RESEARCH PARTICIPANT INFORMATION AND CONSENT FORM
Healthy Control Subjects

TITLE: 5-HT2AR: 5HT2CR Balance in Brain Connectivity in Cocaine Dependence

VCU IRB PROTOCOL NUMBER: HM15289

INVESTIGATOR: Dr. F. Gerard Moeller

Study Sponsor: National Institute on Drug Abuse (NIDA)

ABOUT THIS CONSENT FORM
You are invited to take part in a project called “5-HT2AR: 5HT2CR Balance in Brain Connectivity in Cocaine Dependence” conducted by Dr. F. Gerard Moeller, of Virginia Commonwealth University (VCU) School of Medicine. It is important that you carefully think about whether being in this study is right for you and your situation.

This consent form is meant to assist you in thinking about whether or not you want to be in this study. Please ask the study doctor or the study staff to explain any information in this consent document that is not clear to you. You may take home an unsigned copy of this consent form to think about or discuss with family or friends before making your decision.

Your participation is voluntary. You may decide not to participate in this study. If you do participate, you may withdraw from the study at any time. Your decision not to take part or to withdraw will involve no penalty or loss of benefits to which you are otherwise entitled.

AN OVERVIEW OF THE STUDY AND KEY INFORMATION
The main purpose of this study is to find out how the brain reacts to the medication called mirtazapine and how this is related to cocaine addiction. Mirtazapine is a medicine that is approved by the FDA for the treatment of depression. More information about how the brain reacts to mirtazapine may be helpful to develop new treatments for cocaine addiction. We are inviting you to take part in this research study as a person who does not use cocaine to compare your reactions on computer tasks and brain scans with people who have been using cocaine. Measures of your memory and reaction time while you are in a magnetic resonance imaging (MRI) scanner will be tested to find out whether there are differences in brain scans between people who have been using drugs and people who do not use drugs.

This is a local study. The study will enroll a total of 400 people all at VCU.

What will happen if I participate?
You will also receive one of the following study drugs to take by mouth before you go inside the scanner: Mirtazapine (15 mg) (also called Remeron); or Placebo (inactive pill). The drug you receive first will be assigned randomly, that means the decision is made “by chance”, like the flip of a coin. Using the active study drug and placebo will test whether the medication reacts differently with the brain. You will also be asked to provide a urine sample for complete drug screen analysis, and a breath sample to screen for alcohol. If you are a woman, you will be asked to
provide a urine sample for a pregnancy test. After you take the study medication, you will be monitored for up to 8 hours before being released. If there are any technical problems with the MRI scan during the baseline period, then you may be asked to provide an additional MRI session just like the first one.

You will begin with a baseline period that will last 1 week, you will have a practice MRI session. After the practice MRI session is completed, you will have a real MRI session. One week after your first MRI session, you will have another MRI scanning session just like the first time.

**TIME COMMITMENT**
The study elements and time are:
1. Baseline period (mock MRI scan, real MRI scan, computer tasks,) includes 3 visits per week for 1 week (about 12 hours total time for the baseline period)
2. Follow up MRI scan (real MRI scan, plus preparation time outside the scanner and testing outside the scanner, for total time of about 8 hours)
The total maximum number of hours is about 20 hours.

**What alternative treatments or procedures are available?**
This study is not intended to treat any medical condition. The alternative to taking part in this study is to not take part.

**What are the risks and benefits of participating?**
There are both risks and benefits of participating in research studies. We want you to know a few key risks right now. We will give you more information in the “WHAT RISKS AND DISCOMFORTS CAN I EXPECT FROM BEING IN THE STUDY?” section.

**Risks and Discomforts**

1. The possible psychological effects are that mirtazapine, similar to other antidepressant medicines, can increase thoughts of death and suicide in people who are depressed.

2. Risks associated with the MRI scanner include injury to your body if you have metal in your body, such as a pacemaker, metal shavings, bullet pieces, or surgical clips; feeling anxious or nervous due to being confined in an enclosed space, such as the MRI scanner tube, feeling tired or bored during the MRI testing.

3. There may be some risks that the study doctors do not know about yet, so we will let you know of any new findings.

4. Participation in research might involve some loss of privacy. There is a small risk that

**Benefits to you and others**
This is not a treatment study, and you are not expected to receive any direct medical benefits from your participation in the study. The information regarding your physical status learned during screening procedures. It is also possible that information will be learned about how Mirtazapine affects the brain, which may help develop new treatments for drug abuse in the future. In general, we will not give you any individual results from the study.
someone outside of the research study could see and misuse information about you

**WHAT WILL HAPPEN IF I PARTICIPATE IN THE STUDY?**

You have already completed the screening procedure as per the protocol (HM 20000294, PRE-SCREEN, Keyser-Marcus (PI)) and have qualified to participate in this study. In this study, you will be asked about your possible drug or alcohol use and will also undergo some computer tests that measure attention and reaction time. In addition you will be required to undergo two MRI scans.

If you have ever had any pieces of metal enter your eye, then MRI may be potentially dangerous, and you should tell the study staff about this right away. Also, if you have a pacemaker or other metal such as bullet pieces in your body you need to tell the study staff as these may be dangerous as well. You will also be asked to fill out a standard metal screening check list to make sure that you do not have any metal in your body or conditions that might be dangerous for MRI. If you drink caffeinated beverages we ask that you abstain from these beverages prior to the scan.

You will begin with a baseline period that will last 1 week. During one of your baseline visits, you will have a practice MRI session that will take place in a mock MRI (which is not a real MRI machine) to allow you to practice the same tasks that you will do during the real MRI scan. The total time of the practice MRI session will be around 2 hours. After the practice MRI session is completed, you will have a real MRI session. Each real MRI scanning session consists of two 40-minute periods in which you will be inside the scanner. These periods are separated by a 10-minute bathroom break. While you are in the MRI scanner you will be asked to look at a screen and press some buttons in response to images on the screen, and you will be asked to perform tasks that measure decision-making, impulsivity, and attention. You will be paid $5 per task. As an incentive to do the tasks as accurately as possible, you may earn an additional sum of money that will be a reward based on your accuracy on each task (up to an additional sum of about $5 more per task). During the practice and the real scanning sessions, we will monitor your blood pressure with an automatic blood pressure cuff that is placed on your arm and your pulse with a small clip that is placed on your finger. The total time for the real MRI session, including preparation time outside the scanner, time in the scanner, and testing outside the scanner will take about 8 hours to complete.

For the real MRI session, you will arrive at our building at 7:30 a.m. to have a repeat ECG (heart test) to make sure that there are no medically dangerous changes in your ECG results. You will fill out several forms that ask questions about any problems you may be having. You will also receive one of the following study drugs to take by mouth before you go inside the scanner: Mirtazapine (15 mg) (also called Remeron); or Placebo (inactive pill). The drug you receive first will be assigned randomly, that means the decision is made “by chance”, like the flip of a coin. Using the active study drug and placebo will test whether the medication reacts differently with the brain. You will also be asked to provide a urine sample for complete drug screen analysis, and a breath sample to screen for alcohol. If you are a woman, you will be asked to provide a urine sample for a pregnancy test. After you take the study medication, you will be monitored for up to 8 hours before being released. If there are any technical problems with the
MRI scan during the baseline period, then you may be asked to provide an additional MRI session just like the first one.

One week after your first MRI session, you will have another MRI scanning session just like the first time. If there are any technical problems with this MRI scan, then you may be asked to provide an additional MRI session just like the previous one.

There will be times during the study when the research staff will need to contact you by telephone (for example, to reschedule an appointment).

**WHAT RISKS AND DISCOMFORTS COULD I EXPERIENCE FROM BEING IN THE STUDY?**

There are two kinds of risks; one is physical and the other is psychological (mental). You may have some pain and a bruise when the blood is drawn. There is also a slight chance of infection. Mirtazapine possible side effects include nausea, dry mouth, constipation, dizziness and feeling drowsy. The drowsiness associated with mirtazapine may impair a person’s ability to drive, use machines, or perform hazardous tasks that require alertness. The drowsiness can be worse if you combine this medication with alcohol.

If you have metal in your body, such as a pacemaker, metal shavings, bullet pieces, or surgical clips, the MRI scan can cause injury to you. You will be carefully screened to eliminate this risk by a history, physical examination, and completing a standard MRI screening form.

Possible side effects will be monitored carefully. This will be done through a form that you fill out when you attend study visits.

The possible psychological effects are that mirtazapine, similar to other antidepressant medicines, can increase thoughts of death and suicide in people who are depressed. We will ask you about these thoughts during the study and you should tell us if you have these thoughts. You may experience the potential risk of feeling anxious or nervous due to being confined in a small, enclosed space such as the MRI scanner tube. You may become tired or bored during the memory testing.

**Special note to women:** Additional risks to an unborn child may exist with study drugs and with cocaine use. Therefore, pregnancy testing will be done every week during the study. In addition, pregnancy testing will be done before each MRI scan because of possible unknown risks to the fetus from MRI scanning. Women of childbearing potential must not be pregnant or breast feeding, and if sexually active, must be using acceptable methods of birth control to be enrolled in the study. Acceptable methods include:

a) Oral contraceptives

b) Barrier plus spermicide

c) Hormonal or surgical implants

Risks associated with having an ECG include: discomfort in learning of a potential health condition and the potential for hair removal where the ECG lead is applied to the chest. Some individuals who have allergies to tape adhesive may have a mild allergic reaction to the ECG electrode adhesive.

You will be asked to leave the study if you do not complete the requirements and fulfill your responsibilities to attend, give urine or breath samples, or complete forms. It may also be necessary for you to leave this study if you have other problems or illness during the study. If this happens, appropriate referrals will be given.
HOW WILL MY HEALTH INFORMATION BE USED AND SHARED DURING THE STUDY?

As part of this research study, we will ask you to share identifiable health information with us and/or permit us to access existing information from your healthcare records. New health information may also be created from study-related tests, procedures, visits, and/or questionnaires. This type of information is considered “Protected Health Information” that is protected by federal law.

Who will use or share protected health information about me?
VCU and VCU Health are required by law to protect your identifiable health information. By consenting to this study, you authorize VCU/VCU Health to use and/or share your health information for this research. The health information listed above may be used by and/or shared with the following people and groups to conduct, monitor, and oversee the research.

- Principal Investigator and Research Staff
- Research Collaborators
- Data Safety Monitoring Boards
- Health Care Providers at VCUHS
- Study Sponsor
- Institutional Review Boards
- Government/Health Agencies
- Others as Required by Law

Once your health information has been disclosed to anyone outside of this study, the information may no longer be protected under this authorization.

What type of health information will be used or shared with others during this research?
The following types of information may be used for the conduct of this research: Complete health record

When will this authorization (permission to use) my protected health information expire?
This authorization will expire when the research study is closed, or there is no need to review, analyze, and consider data generated by the research project, whichever is later.

Statement of Privacy Rights
You may change your mind and revoke (take back) the right to use your protected health information at any time. Even if you revoke this Authorization, the researchers may still use or disclose health information they have already collected about you for this study. If you revoke this Authorization you may no longer be allowed to participate in the research study. To revoke this Authorization, you must write to the Principal Investigator: Dr F Gerard Moeller, 203 East Cary Street, Suite 202, Richmond VA 23219.

WHAT ARE THE COSTS?
All medical tests, brain scans, and study drug will be offered at no cost to you. If you receive a bill that you believe is related to your taking part in this research study, please contact Dr. F. Gerard Moeller at (804) 828-3793, with questions.
WILL I BE PAID TO PARTICIPATE IN THE STUDY?
You will be compensated $20 for the practice scan, additional $5 for each task completed (up to $15) and earn bonus money based on task performance (up to $15). You will receive $75 for each scanning session, an additional $5 for each task completed (up to $15) and earn bonus money based on task performance while inside the scanner (up to $15). You will receive $30 for each behavioral task session and earn bonus money based on task performance (up to $10). In addition a study completion bonus of $75 will be provided. Participants traveling more than 20 miles to the study site will be reimbursed for their travel expenses (mileage only at the current VCU rate). Compensation will be provided in the form of vouchers, which can be exchanged for cash or check (to be mailed to you), based on your preference.

If you complete the entire two-week study as per specified procedures, you will receive a maximum amount of $415 dollars. Total payments within one calendar year that exceed $600 will require the University to annually report these payments to the IRS and you. This may require you to claim the money you receive for participation in this study as taxable income. VCU is required by federal law to collect your social security number. Your social security number will be kept confidential and will only be used to process payment.

If you receive payment for taking part in this study please be informed that you will be asked to complete a W-9 form that will be forwarded to the accounting department as a requirement by the Internal Revenue Service. You will also be issued a 1099-Misc form from this study for tax reporting purposes.

WHAT HAPPENS IF I AM INJURED OR BECOME SICK BECAUSE I TOOK PART IN THE STUDY?
If you are injured by, or become ill, from participating in this study, please contact your study doctor immediately. Medical treatment is available at the Virginia Commonwealth University Health System (VCU Health System). Your study doctor will arrange for short-term emergency care at the VCU Health System or for a referral if it is needed.

Fees for such treatment may be billed to you or to appropriate third party insurance. Your health insurance company may or may not pay for treatment of injuries or illness as a result of your participation in this study. To help avoid research-related injury or illness, it is very important to follow all study directions.

CAN I STOP BEING IN THE STUDY?
You can stop being in this research study at any time. Leaving the study will not affect your medical care, employment status, or academic standing at VCU or VCU Health. Tell the study staff if you are thinking about stopping or decide to stop.

You may refuse to answer any questions asked or written on any forms, however answering some questions about your health and safety are necessary to take part in this study.

Your participation in this study may be stopped at any time by the study doctor or the sponsor without your consent. The reasons might include:
- the study doctor thinks it necessary for your health or safety;
• you have not followed study instructions;
• the sponsor has stopped the study; or
• administrative reasons require your withdrawal.

If you withdraw from the study, data that has already been collected about you will remain part of the study database and may not be removed.

**HOW WILL INFORMATION ABOUT ME BE PROTECTED?**

VCU and the VCU Health System have established secure research databases and computer systems to store information and to help with monitoring and oversight of research. Your information may be kept in these databases but are only accessible to individuals working on this study or authorized individuals who have access for specific research related tasks.

Identifiable information in these databases are not released outside VCU unless stated in this consent or required by law. Although results of this research may be presented at meetings or in publications, identifiable personal information about participants will not be disclosed.

Personal information about you might be shared with or copied by authorized representatives from the following organizations for the purposes of managing, monitoring and overseeing this study:

- The study Sponsor, representatives of the sponsor and other collaborating organizations
- Representatives of VCU and the VCU Health System
- Officials of the Department of Health and Human or the Federal Food and Drug Administration
- It will be noted in your protected electronic health record at VCU Health that you are in this study. Information about the study will be included in the record. This information is protected just as any of your other health records are protected.

In general, we will not give you any individual results from the study. If we find something of medical importance to you, we will inform you, although we expect that this will be a very rare occurrence.

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 Website will include a summary of the results. You can search this Web site at any time.

If you tell us that you may hurt yourself or someone else, the law says that we must let people in authority know.

There are no plans to share any money or profits with you if the use of your sample(s) results in inventions or discoveries that have commercial value. In the
future, identifiers might be removed from the information and samples you provide in this study, and after that removal, the information/samples could be used for other research studies by this study team or another researcher without asking you for additional consent.

**Certificate of Confidentiality**

To help us protect your privacy, we have obtained a Certificate of Confidentiality from the National Institutes of Health. A Certificate of Confidentiality helps the researchers keep your information private. For example, researchers can refuse to give out your information in a court case. Researchers may have to give your information if the study is audited, or if the information is required by the Food and Drug Administration (FDA).

However, if we learn about abuse of a child or elderly person or that you intend to harm yourself or someone else, or about certain communicable diseases, we will report that to the proper authorities. The researchers will use the Certificate to resist any demands for information that would identify you, except as explained. The Certificate cannot be used to resist a demand for information from employees of the United States Government that is used for auditing or evaluation of federally funded projects or for information that must be given in order to meet the requirements of the federal Food and Drug Administration (FDA). You should understand that a Certificate of Confidentiality does not prevent you or a member of your family from voluntarily giving information about yourself or your involvement in this research. If an insurer, employer, or other person obtains your written consent to receive research information, then the researchers may not use the certificate to withhold the information.

No one outside the research staff may call and get information about you. If you decide you would like copies of your medical test results, you may sign a form to release information to you or your physician. Only specific medical information, not research information, will be released.

You will not be personally identified in any reports or publications that may result from this study. Any personal information about you that is gathered during this study will remain confidential to every extent of the law. A special number will be used to identify you in the study and only the investigator and research staff will know your name.

You should know that research data or (medical information if applicable) about you may be reviewed or copied by Virginia Commonwealth University.

Although results of this research may be presented at meetings or in publications, identifiable personal information pertaining to participants will not be disclosed.

**GENETIC TESTING**

**Background information:** During this study, if you have not already provided an additional blood sample for genetic testing (as part of the intake screening conducted for HM 20000294, PRE-SCREEN, Keyser-Marcus (PI)), we will draw additional blood (about 2 tablespoons) to test your DNA. This sample will be used to do extra studies to learn more about the biology of drug or alcohol use. DNA is genetic material that we inherit from our parents. This material is made up of elements called genes that may affect our behavior and biology. DNA can be obtained from blood to study the relationship between specific genes and certain medical conditions, including mood disorders and drug dependence.

The research involves genetic testing. Genetic testing may reveal information about the likelihood that a person or his or her relatives may
develop certain diseases. Genetic testing may reveal information about who is related to whom. If known to employers or insurance companies, the results of genetic testing might affect a person's ability to obtain a job or health or life insurance.

Future studies:

1. My blood/tissue samples may be stored and used for future research about drug or alcohol use.

   YES ___________________________ NO ___________________________
   initial initial

2. My blood/tissue samples may be stored and used for future research about other health problems (for example, heart disease, osteoporosis, depression, diabetes, etc.)

   YES ___________________________ NO ___________________________
   initial initial

**Future contact concerning genetic testing results:** Genetic testing in this study may help us understand the way the brain works that can lead to drug or alcohol use. There is no plan to test for genes that are known to cause cancer or other illnesses, so you will not be given the results of the genetic testing.

**Withdrawal of genetic testing consent:** In the future if you decide that you do not want us to use your DNA for genetic research you can tell us this and we will remove your DNA from our research samples for future studies.

**Confidentiality:** All genetic testing in this study is being done for research purposes only and is confidential. We have a certificate of confidentiality from the National Institute on Drug Abuse to help keep all research information in this study confidential.

**WHOM SHOULD I CONTACT IF I HAVE QUESTIONS ABOUT THE STUDY?**

The investigators of this study will be glad to answer any further questions at any time. In the event of problems due to this study you should report them to Dr. Moeller at (804) 828-3793 or 804-828-3686. If you would like to withdraw your permission to use blood or tissue samples, mail address for sending a letter notifying us to 203 East Cary Street, Suite 202, Richmond, VA 23219.

If you have general questions about your rights as a participant in this or any other research, you may contact:

Office of Research
Virginia Commonwealth University
800 East Leigh Street, Suite 3000
P.O. Box 980568
Richmond, VA 23298
Telephone: (804) 827-2157

Contact this number for general questions, concerns, or complaints about research. You may also call this number if you cannot reach the research team or if you wish to talk to someone else. General information about participation in research studies can also be found at http://www.research.vcu.edu/irb/volunteers.htm.

Do not sign this consent form unless you have had a chance to ask questions and have received satisfactory answers to all of your questions.

CONSENT
I have been provided with an opportunity to read this consent form carefully. All of the questions that I wish to raise concerning this study have been answered.

By signing this consent form, I have not waived any of the legal rights or benefits, to which I otherwise would be entitled. My signature indicates that I freely consent to participate in this research study. I will receive a copy of the consent form once I have agreed to participate.

Participant Name, (Printed)

Participant Signature ___________________________ Date __________

Name of Person Conducting Informed Consent Discussion / Witness  (Printed)

Signature of Person Conducting Informed Consent Discussion / Witness ___________________________ Date __________

Principal Investigator Signature (if different from above) ___________________________ Date __________