INSTITUTE: National Cancer Institute

STUDY NUMBER: 13-C-0093 PRINCIPAL INVESTIGATOR: Udai S. Kammula, M.D.

STUDY TITLE: Phase II Study in Patients with Metastatic Ocular Melanoma Using a Non-Myeloablative Lymphocyte Depleting Regimen of Chemotherapy Followed by Infusion of Autologous Tumor-Infiltrating Lymphocytes with or without High Dose Aldesleukin

Continuing Review Approved by the IRB on 10/17/16
Amendment Approved by the IRB on 05/05/17 (L) Date Posted to Web: 05/16/17

INTRODUCTION

We invite you to take part in a research study at the National Institutes of Health (NIH).

First, we want you to know that:

Taking part in NIH research is entirely voluntary.

You may choose not to take part, or you may withdraw from the study at any time. In either case, you will not lose any benefits to which you are otherwise entitled. However, to receive care at the NIH, you must be taking part in a study or be under evaluation for study participation.

You may receive no benefit from taking part. The research may give us knowledge that may help people in the future.

Second, some people have personal, religious or ethical beliefs that may limit the kinds of medical or research treatments they would want to receive (such as blood transfusions). If you have such beliefs, please discuss them with your NIH doctors or research team before you agree to the study.

Now we will describe this research study. Before you decide to take part, please take as much time as you need to ask any questions and discuss this study with anyone at NIH, or with family, friends or your personal physician or other health professional.

If you are signing for a minor child, “you” refers to “your child” throughout the consent document.

Why is this study being done?

We have developed an experimental therapy that involves taking cells called lymphocytes from patients' tumors, growing them in the laboratory in large numbers, and then giving the cells back to the patient. These cells are called Young Tumor Infiltrating Lymphocytes, or Young TIL and the therapy is called cell therapy. Before receiving the cells, the patients receive 2 chemotherapy...
drugs to temporarily suppress the immune system to improve the chances that the tumor fighting cells will be able to survive in the body. The Surgery Branch has pioneered and successfully used this therapy in cancers such as metastatic melanoma, cervical and gastro-intestinal cancers and we now would like to try this therapy in metastatic ocular melanoma. After the cells are given, patients who agree and are medically eligible may receive aldesleukin (IL-2) to help the tumor fighting cells stay alive longer.

The purpose of this study is to see if these tumor fighting cells (young TIL) with or without the administration of aldesleukin can cause metastatic ocular melanoma tumors to shrink and to evaluate the toxicity of this treatment.

**Why are you being asked to take part in this study?**

You are being asked to participate in this study because you have been diagnosed with metastatic ocular melanoma.

**How many people will take part in this Study?**

Up to 54 patients will be enrolled in this study.

**Description of Research Study**

**Research Study Overview**

The Cell protocol has several stages:

<table>
<thead>
<tr>
<th>Stage</th>
<th>Timeframe</th>
<th>Location</th>
<th>Comments &amp; Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Work up</td>
<td>1-2 weeks</td>
<td>Inpatient and out patient</td>
<td>Scans, x-rays, labs leukapheresis other tests as needed</td>
</tr>
<tr>
<td>Chemotherapy (day -7 to -1)</td>
<td>1 week</td>
<td>Inpatient</td>
<td>Receive IV chemotherapy to prepare your immune system for the cells</td>
</tr>
<tr>
<td>Cells and aldesleukin (Day 0-4)</td>
<td>1-5 days</td>
<td>Inpatient and possibly ICU</td>
<td>Receive the TIL cells IV and then high dose aldesleukin about every 8 hours for up to 15 doses</td>
</tr>
<tr>
<td>Recovery</td>
<td>1-2 weeks</td>
<td>Inpatient unit</td>
<td>Recover from the effects of treatment</td>
</tr>
<tr>
<td>Follow-up</td>
<td>Ongoing until disease progression</td>
<td>Outpatient</td>
<td>Return to clinic for physical exam, review of side effects, labs, scans every 1-3 months for the first year.</td>
</tr>
</tbody>
</table>

**PATIENT IDENTIFICATION**

CONTINUATION SHEET for either:

NIH-2514-1 (07-09)
NIH-2514-2 (10-84)
P.A.: 09-25-0099
File in Section 4: Protocol Consent
What will happen if you take part in this research study?

Before you begin the study

Cell harvest and growth

You underwent a procedure, either surgery or a biopsy, to obtain pieces of your tumor so that we can grow the Young TIL from your tumor cells in the laboratory while enrolled on our companion protocol 03-C-0277 (Cell Harvest and Preparation for Surgery Branch Adoptive Cell Therapy Protocols). The surgical procedure may involve liver resection either by laparoscopic or open surgical approaches. In 1 out of 5 patients, we are not able to successfully grow the cells for this procedure. If your cells do not grow, you will not be able to receive the cell infusion. If that happens, we will look at alternative treatments for you (best available care) or return you to the care of your referring physician. We usually know after about 4 weeks whether the cells will grow well enough to be used as an experimental treatment on this protocol. At the time we determine that your cells are not growing, we will inform you and discuss your options with you. Several medications are used during the preparation of your cell product, be sure to tell your doctor if you are allergic to any antibiotics.

Work up

Prior to receiving the experimental treatment, you will undergo many tests. These include imaging procedures, heart and lung function tests, and laboratory tests. If you are a woman, you will undergo a pregnancy test. You will also have a large catheter inserted into a vein and leukapheresis will be performed (see below). You may be admitted to the hospital for these tests. However, you will be allowed to leave on pass on the days that you are not having tests performed.

Catheter insertion

Prior to beginning the experimental treatment, you will have an intravenous (IV) catheter placed in your upper chest. The area will be numbed with an anesthetic before the catheter is put in. Although rare, putting these catheters in can sometimes cause collapse of a lung or cause bleeding. Lung collapse is treated by putting a tube into your chest for a few days to allow your lung to expand. Pressure is placed on any area that might bleed. Other IVs may be needed in one or both of your arms if we need to give you extra fluids, medicines, or nutrition.

Leukapheresis

Leukapheresis is a procedure that allows us to remove certain types of blood cells from you and return the rest of your blood. It is a very common procedure that is done routinely here at the NIH with very few risks. During leukapheresis, blood is removed from you through a needle in your arm, circulated through a machine that divides whole blood into red cells, plasma (the serum part), and lymphocytes (or white cells), and then the plasma and red cells are returned to you through a second needle in your other arm. The procedure takes between 3-4 hours to complete. The white blood cells may be used to help grow the cells. In addition to the
leukapheresis you will undergo as part of your work up, we will also ask you to undergo one additional pheresis procedure between 4 and 6 weeks after your cell treatment to see the impact of this therapy on the immune system and see if the cells we gave you are still active.

**During the Study**

**Chemotherapy Regimen (Day -7 through Day -1)**

After we have grown the Young TIL cells to large numbers in the laboratory, you will be admitted to the hospital to begin your experimental treatment. You will be given two chemotherapy medicines, cyclophosphamide and fludarabine, to make space in your immune system so the Young TIL can work without any interference from the cells in your immune system. These medicines may cause your tumor to shrink some, but this shrinkage is anticipated to be only partial and of small duration. The main purpose of the chemotherapy is to see if we can make the cells more effective in fighting cancer tumors. Animal experiments have indicated that chemotherapy can make the cells more effective in fighting cancer tumors, but it is not known whether this is true in humans. The cyclophosphamide will be given into your catheter over 1 hour for two days (Day -7 and Day -6) and the fludarabine will be given into your catheter for 30 minutes every day for the next five days (Day -5 through Day -1). The side effects of these medicines are described on the following pages.

**Cell Infusion and Aldesleukin Regimen (Day 0 through Day 5)**

All patients will be given the cells through their IV over 20-30 minutes one to four days after the last dose of chemotherapy. Within 24 hours after your cell infusion you will be given high dose aldesleukin through one of the IVs. It will be given as a 15-minute infusion about every 8 hours for up to five days after the cell infusion. Aldesleukin is a cell growth factor and it is thought that it will help the cells live longer in your body. No aldesleukin will be given if you refuse it or are not medically eligible to receive it.

The day after your cells are infused, we will give you G-CSF (filgrastim) as a shot or injection under the skin every day to stimulate your bone marrow to produce blood cells until they increase to a sufficient number. We will watch you closely during this entire time for any side effects of this experimental regimen. We will discuss the side effects below and we will include in your care all the medicines and treatments to prevent as many of these side effects as we can and to make you as comfortable as we can.

**When you are finished taking the drugs (treatment)**

**Recovery**

You will recover in the hospital until you are well enough to go home. This usually takes 7-12 days after you have received cells or your last dose of aldesleukin; however, you may need to stay in the hospital for longer than this before you are well enough to go home. We will continue to give you support medications, do laboratory tests, and watch you closely for any side effects until we feel your condition is stable.
In addition to the laboratory tests to monitor your condition, we will remove between 1 and 9 teaspoons of blood daily to study the effects of this regimen on your immune system. If you experience side effects in your kidneys, we will collect 1 additional teaspoon of blood and about 6 teaspoons of urine to see if we can determine the cause of these side effects. The maximum amount of blood for research is approximately 2.3 cups in 8 weeks.

Follow up and Evaluation of Experimental Regimen

You will need to continue to take Bactrim, an antibiotic, for at least 6 months following your treatment to prevent you from catching a certain type of pneumonia seen in patients who have low white blood cell counts. We will ask you to return to the NIH Clinical Center frequently after you are discharged approximately 6 and 12 weeks following treatment and then if you are responding to the treatment, every 3 months for the first year following treatment, every 6 months for the second year and then as determined by your physician. The first visit will probably take 2 days; the following visits may only take one day. At each visit you will have lab tests, imaging studies and a physical examination. At some of your follow up visits, you may undergo leukapheresis or have about 8 tubes of blood drawn (4 tablespoons) so that we can see the effect this therapy has had on your immune system and if the cells we gave you are still alive. If you are unwilling or unable to travel to the NIH Clinical Center we will contact you by phone or e-mail and we may ask you to send us lab, imaging, and physical exam reports.

Retreatment

If you have sustained stable disease or your tumor shrinks or disappears following the initial treatment and then recurs you may receive one additional treatment if you tolerated the treatment well and if all the side effects have resolved. You will receive the same medications and cell infusion on the same schedule as with the first treatment.

Birth Control

If you are a woman who is breast feeding or pregnant, you may not take part in the study because we don’t know how this medicine would affect your baby or your unborn child. If you are a woman who can become pregnant, or are the partner of a woman who can become pregnant, you will need to practice an effective form of birth control before starting study treatment, during study treatment, and for four months after you finish study treatment. If you think that you or your partner is pregnant, you should tell your study doctor or nurse at once.

- Effective forms of birth control include:
  - Abstinence
  - intrauterine device (IUD)
  - hormonal [birth control pills, injections, or implants]
  - tubal ligation
  - vasectomy
Risks or Discomforts of Participation

What side effects or risks can I expect from being in this study?

The risks and discomforts of this research study can be significant. This experimental treatment can lead to long-term decrease in your immune function. It is also possible that you may lose your fertility following this experimental treatment. It is possible, although unlikely, that this experimental treatment may cause your death.

We will discuss the side effects of this experimental treatment with you. You will be given medicines, transfusions, and treatments to prevent or treat the side effects including drugs to prevent and/or treat different types of infections. We will try to make you as comfortable as possible. You should talk to your study doctor about any symptoms that you experience while taking part in the study.

Leukapheresis

During the leukapheresis procedure, you may have some tingling in your face and lips due to the medicine used to keep your blood from clotting during the procedure. The nurses may give you a calcium-containing antacid to chew that takes away this tingling. Rarely, people may experience lightheadedness or dizziness. We ask that you eat prior to the procedure to prevent this. Rare complications of this procedure are lowered blood pressure, bleeding or bruising where the needles are put in your arms.

Young TIL Cell Infusion

All Young TIL cells will be given through your catheter while we keep you in the patient care unit so we can watch you closely.

Potential risks include:

- Fever, chills and shortness of breath, which are common and may last for a few hours
- Lung congestion causing shortness of breath
- Immune-type reaction, such as pharyngitis and vaginitis. Note: In similar clinical trials with cells targeting a melanoma protein, we have observed the following immune-mediated toxicities: loss of skin pigment (known as vitiligo), and inflammation of the eye (uveitis). The skin and the eye are all sites where that targeted melanoma protein is known to exist.
- As this is a new experimental therapy, side effects that we do not anticipate that may cause your condition to deteriorate may be encountered. Any new information that becomes available during the course of this study will be shared with you.

Medications

The side effects of cyclophosphamide, fludarabine, high dose aldesleukin and some of the other medications you will receive are listed below:
Cyclophosphamide and Fludarabine side effects

<table>
<thead>
<tr>
<th>Common</th>
<th>Less Common</th>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Changes in blood counts including: low</td>
<td>• Bleeding</td>
<td>• Heart damage</td>
</tr>
<tr>
<td>red cell count (causing fatigue and</td>
<td>• Infection</td>
<td>• Lung damage</td>
</tr>
<tr>
<td>shortness of breath), low platelet</td>
<td>• Bladder irritation with bloody urine</td>
<td>• Kidney damage</td>
</tr>
<tr>
<td>count (increasing the risk of bleeding</td>
<td>• Severe allergic reaction (difficulty</td>
<td>• Inflammation of the eye resulting in blindness</td>
</tr>
<tr>
<td>and bruising), decrease in white blood</td>
<td>breathing/swelling)</td>
<td>• Inflammation of nervous system resulting in death</td>
</tr>
<tr>
<td>cells (increasing the risk of infection</td>
<td>• Headache or dizziness</td>
<td>• Epstein Barr Virus Lymphoma. This</td>
</tr>
<tr>
<td>and the need for treatment with</td>
<td>• Sweating</td>
<td>can be fatal (Two patients on other</td>
</tr>
<tr>
<td>antibiotics or other treatment)</td>
<td>• Swelling of arms or legs</td>
<td>studies in the Surgery Branch</td>
</tr>
<tr>
<td>• Loss of appetite, nausea, vomiting,</td>
<td>• Skin changes, rash, blisters</td>
<td>developed EBV lymphoma, and one</td>
</tr>
<tr>
<td>• Diarrhea, stomach pain</td>
<td>• Weakness</td>
<td>died as a result of this disease.)</td>
</tr>
<tr>
<td>• Mouth sores</td>
<td>• Hearing loss</td>
<td>• Loss of fertility</td>
</tr>
<tr>
<td>• Hair loss</td>
<td></td>
<td>• Two out of the first 81 patients</td>
</tr>
<tr>
<td>• Fatigue</td>
<td></td>
<td>enrolled on the initial young TIL study</td>
</tr>
<tr>
<td>• Muscle or joint aches</td>
<td></td>
<td>died from complications resulting from</td>
</tr>
<tr>
<td></td>
<td></td>
<td>suppression of the immune function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>which resulted in a severe infection</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(one of these patients also received</td>
</tr>
<tr>
<td></td>
<td></td>
<td>radiation as part of their treatment regimen).</td>
</tr>
</tbody>
</table>

IL-2 (aldesleukin) side effects

<table>
<thead>
<tr>
<th>Common</th>
<th>Less common</th>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Fever, chills, and fatigue</td>
<td>• Decrease in thyroid function that may</td>
<td>• Bowel perforation (a hole) requiring longer hospitalization or surgery.</td>
</tr>
<tr>
<td>• Lowered platelet and red blood cell levels</td>
<td>require daily thyroid hormone replacement;</td>
<td>• Autoimmune disease, where your immune system attacks cells in organs of your body. Should this occur, you will be treated with steroids to stop the immune response.</td>
</tr>
<tr>
<td>that may require transfusions</td>
<td>Abnormal kidney and liver function that can be severe;</td>
<td>• Damage to the heart muscle or heart attack</td>
</tr>
<tr>
<td></td>
<td>Abnormal heartbeats or low blood pressure that may require treatment in the ICU.</td>
<td>• Loss of blood flow to the extremities due to medicines used</td>
</tr>
<tr>
<td>• Significant fluid retention causing weight</td>
<td>Breathing problems which may need</td>
<td></td>
</tr>
<tr>
<td>gain (as much as 20 pounds).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Low blood pressure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Increased heart rate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Low urine output</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Swelling in your extremities,</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

PATIENT IDENTIFICATION

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- Fluid in your lungs that can require oxygen
- Dry mouth, nausea, vomiting and diarrhea;
- Rash, itching; and changes in skin or hair pigmentation, called vitiligo;
- Changes in mental status, including confusion, difficulty sleeping or vivid dreams; this can be severe and require sedation and monitoring in the ICU

- monitoring in ICU and insertion of a breathing tube.
- to treat very low blood pressure and shock. In one instance a patient had to have her lower arm amputated after treatment with these medicines.
- IL-2 is mixed with human albumin which could cause an allergic reaction or potentially transmit viral infections, although we have not had this occur.

### Support Medications – side effects

<table>
<thead>
<tr>
<th>Common</th>
<th>Less common</th>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filgrastim (To increase production of white blood cells)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| • Bone Pain                 | • Severe headache                                | • Severe breathing problems
|                              |                                                 | • Rupture of your spleen                  |
| Bactrim (To prevent a specific type of pneumonia)      |                                                                 |
| • Fever                     | • Nausea, vomiting,                             |                                           |
|                              | • Skin rash with itching                        |                                           |
|                              | • reduced number of                             |                                           |
|                              | • white blood cells                             |                                           |
|                              | • Allergic reaction                             |                                           |
| Fluconazole: (To prevent fungal infections)            |                                                                 |
| • Headache                  | • A skin disorder called Stevens Johnson Syndrome, which can be fatal |
| • Nausea, vomiting, diarrhea, abdominal pain           | • Liver damage which may be permanent           |
| • Itching                   |                                           |                                           |

**Acyclovir and Valacyclovir**

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**PATIENT IDENTIFICATION**

CONTINUATION SHEET for either:

NIH-2514-1 (07-09)
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P.A.: 09-25-0099
File in Section 4: Protocol Consent
Potential Benefits of Participation

Are there benefits to taking part in this study?

The aim of this study is to see if this new experimental treatment will cause your tumors to shrink. We do not know if you will receive personal medical benefit from taking part in this study. These potential benefits could include shrinking of your tumor or lessening of your symptoms, such as pain, that are caused by the cancer. Because there is not much information about the Young TIL therapy effect on your type of cancer, we do not know if you will benefit from taking part in this study, although the knowledge gained from this study may benefit others in the future who have cancer.

Alternative Approaches or Treatments

What other choices do I have if I do not take part in this study?

If there are effective salvage regimens (regimens used when standard regimens have failed), you will be directed to undergo these regimens before participating in this protocol.

Instead of being on this study, you have these options for treatment:

- Getting treatment or care for your cancer without being in a study
- Taking part in another study
- Getting no treatment; or getting comfort care which is also called palliative care. This type of care helps reduce pain, tiredness, appetite problems, and other problems caused by cancer. It does not treat the cancer directly. Instead it tries to improve how you feel. Comfort care tries to keep you as active and comfortable as possible.

Please talk to your doctor about these and other options.

Research Subject’s Rights

What are the costs of taking part in this study?

If you choose to take part in the study, the following will apply, in keeping with the NIH policy:
• You will receive study treatment at no charge to you. This may include surgery, medicines, laboratory testing, x-rays or scans done at the Clinical Center, National Institutes of Health (NIH), or arranged for you by the research team to be done outside the Clinical Center, NIH if the study related treatment is not available at the NIH.

• There are limited funds available to cover the cost of some tests and procedures performed outside the Clinical Center, NIH. You may have to pay for these costs if they are not covered by your insurance company.

• Medicines that are not part of the study treatment will not be provided or paid for by the Clinical Center, NIH.

• Once you have completed taking part in the study, medical care will no longer be provided by the Clinical Center, NIH.

Will your medical information be kept private?

We will do our best to make sure that the personal information in your medical record will be kept private. However, we cannot guarantee total privacy. Organizations that may look at and/or copy your medical records for research, quality assurance, and data analysis include:

• The National Cancer Institute (NCI) and other government agencies, like the Food and Drug Administration (FDA), which are involved in keeping research safe for people.
• NCI Institutional Review Board
• The study Sponsor (Center for Cancer Research) or their agent(s)
• Lion Biotechnologies

A description of this clinical trial will be available on [http://www.Clinicaltrials.gov](http://www.Clinicaltrials.gov), as required by U.S. Law. This Web site will not include information that can identify you. At most the Web site will include a summary of the results. You can search this Web site at any time.

Stopping the Study

Your doctor may decide to take you off this study under the following circumstances:

• if he/she believes that it is in your best interest
• if your disease comes back during treatment
• if you become pregnant
• if you experience side effects from the treatment that are considered too severe
• if new information becomes available that shows that another treatment would be better for you

In this case, you will be informed of the reason for that decision.

You can stop participating at any time. However, if you decide to stop participating in the study, we encourage you to talk to the study doctor and your regular doctor first. If you refuse to
participate or withdraw from the protocol or at the completion of the protocol, we will attempt to offer you participation in other NIH protocols if these are available, or will refer you to your home physician for further management.

If you decide at any time to withdraw your consent to participate in the trial, we will not collect any additional medical information about you. However, according to FDA guidelines, information collected on you up to that point may still be provided to the Sponsor. If you withdraw your consent and leave the trial, any samples of yours that have been obtained for the study and stored at the NCI can be destroyed upon request. However, any samples and data generated from the samples that have already been distributed to other researchers or placed in the research databases cannot be recalled and destroyed.

**Conflict of Interest**

The National Institutes of Health reviews NIH staff researchers at least yearly for conflicts of interest. This process is detailed in a Protocol Review Guide. You may ask your research team for additional information or a copy of the Protocol Review Guide. Members of the research team who do not work for the NIH are expected to follow these guidelines but they do not need to report their personal finances to the NIH. The NIH and 2 of the investigators on the research team have developed the cell process being used in this research and have a patent pending. This means that it is possible that the results of this study could lead to payments to NIH scientists and to the NIH. By law, government scientists are required to receive such payments for their inventions.

Members of the research team working on this study may have up to $15,000 of stock in the companies that make products used in this study. This is allowed under federal rules and is not a conflict of interest.

**Use of Specimens and Data for Future Research**

Specimens and data collected during the course of this study will be used for future research and will be stored, tracked and disposed of under our companion protocol 03-C-0277, (Cell Harvest and Preparation for Surgery Branch Adoptive Cell Therapy Protocols) on which you have already been enrolled.

In addition, to advance science, it is helpful for researchers to share information they get from studying human samples. They do this by putting it into one or more scientific databases, where it is stored along with information from other studies. A researcher who wants to study the information must apply to the database and be approved. Researchers use specimens and data stored in scientific databases to advance science and learn about health and disease.
We plan to keep some of your specimens and data that we collect and use them for future research and share them with other researchers. We will not contact you to ask about each of these future uses. These specimens and data will be stripped of identifiers such as name, address or account number, so that they may be used for future research on any topic and shared broadly for research purposes. Your specimens and data will be used for research purposes only and will not benefit you. It is also possible that the stored specimens and data may never be used. Results of research done on your specimens and data will not be available to you or your doctor. It might help people who have cancer and other diseases in the future.

If you do not want your stored specimens and data used for future research, please contact us in writing and let us know that you do not want us to use your specimens and/or data. Then any specimens that have not already been used or shared will be destroyed and your data will not be used for future research. However, it may not be possible to withdraw or delete materials or data once they have been shared with other researchers.
OTHER PERTINENT INFORMATION

1. **Confidentiality.** When results of an NIH research study are reported in medical journals or at scientific meetings, the people who take part are not named and identified. In most cases, the NIH will not release any information about your research involvement without your written permission. However, if you sign a release of information form, for example, for an insurance company, the NIH will give the insurance company information from your medical record. This information might affect (either favorably or unfavorably) the willingness of the insurance company to sell you insurance.

The Federal Privacy Act protects the confidentiality of your NIH medical records. However, you should know that the Act allows release of some information from your medical record without your permission, for example, if it is required by the Food and Drug Administration (FDA), members of Congress, law enforcement officials, or authorized hospital accreditation organizations.

2. **Policy Regarding Research-Related Injuries.** The Clinical Center will provide short-term medical care for any injury resulting from your participation in research here. In general, no long-term medical care or financial compensation for research-related injuries will be provided by the National Institutes of Health, the Clinical Center, or the Federal Government. However, you have the right to pursue legal remedy if you believe that your injury justifies such action.

3. **Payments.** The amount of payment to research volunteers is guided by the National Institutes of Health policies. In general, patients are not paid for taking part in research studies at the National Institutes of Health. Reimbursement of travel and subsistence will be offered consistent with NIH guidelines.

4. **Problems or Questions.** If you have any problems or questions about this study, or about your rights as a research participant, or about any research-related injury, contact the Principal Investigator, Udai S. Kammula, M.D., Building 10, Room 3-5930, Telephone: 240-760-6216. If you have any questions about the use of your specimens or data for future research studies, you may also contact the Office of the Clinical Director, Telephone: 240-760-6070. You may also call the Clinical Center Patient Representative at 301-496-2626.

5. **Consent Document.** Please keep a copy of this document in case you want to read it again.
**MEDICAL RECORD**

**CONSENT TO PARTICIPATE IN A CLINICAL RESEARCH STUDY**

- Adult Patient or  
- Parent, for Minor Patient

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**STUDY NUMBER:** 13-C-0093

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**COMPLETE APPROPRIATE ITEM(S) BELOW:**

<table>
<thead>
<tr>
<th>A. Adult Patient's Consent</th>
<th>B. Parent’s Permission for Minor Patient.</th>
</tr>
</thead>
<tbody>
<tr>
<td>I have read the explanation about this study and have been given the opportunity to discuss it and to ask questions. I hereby consent to take part in this study.</td>
<td>I have read the explanation about this study and have been given the opportunity to discuss it and to ask questions. I hereby give permission for my child to take part in this study. (Attach NIH 2514-2, Minor’s Assent, if applicable.)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Signature of Adult Patient/ Legal Representative</th>
<th>Date</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Signature of Parent(s)/ Guardian</th>
<th>Date</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Print Name</th>
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<table>
<thead>
<tr>
<th>C. Child’s Verbal Assent (If Applicable)</th>
</tr>
</thead>
<tbody>
<tr>
<td>The information in the above consent was described to my child and my child agrees to participate in the study.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Signature of Parent(s)/Guardian</th>
<th>Date</th>
<th>Print Name</th>
</tr>
</thead>
</table>

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**THIS CONSENT DOCUMENT HAS BEEN APPROVED FOR USE FROM OCTOBER 17, 2016 THROUGH OCTOBER 16, 2017.**

<table>
<thead>
<tr>
<th>Signature of Investigator</th>
<th>Date</th>
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</table>

<table>
<thead>
<tr>
<th>Signature of Witness</th>
<th>Date</th>
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</table>

<table>
<thead>
<tr>
<th>Print Name</th>
<th>Print Name</th>
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</thead>
</table>

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**PATIENT IDENTIFICATION**

**CONSENT TO PARTICIPATE IN A CLINICAL RESEARCH STUDY (Continuation Sheet)**

- Adult Patient or  
- Parent, for Minor Patient

NIH-2514-1 (07-09)

P.A.: 09-25-0099

File in Section 4: Protocol Consent