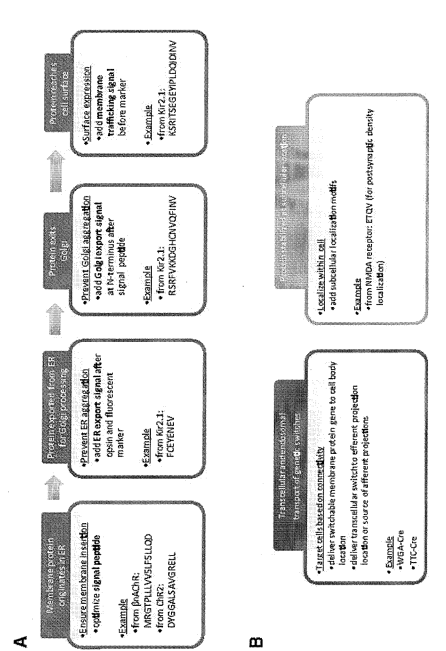


FIG. 29

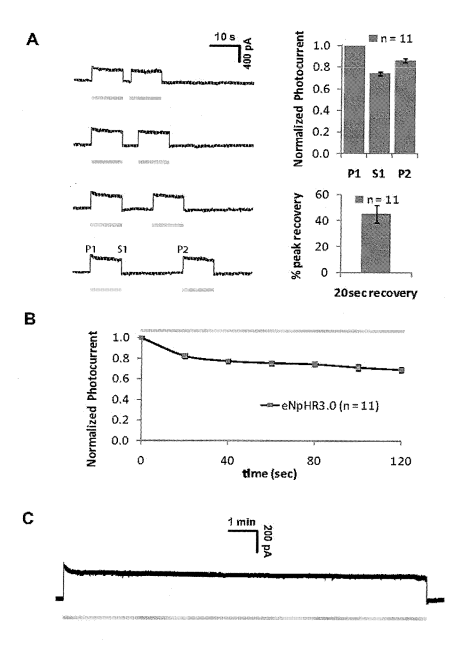


FIG. 30

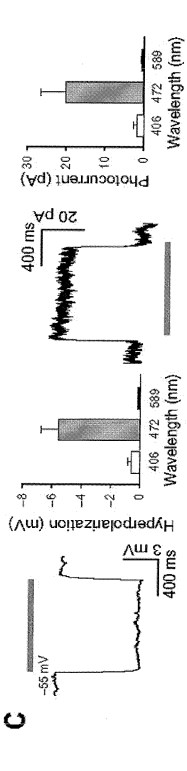


FIG. 31