

Thank you for completing this important information about the way you are conducting outreach for the National ALS Registry. Please submit a form monthly for your organization.

Form Approved OMB No. 0923-0041 Exp. Date xx/xx/201x
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Org Code

Clinic, Support Group, Seminar, Other group activities

Activity	Date	# Registry information kits distributed	# people talked to about the Registry	# people helped sign-up for the Registry	City and State
Total	X				X

Social Media and local mailings (e.g., tweet, email blast, newsletter)

Activity	Date	#	City and State
Total	X		X

Fundraising event (e.g., walk, golf tournament)

Type of event	Date	# attendees	Materials distributed or displayed	City and State
Total	X			X

Outreach calls made where the Registry was discussed

# calls	City and State

Attachment 13B. Outreach Reporting Form for National Offices

Total	

CDC estimates the average public reporting burden for this collection of information as 20 minutes per response, including the time for reviewing instructions, searching existing data/information sources, gathering and maintaining the data/information needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC/ATSDR Information Collection Review Office, 1600 Clifton Road NE, MS D-74, Atlanta, Georgia 30333; ATTN: PRA (0923-0041).