2007 7.013 Problem Set 6
Due before 5 PM on FRIDAY, April 27, 2007.
Turn answers in to the box outside of 68-120.
PLEASE WRITE YOUR ANSWERS ON THIS PRINTOUT.

Question 1.

1a. This is a diagram showing changes in membrane potential before, during and after an action potential. Label the resting potential, action potential, and repolarization period. Label the X and Y axes.

1b. Indicate when and/or where the following are active/open (for each channel/pump, check one or more column for “when” and one or more for “where”):

<table>
<thead>
<tr>
<th>Channel/Pump</th>
<th>resting pot</th>
<th>when</th>
<th>action pot</th>
<th>repolarization</th>
<th>where</th>
</tr>
</thead>
<tbody>
<tr>
<td>voltage gated Na+ channels</td>
<td>X</td>
<td>X</td>
<td>(X)</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>voltage gated K+ channels</td>
<td>(X)</td>
<td>X</td>
<td></td>
<td>(X)</td>
<td></td>
</tr>
<tr>
<td>voltage gated Cl- channels</td>
<td>(X)</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>voltage gated Ca²⁺ channels</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>open Na+ channels</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>open K+ channels</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>open Cl- channels</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Na+/K+ pump</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
</tbody>
</table>

1c. After vigorous exercise, levels of K+ are elevated in the blood. Patients with hyperkalemic periodic paralysis (HPP) have episodes of muscle weakness (paralysis) after exercise. Genetic studies have shown that the disease is caused by a point mutation in the voltage gated Na+ channel found at neuromuscular junctions. This
mutation leads to failure of the Na+ channels to completely inactivate after depolarization.

(i) After depolarization, would the concentration of Na+ ions inside the neuron be higher/lower/the same in HPP patients after exercise than they would have been before exercise? Explain (15 words or fewer).

Higher; Na+ continues to flow in, even after depolarization

(ii) After exercise, neurons from patients with HPP show a resting potential of –40mV (rather than the normal –60mV). At this membrane potential, most voltage gated Na+ channels become inactive. Would an action potential be generated more frequently/less frequently/at the same rate than before exercise? Why? (15 words or fewer)

Less frequently; voltage gated Na+ channels are inactive

(i) Why does the resting potential decrease after exercise in HPP patients (that is, why is the increase K+ concentration a problem)? (15 words or fewer)

K+ outward diffusion is lower because of the higher outside K+ concentration.

OR

A normal K+ gradient cannot be established.

1d. Patients with episodic ataxia exhibit normal neurological function except during periods of mental or physical stress, which can elicit a generalized ataxia (loss of coordinated movements). This disorder is due to a deficit in the neuronal voltage gated K+ channel, which is needed during repolarization.

(i) Would this increase/decrease/not change the resting potential?

Increase: increase in mV, more positive, closer to threshold

(ii) Would this increase/decrease/not change the threshold potential?

not change

(iii) Would this increase/decrease/not alter the tendency of the neuron to fire during periods of stress? Why? (15 words or fewer)

increase; the resting potential is closer to threshold.

Note: In fact, ataxia results from too frequent action potentials.

1e. Myotonia congenita is a neuromuscular disorder that maps to a human chloride channel gene. The channel is a homotetramer, with each subunit having 1000 amino acids. The coding sequence is interrupted by 22 introns and has 12 putative transmembrane domains and a conserved intracellular domain. Sequence analysis of a large number of people, including those with myotonia congenita, as well as healthy controls, has revealed many variants of this gene, including 14 positions where single amino acids may vary, 3 variants where nonsense codons would truncate the protein, and 2 deletions in various exons.
(i) How would you further examine these data, without performing any experiment, to make an initial conclusion as to which variant(s) was most likely to cause myotonia congenita? (15 words or fewer)

*Compare gene sequences of affected individuals to find regions of frequent mutation present in the myotonia patients that are not present in normal controls.*

*Sort variants to find those specifically present in myotonia patients.*

In people with *myotonia congenita*, neuronal open chloride channels conduct chloride ions more slowly than in persons without the disorder.

(ii) In such patients, would you predict that the resting potential of the neuron would return to normal more slowly/more quickly/at the same rate as that of unaffected people? Explain in 15 words or fewer.

*More slowly; Cl- inward movement contributes to repolarization.*

Question 2.

2a. Morphine is the primary bioactive opiate present in the sap isolated from the ripening seed case of the opium poppy. Morphine acts as a neurotransmitter that inhibits the sensation of pain by binding to G-protein coupled receptors. Endogenous ligands for opioid receptors exist, including enkephalins and endorphins. These also function to inhibit response to pain.

(i) To what major class of neurotransmitter receptors does the opioid receptor belong? (1 word – not “G-protein”!)

*Metabotropic*

(ii) How does this receptor class alter ion channel function? (15 words or fewer)

*Activates ion channels indirectly through a second messenger.*

(iii) Where would opioid receptors be localized (presynaptic membrane/postsynaptic membrane/both)?

*Postsynaptic membrane*

(iv) Naloxone is an opioid antagonist, which prevents morphine or enkephalin from binding to opioid receptors. What is the biochemical term for this type of inhibitor? (1 word)

*Competitive*

2b. Morphine increases synaptic transmission between some target neurons, while decreasing synaptic transmission between other neurons.

(i) With morphine treatment, a low level of synaptic activity in some target neurons is increased. Do you expect this increase to be accompanied by an
increased rate of action potential generation in the post-synaptic neuron, or generation of action potentials with larger amplitude than prior to opioid treatment? Explain (15 words or fewer).

*Increased rate of action potential generation because action potentials are binary (on/off). The amplitude never changes.*

(ii) If synaptic transmission were decreased by morphine treatment, would you expect the resting potential in the post-synaptic cell would increase/decrease/stay the same? Explain in 15 words or fewer.

*One would expect the resting potential to be further from threshold (increased difference)*

(iii) Would you expect enkephalin protein to be present at the synapse in the presynaptic/postsynaptic cell? Why (15 words or less)?

*presynaptic cell; enkephalin is released to bind the receptor at the postsynaptic terminal.*

(iv) In which subcellular structures at the synapse would enkephalins be present? (2 words)

*presynaptic vesicle*
Question 3.
3a. (i) What is a “growth cone”? (15 words or fewer)
   migrating tip of a growing nerve cell axon or dendrite
   (ii) What is a pioneer axon? (15 words or fewer)
   the first pathfinding axon

3b. In Drosophila, pioneer axons grow from the sides of the animal towards the midline, due to attractive signals from the midline. However, subsequent to crossing the midline, the axons are repelled by signals from it so they do not recross. slit and robo are loss of function mutants in Drosophila, that have identical phenotypes. In both mutants, axons are attracted to the midline normally, but then cross and recross it multiple times (Fig. 1).

Fig. 1 Axon pathfinding in wild type and slit or robo loss of function mutants (stippled). Initial growth is towards the midline. Arrows indicate direction of axon growth.

It was speculated that one of these genes might encode a ligand secreted by the midline, and one a receptor for an axonal repulsion system. The following assays were performed to determine whether slit acts as a ligand or a receptor.

(i) In the experiment diagrammed below, a slit mutant midline (stippled) replaced the midline cells of a wild type embryo, to make a mosaic slit/wild type embryo. If slit were a ligand, draw on the diagram below how the wild type axons of this embryo would grow as they reached the midline, and explain your result (15 words or fewer).

   like the slit mutant – cross and recross because there is no signal from the midline to prevent re-crossing
(ii) In a similar experiment to that in (i), if slit were a receptor, draw on the diagram below how the wild type axons of this embryo would grow when they reached the midline, and explain your result (15 words or fewer)?

```
Slit mutant midline
in otherwise wildtype embryo

like the wildtype axons; the axons have the WT receptor
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(iii) In a further experiment, you remove the midline from a wild type embryo, and, as diagrammed below, observe that axons are randomly distributed, with no concentration at the midline, as in Fig 1. Explain these results in 15 words or fewer.

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The midline produces attractive signals that direct axon growth.
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3c. In fact, slit is the ligand and robo the receptor in this axon repulsion system (note that these facts will not help you answer 3b!). slit and robo genes are also present in vertebrates. You test whether slit has similar activity in rat as in Drosophila. In the following experiments, explants of lateral spinal cord from rat were placed near a bead coated with slit protein, or with a control protein BSA. Axon outgrowth was monitored from the part of the explant near the bead (proximal, P) or distal (D) to the bead.

(i) Explain the difference in proximal versus distal outgrowth in slit and control bead assays (15 words or fewer).
*Slit inhibits proximal but not distal outgrowth, because it acts in a concentration-dependent way.*

(ii) If the explant was made from a spinal cord that did not contain the robo receptor, what result would you expect and why (15 words or fewer)?
*The result would look like that in the BSA+bead control because without the receptor, there is no inhibition.*

(iii) If the explant was made from a spinal cord defective in lamellipodial and filopodial formation, what would you expect to see, and why (15 words or fewer)?
*No axon growth because the growth cone would be defective (collapsed).*
Question 4.

4a. Crohn’s disease is a poorly understood disorder that involves an aberrant inflammatory response in the intestines of affected individuals. Affected monozygotic (so-called “identical”) twins have a high concordance for Crohn’s disease.

(i) Does this observation suggest a genetic or epigenetic component for Crohn’s disease? Explain in 15 words or fewer.
*Genetic, because a shared trait in twins will generally be due to primary DNA sequence.*

(ii) Do the above observations with twins rule out an environmental component to Crohn’s disease? Explain in 15 words or fewer.
*No, because twins also often share the same environment.*

4b. Given that you can’t do controlled genetic crosses with human subjects, what other evidence would indicate whether there is a genetic basis for Crohn’s disease? Answer in 15 words or fewer.

*Determine whether there is a family history of this disorder ---OR--
Many patients would be expected to have mutations in the same gene(s) if the disease is genetic.*

4c. Ulcerative colitis (UC) is a disorder that also affects the digestive system through an inflammatory response and is often misdiagnosed as Crohn’s disease. Nonetheless, these two diseases can be distinguished. A gene that encodes a protein called NOD2 has been implicated in diseases affecting inflammation of the digestive system. In order to determine whether NOD2 is involved in either Crohn’s or UC, you examine the occurrence of NOD2 mutations in three different populations: individuals affected by Crohn’s disease, individuals affected by UC, and individuals that display neither disorder (the control group). You examine the sequences of the NOD2 gene in each of these individuals. You find the following:

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Controls (n=272)</th>
<th>Crohn’s disease (n=304)</th>
<th>Ulcerative colitis (n=65)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NOD2+/NOD2+</td>
<td>248 (91·2%)</td>
<td>227 (74·7%)</td>
<td>61 (93·8%)</td>
</tr>
<tr>
<td>nod2-/NOD2+</td>
<td>24 (8·8%)</td>
<td>57 (18·8%)</td>
<td>4 (6·2%)</td>
</tr>
<tr>
<td>nod2-/nod2-</td>
<td>0</td>
<td>20 (6·5%)</td>
<td>0</td>
</tr>
</tbody>
</table>

NOD2+ is wildtype, nod2- denotes mutant (Adapted from Hampe et al., 2007, The Lancet 357: 1925)

(i) Based on these observations, which disorder is more strongly associated with mutations in NOD2? (15 words or fewer)
*Crohn’s disease because the frequency of mutation in NOD2 is higher than that in the general population.*

(ii) Based on the above data, can you reasonably infer that mutations in NOD2 are the single strongest genetic determinant for the disease? Explain in 15 words or fewer.
*No, most patients with Crohn’s are WT for NOD2.*