Guide to Submission

Reprogenetics
Submission #2011-ACD-002

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hESC Registry Application Search Results

<table>
<thead>
<tr>
<th>Request #: 2011-ACD-002</th>
<th>Organization: Reprogenetics, LLC</th>
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<tr>
<td>Status: Pending</td>
<td>Org Address: 3 Regent Street, Suite 301, Livingston, NJ 07039</td>
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<td>Review: ACD</td>
<td>DUNS: 148120780</td>
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<tr>
<td>Assurance: Yes (Section II(B))</td>
<td>Grant Number(s): n/a</td>
</tr>
<tr>
<td>Certification: Yes</td>
<td>Signing Official (SO): Santiago Munne / 9734365010 / <a href="mailto:smunne@reprogenetics.com">smunne@reprogenetics.com</a></td>
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<tr>
<td>Authority: Yes</td>
<td>Submitter of Request: Mina Alikani / 9733225043 / <a href="mailto:mina.alikani@embryos.net">mina.alikani@embryos.net</a></td>
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<td>Cell Lines: 1</td>
<td>Submitter Comments: Reprogenetics has previously submitted RNJ7 for inclusion in the registry. This new submission differs from the previous submission in that the patients have been reconsented. The new consent form, IRB approval of the modifications, and IRB approval of reconsenting have been included as supporting documents.</td>
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<td>Document 1: (PDF - 01/25/2011) Initial research consent form signed by donors of embryo for RNJ11; REDACTED - Elements: 1,2,3,4,5,6,7,9,10,11,13</td>
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<td>Document 3: (PDF - 01/25/2011) New revised consent signed by donors of embryo for RNJ11; REDACTED - Elements: 1,2,3,4,5,6,7,8,9,10,11,12,13,14,15</td>
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<td>Document 6: (PDF - 01/25/2011) New consent (2009) in full. - Elements: 1,2,3,4,5,6,7,8,9,10,11,12,13,14,15</td>
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<td>Administrative Comments:</td>
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http://hesregapp.od.nih.gov/login/list.htm?DetailList=yes&id=69

4/21/2011
Redacted Consent for RNJ11 uploaded by D. Hannemann 28 Jan 2011
SO certifications corrected E. Gadbois 1 Feb 2011
Landis approval move to ACD E. Gadbois 8 Feb 2011
Compilation of submission by D. Hannemann 9 Feb 2011; removed on 10 Feb 2011 to update
IIB Assurance submitter email by D. Hannemann 10 Feb 2011
Replaced Redacted Re-Consent for RNJ11 (Doc 2) with addtl redactions by D. Hannemann 10 Feb 2011
Updated submission compilation - by D. Hannemann 10 Feb 2011; removed on 20 Apr 2011 to update
17 Feb Submitter Email with IRB Approval/Renewal Letters - by DHannemann 20 Apr 2011
22 Feb Submitter email re: 2004 Revised Consent and Protocol - by D. Hannemann 20 Apr 2011
2004 Revised Consent and Protocol (22 Feb email attachment) - by DHannemann 20 Apr 2011
Initial Embryo Donation Consent for RNJ11 (22 Feb email attachment) - by D. Hannemann 4 Mar 2011
15 Mar Submitter email - by DHannemann 20 Apr 2011
hESC Line Characterization Chart (15 Mar email attachment) - by DHannemann 20 Apr 2011
13 Apr Submitter email - by DHannemann 20 Apr 2011
15 Apr Submitter email - by DHannemann 20 Apr 2011

Administrative Attachments:
Document 1:  (PDF - 01/28/2011)  28 Jan Submitter Email
Document 2:  (PDF - 02/10/2011)  Redacted Reconsent for RNJ11
Document 3:  (PDF - 02/08/2011)  Landis approval move to ACD
Document 4:  (PDF - 04/20/2011)  17 Feb Submitter Email with IRB Approval/Renewal Letters
Document 5:  (PDF - 02/10/2011)  IIB Assurance submitter email
Document 6:  (PDF - 04/20/2011)  17 Feb Submitter Email with IRB Approval/Renewal Letters
Document 7:  (PDF - 04/20/2011)  22 Submitter Email re: 2004 Revised Consent and Protocol
Document 8:  (PDF - 04/20/2011)  2004 Revised Consent and Protocol
Document 9:  (PDF - 04/20/2011)  Full Initial Embryo Donation Consent for RNJ11
Document 10: (PDF - 04/20/2011)  15 Mar Submitter Email
Document 11: (DOC - 04/20/2011)  hESC Line Characterization Chart (15 Mar email attachment)
Previous ADM Request Number: 2011-ADM-001
Switched from ADM to ACD Date: 02/07/2011

Reason for Switch to ACD Review:
The administrative review group determined that there are two provisions of the Section IIA criteria that are not met in this submission: Element 8: Donor(s) should have been informed that they retained the right to withdraw consent for the donation of the embryo until the embryos were actually used to derive embryonic stem cells or until information which could link the identity of the donor(s) with the embryo was no longer retained, if applicable. Element 15: Whether information that could identify the donor(s) would be available to researchers.

Added By: Commons\smunne1  On: 01/25/2011  |  Last Updated By: NIH\hannemann  On: 04/20/2011  |  Record ID: 69

Total Record Count = 1
Dear Dr. Munne and Dr. Alikani,

We have reviewed this submission under Section IIA of the NIH Guidelines and have found that this submission does not meet all of the requirements under that section. However, it is eligible for review under Section IIB of the Guidelines. Therefore I am sending this submission for analysis by the Working Group on Human Embryonic Stem Cell Eligibility Review under Section IIB. Your submission still appears as pending review on the NIH public website.

For our records, could you please send me an assurance in accordance with Section IIB of the Guidelines? Those criteria are available at http://stemcells.nih.gov/policy/2009guidelines.htm. Essentially we need a statement from Dr. Munne attesting to the following:

I hereby assure that the embryo from which the cell line(s) identified in item 6 of the form was derived was donated prior to July 7, 2009, and the embryo: 1) was created using in vitro fertilization for reproductive purposes and was no longer needed for this purpose; and 2) was donated by individuals who sought reproductive treatment ("donor(s)") who gave voluntary written consent for the human embryo to be used for research purposes.

Please let me know if you have any questions.

Sincerely,

Ellen Gadbois

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From: HESCREGISTRY (NIH/OD)  
Sent: Tuesday, January 25, 2011 4:37 PM  
To: smunne@reprogenetics.com; mina.alikani@embryos.net  
Subject: New hESCs Registry Application Request #2011-ADM-001

To: Santiago Munne (Signing Official)  
Mina Alikani (Submitter)

This is to confirm that the hESC Registry Application request, as detailed below, has just been submitted and is pending NIH Administrative review. You can expect to hear back from us about the status of your application soon.
Stem-cell research using discarded non-viable embryos

A – You are invited to donate any "non-viable" embryos that may result from your current attempt at in-vitro fertilization. Participation will in no way affect the normal progress of your attempt at in vitro fertilization, including the possibility of later replacement of thawed embryos from this attempt, nor will it affect your chances of pregnancy, as the study of non-viable embryos would not commence until completion of each of the steps necessary to identify any individual embryo as non-viable.

B – The Gamete and Embryo Research Laboratory of the Institute for Reproductive Medicine and Science at Saint Barnabas and Reprogenetics are engaged in studies aimed at developing new, more efficient ways of isolating and culturing embryonic stem cells. Stem-cells are cells that are not yet differentiated or changed into specialized cells such as muscle and blood cells. Embryos, even normal ones, can be used as a source of stem cells. Research on such embryonic stem cells is of fundamental importance for the progress of medicine, not least reproductive medicine, in the decades ahead. The reason is that stem cell research not only will allow a better understanding of the causes and mechanisms underlying a host of devastating diseases, but also offers a very real prospect of alleviating, even curing many of these diseases, among them Juvenile Diabetes and Muscular Dystrophy. Physicians and scientists at the Saint Barnabas Medical Center believe that the embryological research for which your non-viable embryos would be donated can play an important, even crucial role in the developments ahead.

Non-viable embryos are selected based on standard criteria used in our laboratory. Embryos are non-viable when eggs fertilize abnormally or fail to divide after fertilization. Others may divide too slowly or appear abnormal. Processes that are included in the evaluation of viability are fragmentation (the blebbing of cells) and the number of nuclei per cell. Fragments are not always considered detrimental, but certain types of fragmentation are known to cause demise of the embryo. Abnormal nuclei do not always cause failure of pregnancy, and the decision to consider such embryos abnormal is very much dependent on the ratio of abnormal cells and the type of abnormal nucleus. Criteria for evaluation are also stage-specific. In other cases embryos are considered non-viable because the embryo appears disorganized and several types of anomalies can be distinguished.

The non-viable embryos donated by you will be used in one or more of the following studies either before or after they are frozen:

- The development of culture techniques that will allow scientists to isolate and grow stem-cells from a single or a few embryonic cells. These studies may involve using animal cells, including cells from animal embryos, for initial growth support. The animal cells would serve as "feeder cells", providing the right environment until the human cells can survive by themselves.

Patient Initials

3/5/2009
- Isolation and analysis of stem-cells from embryos discarded due to abnormal chromosome numbers.
- The development of improved methods for stem-cell storage. Thawed stem-cells would be tested for viability and characteristics as well as their ability to transform into other cell types.
- The development of new methods for transforming stem-cells into other cell types.

Only your "non-viable" embryos, i.e. those that are unsuitable for replacement and would otherwise be discarded, are eligible for donation, and none of the experiments will give rise to normal embryos at any stage. However, the non-viable embryos donated by you may give rise to embryonic stem cells. These stem cells may be genetically normal or genetically abnormal depending, primarily, on whether the isolated cells from which they grew were genetically normal or abnormal to begin with. Both categories of stem cells are of fundamental scientific and biomedical interest.

The methods used in the studies include new processes of cell manipulation and culture of embryonic cells. Any non-viable embryos donated by you may thus be manipulated and cultured, in some instances for an extended period of time, or may be stored, in a frozen state, for later studies. These possibilities also apply to any embryonic stem cells isolated. If embryonic stem cells are obtained from your non-viable embryos, they could give rise to different cell types, such as pancreatic cells or neural cells. Frozen-thawed stem-cells and embryos will be subjected to the same research protocols as described above.

D - Participation in these studies by donation of non-viable embryos will cause no additional physical discomfort whatsoever during the conduct of your procedure, nor will your chances of becoming pregnant be affected in any way. It is also unlikely that you will receive any direct personal benefits from participation at this time; however, it is likely that, in the future, patients will benefit from the knowledge we will gain from these studies.

E - Disclosure of researchers' potential financial interests: In addition to their scientific interests in this research project, the individuals conducting this stem cell study might profit financially from the research. There may be current or potential financial benefits to the researchers, the participating institution(s), and other research institutions or researchers arising from discoveries made through this research project and the stem cells collected from your embryos. If you are undergoing fertility treatment, it is important that your doctor informs you of any personal benefits s/he may gain by your agreement to provide embryos for this project. The person who has been authorized to provide you with information may also have a personal vested interest in this research project. Please feel free to ask your doctor if you have any questions about this.

F - This consent form is being offered to all couples treated with IVF at Saint Barnabas Medical Center. Any information obtained during this study and identified with you will remain confidential. The Food and Drug Administration (FDA) and The Center for Disease Control (CDC) in association with the Society for Assisted Reproduction (SART) may inspect the
records. There are no added costs related to your participation in this study. Your decision whether or not to participate will not prejudice your future relations with Saint Barnabas Medical Center and the treatment you now undergo in this institute. If you decide to participate, you are free to discontinue participation at any time. Your participation is voluntary and your refusal to participate will involve no penalty or loss of benefits to which you are otherwise entitled. If you have any questions regarding your rights as a research subject, please call the office of the IRB chairman, Dr. Goodman at (973) 322-5637. The principle investigator of this protocol is Dr. G. John Garrisi, who can be reached at 973-322-8286. You may request a copy of this form.

G – 1/We hereby attest that we have read the entire consent form, or that it has been read to us, so that we understand it completely. I/we further attest that any and all questions of mine/us regarding this form or this study have been answered to my/our complete satisfaction. We agree that stem-cells may be developed from our non-viable embryos.

Signed: name female partner [redacted] Signature [redacted]

Signed: name male partner [redacted] Signature [redacted]

Witnessed: name Joan Engel Signature Joan Engel

Date: 12/20/2009
Stem-cell research using discarded non-viable embryos

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animal cells, including cells from animal embryos, for initial growth support. The animal cells would serve as “feeder cells”, providing the right environment until the human cells can survive by themselves.

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- The development of new methods for transforming stem-cells into other cell types.

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The methods used in the studies include new processes of cell manipulation and culture of embryonic cells. Any non-viable embryos donated by you may thus be manipulated and cultured, in some instances for an extended period of time, or may be stored, in a frozen state, for later studies. These possibilities also apply to any embryonic stem cells isolated. If embryonic stem-cells are obtained from your non-viable embryos, they could give rise to different cell types, such as pancreatic cells or neural cells. Frozen-thawed stem-cells and embryos will be subjected to the same research protocols as described above.

Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from, the non-viable embryos donated by you, will ever be used to impregnate another woman or to assist any individual or couple in any way to become pregnant. Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from the non-viable embryos donated by you, will ever be used for curing genetic or other disease, or for production of commercial stem-cell lines. Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from, the non-viable embryos donated by you, will ever be donated for use to other individuals, centers, or corporations.

D – Participation in these studies by donation of non-viable embryos will cause no additional physical discomfort whatsoever during the conduct of your procedure, nor will your chances of becoming pregnant be affected in any way. It is also unlikely that you will receive any direct personal benefits from participation at this time; however, it is likely that, in the future, patients will benefit from the knowledge we will gain from these studies.
F – This consent form is being offered to all couples treated with IVF at Saint Barnabas Medical Center. Any information obtained during this study and identified with you will remain confidential. The Food and Drug Administration (FDA) and The Center for Disease Control (CDC) in association with the Society for Assisted Reproduction (SART) may inspect the records. There are no added costs related to your participation in this study. Your decision whether or not to participate will not prejudice your future relations with Saint Barnabas Medical Center and the treatment you now undergo in this institute. If you decide to participate, you are free to discontinue participation at any time. Your participation is voluntary and your refusal to participate will involve no penalty or loss of benefits to which you are otherwise entitled. If you have any questions regarding your rights as a research subject, please call the office of the IRB chairman, Dr. Goodman at (973) 322-5637. The principle investigator of this protocol is Dr. G. John Garrisi, who can be reached at 973-322-8286. You may request a copy of this form.

G – I/We hereby attest that we have read the entire consent form, or that it has been read to us, so that we understand it completely. I/we further attest that any and all questions of mine/us regarding this form or this study have been answered to my/our complete satisfaction. We agree that stem-cells may be developed from our non-viable embryos.

Signed: name female partner __________________ Signature __________________

Signed: name male partner __________________ Signature __________________

Witnessed: name __________________ Signature __________________

Date: 10/16/08
Consent for In Vitro Fertilization/Assisted Reproduction

I / we, ____________________ and ____________________, desire to participate in the Assisted Reproduction Program at The Institute for Reproductive Medicine and Science at Saint Barnabas, P.A. (IRMS). We understand that there are a number of steps to this procedure and that starting this process does not guarantee that we will complete the process, achieve pregnancy or delivery of a healthy child. One of our physicians has discussed with you the etiology of your condition, and alternative therapies, if any, that are available. We understand that the female partner will receive medication to induce the maturation of several eggs and during this period she will undergo a surgical procedure to retrieve her eggs. The steps may be performed by any of the following physicians: Margaret Garrisi, Serena Chen, Patricia Hughes, Natalie Cekleniak. The egg retrieval procedure will be done by needle aspiration, usually under ultrasound guidance or perhaps by laparoscopy. We understand that the eggs will be prepared and inseminated in marked dishes with a sample of the male partner’s sperm after preparation, which removes the sperm from the seminal fluid. The embryos, which may result from fertilization, will be placed into the female partner’s uterus by means of a small catheter, which passes through her cervix under ultrasound guidance. Following this transfer, blood hormone levels will be monitored in the female partner to make sure that there are adequate hormonal levels to support a developing pregnancy and then to determine if a pregnancy has resulted. We understand that each of these steps may fail and carries known risks as well as theoretical concerns as detailed in the following paragraphs.

A - Ovulation induction

We understand that a variety of medications are available for the induction of ovulation including clomiphene citrate (Clomid, Serophene), human menopausal gonadotropins (Pergonal,) pure follicle stimulating hormone (urofollitropin, FSH), human chorionic gonadotropin (hCG), recombinant FSH (Gonal-F or Follistim) and GnRH-agonists (leuprolide acetate, Lupron), and GnRH antagonists (Antagon, Cetrotide). We understand that some of these medications must be given by intra-muscular injection, which may cause bruising or discomfort at the injection site. These medications may cause the ovaries to become over stimulated, leading to a condition called ovarian hyper stimulation syndrome (OHSS). We understand that in its most severe form this condition might require hospitalization for intravenous fluids and monitoring until the syndrome resolves. Worldwide there have been rare reports of death following severe OHSS. We therefore understand the importance of maintaining close contact with the IVF team during the time that these medications are being used and for two weeks afterwards.

We understand that before the start of a cycle, the male partner will be asked to supply a semen sample for analysis by the andrology laboratory. He may be asked to take a specific antibiotic during the first part of the stimulation cycle to treat bacteria that may be present in order to increase chances for a successful fertilization. In certain cases, semen may also be frozen in advance to be certain of its availability at the time of egg retrieval.
B - Monitoring protocol

We understand that while receiving the medications listed above, the female partner will be closely monitored by the IVF team. We understand that this monitoring will include daily blood drawing, which can cause mild discomfort and bruising at the puncture site. We understand that ultrasound examination of the ovarian follicles and the uterus will be performed frequently. These examinations may at times be uncomfortable, but have no known risks of any kind. The female partner may also be asked to collect urine samples for further hormone analysis. We understand that if monitoring suggests a low probability for successful egg retrieval, that the stimulation cycle will be stopped and no egg retrieval will occur. We also understand that we may be given the option of starting the ovarian stimulation procedure again in a subsequent cycle.

C - Egg retrieval

We understand that, at a time determined by the IVF team, the female partner would be admitted to the egg retrieval procedure room and IVF laboratories at Saint Barnabas Medical Center. We understand that in the vast majority of cases, ultrasound directed needle puncture of the follicles will be done. Rarely, the retrieval may be done by laparoscopy under general anesthesia. We understand that the procedure involves the small risk of general anesthesia as well as injury to bowel, bladder, or blood vessels, which might require a large incision (laparotomy) to repair. We understand that a separate informed consent will be obtained for a laparoscopic retrieval if it becomes necessary. With either type of egg retrieval, we understand that in rare cases there could be bleeding from the site where the ovaries were punctured. This may require laparotomy (an incision in the abdomen) if the bleeding cannot be controlled through the laparoscope. The risks of the procedure are similar to the risks of laparoscopy, including general anesthesia.

We understand that we cannot be guaranteed that the number of eggs predicted prior to retrieval will indeed be recovered or that any of the eggs will be normal or ripe. Some follicles may not yield eggs and rarely none of the follicles will yield eggs. The egg retrieval involves equipment such as incubators, suction apparatuses and ultrasound machines that may fail because of technical malfunction. We also understand that once the eggs are isolated in the laboratory, that blood and abnormal nursing cells are removed from around the egg using dissection needles and that although unlikely, some or all of the eggs may be damaged in the process. Eggs may also be damaged because of shock due to differences in conditions.

Initial egg yield numbers are counted once and rapidly in order to place the eggs inside the incubators and stabilize conditions as soon as possible. Eggs themselves are not visualized during egg retrieval, but only the nursing cells surrounding the eggs. The normality of the eggs cannot be assessed at egg retrieval. The exact number of eggs is only determined later on at insemination or ICSI.
D - Insemination, fertilization and embryo growth

Once retrieved, the eggs will be incubated in a special solution (culture medium) and evaluated for timing of insemination by the embryology team of the IVF program. We understand that a sample of semen from the male partner, obtained by masturbation in a private collection room near the laboratory, will be evaluated, prepared, and used for insemination. Semen collection in this way can be unsuccessful and if there are any doubts, a sample can be prepared and frozen in advance for thawing at this time. However, in case of unexpected failure it is possible to obtain spermatozoa from a testicle using a minor operative procedure (testicular sperm retrieval). Separate consent is needed for this procedure.

The seminal fluid that surrounds the spermatozoa must be removed prior to insemination. Sperm processing involves high centrifugal force, washing with an artificial colloidal suspension called Puresperm™ or by swim-up. We understand that the consistency of highly viscous semen will be reduced by an enzyme. The prepared semen may be exposed to substances intended to promote sperm movement or materials intended to remove toxic substances. The zygotes or fertilized eggs are changed over into a culture solution. This solution may be changed every 48 hours or more frequently. Solutions may be specially tailored to embryonic stage. The embryos are checked at least once daily and their development is determined. Embryos will remain in the solution(s) for 48-120 hours and then transferred.

Should a pregnancy occur, we understand that no risk to the fetus is presently known to medical science arising from the materials and methods used in the preparation and handling of eggs, semen and embryos. We understand that not all eggs recovered can be fertilized, and that it is possible that none of the eggs may fertilize. We also understand that not all eggs may be ripe and that ripe eggs may be fertilized multiple times by sperm or even self-fertilize without the sperm participating. Zygotes and later stage embryos may develop abnormally at any time.

E - Blastocyst culture, alternative culture and embryo transfer

We understand that between three to four days after egg retrieval our embryos will be placed into the uterine cavity of the female partner. Alternatively you may be asked to consider having embryos transferred at the blastocyst stage five or six days after egg retrieval, using commercial culture solutions that support growth for a longer period. This protocol may be especially advantageous to couples who are at risk of multiple pregnancy, since the extended culture increases the opportunity for embryologists to select the highest quality embryos. A potential disadvantage is that some embryos may be more sensitive to prolonged presence in the laboratory with the result that cryopreservation and/or embryo transfer may not occur.

We know that some patients have embryos that develop slowly, or show different forms of microscopic anomalies such as fragmentation and multi-nucleation and may not become pregnant. This may be due to a sensitivity of the embryos to the standard culture conditions in the
laboratory. If this is a repeat attempt at assisted reproduction, we may be asked to have our eggs and zygotes cultured in a different culture medium in order to determine whether this sensitivity can be overcome. We realize that alternative culture is only effective in a proportion of the patients. The alternative culture may not have any benefit in our case, and we understand that pregnancy and success is not guaranteed.

For any embryo transfer, a thin catheter will be passed through the cervix and into the uterus so the embryo may be deposited there. We understand that this may involve some cramping and discomfort, and possibly a small amount of bleeding. Infection could be introduced at the time of the catheter insertion into the uterus, requiring antibiotic therapy. We understand there is not guarantee that any of the embryos thus transferred will result in a pregnancy.

We understand that the success of IVF can often relate directly with the number of embryos transferred to the uterus. We also understand that IVF significantly increases the risk for multiple gestation (more than one baby), and that this risk also correlates directly with either the number of embryos transferred, their development, the age of the female partner (or egg donor), the number of prior attempts and other unknown factors. We also understand that in rare cases, embryos may split in two or three, resulting in multiple fetuses; on occasion this can mean that there are more fetuses than embryos transferred. There are distinct obstetric risks to multiple gestations, the most serious of which are pre-term labor and the delivery of premature infants who require intensive care. It is the policy of this program to replace anywhere from one to six embryos in a given cycle all depending on availability and factors such as your age, cycle attempt and embryonic parameters. Any additional viable embryos may be cryopreserved (frozen) for possible replacement in a subsequent cycle. We understand that a separate consent for the cryopreservation must be completed if the embryos are to be cryopreserved.

F - Post-transfer management

We understand that, in conjunction with the transfer of embryos, the female partner may be given natural progesterone by intramuscular injection, vaginal suppository, or oral capsule in an attempt to increase the chances for successful implantation. Sometimes an additional hCG injection may be given instead of the progesterone. Should a pregnancy result, we understand that no harmful effects to the mother of the fetus are presently known to medical science from the use of this natural progesterone or hCG supplementation. During this period, we understand that various blood hormone levels may be evaluated.

G - Disposition of unwanted or unsuitable cells, fluids, spermatozoa, eggs and embryos

Bloods, blood products and cells as well as follicular and seminal fluids and cells contained therein obtained during follicular monitoring, egg or sperm retrieval, may be used for scientific observations. In the event that we have unused or unripe spermatozoa, these may be subjected to scientific observations or discarded without any further observations. Under no circumstances will these spermatozoa be used for fertilization purposes or donation to other individuals, couples, corporations or institutions. In the event that we have unripe, unfertilized or abnormally fertilized
eggs, these may be subjected to scientific observations or discarded without further studies. We understand that these eggs will never be used for fertilization purposes or donation to other individuals, couples, corporations or institutions and that further growth of them will be ceased immediately after the observation. We also understand that these eggs are unwanted and considered abnormal. Embryos that arrest after 1-6 days after egg retrieval, that are partially degenerate or for any other reasons considered unsuitable for embryo transfer or cryopreservation may be observed to determine cellular inclusions, genes, gene mutations, proteins and chromosomes. The studies use protocols that will cease the immediate growth of individual cells. We understand that these embryos or their cells will never be used for purposes other than those described and will never be offered to other individuals, couples, corporations or institutions. We also understand that these embryos or their cells are unwanted and considered abnormal.

A separate research consent form is being offered to you, by which you can grant permission for the use of your unwanted and unsuitable spermatozoa, eggs, embryos and their cells in research studies for the development of new reproductive technologies. This separate research protocol is being supervised by the Internal Review Board (IRB) of Saint Barnabas Medical Center, and you are not in any way required to give your permission. Your further and ongoing treatment at The Institute for Reproductive Medicine (IRMS) will not be affected if you do not wish to participate in that or any other IRB-reviewed protocol.

H - Use of blood products

We understand that maternal blood serum, derived from blood collected from the female partner shortly before egg collection will be prepared for culture with your eggs and embryos in order to promote growth. Sometimes we will not be able to use your serum. It has been shown in rare instances that maternal serum may inhibit embryos developing in-vitro. Human serum albumin, a commercially prepared blood product for clinical laboratory use, is added to the egg collection fluid, micromanipulation, and semen preparation fluids. Careful screening is done by the manufacturers to reduce the likelihood of transmission of infectious diseases such as HIV, Hepatitis B and C. To date there have been no documented cases of disease transmission linked to human serum albumin usage in our Center. We understand and accept the risk that use of these blood products could result in the transmission of HIV, Hepatitis and/or other viral or possibly as yet unknown non-viral diseases.

I - Use of chemical substances, disposable items and mechanical devices during the procedures

A large number of chemical substances (sugars, salts, enzymes, proteins), mechanical devices (incubator chambers, microscopes, air handling systems, filters, standard laboratory equipment) and disposable items (pipettes, petri dishes, flasks, microtools) are used during the laboratory procedures. There may be unknown risks associated with the use of any of these items that cause your procedure to fail, even though checks and quality control measures are performed on a regular basis. Thus far we do not know of any association between the use of these materials and anomalies of pregnancy and fetal development, but underlying unidentified problems may nevertheless exist. An enzyme made from cow testis called hyaluronidase is routinely used to
remove nursing cells from around the eggs, and there is a chance that this enzyme may inadvertently remove the zona pellucida (the layer surrounding eggs) and cause your procedure to fail. Another enzyme called chymotrypsin made from cow pancreas is used to reduce the viscosity of seminal fluid. This enzyme may also in very rare instances inadvertently remove the zona pellucida and also may cause your procedure to fail.

J - Risks associated with procedures

Based on current medical knowledge, we understand there does not appear to be a higher incidence of birth defects associated with IVF procedures. However, there is not at present sufficient statistical data available to definitively conclude that this is so. Therefore, we understand that IVF may impose risks to the fetus during development. We also understand that because more than one embryo or egg may be transferred, there may be a higher incidence of multiple births. An embryo may split when inside the uterus, forming monozygotic twins and there may be other associated anomalies. In certain cases, fetal reduction may be considered if more embryos implant that can be medically (or personally) deemed advisable to carry through a pregnancy. We also understand that ectopic or tubal pregnancies may occur in the procedure. These associated procedures can also produce increased financial and emotional burdens.

We understand and accept that the use of ovarian fertility drugs may be associated with an increased risk of ovarian diseases in later life, including cancer. We recognize that the exact risk, if any, has yet to be established and may not be known for many years.

K - Success rate and outcome

We understand that failure to obtain a pregnancy may result from many reasons, including the following:

- Maturation of the egg(s) may not occur, or the time of the egg maturation may be misjudged, may not be predictable or may not take place in the monitored cycle.
- Pelvic adhesions may prevent access to the ovary with the follicles, thus causing the procedure to obtain the egg from the ovary to fail.
- The egg(s) obtained from the wife may be abnormal.
- Normal spermatozoa may not be available.
- Normal fertilization of the egg(s) by the sperm may not occur.
- Cleavage or growth of the embryo(s) may not occur at any day of development or the embryo(s) may not develop normally.
- The embryo(s) may become infected in the laboratory or an unforeseen laboratory accident may result in loss or damage to the eggs, sperm, or embryo(s).
- The embryo(s) may become contaminated by infection in the semen or bacteria from the vagina.
- Some embryos may not develop well in approved commercial culture medium despite standard testing.
- Implantation of the embryo(s) in the uterus after embryo transfer may not occur or an early pregnancy may be lost after an initial positive result.
- Even if a pregnancy is established, we understand that delivery of a child may not occur due to miscarriage, ectopic pregnancy (outside the uterus), stillbirth, or other complications associated with pregnancy and delivery.
- There may be unknown side-effects from any of the procedures used resulting in abnormal pregnancy or abnormal fetal development.

We understand that the members of the IVF team cannot guarantee that a pregnancy will result from this procedure. Even in normally fertile couples, the chance of pregnancy is approximately 25% in a given natural cycle. If no pregnancy occurs, we may be offered participation in future cycles when assessment by the IVF team reveals no contra-indications. We understand that the IVF team cannot guarantee the normality of any infant that results from this procedure.

We understand that we may at any time decide to withdraw from participation in this program without prejudice. Any information obtained during this procedure and identified with us will remain confidential and will be disclosed only with our permission. Any publication resulting from this procedure will not identify us individually. Representatives of The Food and Drug Administration (FDA), The Center for Disease Control (CDC) and The Department of Health of New Jersey may inspect the records.

We have been encouraged to ask questions and any that we have asked have been answered to our satisfaction. A member of the IVF team will answer future questions.

(Signature of patient)  
Date

(Signature of Partner, if not applicable write NA)  
Date

(Signature of Witness)  
Date

The Institute for Reproductive Medicine and Science at Saint Barnabas, is not owned by, and its physicians and other personnel are not employed by, Saint Barnabas Health Care System, Saint Barnabas Medical Center, or any of their affiliates.
August 18, 2004

Robert L. Goodman, MD
Chairman, Institutional Review Board
Saint Barnabas Medical Center
Old Short Hills Road
Livingston, NJ 07039

Dear Dr. Goodman:

I would like to submit the attached revised consent form entitled *Stem-cell research using discarded non-viable embryos* to the Institutional Review Board of Saint Barnabas Medical Center to be considered for expedited IRB review.

There have been some minor changes to the consent form, which have been highlighted. In reviewing the consent, it was also noticed that the telephone number for the IRB office has not been updated. I have highlighted this change as well.

With warmest regards,

Santiago Munné, Ph.D.

SM/laf
Enclosure
APPLICATION TO UNDERTAKE RESEARCH INVOLVING HUMAN SUBJECTS

Checklist

TITLE OF PROPOSED RESEARCH:
Stem-cell research using discarded non-viable embryos

INVESTIGATOR(S) AND DEPARTMENT(S):
Steen Willadsen, DVM, Jacques Cohen, Ph.D., Mina Alikani, M.Sc., Margaret Garrisi, M.D.,
Santiago Munné, Ph.D. Institute for Reproductive Medicine and Science, Department of
Gynecology and Obstetrics

THIS PROTOCOL IS TO BE CONSIDERED FOR
(check only one box)  
______ EXEMPTION FROM REVIEW
______ EXPEDITED REVIEW
______ FULL IRB REVIEW

The proposed research will be carried out in cooperation with the following institution
Reprogenetics (if applicable)
Date of approval by cooperating institution's IRB

Human subjects would be involved in the proposed research as either:

______ None of the following

 or including (Check all applicable boxes)

_____ Minors
______ Fetuses
______ Abortuses
______ Pregnant women
______ Prisoner
______ Mentally retarded
______ Mentally disabled
______ Non-English-speaking subjects
APPLICATION TO UNDERTAKE RESEARCH INVOLVING HUMAN SUBJECTS

Part A (to be completed in typescript by all applicants)

1). Date 7/31/2004

2). IRB No. (leave blank)

3). Title of Project

Stem-cell research using discarded non-viable embryos

4). Responsible Investigator(s) Full Name(s) and Department(s)

Steen Willadsen, DVM, Jacques Cohen, Ph.D., Mina Alikani, M.Sc., Margaret Garrisi, M.D., Institute for Reproductive Medicine and Science of Saint Barnabas, Department of Gynecology and Obstetrics; Gamete and Embryo Research Laboratory at the Institute for Reproductive Medicine and Science of Saint Barnabas
Santiago Munne, Ph.D., Reprogenetics LLC

5). Personal Signature of Investigator(s)

Steen Willadsen

Jacques Cohen

Mina Alikani

Margaret Garrisi

Santiago Munné

6). Personal Signature of Department Chairman

7). Source of Funds for Study

Departmental

Number of Subjects to be Studied

1500

Dates of Study September 1, 2002 to August 31, 2005

Is this an multi-institutional study? ☑ Yes _____ No
APPLICATION TO UNDERTAKE RESEARCH INVOLVING HUMAN SUBJECTS

Part A (to be completed in typescript by all applicants)

1). Date 9/31/2004

2). IRB No. (leave blank)

3). Title of Project

Stem-cell research using discarded non-viable embryos

4). Responsible Investigator(s) Full Name(s) and Department(s)

Steen Willadsen, DVM, Jacques Cohen, Ph.D., Mina Alikani, M.Sc., Margaret Garrisi, M.D., Institute for Reproductive Medicine and Science of Saint Barnabas, Department of Gynecology and Obstetrics; Gamete and Embryo Research Laboratory at the Institute for Reproductive Medicine and Science of Saint Barnabas, Santiago Munné, Ph.D., Reprogenetics LLC

5). Personal Signature of Investigator(s)

Steen Willadsen
Mina Alikani
Santiago Munné

6). Personal Signature of Department Chairman

Veronica Ravniker, M.D.

7). Source of Funds for Study

Departmental
Number of Subjects to be Studied
1500

Dates of Study September 1, 2002 to August 31, 2005

Is this an multi-institutional study? X Yes No
If yes, under what sponsorship? Institute for Reproductive Medicine and Science at Saint Barnabas, and Reprogenetics
8). Background of Proposed study (State briefly the reason for doing the study. What question(s) is it designed to answer and why is the question being asked? Include references to no more than three key literature citations, if appropriate.)

Abnormal gametes and embryos are an excellent source of material for studying anomalies that can occur during gametogenesis and embryogenesis. Discarded non-viable material resulting from in-vitro fertilization (IVF) can also be used to develop new technology applicable to the treatment of infertile patients (protocol #769). Protocol 769 was revised in February 2002 and allowed us to extend the study of non-viable embryos into the field of embryonic stem-cell research. We have developed several technologies that could almost certainly be used to make producton of human embryonic stem-cells from embryos simpler and more efficient. The fact that these technologies primarily use non-viable embryos as the source of embryonic stem-cells is of particular relevance in the current funding climate which does not allow the use of viable human embryos for that purpose (Alikani and Willadsen; Reproductive Medicine Online; 5, 56-58, 2002). However, we have not yet developed stem-cell lines from such embryos. Here we propose to develop the technology further with the endpoint of stem-cell formation. The isolation of chromosomally normal and abnormal stem-cell lines from non-viable embryos is important. For one thing it avoids the dilemma posed by the use of viable human embryos for stem-cell production, and may therefore allow government funding. The reason for making the present protocol separate from protocol 769 using non-viable embryos is the need for patients’ permission to use discarded non-viable embryos for isolation of stem-cells. Inherent in this kind of research is the unpredictability of the duration of individual experiments, particularly since abnormal embryos and stem-cells can be frozen. The isolation and experimental culture of embryonic stem cells is considered to be of fundamental importance for the progress of medicine. Stem cell based research offers the prospect of cures to many devastating diseases such as Juvenile Diabetes, Muscular Dystrophy and Alzheimer’s Disease.

The reason of the current update of this application is because of our long relationship with Dr. Munne, before an employee of Saint Barnabas, and now directing Reprogenetics, we want to expand the study to develop stem cells from chromosomally normal embryos detected from Preimplantation Genetic Diagnosis cycles.

9). Outline of Proposed Study (State briefly but precisely what is to be done. If the study is to be conducted according to a detailed protocol of a pharmaceutical company or outside agency, include a precis here and attach the full protocol as an appendix. If the study involves the use of a questionnaire or structured interview, attach the text of such instruments as an appendix. Note: IRB review focuses on the scientific merit and adequacy of experimental design as well as on issues of safety and protection of confidentiality).

Non-viable embryos donated by patients will be selected based on criteria used in our program. These are embryos that contain too few normal cells – often no normal cells at all – for the
formation of a viable/ normal conceptus. Non-viable embryos include those that have
detrimental forms of fragmentation or multi-nucleation, or that stopped growing during serial
observation. Other embryos may seem viable following preimplantation genetic diagnosis
(PGD) (that is they divide) but are diagnosed as trisomic or monosomic. Yet other abnormal
embryos are polyploid, haploid or chaotic mosaic.

The zona pellucida of the embryos will be removed, and individual mono-nucleate cells will be
isolated. They will be grown together from cohorts of embryos in order to form pseudo-
blastocysts (Alikani and Willadsen, 2002) or cultured individually with mouse or cow
embryonic cells or embryos that have been manipulated to support the growth of human
embryonic cells to develop into stem-cells. Animal and human cells will be immunologically or
mechanically separated, but in some models the cells can be isolated by varying culture
conditions. The stem-cell development will be studied by analyzing the expression of Oct-4, a
gene that only is expressed in early embryos, and the inner cell mass and embryonic stem cells.
Ploidy will be studied by fluorescence in-situ hybridization (FISH).

10) Subject (State the kind(s), ages, sex, and appropriate numbers of subjects to be
studied. Indicate the criteria for the selection of the proposed kinds and number of
subjects. If populations at special risk (cf. checklist) are to be studied, provide the
reason(s) for their inclusion. Specify population groups to be excluded (e.g.
pregnant women). Indicate composition of control groups.)

The investigations will be conducted in 500 patients annually. There are no risks to the couple
and no specific groups will be excluded.

11). Drugs (Provide specific information on proposed dosage levels and schedules.
Include both the generic and commercial name of each drug and summarize
available information on efficacy and side effects. If none are to be administered,
enter this statement: "No drugs will be given.")

No additional drugs are needed for the completion of the proposed studies other than those
used for assisted reproduction procedures.

12). Blood Sampling (State the volume to be drawn on each occasion and the frequency
of sampling from the same subject. If none is to be drawn, enter this statement:
"No blood will be drawn.")

No blood will be drawn.

13). Tissue Sampling (State the type of tissue to be collected, volume (if applicable), and
frequency of sampling from the same subject (if applicable). If no other tissues will
be collected, enter this statement: "No other tissues will be collected.")

No extra tissues will be collected.

14). Radioactive Isotopes (Give the identity and dose of each isotope. If none are to be
administered, enter this statement: "No radioactive isotopes will be given").
No radioactive isotopes will be given.

15). Safety (State in adequate detail any anticipated physical, mental, or emotional risk to the subjects of this research activity and the degree of likelihood that it may occur. If no such risk is anticipated, state why this is so). There is no additional risk to the patients other than those involved with regular IVF.

16). Confidentiality (Describe the adequate detail what measures will be taken to protect the confidentiality of the data to be obtained and the subject's rights to privacy). Patient names, IDs, etc. will be kept strictly confidential. Files and databases will be locked or only made accessible to privileged investigators. Anonymous case reports will only be published after the patients grant permission.

17). Informed Consent (State whether it is planned to obtain informed consent from each subject of the research activity. If informed consent will be obtained, complete the attached Part B. It is not planned to do so, provide in this space, thorough justification for its omission. Note: Federal regulations provide that written informed consent be obtained from each subject of a research activity, but this requirement may be waived by the IRB under specific circumstances (cf. attachment to Part B). We will ask each couple to provide written consent after fully informing them in a private consultation. The IRB consent form is attached.
Stem-cell research using discarded non-viable embryos

A – You are invited to donate any “non-viable” embryos that may result from your current attempt at in-vitro fertilization. Participation will in no way affect the normal progress of your attempt at in vitro fertilization, including the possibility of later replacement of thawed embryos from this attempt, nor will it affect your chances of pregnancy, as the study of non-viable embryos would not commence until completion of each of the steps necessary to identify any individual embryo as non-viable.

B – The Gamete and Embryo Research Laboratory of the Institute for Reproductive Medicine and Science at Saint Barnabas and Reprogenetics are engaged in studies aimed at developing new, more efficient ways of isolating and culturing embryonic stem cells. Stem-cells are cells that are not yet differentiated or changed into specialized cells such as muscle and blood cells. Embryos, even abnormal ones, can be used as a source of stem cells. Research on such embryonic stem cells is of fundamental importance for the progress of medicine, not least reproductive medicine, in the decades ahead. The reason is that stem cell research not only will allow a better understanding of the causes and mechanisms underlying a host of devastating diseases, but also offers a very real prospect of alleviating, even curing many of these diseases, among them Juvenile Diabetes and Muscular Dystrophy. Physicians and scientists at the Institute for Reproductive Medicine and Science and Reprogenetics believe that the embryological research for which your non-viable embryos would be donated can play an important, even crucial role in the developments ahead.

Non-viable embryos are selected based on standard criteria used in our laboratory. Embryos are non-viable when eggs fertilize abnormally or fail to divide after fertilization. Others may divide too slowly or appear abnormal. Processes that are included in the evaluation of viability are fragmentation (the blebbing of cells) and the number of nuclei per cell. Fragments are not always considered detrimental, but certain types of fragmentation are known to cause demise of the embryo. Abnormal nuclei do not always cause failure of pregnancy, and the decision to consider such embryos abnormal is very much dependent on the ratio of abnormal cells and the type of abnormal nucleus. Criteria for evaluation are also stage-specific. In other cases embryos are considered non-viable because the embryo appears disorganized and several types of anomalies can be distinguished. Finally, embryos classified by Preimplantation Genetic Diagnosis (PGD) as being genetically abnormal were also considered non-viable.
The non-viable embryos donated by you will be used in one or more of the following studies either before or after they are frozen:

- The development of culture techniques that will allow scientists to isolate and grow stem-cells from a single or a few embryonic cells. These studies may involve using animal cells, including cells from animal embryos, for initial growth support. The animal cells would serve as “feeder cells”, providing the right environment until the human cells can survive by themselves.
- Isolation and analysis of stem-cells from embryos discarded due to abnormal chromosome numbers.
- The development of improved methods for stem-cell storage. Thawed stem-cells would be tested for viability and characteristics as well as their ability to transform into other cell-types.
- The development of new methods for transforming stem-cells into other cell types.

Only your “non-viable” embryos, i.e. those that are unsuitable for replacement and would otherwise be discarded, are eligible for donation, and none of the experiments will give rise to normal embryos at any stage. However, the non-viable embryos donated by you may give rise to embryonic stem cells. These stem cells may be genetically normal or genetically abnormal depending, primarily, on whether the isolated cells from which they grew were genetically normal or abnormal to begin with. Both categories of stem cells are of fundamental scientific and biomedical interest.

The methods used in the studies include new processes of cell manipulation and culture of embryonic cells. Any non-viable embryos donated by you may thus be manipulated and cultured, in some instances for an extended period of time, or may be stored, in a frozen state, for later studies. These possibilities also apply to any embryonic stem cells isolated. If embryonic stem-cells are obtained from your non-viable embryos, they could give rise to different cell types, such as pancreatic cells or neural cells. Frozen-thawed stem-cells and embryos will be subjected to the same research protocols as described above.

Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from, the non-viable embryos donated by you, will ever be used to impregnate another woman or to assist any individual or couple in any way to become pregnant. Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from the non-viable embryos donated by you, will ever be used for curing genetic or other disease, or for production of commercial stem-cell lines. Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from, the non-viable embryos donated by you, will ever be donated for use to other individuals, centers, or corporations.

D – Participation in these studies by donation of non-viable embryos will cause no additional physical discomfort whatsoever during the conduct of your procedure, nor will
your chances of becoming pregnant be affected in any way. It is also unlikely that you will receive any direct personal benefits from participation at this time; however, it is likely that, in the future, patients will benefit from the knowledge we will gain from these studies.

F – This consent form is being offered to all couples treated with IVF at Saint Barnabas Medical Center. Any information obtained during this study and identified with you will remain confidential. The Food and Drug Administration (FDA) and The Center for Disease Control (CDC) in association with the Society for Assisted Reproduction (SART) may inspect the records. There are no added costs related to your participation in this study. Your decision whether or not to participate will not prejudice your future relations with Saint Barnabas Medical Center and the treatment you now undergo in this institute. If you decide to participate, you are free to discontinue participation at any time. Your participation is voluntary and your refusal to participate will involve no penalty or loss of benefits to which you are otherwise entitled. If you have any questions regarding your rights as a research subject, please call the office of the IRB chairman, Dr. Goodman at (973) 322-5548. The principle investigators of this protocol are Dr. Steen Willadsen and Santiago Munne, who can be reached at 973-3226236, respectively. You may request a copy of this form.

G – I/We hereby attest that we have read the entire consent form, or that it has been read to us, so that we understand it completely. I/we further attest that any and all questions of mine/us regarding this form or this study have been answered to my/our complete satisfaction. We agree that stem-cells may be developed from our non-viable embryos.

Signed: name female partner ________________  Signature __________________

Signed: name male partner ________________  Signature __________________

Witnessed: name __________________________ Signature __________________

Date: ________________________________

IRB Consent Form 3 Revised 8/18/2004
SAINT BARNABAS MEDICAL CENTER
AUTHORIZATION TO USE AND DISCLOSE HEALTH INFORMATION FOR RESEARCH
(HIPAA 45 CFR PART 164)

Patient Name: ___________________________ Patient MRN: ___________________________


The purpose of this research is as follows: To develop chromosomally normal stem cells from chromosomally abnormal and unviable embryos diagnosed as such by PGD.

A representative of Saint Barnabas Medical Center or principal investigator must answer these questions completely before providing this authorization form to you. PLEASE DO NOT SIGN A BLANK FORM. You or your personal representative should read the descriptions below before signing this form.

This document specifically relates to the uses and disclosures of your “protected health information” or “PHI” as referred to in the federal law. For study purposes, your PHI may include records of your blood samples, physical examinations, test results, medical history and any other data collected or reviewed during the course of the study.

By signing this authorization, you are agreeing that your physicians and other health care providers may provide Saint Barnabas Medical Center and the investigators with the PHI they request for purposes of the study. You also are agreeing that Saint Barnabas Medical Center and the investigators may, for purposes of the study, use your PHI collected or created as part of the study and share this information with the parties described below. Additionally, you are agreeing that, during the study, you may not have access to the PHI obtained or created as part of this study, although you will have access to this information once the study is finished.

This information may be shared, with the individuals and/or organization described on this form, these parties may not required to protect the privacy of your information by law. It is your right to refuse to sign this authorization form. If you do not sign this form, you will not be able to participate in the research study. You will not receive the research treatment that was described to you. Your health care outside the study will not be affected. The payment for your health care or your health care benefits will not be affected.

If you sign this authorization form, you will have the right to withdraw it at any time. To withdraw, the authorization will prohibit further use or disclosure of your health information. If the hospital has already used your health information approved by your authorization or needs the information to fulfill an obligation or analyze the data, the use or disclosure can not be stopped. This authorization form will not expire unless you withdraw it. If you want to withdraw this authorization, please write to Robert Goodman, MD, IRB Chairman, 94 Old Short Hills Road, Livingston, New Jersey 07039

1. Who will have access to and/or use your health information?

The following individuals and/or organization(s) may have access to use, disclose or receive my identifiable health information. They may only share the information to the individuals/parties indicated on this list. This information must be shared with you, the research subject and/or your personal representative, as required by law.

☐ Every research site for this study, including Saint Barnabas Medical Center and the research support staff (for example, research study assistant) at each study location: Reprogenetics, LLC (West Orange, NJ).

☐ Every health care personnel who provides services to you in connection with this study.

☐ Any laboratories, other individuals/organizations that analyze your health information in connection with this study.

☐ The following research sponsors: Reprogenetics, LLC (West Orange, NJ).

☐ The National Cancer Institute and/or the National Institute of Health.

☐ The United States Food and Drug Administration, or other governmental surveillance organization.

☐ The members and staff of the hospital’s Institutional Review Board and Privacy Board.

☐ Principal Investigator and Co-Principal Investigation(s): Santiago Munné, PhD, Mercedes Garcia Bermudez, PhD, Pere Colls, PhD.

☐ Members of the Research Team including the participating investigators, research assistants, clinical nurses, and clerical support staff.

☐ Employees of the Contract Research Organization (CRO):

☐ Others (as described below):
II. What information will be used or disclosed?

The boxes checked below should provide you with a description of the information to be used or disclosed in the research.

☐ Your entire research record
☐ Any part of your medical records held by the hospital
☐ The following information: The genetic diagnosis of your oocytes, embryos or PGD procedure, if stem cells were obtained from those embryos and the characteristics of those stem cells, female age.

SPECIFIC UNDERSTANDINGS

III. Research Participant or Research Participant’s Representative Must Read and Initial the Following:

1. I understand that my participation in the research discussed by this authorization is conditioned upon my signing of this research authorization.

For Clinical Trials:

2. I understand that, despite any statements made in other places in this authorization, while the clinical trial is ongoing, Investigator or Hospital may refuse to permit me access to individually identifiable health information obtained in the course of the clinical trial. I also understand that I will be granted access to this health information following the completion of the clinical trial.

SIGNATURE

Signature of Subject or Personal Representative

Initials

Date

Print Name of Subject or Personal Representative

Description of Personal Representative’s Authority

Expiration Date:

☐ None
☐ End of Research 3/30/2006
☐ Date:__________________

CONTACT INFORMATION

The contact information of the subject or personal representative who signed this form should be filled in below.

Address: ___________________________ Telephone: _________________________(daytime)

_____________________________ _________________________(evening)

THE SUBJECT OR HIS OR HER PERSONAL REPRESENTATIVE MUST BE PROVIDED WITH A COPY OF THIS FORM AFTER IT HAS BEEN SIGNED.
Dear Dr. Gadbois,

Attached, please find all the documentation related to the protocol in question. You will note that this IRB-approved protocol (02-33) was in place and valid at the time of these donations. It was renewed and reapproved on a yearly basis, as we are required to submit a report to the IRB and state whether we wished to continue the study.

Please let me know if this answers the WG's concerns.

Best wishes,

Mina Alkani, Ph.D.

On 2/16/11 12:50 PM, "HESCREGISTRY (NIH/OD)" <hesregistry@mail.nih.gov> wrote:

Thank you. To go back to the protocol: the protocol you submitted with RN17 is attached, so can you check whether this is the correct protocol for RN11 and 12? The protocol was approved for use from September 1, 2002-August 31, 2005. We note that RN11 and 12 came from embryos donated in 2008 (as did RN17), so please confirm we have the protocol that was in effect during the donations of all of those embryos.

Sincerely,
Ellen Gadbois

On 2/15/11 1:09 PM, "HESCREGISTRY (NIH/OD)" <hesregistry@mail.nih.gov> wrote:

Thank you Dr. Alkani. Can you also confirm whether the new consent document signed by the donors of the embryos from which RN11 and RN12 were derived are identical to the reconsent document that was included in the RN17 submission (but not signed by those donors)?

Sincerely,
Ellen Gadbois

On 2/14/11 2:22 PM, "HESCREGISTRY (NIH/OD)" <hesregistry@mail.nih.gov> wrote:

Certainly. Please don't forget the other request (attached) regarding the two submissions that have been moved to the ACD Working Group for review—we need to have the protocol that was in effect at the time of donation of those embryos. It may be the same version as the protocol you submitted with your first submission, but we don't assume anything.
September 24, 2002

Jacques Cohen, Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange, NJ. 07052

RE: IRB Study # 02-33

Dear Dr. Cohen:

Meeting Date: 8/20/2002 At: Saint Barnabas Med Ct

Protocol Title:
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

To advise you that the above referenced Study has been presented to the Institutional Review Board identified above, and the following action taken subject to the conditions and explanation provided below.

Internal #: New Appl
Expiration Date: 8/19/2003
On Agenda For: initial Submission
Reason 1:
Description Date Received- 7/31/2002
IRB ACTION: Approved
Action Protocol (dated 8/26/02) and consent form (dated 8/23/02) approved
Explanation: IRB regulations require submission of an annual report and prompt notification of untoward events. At the completion of your study a final report should be submitted.

Sincerely,

[Signature]
Robert L. Goodman, M.D.
Chairman, IRB
November 02, 2004

Jacques Cohen, Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange, NJ. 07052

RE: IRB Study # 02-33  At: Saint Barnabas Med Ct

Dear Dr. Cohen:

Protocol Title:
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

This letter is to acknowledge the receipt of the information identified below.

Expiration Date: 8/19/2003
Our Internal #: 1072
Type of Change: Consent Modification Approved
Expedited?: 
Date of Change: 11/2/2004
Date Received: 11/2/2004
On Meeting Date: 11/16/2004
Description: Revised informed consent form dated 8/18/04
- Addition of Reprogenetics
- Update of contact information for Investigators and IRB
- Addition of sentence "Finally, embryos classified by Preimplantation Genetic Diagnosis (PGD) as being genetically abnormal were also considered non-viable."

Updated IRB application - Dr. Múnee previously employed by Saint Barnabas is now the Director of Reprogenetics

Respectfully yours,

N. Peter Zauber, M.D.
Co-Chairman, IRB
Friday, November 19, 2004

Jacques Cohen, Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange, NJ. 07052

RE: IRB Study # 02-33

Dear Dr. Cohen:

Meeting Date: 11/16/2004 At: Saint Barnabas Med Ct

Protocol Title:
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

To advise you that the above referenced Study has been presented to the Institutional Review Board identified above, and the following action taken subject to the conditions and explanation provided below.

Internal #: 1090
Expiration Date: 11/15/2005
On Agenda For: Renewal
Reason 1: Progress Report Reason 2:
Description: HIPAA Tool #1
IRB ACTION: Approved
Action Please make HIPAA Tool #1 available to all patients you are planning to enroll.
Examination: IRB regulations require submission of an annual report as long as patients are enrolled in the study and prompt notification of untoward events. At the completion of your study a final annual report should be submitted.

Sincerely,

Robert L. Goodman, M.D.
Chairman, IRB
January 18, 2006

Jacques Cohen, Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange, NJ. 07052

RE: IRB Study # 02-33

Dear Dr. Cohen:

**Meeting Date:** 1/17/2006  
**At:** Saint Barnabas Med Ct

**Protocol Title:**
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

To advise you that the above referenced Study has been presented to the Institutional Review Board identified above, and the following action taken subject to the conditions and explanation provided below.

**Internal #:** 1660  
**Expiration Date:** 1/16/2007  
**On Agenda For:** Renewal  
**Reason 1:** Progress Report  
**Description:**  
**IRB ACTION:** Approved  
**Action**  
**Explanation:** IRB regulations require submission of an annual report and prompt notification of untoward events. At the completion of your study a final report should be submitted.

Sincerely,

Robert L. Goodman, M.D.
Chairman, IRB
February 21, 2007

Jacques Cohen Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange NJ 07052

RE: IRB Study # 02-33

Dear Dr. Cohen:

Meeting Date: 2/20/2007 At: Saint Barnabas Med Ctr

Protocol Title:
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

This is to advise you that the above referenced Study has been presented to the Institutional Review Board, and the following action taken subject to the conditions and explanation provided below.

Internal #: 2201
Expiration Date: 2/19/2008
On Agenda For: Renewal
Reason 1: Progress Report Reason 2: Revised informed consent form dated 1/25/07 - updates Principal Investigator's current contact information
IRB ACTION: Approved
Condition 1: Approval
Explanation: IRB regulations require submission of an annual report and prompt notification of untoward events. At the completion of your study a final report should be submitted.

Sincerely,

Robert L. Goodman, M.D.
Chairman, IRB
April 9, 2009

Jacques Cohen Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange NJ 07052

Dear Dr. Cohen:

This letter is to acknowledge the receipt of the information identified below.

Our Study # 02-33

Protocol Title: STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

Expiration Date: 3/17/2009
Our Internal #: 3229
Type of Change: Consent Modification
Expedited ?: Yes
Date of Change: 4/9/2009
Date Received: 4/9/2009
On Meeting Date: 4/21/2009
Description: Revised informed consent form dated 3/13/09

Page 2, paragraph 3, has been deleted: "Neither the non-viable embryos donated by you, nor any cells isolated from, or developing from the non-viable embryos donated by you, will ever be used for curing genetic or other disease, or for production of commercial stem-cell lines." The reasons for this are obvious. Stem-cell lines once obtained are rare commodities and clinically important. They can not be shared with universities or other institutions by the self-imposed limitations of this paragraph. This paragraph has been replaced with Section E which was verbatim obtained from guidelines provided by NIH regarding this topic.

Section E states 'Disclosure of researchers' potential financial interests: In addition to their scientific interests in this research project, the individuals conducting this stem cell study might profit financially from the research. There may be current or potential financial benefits to the researchers, the participating institution(s), and other research institutions or researchers arising from discoveries made through this research project and the stem cells collected from your embryos. If you are undergoing fertility treatment, it is important that your doctor informs you of any personal benefits s/he may gain by your...
agreement to provide embryos for this project. The person who has been authorized to provide you with information may also have a personal vested interest in this research project. Please feel free to ask your doctor if you have any questions about this.

Respectfully yours,

[Signature]
N. Peter Zauber, M.D.
Co-Chairman, IRB
April 21, 2009

Jacques Cohen Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange NJ 07052

RE: IRB Study # 02-33

Dear Dr. Cohen:

Meeting Date: 4/21/2009 At: Saint Barnabas Med Ct

Protocol Title:
STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS
This is to advise you that the above referenced Study has been presented to the Institutional
Review Board, and the following action taken subject to the conditions and explanation
provided below.

Internal #: 3224
Expiration Date: 4/20/2010
On Agenda For: Renewal
Reason 1: Progress Report Reason 2:
Description:
IRB ACTION: Approved
Condition 1:
Action Explanation: IRB regulations require submission of an annual report and prompt notification
of untoward events. At the completion of your study a final report should be
submitted.

PLEASE NOTE: The Saint Barnabas Medical Center IRB Policy #29 states
under Investigator Responsibilities, Item #9, “On-Site adverse events and
unanticipated outcomes will be reported to the IRB by the Principal
Investigator within seventy-two hours of the Investigator becoming aware that
there has been an on-site adverse event.”

EXPIRATION DATE: 4/20/10: A REQUEST FOR RENEWAL MUST BE
SUBMITTED TO THE IRB OFFICE AT LEAST 30 DAYS PRIOR TO THE

OLD SHORT HILLS ROAD ■ LIVINGSTON, NEW JERSEY 07039 ■ (973) 322-5000

Saint Barnabas Medical Center is a major teaching affiliate of UMDNJ-New Jersey Medical School
ABOVE EXPIRATION DATE. AN ANNUAL REPORT FORM IS ENCLOSED.

Sincerely,

Robert L. Goodman, M.D.
Chairman, IRB
October 8, 2009

Jacques Cohen Ph.D.
Saint Barnabas Medical Center
Institute For Reproductive Medicine & Science
101 Old Short Hills Road, Suite 501
West Orange NJ 07092

Dear Dr. Cohen:

This letter is to acknowledge the receipt of the information identified below.

Our Study # 02-33

Protocol Title: STEM CELL RESEARCH USING DISCARDED NON-VIABLE EMBRYOS

Expiration Date: 4/20/2010
Our Internal #: 3445
Type of Change: (Other) Pre Meeting Action: Acknowledged
Expedited ?: No
Date of Change: 10/8/2009
Date Received: 10/8/2009
On Meeting Date: 10/20/2009
Description:

Receipt of notification (letter dated September 15, 2009) that the investigator will re-consent patients with revised informed consent form dated 3/13/09. The revised consent form was reviewed and granted expedited approval on April 9, 2009.

Respectfully yours,

Chairman, IRB
From: Mina Alikani [mailto:mina.alikani@embryos.net]
Sent: Friday, January 28, 2011 12:46 PM
To: Gadbois, Ellen (NIH/OD) [E]
Cc: Santiago Munne
Subject: Additional cell lines submitted

Dear Dr. Gadbois,

I have just completed submission of five additional cell lines on behalf of Reprogenetics for administrative review. I would like to add the following:

First, cell lines RNJ11 and RNJ12 were derived from donations by IRMS patients (like RNJ7) but these donor patients agreed with and signed a new consent form that we believe fulfills the requirements of section IIA. I submitted the redacted signed reconsent forms but only included the signature page. I am attaching the full consent here in case that is a problem.

Secondly, cell lines RNJ18-20 are from a donation made by a patient undergoing treatment in a collaborating IVF center, Northwest Center for Reproductive Sciences, in Kirkland, Washington. The collaboration was acknowledged and approved through WIRB and all the supporting paperwork was submitted. This submission should also fulfill section IIA requirements.

As before, if there are any questions or concerns regarding these submissions, please do not hesitate to contact me.

Best wishes,

Mina Alikani, Ph.D.
I agree with this recommendation.

Sent from my iPad

On Feb 4, 2011, at 1:46 PM, "Gadbois, Ellen (NIH/OD) [E]" <gadboisel@od.nih.gov> wrote:

Dr. Landis,

Two new submissions from Reprogenetics were reviewed yesterday by the NIH administrative review group: 2011-ADM-001 and 2011-ADM-002. The administrative review group determined that there are two provisions of the Section IIA criteria that are not met in either submission:

Element 8: Donor(s) should have been informed that they retained the right to withdraw consent for the donation of the embryo until the embryos were actually used to derive embryonic stem cells or until information which could link the identity of the donor(s) with the embryo was no longer retained, if applicable.

Element 15: Whether information that could identify the donor(s) would be available to researchers.

Therefore the NIH administrative review group recommends referring these submissions to the ACD Working Group for consideration under Section IIB of the Guidelines. (The embryos were donated prior to the effective date of the Guidelines.) In prior administrative review of an earlier submission from Reprogenetics (2010-ACD-006), we found the same elements to be lacking and moved that submission to the ACD Working Group for review.

Please let us know if you concur with this recommendation or need further information.

Ellen
Dear Dr. Hannemann,

Thank you for your e-mail. In response to your question, RN7, 11, and 12 were not derived from PGD embryos and as far as we know, they do not contain genetic mutations. - though they have not been specifically tested for such abnormalities.

For further clarification, I have enclosed here a table that lists our cell lines, their origin, and the extent of characterization done for each. I hope this will be helpful.

The wording in the highlighted question 8 of the application to which you refer is actually puzzling to me — I have the distinct impression that the word “normal” should have actually been “abnormal.” At the time this revision was made, I was not involved directly in this project. The project really started back in 2002 with a series of hypothetical aims and goals and evolved to what it is today with more specific goals — each stage of this evolution having been managed by different team members. I have offered this previously to Dr. Gadbois as a way of explaining some confusions and uncertainties. At the same time, I have and will continue to emphasize that our work has never crossed any ethical boundaries and we have and continue to adhere to basic principles of ethical research conduct.

Best wishes,

Mina Allikani, Ph.D.

On 3/15/11 10:50 AM, "HESCREGISTRY [NH/OD]" <hescregistry@mail.nih.gov> wrote:

Dear Dr. Allikani,

Thank you for the additional information. Looking at the 2004 application (attached), we note that item 8 states, “... we want to expand the study to develop stem cell from chromosomally normal embryos detected from Pre-implantation Genetic Diagnosis cycles.”

Please let us know whether any the cell lines in review (RN7, RN11, or RN12) was derived from embryos that were tested by PGD and if any of the lines contain genetic mutations.

We will also be sending 3 follow-up emails to confirm documentation on each of the cell lines. Please respond to each email separately to help us with our records. Again, thank you for your on-going efforts to answer our questions.

Sincerely,
Diane Hannemann

Diane E. Hannemann, Ph.D.
Office of Science Policy Analysis
National Institutes of Health
tel: 301-594-2567
fax: 301-402-0280

From: Mina Allikani [mailto:mina.allikani@embryos.net]
Sent: Tuesday, February 22, 2011 11:27 AM
To: HESCREGISTRY [NH/OD]
Subject: Re: New hESC Registry Application Request #2011-ADM-004 FURTHER CLARIFICATION
Importance: High

Dear Dr. Gadbois,

I have now looked into the question regarding the IRB letter of 2 November, 2004 further and obtained more documents that clarify the situation a little more. I have concluded that while a revision to the consent and the protocol went through the IRB on 2 November, 2004, as you will note in the attached document, the IRMS clinical staff who normally oversees the consenting process did not put this latest version of the consent into use in 2004. Instead, they continued to use, by error, an earlier version of the consent — i.e., the version that was signed by the donors of RN7, 11, and 12.

Again, I do apologize for the confusion this has caused. There are many entities and individuals involved here, including the IRMS clinical staff, IRB staff, and one staff person at Reprogenetics who overlooks the submission of protocols to the IRB, renewals, reports, etc. This arrangement is evidently error-prone. I think the fact that the 2004 version of the consent did not go into circulation at IRMS
Table 1. Summary of Characteristics of new hES Cell Lines Derived by Reprogenetics.

<table>
<thead>
<tr>
<th>Cell line Code</th>
<th>RNJ7</th>
<th>RNJ8</th>
<th>RNJ9</th>
<th>RNJ10</th>
<th>RNJ11</th>
<th>RNJ12</th>
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<tbody>
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<td><strong>Embryo Source</strong></td>
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<td>IVF Clinic</td>
<td>IVF Clinic</td>
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<td>IVF Clinic</td>
<td>IVF Clinic</td>
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</tr>
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<td>Discarded</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Cell Line Characteristics</strong></td>
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<td></td>
<td></td>
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<td>25</td>
<td>24</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Human Feeder Cells</td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
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<td>Normal (46,XX)</td>
<td>Normal (46,XY)</td>
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<td>FISH (XX)</td>
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<td></td>
</tr>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
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<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>+</td>
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<td>+</td>
<td>+</td>
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Table 2. Summary of Characteristics of New hES Cell Lines Derived by Reprogenetics.

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<thead>
<tr>
<th>Cell Line Code</th>
<th>RNJ13</th>
<th>RNJ14</th>
<th>RNJ15</th>
<th>RNJ16</th>
<th>RNJ17</th>
<th>RNJ18</th>
</tr>
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<td>Embryo Source</td>
<td>IVF Clinic</td>
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<td>IVF Clinic</td>
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<td>Fresh Discarded</td>
<td>Fresh Discarded</td>
<td>Fresh Discarded</td>
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<tr>
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<td>N/A</td>
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<td>Trisomy 22 (XY)</td>
<td>Trisomy 15, 22 (XX)</td>
<td>Complex Single Gene Abnormal Defect (XXX)</td>
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</table>

<table>
<thead>
<tr>
<th>Cell Line Characteristics</th>
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<th></th>
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<td>Human Feeder Cells</td>
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<td>Yes</td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
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<td>FISH Normal (XY)</td>
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<td>FISH SGD</td>
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<td></td>
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</tbody>
</table>
------ Forwarded Message

From: Mina Alikani <mina.alikani@embryos.net>
Date: Wed, 13 Apr 2011 17:49:33 +0100
To: "HESCREGISTRY (NIH/OD)" <hescregistry@mail.nih.gov>
Conversation: RNJ11 - hESC Registry Application #2011-ACD-002
Subject: Re: RNJ11 - hESC Registry Application #2011-ACD-002

Dear Dr. Hannemann:

The answer to your first question is YES: the consent version signed by the patients was dated 3/27/2008. The re-consent version signed by the donors is dated 10/2009. That date must have been placed on the form by the IRMS staff to indicate that it was the latest version.

The answer to your second question is NO for the reasons explained previously for RNJ7: The documentation attached to the IRB letter dated 2 November 2004 shows that a paragraph was added to question number 8 of the application/protocol. The paragraph is highlighted in the document. The IRB had approved the protocol revisions submitted on that date (and in August of 2004) but the consent form associated with those revisions was never put into circulation by IRMS.

I hope this clarifies the issues of concern. However, if you wish to discuss this matter over the telephone, please let me know if this Friday, 15 April, or the following Monday would be a possibility.

Sincerely,

Mina Alikani, Ph.D.

On 3/25/11 1:21 PM, "HESCREGISTRY (NIH/OD)" <hescregistry@mail.nih.gov> wrote:

Dear Dr. Alikani,

In follow-up to my earlier email, please confirm the following information for hESC line RNJ11:

**RNJ11**
- Embryo Donation Consent, version dated 3/27/2008:
- Regarding the Embryo Donation Re-Consent for RNJ11, NIH has received a version from you dated 10/2009, while the IRB approved use of the re-consent version dated 3/13/2009. We are not asking if there is a text difference between the two versions. Instead, please confirm which embryo donation re-consent version was signed by the donors for the donation of embryo(s) from which RNJ11 was derived.

- Protocol version used at the time of embryo donation was the version dated 6/30/2002:
  - Yes
  - No
  - If no, please clarify which version was actually used at the time of donation.

From the information NIH has received from you, we understand that the protocol dated 6/23/2002 was modified and approved by the IRB in 2004, but that this modified version of the protocol was not implemented by the clinic when the embryos were donated for the derivation of RNJ11. As such, please confirm above if the protocol actually used in relation to RNJ11 was the version dated 6/30/2002.

Sincerely,
Diane Hannemann

Diane E. Hannemann, Ph.D.
Office of Science Policy Analysis
National Institutes of Health
tel: 301-594-2567
fax: 301-402-0280

From: Mina Alikani [mailto:mina.alikani@embryos.net]
Sent: Friday, March 18, 2011 12:26 PM
To: HESCREGISTRY (NIH/OD)
Subject: Re: RNJ11 - hESC Registry Application #2011-ACD-002
Dear Dr. Hannemann,

Please see my responses in red below.

Best wishes,

Mina Alikani, Ph.D.

On 3/15/11 11:00 AM, "HESCREGISTRY (NIH/OD)" <hescregistry@mail.nih.gov> wrote:

Dear Dr. Alikani,

We’re writing to confirm documentation for RNJ11.

1) CONSENT – Please confirm that for cell line RNJ11, the embryo donation consent form version 3/27/2008 was used (attached).

I am not sure I understand this question, but as far as I can see, the date on the initial consent document is noted as 3/27/2008.

2) RE-CONSENT – We note that on October 8, 2009, the IRB stated, “that the investigator will re-consent patients with revised informed consent form dated 3/13/2009.” However, we understand that the re-consent form version 10/2009 (attached) was used to re-consent for embryos used to derive cell line RNJ11. Please confirm that the 10/2009 version was used.

As far as I can see the two reconsent documents dated 10/2009 and 3/13/2009 are the same. If there is a difference between the two, perhaps that can be pointed out to me so I can investigate.

Please let me also emphasize that the consent process is entirely handled by the IRMS staff. We do submit annual reports to the IRB and make revisions as necessary but we do not have control over what is handed to the patient at the time of consenting.

3) PROTOCOL:

a. For RNJ11, please confirm which protocol version was followed.

The original protocol was approved in 2002; it was modified and reapproved in 2004, but we discovered much later that the consent associated with the revised protocol was not circulated at the time (hence the discrepancy in RNJ 7 documents). The original protocol did not explicitly state the standard method of hESC isolation, although this was the only means by which the cell lines were derived. For that reason, we have amended the protocol during the 2010 annual review through an
amendment that states all the cell lines were derived through either whole blastocyst
culture or isolation and culture of the ICM of the blastocyst.

b. For RNJ11, please confirm that the protocol used was IRB-approved.

Please see 'a' above.

Sincerely,
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