Department of Veterans Affairs	Authorization for Use & Release of Individually Identifiable Health Information for Veterans Health Administration (VHA) Research			
Subject Name (Last, First, Middle Initial):		Subject SSN (last 4 only):	Date of Birth:	
VA Facility (Name and Address):				
VA Principal Investigator (PI):		PI Contact Information:		
Study Title:				
Purpose of Study:				
USE OF YOUR INDIVIDUALLY IDENTIFIABLE HEALTH INFORMATION (IIHI): Your individually identifiable health information is information about you that contains your health information and information that would identify you such as your name, date of birth, or other individual identifiers. VHA is asking you to allow the VA Principal Investigator (PI) and /or the VA research team members to access and use your past or present health information in addition to new health information they may collect for the study named above. The investigators of this study are committed to protecting your privacy and the confidentiality of information related to your health care.				
Signing this authorization is completely voluntary. However, your authorization (permission) is necessary to participate in this study. Your treatment, payment, enrollment, or eligibility for VA benefits will not be affected, whether or not you sign this authorization. Your individually identifiable health information used for this VA study includes the information marked below: Information from your VA Health Records such as diagnoses, progress notes, medications, lab or radiology findings, etc.				
 □ Specific information concerning: □ alcohol abuse □ drug at a drug a	e, age, race, etc. tapes of you Diary	anemia □ HIV		

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Subject Name (Last, First, Middle Initial):	Subject SSN (last 4 only):	Date of Birth:		
USE OF YOUR DATA OR SPECIMENS FOR OTHER RESEARCH banking is a required component of this study. When banking is an of this form in lieu of this section.)	`	•		
☐ Not Applicable - No Data or Specimen Banking for Other Res	search			
An important part of this research is to save your				
☐ Data				
☐ Specimen				
in a secure repository/bank for other research studies in the future. and/or specimen for future studies approved by the required commit will not be able to participate in this study.	•	•		
DISCLOSURE: The VA research team may need to disclose the infinstitutions that are not part of VA. VA/VHA complies with the require Accountability Act of 1996 (HIPAA), Privacy Act of 1974 and all other protect your privacy. The VHA Notice of Privacy Practices (a separative protect your information. If you do not have a copy of the Notice, Giving your permission by signing this authorization allows us to discover your permission by signing this authorization allows us to discover your permission by signing this authorization allows us to discover your permission by federal laws and regulations and might be retained information. These non-VA/VHA institutions or persons included.	ements of the Health Insurance rapplicable federal laws and of the document) provides more in the research team will provide close your information to other in has been disclosed outside disclosed by the persons or in	re Portability and regulations that information on how e one to you. Institutions or VA/VHA, it may no		
☐ Non-VA Institutional Review Board (IRB) at who will monitor the study				
Study Sponsor (name):				
Person or entity who takes responsibility for and initiates a clinical	al investigation			
Academic Affiliate (institution/name/employee/department): A relationship with VA in the performance of this study				
Compliance and Safety Monitors:	-A d			
Advises the Sponsor or PI regarding the continuing safety of this	-			
Other Federal agencies required to monitor or oversee research	(such as FDA, OHRP, GAO):			
☐ A Non-Profit Corporation (name and specific purpose):				
☐ Other (e.g. name of contractor and specific purpose):				

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Note: Offices within VA/VHA that are responsible for oversight of VA research such as the Office of Research Oversight (ORO), the Office of Research and Development (ORD), the VA Office of Inspector General, the VA Office of General Counsel, the VA IRB and Research and Development Committee may also have access to your information in the performance of their VA/VHA job duties.				
Access to your Individually Identifiable Health Information create While this study is being conducted, you	ed or obtained in the cours	e of this research:		
☐ will have access to your research related health records				
☐ will not have access to your research related health records				
This will not affect your VA healthcare including your doctor's ability to see your records as part of your normal care and will not affect your right to have access to the research records after the study is completed.				
REVOCATION: If you sign this authorization you may change your n any time. You must do this in writing and must send your written require following address:				
If you revoke (take back) your permission, you will no longer be able to participate in this study but the benefits to which you are entitled will NOT be affected. If you revoke (take back) your permission, the research team may continue to use or disclose the information that it has already collected before you revoked (took back) your permission which the research team has relied upon for the research. Your written revocation is effective as soon as it is received by the study's Principal Investigator.				
EXPIRATION: Unless you revoke (take back) your permission, your your information will:	authorization to allow us to us	se and/or disclose		
☐ Expire at the end of this research study				
☐ Expire on the following date or event:				
☐ Not expire				
Expires at the end of this research study unless you have: (1) provand/or biological specimens in a research data repository or (2)wh has been completed				

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Subject Name (Last, First, Middle Initial):	Subject SSN (last 4 only):	Date of Birth:		
TO BE FILLED OUT BY THE S	SUBJECT			
Research Subject Signature. This permission (authorization) has be opportunity to ask questions. If I believe that my privacy rights have facility Privacy Officer to file a verbal or written complaint.				
I give my authorization (permission) for the use and disclosure of my described in this form. I will be given a signed copy of this form for m		th information as		
Signature of Research Subject	Date			
Signature of Legal Representative (if applicable)	 Date			
To Sign for Research Subject (Attach authority to sign: Health Care Power of Attorney, Legal Guardian appointment, or Next of Kin if authorized by State Law)				
Name of Legal Representative (please print)	Date			