#### **Page Layout: Browse RCTs Introduction Page**

<< What Works Clearinghouse Registries Page Header>>

**Browse RCTs** 

The Registry of Randomized Controlled Trials (RCTs) website provides information about various ongoing and completed education-related RCTs in the United States. Any visitor can browse the RCTs based on the title of the study, the study's sponsor organization, or its unit of randomization. All registered RCTs receive a WWC Registration Number, and those registered prior to trial completion are indicated by a WWC Star Rating.

Browse by Title of Study

Browse by Sponsor Organization

Browse by Unit of Randomization

#### Page Layout: Registries Log-In Page

<< What Works Clearinghouse Registries Page Header>>

#### WWC Log-In

In order to submit an evaluation researcher profile or register a randomized controlled trial (RCT), researchers must have secure log-in credentials.

Researchers registering with the Registry of Evaluation Researchers are required to submit:

- Contact information
- Examples of project experience by study design
- Examples of project experience by publication
- A signed letter of commitment to the WWC Evidence Standards

Log In	
O Evaluation Researchers	○ RCT
User Name:	
Password:	
Log In	
Please Sign Up for New User	
Forgot Your Password?	

We encourage you to print a paper copy of the <u>Registry of Evaluation Researchers form</u> in order to review all the required information before completing the online registration. If you have questions about completing this form, please contact us.

Researchers registering RCTs are required to submit study details, including:

- Principal investigator
- Start and anticipated end dates
- Setting and sample details
- Primary and secondary outcomes

RCTs may be registered once the study receives funding and is in progress. We encourage you to print a paper copy of the <u>RCT Registry form</u> in order to review all the required information before completing the online registration. If you have questions about completing this form, please <u>contact us</u>.

#### **Paperwork Burden Statement**

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless such collection displays a valid Office of Management and Budget (OMB) control number. The valid OMB control number for this information collection is 1850-0788, which is valid through June 30, 2010. The time required to complete this information collection is estimated to average 30 minutes for individuals and organizations per response, including the time to review instructions, gather the data needed, and complete and review the information collection. If you have any comments concerning the accuracy of the time estimate(s) or suggestions for improving this form, please write to: U.S. Department of Education, Washington, DC 20202-4651. If you have comments or concerns regarding the status of your individual submission of this form, contact the What Works Clearinghouse.

#### **Page Layout: Contact Information**

(Details regarding each field are available in Form 07 Details.)

<<What Works Clearinghouse Registries Page Header>>

**Registry of Randomized Controlled Trials** 

Logout

**Contact Information** 

First Name [Text Box]
Last Name [Text Box]

Prefix [Pre-populated drop-down menu]

Title [Text Box]

Type [Pre-populated drop-down menu]

Organization [Text Box] Phone Number [Text Box] Fax Number [Text Box] No Fax Number [Check Box] Email [Text Box] Confirm Email [Text Box] Address Line 1 [Text Box] Address Line 2 [Text Box] City [Text Box] Foreign State [Check Box]

State [Pre-populated drop-down menu]
Country [Pre-populated drop-down menu]

ZIP Code [Text Box]

Save [Button] Reset [Button]

## **Page Layout: Study Information Summary Page**

<<What Works Clearinghouse Registries Page Header>>

Registry of Randomized Controlled Trials

Logout

Studies by Study Title

Below is a list of studies you have submitted. To modify or delete one of these study records, click on the WWC ID number. To enter information about another study, click on the appropriate "Add New Study" link.

WWC ID Study Title

000000 Sample Study Title Here

<u>Add New Study – Funded and In Progress</u> <u>Add a New Study – Completed with Available Final</u> Report

## Page Layout: Study Information Entry (Add New Study – Funded and In Progress) – PAGE 1

(Details regarding each field are available in Form 07 Details.)

<< What Works Clearinghouse Registries Page Header>>

**Registry of Randomized Controlled Trials** 

Logout

Study Information Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is funded and in progress, but not yet completed with an available final report. After review, the WWC will assign a unique ID number to each RCT.

#### Study Information – For Studies Funded and In Progress

Study Status [Radio Button option]

RCT Title [Text Box]
Principal Investigator First Name [Text Box]
Principal Investigator Last Name [Text Box]

Principal Investigator Prefix [Pre-populated drop-down menu]

Principal Investigator Title [Text Box]
Start Date [Text Box]
Anticipated End Date [Text Box]
Sponsor [Text Box]

#### Study Abstract and Details - For Studies Funded and In Progress

- 1. What is the study setting? [Text Box]
  Include settings and locations where the data will be collected, as well as relevant eliqibility criteria for participants.
- 2. What is the intervention? [Text Box] Include intervention type, name(s), and description. Describe how the intervention is designed to be delivered to each group.
- 3. What is the randomized unit? [Radio Button option]

<sup>\*\*</sup>Continued on Next Page\*\*

# Page Layout: Study Information Entry (Add New Study – Funded and In Progress) – PAGE 2 (Details regarding each field are available in Form 07 Details.)

<<What Works Clearinghouse Registries Page Header>>

**Registry of Randomized Controlled Trials** 

Logout

Study Information (continued)
<a href="mailto:Back to Study List">Back to Study List</a>

Please complete the following information related to the randomized controlled trial (RCT) that is funded and in progress, but not yet completed with an available final report. After review, the WWC will assign a unique ID number to each RCT.

#### Study Abstract and Details (continued) – For Studies Funded and In Progress

4. Study Sample Characteristics

Below is a list of target populations identified by the WWC.

Select all that apply to the study sample for this RCT.

Gender

Gender	
Male	[Check Box]
Female	[Check Box]
Not Applicable	[Check Box]
Student Race/Ethnicity	
American Indian or Alaska Native	[Check Box]
Asian	[Check Box]
Black or African American	[Check Box]
Caucasian	[Check Box]
Hispanic or Latino	[Check Box]
Native Hawaiian or Other Pacific Islander	[Check Box]
Not Applicable	[Check Box]
Student Level(s) of Education	
Adult/Continuing Education	[Check Box]
Elementary School	[Check Box]
High School	[Check Box]
Kindergarten	[Check Box]
Middle School	[Check Box]
Preschool	[Check Box]
Postsecondary Education	[Check Box]
Vocational/Career Education	[Check Box]
Not Applicable	[Check Box]

<sup>\*\*</sup>Continued on Next Page\*\*

## Page Layout: Study Information Entry (Add New Study – Funded and In Progress) – PAGE 3 (Details regarding each field are available in Form 07 Details.)

<< What Works Clearinghouse Registries Page Header>>

Registry of Randomized Controlled Trials

Logout

Study Information (continued) Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is funded and in progress, but not yet completed with an available final report. After review, the WWC will assign a unique ID number to each RCT.

## Study Abstract and Details (continued) – For Studies Funded and In Progress

4. Study Sample Characteristics (continued) Student Disability

**Autism** [Check Box] **Deaf-Blindness** [Check Box] **Deafness** [Check Box] Emotional Disturbance (ED) [Check Box] **Hearing Impairment** [Check Box] Mental Retardation [Check Box] **Multiple Disabilities** [Check Box] Orthopedic Impairment [Check Box] Other Health Impairment (OHI) [Check Box] Specific Learning Disability (LD) [Check Box] Speech or Language Impairment [Check Box] Traumatic Brain Injury [Check Box] Visual Impairment [Check Box] Not Applicable [Check Box]

- 5. Additional Study Sample Information [Text Box]
  Include the expected size of the sample and any additional characteristics
  not described in question 4. Describe attrition that has occurred at this point,
  as well as any interim analyses.
- 6. What research design and methods will be used? [Text Box] Include a description of any alterations to the sample after random assignment. Also note the statistical methods intended to compare compatibility of groups on primary outcome(s) and methods for additional analyses.
- 7. What is the control (counterfactual) condition? [Text Box]

<sup>\*\*</sup>Continued on Next Page\*\*

Page Layout: Study Information Entry (Add New Study – Funded and In Progress) – PAGE 4 (Details regarding each field are available in Form 07 Details.)

<< What Works Clearinghouse Registries Page Header>>

Registry of Randomized Controlled Trials

Logout

Study Information (continued) Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is funded and in progress, but not yet completed with an available final report. After review, the WWC will assign a unique ID number to each RCT.

#### Study Abstract and Details (continued) – For Studies Funded and In Progress

- 8. What are the intended primary outcomes? [Text Box] Include all relevant measures, subgroups, and time periods (e.g., Reading test scores for all female students in second grade one year after intervention), and list any planned subgroup analyses.
- 9. What are the intended secondary outcomes? [Text Box] *See question 8 for instructions.*
- 10. What is the data analytic strategy? [Text Box] Include information about power level analysis and plans for estimating effect of the intervention, including calculation of effect sizes.
- 11. Summary of the Study/Abstract [Text Box] *State specific objectives and hypotheses.*

Save [Button] Reset [Button]

## Page Layout: Study Information Entry (Add New Study – Completed with Available Final Report) – PAGE 1

(Details regarding each field are available in Form 07 Details.)

<< What Works Clearinghouse Registries Page Header>>

**Registry of Randomized Controlled Trials** 

Logout

Study Information Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is completed and has an available final report. After review, the WWC will assign a unique ID number to each RCT.

#### Study Information – For Studies Completed with an Available Final Report

Study Status [Radio Button option]
RCT Title [Text Box]
Principal Investigator First Name [Text Box]
Principal Investigator Last Name [Text Box]

Principal Investigator Prefix [Pre-populated drop-down menu]

Principal Investigator Title [Text Box]
Start Date [Text Box]
End Date [Text Box]
Sponsor [Text Box]
Web Address for Report [Text Box]
Published Report Citation (APA Format) [Text Box]

#### Study Abstract and Details - For Studies Completed with an Available Final Report

- 1. What was the study setting? [Text Box] Include settings and locations where the data were collected, as well as relevant eligibility criteria for participants.
- 2. What was the intervention? [Text Box]

  Include intervention type, name(s), and description. Describe how and when the intervention was delivered to each group.
- 3. What was the randomized unit? [Radio Button option]

<sup>\*\*</sup>Continued on Next Page\*\*

## Page Layout: Study Information Entry (Add New Study – Completed with Available Final Report) – PAGE 2

(Details regarding each field are available in Form 07 Details.)

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**Registry of Randomized Controlled Trials** 

Logout

Study Information (continued)
<a href="mailto:Back to Study List">Back to Study List</a>

Please complete the following information related to the randomized controlled trial (RCT) that is completed and has an available final report. After review, the WWC will assign a unique ID number to each RCT.

# Study Abstract and Details (continued) – For Studies Completed with an Available Final Report

4. Study Sample Characteristics

Below is a list of target populations identified by the WWC.

Select all that apply to the study sample for this RCT.

Gender

Gender	
Male	[Check Box]
Female	[Check Box]
Not Applicable	[Check Box]
Student Race/Ethnicity	
American Indian or Alaska Native	[Check Box]
Asian	[Check Box]
Black or African American	[Check Box]
Caucasian	[Check Box]
Hispanic or Latino	[Check Box]
Native Hawaiian or Other Pacific Islander	[Check Box]
Not Applicable	[Check Box]
Student Level(s) of Education	
Adult/Continuing Education	[Check Box]
Elementary School	[Check Box]
High School	[Check Box]
Kindergarten	[Check Box]
Middle School	[Check Box]
Preschool	[Check Box]
Postsecondary Education	[Check Box]
Vocational/Career Education	[Check Box]
Not Applicable	[Check Box]

<sup>\*\*</sup>Continued on Next Page\*\*

## Page Layout: Study Information Entry (Add New Study – Completed with Available Final Report) – PAGE 3

(Details regarding each field are available in Form 07 Details.)

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**Registry of Randomized Controlled Trials** 

Logout

Study Information (continued) Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is completed and has an available final report. After review, the WWC will assign a unique ID number to each RCT.

## Study Abstract and Details (continued) – For Studies Completed with an Available Final Report

4. Study Sample Characteristics (continued) Student Disability

**Autism** [Check Box] **Deaf-Blindness** [Check Box] Deafness [Check Box] Emotional Disturbance (ED) [Check Box] **Hearing Impairment** [Check Box] Mental Retardation [Check Box] Multiple Disabilities [Check Box] Orthopedic Impairment [Check Box] Other Health Impairment (OHI) [Check Box] Specific Learning Disability (LD) [Check Box] Speech or Language Impairment [Check Box] Traumatic Brain Injury [Check Box] Visual Impairment [Check Box] Not Applicable [Check Box]

- 5. Additional Study Sample Information [Text Box]
  Include the sample size and any characteristics not described in question 4.
  Describe attrition that occurred, as well as any interim analyses.
- 6. What research design and methods were used? [Text Box] Include a description of any alterations to the sample after random assignment. Also note the statistical methods used to compare compatibility of groups on primary outcome(s) and methods for additional analyses.
- 7. What was the control (counterfactual) condition? [Text Box]
- 8. What were the primary outcomes? [Text Box]
  For each outcome, include a summary of results of each group and the estimated effect size and level of power analysis. Include all relevant measures, subgroups, and time periods, as well as the number of participants in each group included in each analysis.

<sup>\*\*</sup>Continued on Next Page\*\*

## Page Layout: Study Information Entry (Add New Study – Completed with Available Final Report) – PAGE 4

(Details regarding each field are available in Form 07 Details.)

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**Registry of Randomized Controlled Trials** 

Logout

Study Information (continued)
Back to Study List

Please complete the following information related to the randomized controlled trial (RCT) that is completed and has an available final report. After review, the WWC will assign a unique ID number to each RCT.

## Study Abstract and Details (continued) – For Studies Completed with an Available Final Report

- 9. What were the secondary outcomes? [Text Box] See question 8 for instructions.
- 10. What was the data analytic strategy? [Text Box] *Include information about power level analysis and effect size.*
- 11. Summary of the Study/Abstract [Text Box] *State specific objectives and hypotheses.*
- 12. Interpretation of Results/Discussion [Text Box]

  Provide a brief discussion of findings. Clearly state any limitations and sources of bias.

Save [Button] Reset [Button]