Instructions: Using the information that you have learned in class and from your textbook, complete the questions in the case. Internet sites and resources are cited below in the case study for you to use to look up key information.

Part I – Genetic Testing
You are a second-year medical student in an innovative medical school that allows you to get in-depth clinical experience early in your medical education. You are currently assigned to work with Dr. Aikenhed, a pediatric oncologist. She has done a biopsy on a tumor-like growth in the adrenal gland of her 17-year-old patient, Lee F. You record the following notes while Dr. Aikenhed takes a detailed family history from the patient’s parents:

- Lee has a sister, Leah (age 10), and a brother, Luke (age 6). Both are healthy.
- Lee’s mother, Grace, was diagnosed with bilateral breast cancer last year at age 35. Lee’s father, Brian, and Lee’s paternal grandparents have no history of cancer.
- Grace is the youngest of four children. Her eldest brother, Greg (age 42), and sister, Greta (age 40), have never had any signs or symptoms of cancer. She had a brother, Geoff, who died of leukemia at age 8. He was the third child of Grace’s parents.
- Greg’s two fraternal twin daughters have no signs or symptoms of cancer.
- Grace’s father, Roger, died of a soft tissue sarcoma at age 35. Grace’s mother, Renee, is still living and in excellent health.
- Roger’s mother died of a brain tumor at age 30. Roger was her only child.

Questions
1. Draw the pedigree for this family in the space provided below. Make sure to indicate Lee, the proband, with an arrow and use the proper symbol for individuals who have died from cancer. Remember to also label each generation with roman numerals.
It is clear from the pedigree that there is a pattern. You strongly suspect that the high incidence of cancer in this family is caused by an inherited genetic defect. Due to the severity of the phenotype, the defective allele is probably rare in the population.

2. Complete the following table to help organize your thoughts. Indicate in the table if a mode of inheritance is definitely possible, possible but unlikely, or not possible based on the pedigree. Also briefly explain your rationale behind these choices in the space provided.

Table 1 – Determining Mode of Inheritance

<table>
<thead>
<tr>
<th>Mode of Inheritance</th>
<th>Possible</th>
<th>Possible but Unlikely</th>
<th>Not possible</th>
<th>Brief Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autosomal recessive</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Autosomal dominant</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-linked recessive</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-linked dominant</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y-linked</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mitochondrial</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

3. What mode of inheritance pattern best fits this pedigree in your opinion?

4. In order to impress Dr. Aikenhed, you begin to research genetic disorders that can cause adrenal cancer. You search the Genetics Home Reference (http://ghr.nlm.nih.gov/) with the key words “autosomal dominant adrenal cancer” and you discover seven disorders. You rule out two immediately since they do not usually cause malignant tumors. Write down a summary of the clinical symptoms for the other five in Table 2 below:
Table 2 – Clinical & Genetic Characteristics of Selected Syndromes Associated with Adrenal Cancer

<table>
<thead>
<tr>
<th>Genetic Disorder</th>
<th>Clinical manifestations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiple endocrine neoplasia</td>
<td></td>
</tr>
<tr>
<td>Carney complex</td>
<td></td>
</tr>
<tr>
<td>Li Fraumeni syndrome</td>
<td></td>
</tr>
<tr>
<td>Neuroblastoma</td>
<td></td>
</tr>
<tr>
<td>Von-Hippel-Lindau syndrome</td>
<td></td>
</tr>
</tbody>
</table>

5. Comparing this information with information from both your patient and his family, what genetic disorder should you most suspect?

6. What is the OMIM number (www.ncbi.nlm.nih.gov/omim) for this disorder?

7. What kind of information can you look up using the OMIM number for a genetic disorder?

8. OMIM lists which gene as being defective in this genetic disorder?

9. When you discuss the pedigree and your suspicions with Dr. Aikenhed, she shares with you her concerns about Leah and Luke, despite the fact that they do not appear to have cancer. Why is she concerned about them?

10. With respect to this genetic disorder, would it be possible for a person with cancer to have two parents who do not have cancer? Explain.
You are worried about sharing your suspicions with Lee and Grace, but for different reasons. Grace, who is still recovering from breast cancer surgery and chemotherapy, seems very fragile as well as distraught that her son has a tumor-like growth. If Lee did, in fact, inherit a faulty gene from her that will predispose him to cancer, you are worried about how Grace will handle the news. She seems at great risk for deep feelings of guilt and serious depression. Lee, on the other hand, is a very upbeat young man with a great sense of humor. He seems mostly concerned about improving his mother’s spirits. You would dread being the one to tell him if it turns out that he has a genetic condition that will predispose him to various types of cancers for his entire life. In order to best communicate the pros and cons of genetic testing to them, you review the different types available. You decide to use this reference to understand more: ghr.nlm.nih.gov/handbook/testing.

11. List at least four general categories of genetic tests along with a specific example (e.g., Category: Carrier testing — Example: Assess a couple’s risk for having a child with Tay Sachs disease).

12. What types of issues come up when determining whether to test someone for a genetic illness?

13. If it turns out that Lee has inherited a gene from his mother that predisposes him to cancer, would you recommend that the family test his younger siblings at this point? Why or why not?
Part II – Applications of DNA Technology

While you are waiting for Lee’s biopsy results, you decide to do some concentrated studying for the USMLE Step 1, the exam that all second-year medical students must take to progress in medical school. Because of your interest in Lee’s case, you review material related to genetic diagnosis, prevention, and treatment. You use a more in-depth scientific source on genetic testing (www.ncbi.nlm.nih.gov/books/NBK7586/) and your own knowledge of molecular genetics to answer the following questions.

Questions

1. In order to test Lee for a genetic disorder, Dr. Aikenhed will probably take a sample of cells containing his genomic DNA by swabbing his cheek or taking a blood sample. Why is it important to use non-cancerous cells to test Lee for a genetic disorder instead of the biopsy tissue?

2. Describe the main technique for amplifying a segment of DNA (like the one you suspect is involved in Lee’s cancer) from a complex mixture of genomic DNA. Remember that the entire human genome sequence is known. (*Hint:* This is a technique that is commonly used by laboratories that do genetic testing and various other applications of molecular biology.)

3. If Dr. Aikenhed wanted to see if there was mutation within the protein-coding sequence of the gene implicated in this disorder (as opposed to mutations affecting regulatory elements), what technique involving dideoxynucleotides could be used? Briefly describe this technique.

Lee and his family have inspired in you a desire to find out more about how to prevent and treat this disorder. In your medical genetics textbook, you read that some genetic disorders have specific treatments and/or prevention strategies. The symptoms and progression of phenylketonuria (PKU), for example, can be mitigated with a strict diet that limits phenylalanine intake. Cystic fibrosis is treated with palpitations (tapping) of the chest to loosen mucus and with antibiotics to help prevent infection, among other strategies. Most genetic disorders, however, do not have such clear cut recommendations.
4. Frequent screening/check-ups and avoidance of risk factors (e.g., smoking) are often advised in cases involving inherited genes which predispose a person to cancer. Do you think that this advice will make a difference in morbidity or mortality related to this particular disorder? Why or why not?

5. What is the name of the genetic technique that would help a person who inherited this genetic disorder to have a child without the defective allele?
Part III – Gene Expression and Disease

Unfortunately, the results from Lee's biopsy are in and it appears that he has adrenocortical carcinoma (ACC). Fortunately, Dr. Aikenhed (who, in addition to her clinical practice in the area of oncology also conducts research) is on the cutting edge of developing treatments for this condition. She is part of a consortium of researchers that are trying to catalog all of the different genes that may be involved in adrenal cancer. This will help physicians and scientists to discover drugs to target specific cancer-contributing gene products.

Cancer genes can be categorized in many different ways. **Proto-oncogenes** produce proteins that are involved in promoting the cell cycle. Mutations in these genes tend to cause either an over-production of protein (over-expression) or a protein to be constitutively active (pushing the cell cycle forward even in the absence of a signal to divide (e.g. growth factor)).

**Tumor suppressor genes**, on the other hand, put the brakes on the cell cycle, usually at key checkpoints. Mutations in tumor suppressor genes that contribute to carcinogenesis can cause either abnormally low levels of protein or no functional protein (under-expression) at all. A mutation like this could allow a damaged or rogue cell to divide when it shouldn’t.

**Genome-maintenance genes** can also play a role in cancer and they include the telomerase gene (which helps extend the life of a chromosome and, consequently, the number of cell division cycles that it can undergo) and DNA repair genes (which keep the level of mutations in all genes low).

Dr. Aikenhed and her colleagues use microarrays containing the most common genes implicated in ACC to analyze gene expression (both over- or under-expressed genes) in cancer samples compared to normal tissue. To understand more about this technique, you decide to review the basics of microarray analysis by going through slides 1-9 at this website: www.hhmi.org/biointeractive/genomics/microarray_analyzing/01.html. It never ceases to amaze you how they can put thousands of tiny pieces of single-stranded DNA on these microarray chips in an organized way, creating all sorts of different sequences for samples to hybridize (bind) to.

Several microarrays were made with the most common genes implicated in ACC. DNA samples from six patients with ACC were analyzed. Lee F. is Patient #3. The last row represents a control using normal adrenal tissue from a healthy individual.

**Table 3 – Different Genes* Implicated in ACC**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
<th>Patient 4</th>
<th>Patient 5</th>
<th>Patient 6</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>IGF2</td>
<td>R</td>
<td>R</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>TP53</td>
<td>Y</td>
<td>Y</td>
<td>G</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>RAS</td>
<td>Y</td>
<td>G</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>BCLXL</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>CHRBL</td>
<td>G</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>APC</td>
<td>R</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>DDB1</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>MRPL48</td>
<td>G</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
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<td>Y</td>
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<tr>
<td>EGRF1</td>
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<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>CTNNB1</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
</tbody>
</table>

*gene symbols are consistent with HGNC & OMIM
R=red (high expression in diseased compared with normal cells)
G=green (low expression in diseased compared with normal cells)
Y=yellow (the same expression in both cell types)

**Questions**

1. What can you conclude about gene expression in your patient’s cells?
You decide to search the following World Health Organization website, which contains important statistics on this gene: www-p53.iarc.fr/Statistics.html.

2. The most common type of germ line mutation (occurring in ~72% of cases) is a ___________ mutation.

3. Describe how this type of mutation changes the amino acid sequence of a polypeptide made from this gene.

4. There are two other types of common point mutations (a change, insertion or deletion of a nucleotide) that affect the coding sequence of various proteins. Briefly describe all three types of mutations in Table 4 and include whether you predict that the level of mRNA and protein might be affected (increased or decreased).

Table 4 – How Different Mutations Might Affect Gene Expression

<table>
<thead>
<tr>
<th>Name of mutation</th>
<th>How does mutation alter mRNA sequence?</th>
<th>Could this mutation affect the level (increase or decrease) of functional protein in the cell? Why?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tbody>
</table>

5. Sometimes mutations involve the deletion of large regions of genetic information. Other mutations affect gene expression by altering the DNA sequence of regulatory elements like promoter regions. Further analysis reveals that your patient appears to have a mutation that affects mRNA splicing (removal of introns and splicing together of various exons), which occurs approximately ________% of the time in this condition. This mutation reduces the level of mature mRNA in his cells. This is why the level of the affected gene shows a lower expression level than controls in your patient, Lee.
Part IV – Gene Therapy

Despite trying to maintain a professional level of emotional distance, you have become attached to Lee and his good natured family. You spend evenings searching the medical literature for potential treatments, including clinical trials, to help them. You remember learning about gene therapy trials for SCID (Severe Combined ImmunoDeficiency) and LCA (Leber’s Congenital Amaurosis—a type of hereditary blindness) in the *New England Journal of Medicine.*

Questions

1. How would you describe the main goal of gene therapy to this family?

2. Several different types of viruses, including adenoviruses, have been used as vectors for gene therapy in humans. Why are viruses used for gene therapy?

3. The two basic types of gene therapy are _________ and _________. What is the main difference between them?

4. Which type would best help this patient and his family? Why?

5. What types of risks/problems are currently associated with gene therapy?

You discover that there have been gene therapy trials involving *TP53*. For a variety of reasons, however, the treatment has not been approved by the FDA (U.S. Food & Drug Administration) and the trials stalled. The main problem appeared to be that the company developing the treatment did not provide enough evidence to the FDA that the treatment was effective.
6. If there were a current TP53 gene therapy clinical trial, would you recommend that Lee enroll in it? Why or why not?

7. Another possible strategy to treat this disorder involves “virotherapy.” Watch the following animation (www.hhmi.org/biointeractive.cancer/using_p53.html) and then briefly describe how this process works.

8. Write a brief, encouraging letter addressed to Lee and his family that highlights the hope of a future gene therapy cure for the disorder affecting them.

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For more information on gene therapy:

