



National Registry of Myotonic Dystrophy and Facioscapulohumeral Muscular Dystrophy Patients and Family Members

Dear Registry Applicant,

Thank you for your interest in the **National Registry**! The Registry connects people with Myotonic Dystrophy (DM) and FSHD with researcher opportunities. Anyone with DM or FSHD is eligible to join, as well as family members.

Please complete the following enclosed forms to join the Registry:

- 1. Consent Form Please sign and return one copy. The second copy is for you to keep.
- 2. Assent Form Completed if the enrollee is a child between the ages of 13-17 years old.
- 3. Patient Information Form
- 4. Medical Information Form This form gives us permission to request your medical records from your neurologist or primary care physician.

Please return the completed forms to us in the enclosed prepaid envelope. If you have any questions, please contact us at 1-888-925-4302 or at dystrophy registry@URMC.rochester.edu.

Digleto a Luella

Elizabeth Luebbe

We appreciate your support of research for DM and FSHD!

Sincerely,

James Il

James Hilbert, MS
Health Project Coordinator

Health Project Coordinator Health Project Coordinator





CONSENT FORM

Study title: National Registry of Myotonic Dystrophy and Facioscapulohumeral Muscular Dystrophy Patients and Family Members

Principal Investigator: Rabi Tawil, MD

This consent form describes a research study, what you may expect if you decide to take part, and important information to help you make your decision. Please read this form carefully and ask questions about anything that is not clear before you agree to participate.

A person who takes part in a research study is called a research subject, or research participant. In this consent form, "you" generally refers to the research subject. If you are a parent/legal guardian for the potential subject, "you" in the rest of this form generally means your child or the adult who will be the research subject.

Key Information

- Being in this research study is voluntary it is your choice.
- You are being asked to take part in this study because you or a family member has myotonic dystrophy (DM) or facioscapulohumeral muscular dystrophy (FSHD).
- The purpose of the National Registry is to collect information about the symptoms of DM and FSHD and to connect patients with researchers.
- Your participation in this study will last for the next 5-10 years or longer.
- Procedures include completing a questionnaire and providing updates to your information each year. You will also receive information about studies related to DM and FSHD and information on how to participate. You may also receive email and newsletters related to Registry activities.
- There are risks from participating.
 - o The most common risk is that you may feel uncomfortable answering certain questions about your symptoms. You do not have to share any information that you do not want to.
 - One of the most serious risks is a possible loss of confidentiality due to the unauthorized release of medical information. See the "Risks of Participation" section in this consent form for more information. You should discuss these risks in detail with the study team if you have any questions.
- You might not benefit from being in this research study. A potential benefit is receiving information about studies that you may want to join and receiving updates on advances in DM and FSHD research and clinical care.

PURPOSE

The goals of this Registry are to:

- Help researchers collect and study information on how DM and FSHD affect people;
- Help researchers recruit patients with DM and FSHD into clinical studies and trials;
- Share information about opportunities and advances in DM and FSHD research with you, care providers, and researchers.

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DESCRIPTION OF PROCEDURES

The forms for the Registry will take about 20 minutes to read and complete. You can complete the forms by paper or online through Research Electronic Data Capture (REDCap). REDCap is a secure, HIPAA-compliant, web-based application used for electronic capture and management of research and clinical study data. The following is requested to participate in the Registry:

- Complete the "Patient Information Form" questionnaire. This form will ask for your contact information as well as information about your muscle strength, general health, and how your muscular dystrophy affects your daily life. Unaffected family members will complete a shortened version of this form.
- Complete Authorization for Release of Medical Information form. Please provide the complete name, address, and phone number of one or two of your doctors on this form. This form gives us permission to request medical records about your muscular dystrophy and how it was diagnosed. This form permits your physician(s) to send test results such as the results of muscle biopsies, genetic testing, heart tracing (e.g., EKG), electromyography (EMG), as well as records that pertain to your muscular dystrophy. If you are an unaffected family member, we will only request this information if you have received a genetic test or other exams that show that you do not have muscular dystrophy.

If you complete the forms on paper, please mail all completed forms to us in the enclosed, prepaid envelope. If you complete the forms online, you have the option to save and return later. When you click "save," you will receive an individualized Return Code to return and complete your application at a later time, if you choose.

Once we receive your application through the mail or online, we will review your forms and may contact you if additional information is needed. You will receive a notification in the mail or email that all of your forms have been reviewed and that you are enrolled in the Registry.

After joining the Registry

- Once you are enrolled in the Registry, we may contact you through the mail or email about opportunities to participate in research studies. Some studies involve filling out questionnaires at home about your quality of life. Other studies involve collecting blood or tissue samples, testing your muscle strength, or testing new treatments. Each study is voluntary and requires your agreement (consent).
- If you are interested in such studies, you can contact the researcher for more information about the study. The Registry will not provide any information that could identify you to the researcher.
 All research studies are reviewed and approved by the researcher's human subjects institutional review board and by the Scientific Advisory Committee of this Registry.
 - Once a year, we will send you a form through the mail oremail to update your address, phone number, and information about your health and/or any symptoms of your muscular dystrophy. It should take about 15 minutes to review and complete this form. Completion of the form is voluntary.
- We ask that you contact us if there are changes to your home address, phone number, or email address so that we are able to update your contact information.

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- Participation of family members is strongly encouraged. No information about you will be shared
 with members of your family. Each family member is encouraged to enter the Registry and to
 complete the forms themselves, if interested and able.
- Scientists, researchers, and clinicians will be allowed to see and study Registry data that is deidentified or anonymous (information that cannot identify you). Researchers need to submit an
 application to the Registry team to get approval and receive data. They can analyze this deidentified information to study the symptoms in DM and FSHD, learn how symptoms progress over
 time, and other topics to better understand these diseases and to develop new treatments.
- A subset of de-identified information collected from you may be shared with certain other
 databases. We may share de-identified information with other national or international registries that
 collect information on multiple rare disease and registries that are specific to DM or FSHD. We may
 share de-identified information with other databases in order to increase global knowledge of DM
 and FSHD that may lead to new research studies, clinical trials, and clinical treatments. No
 information will be shared that could identify you.

NUMBER OF SUBJECTS

We expect 3,500 subjects or more to participate in this Registry.

BENEFITS OF PARTICIPATION

You might not benefit from being in this Registry. A potential benefit to you from being in the Registry is receiving information about other studies you may want to join. You will receive information about Registry activities and research advances in myotonic dystrophy, FSHD, and related diseases.

Researchers may benefit by using the Registry to study why individuals have different symptoms, learn about how certain treatments work, help medical professionals improve how they manage care for individuals with DM and FSHD, and advance research in DM and FSHD by analyzing de-identified Registry data.

RISKS OF PARTICIPATION

There is minimal risk in taking part in this Registry. Participation includes questions that can be sensitive and that may make you may feel uncomfortable. You do not have to share any information that you do not want to. Another risk of participation is the possible loss of confidentiality due to an unauthorized release of medical information.

SPONSOR SUPPORT

The University of Rochester is receiving payment from the National Institutes of Health (NIH) for conducting this research.

COSTS

There will be no cost to you to participate in this Registry.

PAYMENTS

You will not be paid for participating in this Registry.

CERTIFICATE OF CONFIDENTIALITY

To help us protect your privacy, we have a Certificate of Confidentiality from the National Institutes of Health (NIH). With this Certificate, the investigators cannot be forced (for example, by court subpoena)

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to disclose research information that may identify you in any Federal, State, or local civil, criminal, administrative, legislative, or other proceedings. Disclosure will be necessary, however, upon request of DHHS for audit or program evaluation purposes.

You should understand that a Certificate of Confidentiality does not prevent you or a member of your family from voluntarily releasing 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 investigator may not use the Certificate of Confidentiality to withhold this information. This means that you and your family must also actively protect your own privacy.

Finally, you should understand that the researcher is not prevented from taking steps, including reporting to authorities, to prevent serious harm to yourself or others.

<u>Confidentiality of Records and Authorization to Use and Disclose Information for Research Purposes</u>

The University of Rochester makes every effort to keep the information collected from you private. In order to do so, we have sophisticated computer safeguards, such as firewalls, virus checking, network/workstation access passwords, and backup and disaster recovery. Paper forms are stored by unique Registry identification numbers, double locked, and maintained by other University safeguards. Sometimes, however, researchers need to share information that may identify you with people that work for the University, regulators or the study sponsor.

If you have never received a copy of the University of Rochester Medical Center (URMC) and Affiliates Notice of Privacy Practices, please ask the investigator for one.

What information may be used and given to others?

The study doctor will get your personal and medical information. For example:

- Research records
- Records about phone calls made as part of this research

Who may use and give out information about you?

- The study doctor and the study staff
- URMC and Affiliates

Your information may be given to:

- The Department of Health and Human Services
- The University of Rochester
- The Registry's Scientific Advisory Committee, the National Institutes of Health, other government agencies, and foreign government regulatory agencies.

Why will this information be used and/or given to others?

- To do the research
- To study the results

Registry Number:

To see if the research was done correctly

If the results of this study are made public, information that identifies you will not be used.

What if I decide not to give permission to use and give out my health information? Then you will not be able to be in this research study.

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May I review or copy my information? Yes, but only after the research is over.

How long will this permission be valid? This permission will last indefinitely.

May I cancel my permission to use and disclose information?

Yes. You may cancel your permission to use and disclose your health information at any time. You do this by sending written notice to the study doctor. Upon receiving the written notice, the study team will no longer use or disclose your health information and you will not be able to stay in this study. Information that has already been gathered may need to be used and given to others for the validity of the study.

May I withdraw from the study?

Yes. If you withdraw your permission to be in the study, no new health information identifying you will be gathered after that date. Information that has already been gathered may still be used and given to others.

Is my health information protected after it has been given to others?

No. There is a risk that your information will be given to others without your permission.

Use of Email for Communication in Research

When using e-mail to communicate with you in this study, the researcher cannot guarantee, but will use reasonable means to maintain security and confidentiality of e-mail information sent and received. You and the researcher should understand the following conditions, instructions and risks of e-mail use:

Conditions for e-mail use:

- a) E-mail is not appropriate for urgent or emergency situations. The researcher cannot guarantee that any particular e-mail will be read and responded to.
- b) E-mail must be concise. You should schedule an appointment if the issue is too complex or sensitive to discuss via e-mail.
- c) E-mail communications between you and the researcher will be filed in your research record.
- d) Your messages may also be delegated to any member of the study team for response.
- e) The researcher will not forward subject-identifiable e-mails outside of URMC and Affiliates without your prior written consent, except as authorized or required by law.
- f) You should not use e-mail for communication regarding sensitive medical information.
- g) It is your responsibility to follow up and/or schedule an appointment if warranted.

Instructions for e-mail use:

- a) Avoid use of your employer's computer.
- b) Put your name in the body of the e-mail.
- c) Put the topic (e.g., study question) in the subject line.
- d) Inform the researcher of changes in your e-mail address.
- e) Take precautions to preserve the confidentiality of e-mail.
- f) Contact the researcher's office via conventional communication methods (phone, fax, etc.) if you do not receive a reply within a reasonable period of time.

Risks of e-mail use:

Sending your information by e-mail has a number of risks that you should consider. These include, but are not limited to, the following:

a) E-mail can be circulated, forwarded, stored electronically and on paper, and broadcast to unintended recipients.

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Registry Number: _____ RSRB Approval Date: 9/5/2023 Expiration Date: 9/4/2024

- b) E-mail senders can easily misaddress an e-mail.
- c) Backup copies of e-mail may exist even after the sender or the recipient has deleted his or her copy.
- d) Employers and on-line services have a right to inspect e-mail transmitted through their systems.
- e) E-mail can be intercepted, altered, forwarded, or used without authorization or detection.
- f) E-mail can be used to introduce viruses into computer systems.

CONTACT PERSONS

For more information about this research study, please contact:

James Hilbert, MS or Elizabeth Luebbe, MS
University of Rochester, Department of Neurology
601 Elmwood Ave, Box 673
Rochester, NY 14642

Email: dystrophy_registry@urmc.rochester.edu Telephone: (888) 925-4302 or (585) 276-0004.

Please contact the University of Rochester Research Subjects Review Board at 265 Crittenden Blvd., CU 420628, Rochester, NY 14642, Telephone (585) 276-0005 or (877) 449-4441 for the following reasons:

- You wish to talk to someone other than the research staff about your rights as a research subject;
- To voice concerns about the research;
- To provide input concerning the research process;
- In the event the study staff could not be reached.

VOLUNTARY PARTICIPATION

Taking part in this study is voluntary. You are free not to take part or to withdraw at any time, for whatever reason. No matter what decision you make, there will be no penalty or loss of benefit to which you are entitled. In the event that you do withdraw from this study, the information you have already provided will be kept in a confidential manner.

SIGNATURE/DATES

After reading and discussing the information in this consent form, you should understand:

- Why this study is being done;
- What will happen during the study;
- Any possible risks and benefits to you;
- How your personal information will be protected;

What to do if you have problems or questions about this study.

Please complete section 1 **OR** section 2.

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1.) SUBJECT CONSENT (For participants 18 year I have read (or it has been read to me) the contents ask questions. If I had any questions, I have asked to my questions. I agree to participate in this study.	of this consent form and have been encouraged to
If completing these forms on paper, I have received to the study team and the other copy for my records online, I will receive an email with a copy of this form	and future reference). If completing these forms
Subject Name (Printed by Subject)	
Signature of Subject	Date
2.) CONSENT FROM PARENT, LEGAL GUARDIAN REPRESENTATIVE (LAR) I have read (or it has been read to me) the contents ask questions. If I had any questions, I have asked to	of this consent form and have been encouraged to
my questions. I agree to allow the subject to participal of the study team and the other copy for my records	two copies of this consent form (one copy to return
online, I will receive an email with a copy of this form	
Subject Name (Printed by parent, guardian, or LAR)	
Name of Parent, Guardian, or LAR (Printed)	
Signature of Parent, Guardian, or LAR	Date
PERSON OBTAINING CONSENT The subject has been given adequate opportunity to provided with a copy of the consent form for his/her	records.
REGISTRY COORDINATOR PRINTED NAME: REGISTRY COORDINATOR'S SIGNATURE:	
MEGIOTICI GONDINATOR G GIGNATORE:	DATE:
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ASSENT FORM (Adolescents ages 13-17 years)

Study title: National Registry of Myotonic Dystrophy and Facioscapulohumeral Muscular Dystrophy Patients and Family Members

Principal Investigator: Rabi Tawil, M.D.

What are some things you should know about research studies?

You are being asked to take part in a study. Your parent or guardian needs to give permission for you to be in this study. You do not have to be in this study if you don't want to, even if your parent has given permission. You can choose whether or not to be in this study. You may decide not to join. If you join, you may decide to stop being in the study, at any time, for any reason.

What is the purpose of this study?

Research is how we often learn new things. The purpose of this study is to join a Registry that may help doctors and scientists learn about ways to help people with two muscle problems. The two muscle problems are myotonic dystrophy and facioscapulohumeral muscular dystrophy (or FSHD). A registry is a place where medical information is collected and studied for medical research.

You are being asked to join because you or somebody in your family has one of these muscle problems. The goals of the Registry are to:

- To keep track of people with muscle problems.
- To share information with doctors and scientists so that they can learn more about the
 cause of muscle problems and develop better treatments. We won't share your name or any
 information that could identify you.
- To help doctors and scientists find people with muscle problems to participate in their studies. You and your parents can choose whether or not to join any other studies. You don't have to join any other studies.
- To learn more about families with muscle problems.

What will happen if you take part in the study?

If you decide to take part in this study, you will be asked to help your parents answer questions about your symptoms or problems. People without these muscle problems will answer a few questions about their family. We will collect information from your doctor to learn more about your symptoms if you have a muscle problem. We will also collect information from your doctor if you had test that says you don't have a muscle problem.

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Registry Number:	

If you decide to join the Registry, you may be asked at a later time if you would like to help with other studies about these muscle problems. We will send a letter through the mail, email, or online to describe these studies. You can review the information with your parents and decide if you want to help with these studies too. No other doctor or research will know you are in the Registry. It will be up to you and your parents to talk to the other doctors or researchers. We keep your name private and let you decide about what other studies to join.

We will also send you a newsletter through the mail, email, or online with new information about research and muscle problems.

How long will you be in this study?

Your participation in this study may last for several years. We will send you a new questionnaire each year to see if you have any changes (new address, new phone number, or new symptoms if you have a muscle problem). These forms help us keep track of how muscle problems change over time.

Who will be told the things we learn about you in this study?

The information we collect about you will be kept private. Some of your information may be shared with other researchers, but this information won't include your name or anything that could identify you.

What are the possible risks or discomforts involved from being in this study?

The Registry includes questions that may make you feel uncomfortable. You do not have to share any information you do not want to. There may also be an accidental release of your information to other groups. We have many rules to help prevent such accidents.

The University of Rochester makes every effort to keep the information collected from you private. In order to do so, we follow governmental laws about privacy, lock our computers and files, and have other safety tools. Sometimes, however, researchers need to share information that may identify you with people that work for the University, the government or the study sponsor. If this does happen we will take steps to protect the information that you have provided. Results of the research may be presented at meetings or in publications, but your name will not be used.

What are the possible benefits from being in this study?

The potential benefit to you from being in the Registry is receiving information about studies you may want to join. You will also receive newsletters and other information about muscle problems.

What if you or your parents don't want to be in this study?

You do not have to sign this form if you don't want to be in the Registry. Even if your parents say yes, you do not have to. You can change your mind at any time. If some day you decide you want your name taken off the Registry list, just tell your parents or call us and we will remove your name. No one will be upset with you.

Will you get any money or gifts for being in this study?

You will not be paid or given anything for being in this study.

What if you have questions about this study?

For more information concerning this research or if you feel that being in the study has resulted in any research related injury, emotional or physical discomfort, please contact:

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For office use only: Name:	Registry Number:	

James Hilbert, MS or Elizabeth Luebbe, MS
University of Rochester, Department of Neurology
601 Elmwood Ave, Box 673
Rochester, NY 14642

Telephone: (888) 925-4302 or (585) 276-0004.

What if you have questions about your rights as a research subject?

Please contact the University of Rochester Research Subjects Review Board at 265 Crittenden Blvd., CU 420628, Rochester, NY 14642, Telephone (585) 276-0005 or (877) 449-4441 for the following reasons:

- You wish to talk to someone other than the research staff about your rights as a research subject;
- To voice concerns about the research;
- To provide input concerning the research process;
- In the event the study staff could not be reached.

Do I have to be in this study?

Taking part in this study is your choice. You are free not to take part or to stop at any time, for whatever reason. No matter what decision you make, there will be no penalty to you. In the event that you do stop this study, the information you have already provided will be kept private.

SIGNATURE/DATES

SUBJECT ASSENT

I have read (or it has been read to me) the contents of this consent form and have been encouraged to ask questions. If I had any questions, I have called the study team and have received the answers to my questions. I agree to participate in this study.

If completing these forms on paper, I have received two copies of this consent form (one to return to the study team and the other copy for my records and future reference). If completing these forms online, I will receive an email with a copy of this form for my records and future reference.

CHILD'S PRINTED NAME:		
CHILD'S SIGNATURE:		
	DATE:	
PERSON OBTAINING CONSENT The subject has been given adequate opportunity to read the provided with a copy of the consent form for his/her records.	consent before signing a	nd will be
REGISTRY COORDINATOR PRINTED NAME:		_
REGISTRY COORDINATOR'S SIGNATURE:		
	DATE:	
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For office use only: Name:	Registry Number:	



University of Rochester Department of Neurology National Registry for DM and FSHD

601 Elmwood Ave, Box 673

Rochester, NY 14642

Telephone: (888) 925-4302 Fax: (585-276-1947)

& Affiliates Telephone: (888) 925-4302 Fax: (585-276-1947) SH 48 Authorization for Release/Disclosure of Medical and/or Behavioral Health Information

PLEASE PRINT. Patient name:	Date of Birth:
Patient name:	Patient's phone#:
City/State/Zip:	. diente prienen.
•	
This Authorization allows URMC & Affiliates to: (c	heck one or both)
·	
☐ SEND copies of your record to (or discuss your infor	mation with) the provider/person/facility bel*""
☐ RECEIVE copies of your record from (or discuss you	ur information with) the provider/person/facility bel°""
Name of Provider/ Person/Facility	Address
City, State, Zip Code	Phone #/Fax# include area code
	am morado aroa codo
PURPOSE FOR THIS REQUEST: ☐ Healthcare or App	ointment (date) ☐ Insurance ☐ Other
TYPE OF RECORDS or INFORMATION REQUESTED:	Check all that apply:
The records requested are to include: □Mental Health Treatme	nt Records ⊟Alcohol/Drug Treatment Records
Release/disclosure of HIV-related information requires addi	5
•	,
☐ Inpatient admission(s)/date(s): (Check only <u>one</u> of the following 3 choices if requesting inpatier	nt records)
☐ Treatment summary (includes discharge summary, hi	story/physical, laboratory tests, x-ray reports, operative reports,
pathology) Specific information or reports (describe):	
Other (describe):	
□ Outpatient/Officevisitsdate(s):	and/or specific illness/injury:
(Check type of outpatient visit to be released)	
 ☐ Clinic/doctor/dental visit ☐ Ambulatory Surgery visit ☐ Radiology report(s) ☐ Laboratory test results ☐ Immu 	: ☐ Emergency Department Record nizations ☐ Physical/occupational therapy record(s)
☐ Other (describe):	
AUTHORIZATION VALID FOR: (If nothing is checked bel	ow, this authorization is valid for this request only.)
☐ This requestonly	
One year from the date of this authorization OR	
 ☐ This request and for medical records of any future treatment 	
l understand that:	·
 My right to healthcare treatment is not conditioned circumstances (e.g. non-emergent mental health 	
 I may cancel this authorization at any time by subm 	nitting a written request to the address provided at the
top of this form, except where a disclosure has alre- If the person or facility receiving this information	eady <u>been made</u> in reliance on my prior authorization. is not a health care or medical insurance provider
covered by privacy regulations, the information sta	ated above could be redisclosed, <u>except that</u>
chemical dependency treatment records protected not be disclosed without my written authorization i	d by Federal Confidentiality Rules 42C R Part 2 may unless otherwise provided for in the regulations.
 There may be a charge for the requested records The medical records requested above may be fa 	S
- The medical records requested above may be la	Aed III Cases Of Medical Necessity.
Signature of Patient or Representative	Date
Relationship to Patiend(\$AÜ^1 ¦^•^} carãc^DÁ	

National Registry of Myotonic Dystrophy and Facioscapulohumeral Muscular Dystrophy Patients and Family Members

Patient Information Form for individuals with Myotonic Dystrophy or Related Diseases

The purpose of this form is to collect information from individuals who have myotonic dystrophy or a related disease. <u>Please return this form within three weeks if at all possible.</u> If you have any questions about this form, please call Local: (585) 506-0004, in Rochester NY or Toll Free: (888) 925-4302 for assistance.

		write your diagnosis here _ons may not apply to your co	
Date:			
NAME:			
NAME: First	Middle	(Maiden)	Last
ADDRESS:			
Street			
City		State Zip Code	
·	:()	•	
TELEPHONE: Home	: () Area Code Number	State Zip Code Work: () Area Code	
TELEPHONE: Home		Work: ()	Number
TELEPHONE: Home		Work: ()Area Code	Number
TELEPHONE: Home	Area Code Number	Work: ()Area Code	Number
TELEPHONE: Home	Area Code Number	Work: ()Area Code	Number
TELEPHONE: Home	Area Code Number	Work: ()Area Code	Number
TELEPHONE: Home	Area Code Number	Work: ()Area Code	Number
TELEPHONE: Home EMAIL ADDRESS: _ Date of Birth:/_ Mo Da	Area Code Number Sex	Work: ()Area Code	Number
TELEPHONE: Home EMAIL ADDRESS: _ Date of Birth:/_ Mo Da Where did you learn al Your doctor	Area Code Number / Ser bout the Registry? □ Internet	Work: ()Area Code x: Male Female	Number
TELEPHONE: Home	Area Code Number / Ser bout the Registry? □ Internet	Work: ()Area Code x: □ Male □ Female	Number

INFORMATION ABOUT YOUR DIAGNOSIS OF MYOTONIC DYSTROPHY:

1. —	What was the first symptom of myotonic dy	strophy	/?					
2.	. How old were you when you had your first symptom of myotonic dystrophy? (Give your bes estimate even if you are not sure.) years old.							
3.	How old were you when your myotonic dysteven if you are not sure.) years old		was dia	agnosed? (Gi	ve your best estimate			
4.	Did you have any of these tests?							
	Examination by a neurologist		Yes	□ No	□ Not sure			
	Electromyography (EMG, needle inserte into muscles to check electrical activity)		Yes	□ No	□ Not sure			
	Muscle biopsy		Yes	□ No	□ Not sure			
	DNA test (blood test) for myotonic dystrophy		Yes	□ No	□ Not sure			
6.	☐ family member ☐ y ☐ a specialist in a neuromuscular clinic of Were you the first person in your family to h	nave the	cular D					
7.		YES	NO	Not Sure	T			
	anyone else in your family affected with	1 ES	NO	Not Sure				
	votonic dystrophy? If yes, please indicate							
wi	th a check in the appropriate boxes below.	*******	710	N G	N. 1 400 . 1			
D۳	others and sisters	YES	NO	Not Sure	Number Affected			
	ildren							
	(If yes, are any affected children							
	under the age of 18? ☐ yes ☐ no)							
	other							
	ther							
	andparents							
	ints or uncles							
Co	ousins or other relatives							
8.	Are any other members of your family in the	_	try? Yes	□ No	□ Not sure			
For	office use only. Name:			Registry Number				
		2 of 7		-8 / 1 (4111001)	Revised: 09/30/10			

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OCCUPATION AND EMPLOYMENT

What is your current occupation			
☐ Employed (describe your j	ob)		
☐ Homemaker	□ Stude	nt	□ Retired
☐ Disabled because of myoto	onic dyst	rophy	☐ Disabled (not due to myotonic dystrophy)
☐ Unemployed (not due to d	•		Bisacica (not due to myotome dystropmy)
Comments			
Comments			
Has myotonic dystrophy affec	ted your	employmo	ent?
If yes, how (check boxes)	J	1 2	
☐ Lost job			☐ Forced to go on disability
☐ Job modified to accommod	ate vour	nhysical li	•
500 modified to decommod	ate your	physicar n	Larry retirement
EDUCATION			
Highest level of education con	mpleted:		
☐ No formal education			College
☐ Grade school			Graduate school
☐ High school			Other
☐ Technical school			Oon't know
USE OF ASSISTIVE DEVICE	CES		Your age when you started using the
			device (give your best estimate even if
			you are not sure).
	YES	NO	Years old
Use ankle braces			Years old
Use long leg braces			Years old
Use a cane at times			Years old
Use a walker at times			Years old
Use a wheelchair.			Years old
If yes, circle one:			1 3425 314
1. For long distances only			
2. Usually			
3. Always			
Use of CPAP or BIPAP for			Years old
			Tears old
Use ventilator	1		Years old
Have a pacemaker			Years old
Other			Years old

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SIGNS AND SYMPTOMS

Do you have any of the following?	o you have any of the following?			Your age when the		
	Yes	No	Not	problem began (give your		
			sure	best estimate even if you		
				are not sure).		
Trouble with your hands/grip locking up,						
or hand stiffness						
Difficulty making a tight fist, loss of grip						
strength or difficulty opening jars						
Trouble speaking clearly						
Trouble with swallowing						
Weakness of face						
Difficulty walking on your toes or heels, or						
ankle weakness						
Difficulty getting up from the floor, rising						
from a chair, or climbing stairs						
Trouble with breathing or shortness of						
breath						
Cataracts						
Racing heart beat, irregular heart beat,						
palpitations, or pacemaker						
Baldness						
Have you ever had a broken bone or operation If yes, please list them and the date they occur If yes, please list them and the date they occur						
this form.			. •.			
Broken bone or operation	Y	ear th	at it oc	curred		

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MEDICATIONS

<u>MEDICATIONS</u>						
Do you take medications? Yes If yes, please give the name of each medications and herbal remedies.					d non-p	prescription
Codes: 1 Have taken for less than one						
2 Have taken for one month to						
3 Have taken for more than on						
Name of medication	C ₁	rcle o	ne	_	Dosage	
		1 _	I _	Milligrams/Tablet	T	ablets/Day
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
	1	2	3			
What is your current height:feet_ ALLERGIES Please list any foods or drugs to which you smoke tobacco? TREATMENTS OR COUNSELING				u weight	_pounds	
Have you ever received any of the follow	ving?			100		T
DI 1.1.1				Yes	No	Not sure
Physical therapy						
Genetic counseling						
Emotional or psychological counseling						
Speech therapy						
Occupational therapy						
Vocational rehabilitation						
Other						
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OTHER MEDICAL PROBLEMS

Have you ever had or do you have any of these con ☐ Diabetes	nditions:				
☐ High blood pressure	☐ Kidney trouble				
□ Asthma	☐ Thyroid trouble				
☐ Rheumatoid arthritis	☐ Stomach ulcers				
□ Emphysema	☐ Gall bladder trouble				
□ Pneumonia	☐ Prostate trouble				
☐ Heart disease or heart beat irregularity	☐ Liver trouble				
☐ Cancer or tumor, type	☐ Chronic infection				
☐ High cholesterol	☐ Trouble with sexual function				
☐ Miscarriage	☐ Acid reflux or "heartburn"				
□ Stillbirth	□ Constipation				
☐ Child showing signs of myotonic dystrophy within the 1 st four weeks of life	-				
☐ Psychological problems such as depression or	anxiety				
□ Other	•				
ETHNICITY/RACE					
Are you Hispanic or Latino?	\square Yes \square No				
How would you describe your race? Select one or	more of the following categories:				
☐ American Indian or Alaskan Native	□ Asian				
☐ Black or African American	□ White				
☐ Native Hawaiian or other Pacific Islander					
SLEEP PROBLEMS					
How likely are you to doze off or fall asleep in the f					
tired? This refers to your usual way of life in recent					
these things recently try to work out how they would	•				
to choose the <i>most appropriate number</i> for each situ 0 = would <i>never</i> doze	eation:				
1 = slight chance of dozing					
2 = moderate chance of dozing					
3 = high chance of dozing					
Situation	Chance of dozing				
and the					
Sitting and reading					
Sitting, inactive in a public place (such as a theater or a meeting					
As a passenger in a car for an hour without a break					
Lying down to rest in the afternoon when circumstances permi	t				
Sitting and talking to someone					
In a car, while stopped a few minutes in traffic					
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Have you ever participated in a research study for myotonic dystrophy ? $ \ \ \Box \ \ Yes \Box \ \ No \ \ \Box \ \ Not \ sure$
Have you ever received an experimental treatment for myotonic dystrophy? \Box Yes \Box No
If yes, what was that treatment:
In case you needed help filling out this form, who was your helper? (state below) Name of individual filling out the form: Relationship to applicant:
Please provide the name, address, and telephone number of a family member or friend we can contact in case you move or change your phone number.
NAME:RELATIONSHIP:
ADDRESS:
PHONE NUMBER:
Medical records, which confirm your diagnosis, must be sent to us for review. Attached is a Request for Information form. If you sign it and return it to us, we can contact your doctor for any test results and they can send them directly to us.
IMPORTANT Please read, sign and return the attached Consent Form. Without it we cannot consider you for entry into the Registry.
Thank you for your help with the Registry.
Local: (585) 506-0004, Rochester NY Toll Free: (888) 925-4302
FAX: (585) 273-1255
Address: 601 Elmwood Avenue, Box 673, Rochester, NY 14642-8673
The information for this Registry is collected under the authority of Sections 435-442 of the PHS Act (285d-285d-7 of Title 42, USC). The data will be maintained in accordance with the Privacy Act 42 United States Code 241.
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