

# Antibacterial assays of *Montipora capitata* organic extracts collected in Kaneohe Bay, Oahu, Hawaii during 2014

**Website:** <https://www.bco-dmo.org/dataset/544862>

**Data Type:** experimental

**Version:** 2014-08-28

## Project

» [Host-environment-pathogen interactions in a model coral disease system](#) (coral-pathogen interaction)

Contributors	Affiliation	Role
<a href="#">Gochfeld, Deborah J.</a>	University of Mississippi (UM-NCNPR)	Principal Investigator
<a href="#">Copley, Nancy</a>	Woods Hole Oceanographic Institution (WHOI BCO-DMO)	BCO-DMO Data Manager

## Table of Contents

- [Dataset Description](#)
  - [Methods & Sampling](#)
  - [Data Processing Description](#)
- [Data Files](#)
- [Parameters](#)
- [Deployments](#)
- [Project Information](#)
- [Funding](#)

## Dataset Description

2018-11-12: Contacted Gochfeld about unrestricting this dataset. She requested further restriction till after the new year. Changed release date from 2018-10-01 to 2019-02-01.

Data access is restricted. Please contact the PI for further information.

### Related datasets:

[Montipora antibacterial-aqueous](#)

[Montipora antibacterial-mucus](#)

[Montipora chemical fingerprints](#)

[MWS accession numbers](#)

[MWS lesion progression](#)

### Related references:

Deborah J. Gochfeld & Haidy N. Kamel & Julie B. Olson & Robert W. Thacker. 2012. Trade-offs in defensive metabolite production but not ecological function in healthy and diseased sponges. *J Chem Ecol* 38:451-462. DOI 10.1007/s10886-012-0099-5.

## Methods & Sampling

**Preparation of extracts:** Coral samples of orange and red colonies of *Montipora capitata* were collected in Kaneohe Bay, Oahu. Frozen samples were extracted in 2:1 dichloromethane:methanol, which was replaced daily for 3 d. Extracts were filtered, dried, and weighed. We tested extracts at concentrations approximating those naturally found in the coral tissues (Gochfeld & Aeby 2008). To determine the volumetric concentrations of extracts, the surface area of each piece of coral was calculated using the wax technique (Gochfeld 1991). Tissue volume was determined by multiplying surface area by tissue depth measured from replicate decalcified pieces of *M. capitata*. Extract concentrations were determined as g dried extract ml<sup>-1</sup> of coral tissue. Extracts were re-suspended to these concentrations in 2:1 dichloromethane:methanol for use in disk diffusion assays.

**Bacterial strains tested:** The strains used for our antibacterial assays were selected as model systems to represent a range of potential bacterial pathogens from the marine environment, including known coral pathogens (*Aurantimonas coralicida*, *Serratia marcescens*, *Vibrio coralliilyticus*, *Vibrio shiloi*) and human enteric bacteria that have the potential to enter near-shore waters and can survive in the marine environment (*Yersinia enterocolitica*), and strains previously isolated from the surfaces of Hawaiian corals (*Klebsiella pneumonia*, *Vibrio agarivorans*) (Gochfeld & Aeby 2008). In addition, bacterial strains isolated from surfaces of *Montipora capitata* (OCN001, OCN002, OCN003, OCN008, O1-1, O2-8, O2-12, R1-13, R5-5, R5-13, R5-29) by Dr. Sean Callahan's lab at University of Hawaii were also used. Two of these, OCN002 and OCN008, are pathogens associated with *Montipora* White Syndrome (Ushijima et al. 2012, 2014).

**Disk Diffusion Assay** (Gochfeld et al. 2012): Extracts from orange and red colonies (n = 4 replicates of each color morph) were added to 4-mm diameter paper disks at natural volumetric concentrations. Assays were run on triplicate petri plates. Twenty-four hour bacterial cultures were plated onto appropriate medium, and disks impregnated with the organic extracts were placed on the plate. Following a 24-hr incubation period, the zones of inhibition surrounding each disk were measured.

## Data Processing Description

Data represent mean zones of inhibition in millimeters for the three replicate disks of each sample against each bacterium.

### BCO-DMO Processing:

added conventional header with dataset name, PI name, version date, reference information  
renamed parameters to BCO-DMO standard  
replaced blanks and / with underscores

[ [table of contents](#) | [back to top](#) ]

---

## Data Files

File	
<b>assays.csv</b>	(Comma Separated Values (.csv), 1.12 KB) MD5:f0a329f644dad5cec74e0dab61c1384a
Primary data file for dataset ID 544862	
<b>assays_Aeby_2014.csv</b>	(Comma Separated Values (.csv), 1.12 KB) MD5:f0a329f644dad5cec74e0dab61c1384a
Antibacterial assays of <i>Montipora capitata</i> extracts D. Gochfeld (UMiss) version: 2014-08-28	

[ [table of contents](#) | [back to top](#) ]

---

## Parameters

Parameter	Description	Units
assay	coral extract material examined	unitless
color_morph	color morph of coral	unitless
coral	coral fragment identification	unitless

Aurantimonas_coralicida	mean zone of inhibition against Aurantimonas coralicida	millimeters
Klebsiella_pneumoniae	mean zone of inhibition against Klebsiella pneumoniae	millimeters
Pseudomonas_nautica	mean zone of inhibition against Pseudomonas nautica	millimeters
Serratia_marcescens	mean zone of inhibition against Serratia marcescens	millimeters
Vibrio_agarivorans	mean zone of inhibition against Vibrio agarivorans	millimeters
Vibrio_corallyliticus	mean zone of inhibition against Vibrio corallyliticus	millimeters
Vibrio_shiloi	mean zone of inhibition against Vibrio shiloi	millimeters
Yersinia_enterocolitica	mean zone of inhibition against Yersinia enterocolitica	millimeters
OC_N001	mean zone of inhibition against OC-N001	millimeters
OC_N002	mean zone of inhibition against OC-N002	millimeters
OC_N003	mean zone of inhibition against OC-N003	millimeters
OC_N008	mean zone of inhibition against OC-N008	millimeters
O1_1	mean zone of inhibition against O1-1	millimeters
O2_8	mean zone of inhibition against O2-8	millimeters
O2_12	mean zone of inhibition against O2-12	millimeters
R1_13	mean zone of inhibition against R1-13	millimeters
R5_5	mean zone of inhibition against R5-5	millimeters
R5_13	mean zone of inhibition against R5-13	millimeters
R5_29	mean zone of inhibition against R5-29	millimeters

---

## Deployments

### Aeby\_2014

<b>Website</b>	<a href="https://www.bco-dmo.org/deployment/544868">https://www.bco-dmo.org/deployment/544868</a>
<b>Platform</b>	Hawaii_reef
<b>Start Date</b>	2010-06-01
<b>End Date</b>	2014-05-31
<b>Description</b>	Coral reef pathogen studies.

[ [table of contents](#) | [back to top](#) ]

---

## Project Information

### Host-environment-pathogen interactions in a model coral disease system (coral-pathogen interaction)

**Coverage:** Kaneohe Bay, Oahu, Hawaii (21 26' N, 157 47' W)

*Extracted from the NSF award abstract:*

Diseases of marine organisms have emerged as a serious problem contributing to the decline of coral reef resources worldwide. Loss of coral reef habitats carry social and economic implications especially in island states, such as Hawaii, which depend on reefs for food, shoreline protection and tourism. Our ability to manage coral diseases is hampered by a lack of knowledge of which environmental variables affect disease, mechanisms of host defense, and the etiology of most of the numerous described coral diseases. The PIs of this project discovered a coral disease system that can be used as a model to explore many components of the host-environment-pathogen triangle of disease causation. Montipora white syndrome (MWS) is an infectious disease that results in progressive tissue loss on colonies of Montipora capitata, and has been found on reefs throughout the Hawaiian archipelago. It is particularly prevalent in Kaneohe Bay, Oahu, which has a long history of reduced water quality, and this suboptimal environment sets the stage where host-pathogen interactions occur. In Kaneohe Bay, M. capitata is a major reef-building species, and is found in two color morphs (red and orange) that harbor different clades of zooxanthellae. During preliminary surveys, the PIs discovered intraspecific variability in response to MWS between color morphs. Although the red morph was dominant within survey transects (80% of the colonies), the orange morph was disproportionately affected by MWS (70% of the affected colonies). Microbial studies found a shift in bacterial communities on MWS-affected and healthy M. capitata and allowed identification of potential pathogens. Numerous bacterial strains were cultured and screened for pathogenicity and three strains, which produced lesions, were identified as potential pathogens. Two of the putative pathogens (Vibrio spp.) produced diffuse tissue whereas the other bacterial strain (Pseudoalteromonas sp.) produced acute tissue loss.

In the field, the PIs also observed two patterns of tissue loss on M. capitata; a slow, chronic pattern of tissue loss, which they followed through time with tagged colonies (chronic MWS), but also a rapid onset of acute tissue loss (acute MWS). Thus they discovered an infectious coral disease that results in significant coral mortality that has the unique component of differences in disease susceptibility among color morphs. The PIs identified three potential bacterial pathogens that will be used to investigate underlying factors affecting the coral-environment-pathogen triad of disease causation. The Hawaii Institute of Marine Biology (HIMB) is located within Kaneohe Bay allowing year-round access to reefs for research on Montipora white syndrome. The goal of this project is to investigate the host- environment-pathogen triangle of disease causation for Montipora white syndrome. The objectives of this research will be to: 1) investigate mechanisms contributing to differential disease resistance in red (less susceptible) vs. orange (more susceptible) morphs of M. capitata. The PIs will compare antimicrobial activity in the holobiont, mucus and mucus-associated bacteria of the two color morphs of M. capitata, and will compare the natural coral-associated microbial flora between the two color morphs; 2) use manipulative aquarium studies to determine whether environmental stressors (elevated temperature, nutrient stress) differentially affect the progression or transmission efficiency of MWS in red vs. orange morphs of M. capitata; 3) use challenge experiments to confirm the role of bacterial pathogens as

causative agents of MWS, and to determine the response of red vs. orange morphs of *M. capitata* to three putative pathogens. This project will involve a multidisciplinary team to provide a broader perspective of coral disease processes. This will be the first comprehensive study conducted on a coral disease in Hawaii.

*Related Publications:*

Ushijima, B, Videau, P, Burger, A, Shore-Maggio, A, Runyon, C, Sudek, M, Aeby, G and S. Callahan. 2014. *Vibrio coralliilyticus* strain OCN008 is an etiological agent of acute Montipora white syndrome. *Applied & Environ Microbiology* doi:10.1128/AEM.03463-13.

Ushijima B, Videau P, Aeby GS, Callahan SM. 2013. Draft Genome Sequence of *Vibrio coralliilyticus* Strain OCN008, Isolated from Kane'ohe Bay, Hawai'i. *Genome Announc.* 2013 Oct 3;1(5). doi:pii: e00786-13. 10.1128/genomeA.00786-13. PMID: 24092784

Ushijima B, Smith A, Aeby GS, Callahan SM (2012) *Vibrio owensii* Induces the Tissue Loss Disease Montipora White Syndrome in the Hawaiian Reef Coral *Montipora capitata*. *PLoS ONE* 7: e46717.

[ [table of contents](#) | [back to top](#) ]

---

## Funding

Funding Source	Award
<a href="#">NSF Division of Ocean Sciences (NSF OCE)</a>	<a href="#">OCE-0961384</a>

[ [table of contents](#) | [back to top](#) ]