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RPPR Final Report
as of 12-Jun-2018

Agency Code:

Proposal Number: 65324CH

Agreement Number: W911NF-14-1-0208

INVESTIGATOR(S):

Name: Joel M Bowman
Email: jmbowma@emory.edu
Phone Number: 4047276592
Principal: Y

Organization: **Emory University**

Address: 1599 Clifton Road NE, 4th Floor, Atlanta, GA 303224250

Country: USA

DUNS Number: 066469933

EIN: 580566256

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Final Report for Period Beginning 15-May-2014 and Ending 14-May-2017

Title: Theoretical Chemistry: Theoretical Studies of "Roaming" Chemical Reactions

Begin Performance Period: 15-May-2014

End Performance Period: 14-Nov-2017

Report Term: 0-Other

Submitted By: Joel M Bowman

Email: jmbowma@emory.edu

Phone: (404) 727-6592

Distribution Statement: 1-Approved for public release; distribution is unlimited.

STEM Degrees: 0

STEM Participants: 0

Major Goals: Theoretical reaction dynamics including studies of roaming

Accomplishments: Numerous publications and the training of graduate students and postdocs

Training Opportunities: Training of postdocs and graduate students

Results Dissemination: Numerous publications

Honors and Awards: Nothing to Report

Protocol Activity Status:

Technology Transfer: Nothing to Report

PARTICIPANTS:

Participant Type: PD/PI

Participant: Joel Bowman

Person Months Worked: 12.00

Funding Support:

Project Contribution:

International Collaboration:

International Travel:

National Academy Member: N

Other Collaborators:

Participant Type: Faculty

Participant: Paul Houston

Person Months Worked: 12.00

Funding Support:

Project Contribution:

International Collaboration:

International Travel:

National Academy Member: N

RPPR Final Report
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Other Collaborators:

Participant Type: Graduate Student (research assistant)

Participant: Kee Wang

Person Months Worked: 12.00

Funding Support:

Project Contribution:

International Collaboration:

International Travel:

National Academy Member: N

Other Collaborators:

Participant Type: Postdoctoral (scholar, fellow or other postdoctoral position)

Participant: Yimin Wang

Person Months Worked: 3.00

Funding Support:

Project Contribution:

International Collaboration:

International Travel:

National Academy Member: N

Other Collaborators:

Participant Type: Graduate Student (research assistant)

Participant: Chen Qu

Person Months Worked: 6.00

Funding Support:

Project Contribution:

International Collaboration:

International Travel:

National Academy Member: N

Other Collaborators:

Publications 2015-2017

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