

Version 1

Jun 22, 2021

Gates lab qPCR protocol for *Cladocopium* and *Durusdinium* Symbiodiniaceae detection V.1

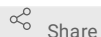
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ABSTRACT

The purpose of this protocol is to quantify *Cladocopium* and *Durusdinium* Symbiodiniaceae cell ratio densities (symbiont to host cell ratios) from individual coral DNA samples using the StepOnePlus™ Real-Time PCR System. It allows for a quick assessment of the symbiont community composition and ratio using targeted primers.

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KEYWORDS

qPCR, Symbiodiniaceae, coral , Quantitative PCR, Real Time PCR

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Apr 14, 2020

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35702

MATERIALS TEXT

MATERIALS

 [qPCR primers](#) **Contributed by users**

 [MicroAmp™ Optical Adhesive Film](#) **Thermo**

Fisher Catalog #4311971

 [qPCR Mastermix \(GTMM\)](#) **Fisher**

Scientific Catalog #4346907

 [qPCR 96-well](#)

[plate](#) **Qiagen Catalog #209002**

StepOnePlus™ Real-Time PCR System

ThermoFisher 4376600

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BEFORE STARTING

Introduction

The purpose of this quantitative PCR protocol is to quantify *Cladocopium* and *Durussdinium* symbiont cell ratio densities (symbiont to host cell ratios) from each DNA sample. It measures the abundance of symbionts relative to the total number of host cells at the tissue or colony level. This technique amplifies specific target gene loci in both the symbiont and the host to calculate a ratio of the total number of symbiont cells to host cells (S/H cell ratio).

The primers and probes presented here have been developed for two coral species commonly found in Hawaii: *Montipora capitata* (Cunning et al. 2016) and *Pocillopora damicornis* (Cunning and Baker, 2013). *Cladocopium* (former clade C) and *Durussdinium* (former clade D) are the most abundant symbiont species found in Hawaii, and the ones expected to be found in those coral species. Additional actin primers have been developed for detection of other Symbiodiniaceae species (see Cunning et al. 2015 for *Breviolum* (former clade B) and Winter, R. N. 2017 for *Symbiodinium* (former clade A)).

The same protocol can be applied to other coral species. In the absence of a specific primer and probe for the coral species of interest, the protocol can be run without host (in this case, qPCR results will not provide S/H cell ratio, but only symbiont relative proportions).

Make sure you have the DNA extracted and quality checked.

Any mistakes?

This protocol is always changing to move with the times, so if you notice any mistakes or issues, please feel free to let us know so we can improve it.

Primer and Probe Dilutions

Primers usually come in lyophilized form. We recommend diluting them to a 100µM stock, then diluting the stock to get the desired concentration for the reactions. For the reactions, Forward primers should be 1µM and Reverse primers should be 1.5 µM. Probes should be diluted to 2µM (**Table 1**).

Stock should be kept at -20 °C and when in use, kept at 4 °C (avoid freezing and thawing). Mastermix should be kept at 4 °C at all times.

Probes for the detection of Symbiodiniaceae *Cladocopium* (former clade C) and *Durusdinium* (former clade D) were labeled by [Cunning et al. \(2013\)](#) with different fluorescent dyes (VIC and FAM respectively), allowing those reactions to be multiplexed. Primers and probes for two coral hosts (*Montipora capitata* and *Pocillopora damicornis*) have been developed by [Cunning et al. \(2017\)](#) and [Cunning & Baker \(2013\)](#) respectively.

Table 1. Primers and probes for symbionts and coral host.

A	B	C	D	E
Target	Name		Working Concentration	Sequence
Symbiodiniaceae				
<i>Cladocopium (C)</i>	CActF	Forward	1µM	5'-CCAGGTGCGATGTCGATATTC- 3'
	CActR	Reverse	1.5 µM	5'-TGGTCATTGCTCACCAGT- 3'
	CActProbe	Probe	2 µM	5'-VIC-AGGATCTCTATGCCAACG-MGB- 3'
<i>Durusdinium (D)</i>	DActF	Forward	1µM	5'-GGCATGGGGTAAGCACTTCTT- 3'
	DActR	Reverse	1.5 µM	5'-GATCCTTGAAGTAGCCTTGGAAAC- 3'
	DActProbe	Probe	2 µM	5'-6FAM-CAAGAACGATACCGCC-MGB- 3'
Coral host				
<i>Montipora capitata</i>	PaxC-For	Forward	1µM	5'-GTGCAGGTGAGATTGAGTCTTATAACA-3'
	PdAct-Rev	Reverse	1.5 µM	5'-CGGTTGAGCTTCGCTAAACAG-3'
	PaxC-Probe	Probe	2 µM	5'-FAM-CAGTCTTCCAACAATG-MGB-3'
<i>Pocillopora damicornis</i>	PdAct-For	Forward	1µM	5'-GAGAAGCTCTGCTATGTTGCCA- 3
	PdAct-Rev	Reverse	1.5 µM	5'-TCCACGGAATCGCTCGTT- 3'
	PdActProbe	Probe	2 µM	5'-NED-AGACTGCTGCCTCAAC-MGB- 3'

Mastermix preparation

Note:

- Always run duplicates or triplicates of each sample
- The Micro Amp Fast plate has 96 wells, but in our machine, the 4 corners of the plate don't provide a good read, so in a 96 well plate, we only use 92 wells. You might need to test that for the machine you are using.
- Print a 96 well plate template and write the sample ID and the reaction it will be running (S for Symbiodiniaceae and

H for coral host) on the printed wells .

- To minimize the time the mastermix will sit on the bench at room temperature, separate the DNA samples you will use in the order they will be plated before you start preparing the mastermix. Leave them in order in a microcentrifuge rack in the freezer.
- It is always recommended to do the whole protocol with the reagents and the samples on ice.

Calculation

Each well should have 9 µl of your qPCR mastermix and 1 µl of DNA sample. Prepare two separate mastermixes, one for the host and one for the Symbiodiniaceae .

qPCR Symbiodiniaceae C and D mastermix (amount per well)

- 5 µl GTMM
- 0.5 µl Symb C primer (F), 0.5 µl Symb C primer (R), 0.5 µl, 0.5µl Symb C probe
- 0.5 µl Symb D primer (F), 0.5 µl Symb D primer (R), 0.5 µl Symb D probe
- 1 µl Hi clone ddH2O

qPCR Coral Host master mix (amount per well)

- 5 µl GTMM
- 0.5 µl Host primer (F), 0.5 µl Host primer (R), 0.5 µl Host probe
- 2.5 µl Hi clone ddH2O

- Remember to have 3 Controls (c) :

Positive control: coral with Symb C and Symb D (a sample that has been confirmed to have both symbionts you are testing)

Negative control 1: Hi clone ddH2O (will test for external source of DNA)

Negative control 2: control TE buffer (extraction control, replace with the reagent your DNA samples had been eluted with)

- Calculate the quantity of mastermix according to the number of samples you are running and make each mix in a 1.5 µl centrifuge tube, one for the Symbiodiniaceae and one for the coral host.

If doing a full plate, here are the calculations (92 samples):

qPCR Symbiodiniaceae C and D (46 samples)

- 230 µl GTMM
- 23 µl Symb C primer (F), 23 µl Symb C primer (R), 23 µl Symb C probe
- 23 µl Symb D primer (F), 23 µl Symb D primer (R), 23 µl Symb D probe
- 46 µl Hi clone ddH2O

qPCR Coral Host master mix (46 samples)

- 230 µl GTMM
- 23 µl Host primer (F), 23 µl Host primer (R), 23 µl Host probe
- 115 µl Hi clone ddH2O

2 Plating mastermix

IMPORTANT: Vortex each mix thoroughly before plating.

- Add 9 µl of mastermix to the bottom of each well (remember to check which ones are Symbiont wells and which ones are Host wells). Change pipettes between mastermixes.
- Flick each DNA sample before use, for better mixing
- Add 1 µl of DNA sample to the top side of each well. Change pipette between each DNA sample. As 1 µl is such a small volume, adding to the side makes it easy to track which samples have been loaded already. The droplets will be visible on the side. Plus, adding to the side instead of directly into the mix in the bottom of the well helps prevent bubbles.
- Seal the plate with qPCR sealing tape and use roller to ensure it is well sealed.

- Put plate in centrifuge to spin DNA into mastermix at 10,000 g for about 10s.
- Check that the liquids are at the bottom of each well, that all wells have same amount of liquid, and that there are no bubbles. If there are bubbles, flick the wells and centrifuge again.

3 Running the samples in the Step One Plus machine

Put samples in thermocycler (StepOnePlus machine), and check that the samples are placed in the same orientation you loaded them.

Thermocycler settings have previously been saved with the name TaqMan_Std (**Figure 1**).

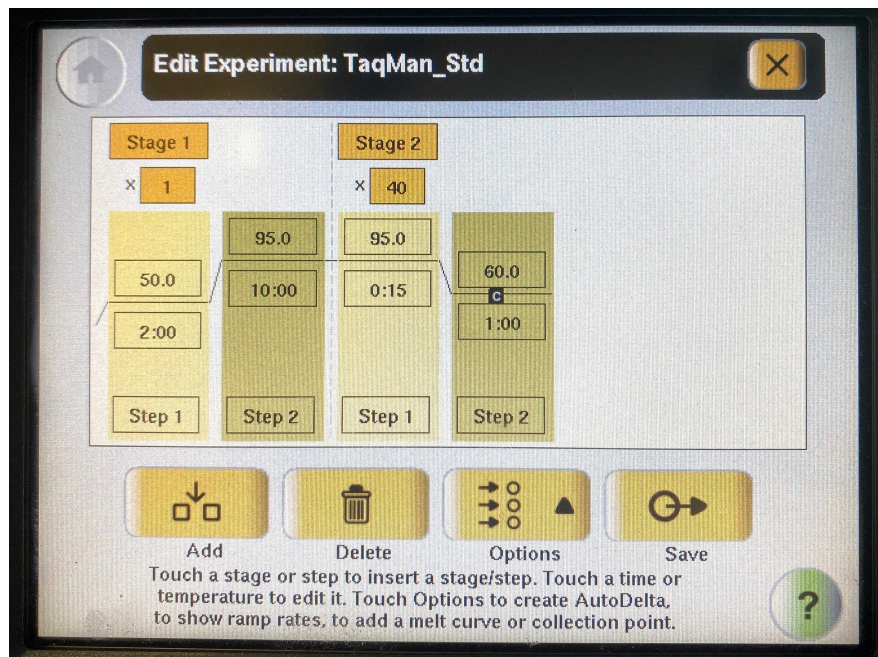


Figure 1. Thermocycler settings

- Check that the settings are correct and press save!
- Change volume to your sample volume (10ul)
- Select Run and wait for reaction to start to make sure it is running

4 Collecting results and visualizing results on computer

- When reaction is finished, insert a USB thumbdrive into the Step One Plus Machine and select "Save Results", once the file has been saved, remove the USB thumbdrive. Results can also be taken directly from the computer connected to the Step One Machine by selecting "Collect experimental results" on the computer.
- Open your Taq file on the computer (computer should have StepOne software installed).
- Click on "Setup" on the left hand column. In "Define Targets" (your C and D and host) and your samples. Click "Add New Target" into "Define Targets" to get a new row. For reporter, select the fluorescent dye for each reaction (**Table 2**). Choose also the color you would like to display each reporter.

Table 2. Target and fluorescent dyes

A	B
Target	Fluorescent dye
<i>Cladocopium (C)</i>	VIC
<i>Durusdinium (D)</i>	FAM
<i>Montipora capitata</i>	FAM
<i>Pocillopora damicornis</i>	NED

- Now on "Define Samples", on the right side, list all your sample names (**Figure 2**).

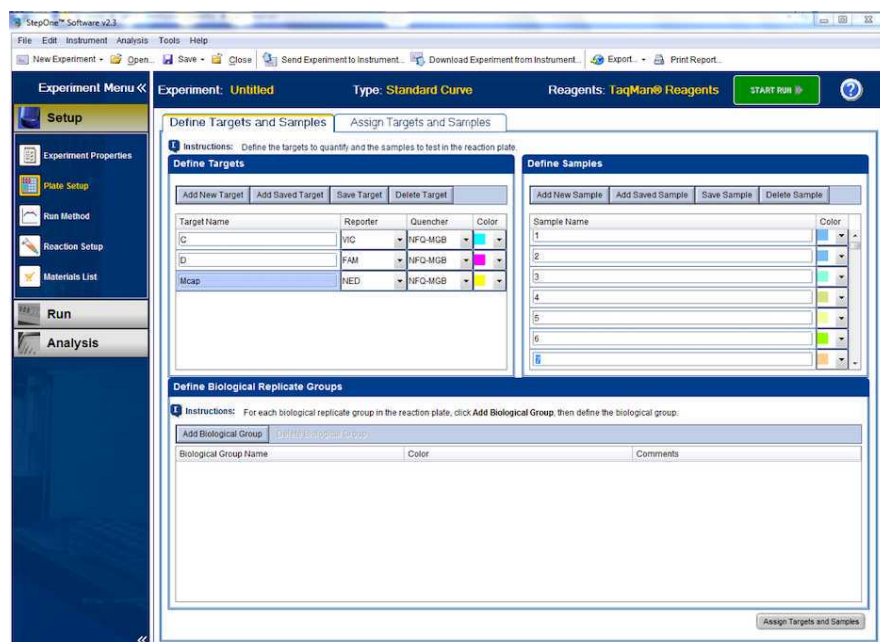


Figure 2. Defining samples

- Now on "Define Samples", on the right side, list all your sample names (**Figure 2**).
- Now click on the tab "Assign Targets and Samples". You will assign targets and samples to each well (**Figure 3**). Highlight the well with your mouse and then check the box next to the target (both) and the sample name. You can highlight all the wells and assign targets at once, and do the sample names individually.

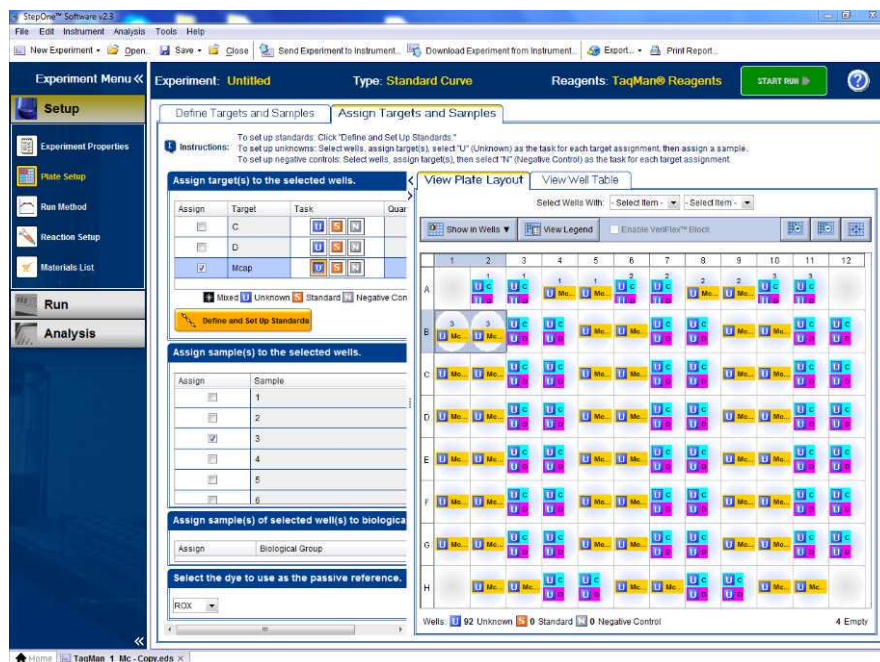


Figure 3. Assigning targets and samples.

- Now click on "Analysis" on the left hand column, then click "Analyze" on the top right corner.
- On left, click "Multicomponent Plot" (**Figure 4**). Here we are looking for the number of cycles before amplification. We want to tell the machine what our baseline is. We want it to be large, but remember, it needs to be the same for all of your runs. Let's do 6-23.
- On the top right, click on "Analysis settings" (**Figure 5**). Enter the threshold (usually 0.01) and the baseline for both C and D (usually 6-23). Then hit "Apply Analysis Settings" (we tend not to use the "auto" settings and instead edit them ourselves).

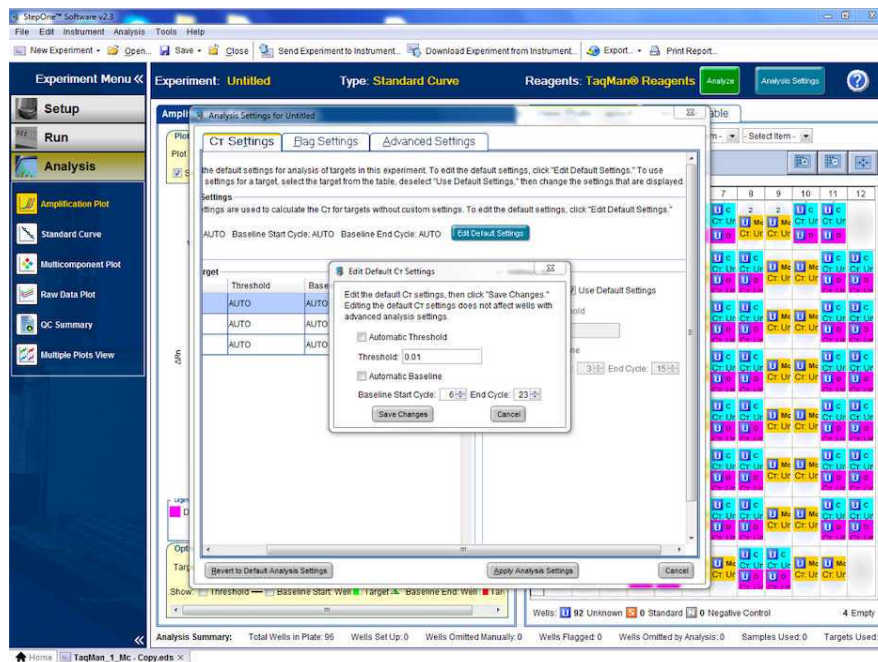


Figure 5. Editing default settings.

- Now the screen should show you the amplification plot (if you don't see the lines, make sure the "Show baseline" and "Show threshold" buttons are clicked at bottom). Also, select "Well" or Target into the Plot Color. Note, you can drag to make the wells larger so you can read the numbers as you would in an Excel sheet. You can also select specific wells, or row or wells. (**Figure 6**).

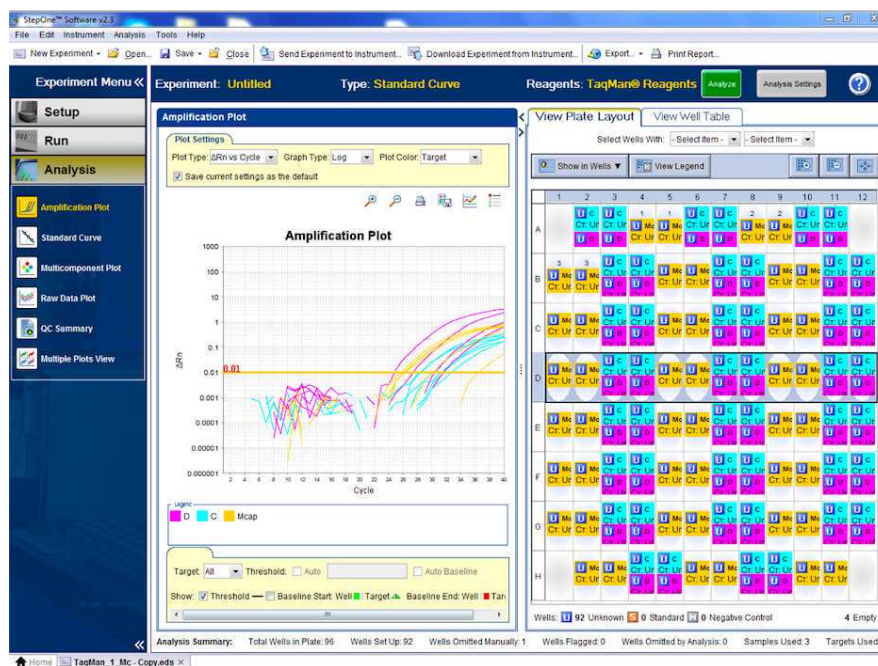


Figure 6. Amplification plot.

- Now we want to export the data as an Excel sheet. Under "File", select "Export". Select "Results" as the data to export. Name your file, and be sure it is exported as an Excel file.
- Also save the StepOne settings and set up (in case you find a mistake, you won't have to re-do everything), by going into "saving as". Give a name for the settings and this run.
- Then you can copy it to your USB and save to your laptop.

5 Bioinformatics

- You need to run your data through an R script to see what the values mean. Ross Cunning has developed calculations (Cunning et al. 2012, 2017) and an R script to do so [GitHub](#). A modified version is below.
- The file "qPCR1.xls" referred to in the script is the Excel file exported from the StepOne machine after doing the qPCR.

```
library(tidyverse)
library(janitor)
library(readxl)
library(ResourceSelection)

#Cunning et al. 2012 - Excess algal symbionts increase the susceptibility of reef corals to bleaching
#Cunning et al. 2017 - Patterns of bleaching and recovery of Montipora capitata in Kaneohe Bay, Hawaii, USA

#-----#qPCR1 #-----#

##### Uploading data and cleaning #####
setwd("~/Desktop/qPCR_results/")
rawdata<-read_excel("qPCR1.xls")
rawdata<-rawdata[-(1:6),] #####deleted first six rows of header information
colnames(rawdata) <- rawdata[1,] #####set first row as column names
rawdata <- rawdata[-1,] #####delete first row
format_data<-clean_names(rawdata); colnames(format_data)[7] <- "ct"; colnames(format_data)[8] <- "ct_mean";
colnames(format_data)[9] <- "ct_sd" ### this renames columns manually, there are some weird capital subscripts in
column names, so this fixes them
format_data$sample_name <- gsub("\\\\','-", format_data$sample_name); head(format_data)

##### Set arguments #####
copy.n.C<-33 #####from Cunning et al. 2017 Supplemental Materials
copy.n.D<-3 #####from Cunning et al. 2017 Supplemental Materials
copy.n.Mcap<-1 #####from Cunning et al. 2017 Supplemental Materials
copy.n.ratioCD<-(copy.n.C/copy.n.D)
copy.n.ratioCM<-(copy.n.C/copy.n.Mcap)
copy.n.ratioDM<-(copy.n.D/copy.n.Mcap)
fluo.C<-2.26827 #####from Cunning et al. 2017 Supplemental Materials
fluo.D<-0 #####from Cunning et al. 2017 Supplemental Materials
fluo.Mcap<-0.84815 #####from Cunning et al. 2017 Supplemental Materials

##### Process data #####
data<-format_data%>%
dplyr::select(sample_name,target_name,ct_mean, ct_sd)%>% ###take only the columns we care about
mutate_at(vars(starts_with("ct")),funs(as.numeric))%>% ###set all the ct columns to numeric values (they are
originally stored as characters)
group_by(sample_name,target_name)%>%###group technical replicates
summarise_all(funs(mean(., na.omit = TRUE)))%>% ###take mean of technical replicates
filter(!grepl("con",sample_name))%>%
filter(sample_name!="NA")

C<-filter(data,target_name=="C")
D<-filter(data,target_name=="D")
Mcap<-filter(data,target_name=="Mcap")
```

```

final_data1<-left_join(left_join(C,D,by="sample_name"),Mcap,by="sample_name")%>%
dplyr::select(-target_name.x,-target_name.y,-target_name)%>%
dplyr::rename(d_mean=ct_mean.y)%>% ###rename columns from rbind
dplyr::rename(d_sd=ct_sd.y)%>%
dplyr::rename(c_mean=ct_mean.x)%>%
dplyr::rename(c_sd=ct_sd.x)%>%
dplyr::rename(mcap_mean=ct_mean)%>%
dplyr::rename(mcap_sd=ct_sd)%>%
mutate(c_mean=c_mean-fluo.C)%>%#from Cunning et al. 2017
mutate(mcap_mean=mcap_mean-fluo.Mcap)%>% #from Cunning et al. 2017
mutate(presence=case_when(d_mean>0~"CD",is.na(d_mean)~"C"))%>% ###set presence/absence in new column
based on ct values
mutate(dm_ratio=(2^(mcap_mean-d_mean))/copy.n.ratioDM)%>%#from Cunning et al. 2012
mutate(cm_ratio=(2^(mcap_mean-c_mean))/copy.n.ratioCM)%>%#from Cunning et al. 2012
mutate(cd_ratio=(cm_ratio/dm_ratio))%>%#from Cunning et al. 2017 ###calculate ratio, log of 2^(difference in ct
values)
mutate(sh_ratio=sum(cm_ratio,dm_ratio,na.rm=TRUE))%>%#from Cunning et al. 2012 ###add C:host ratio + D:host
ratio for total symbiont:host ratio###set even numbers as nonbleached, odd as bleached. this works by using the
remainder command (%), if 0 it must be an even number
mutate(prop_c=cd_ratio/(cd_ratio+1))%>%#from Cunning et al. 2012
mutate(prop_c=case_when(is.na(prop_c)~"1",TRUE ~ as.character(prop_c)))%>% #from Cunning et al. 2012
mutate(prop_c=as.numeric(prop_c))%>%
mutate(prop_d=1-prop_c)%>% ###set proportion D off of proportion C
mutate(abundance=case_when(prop_d>prop_c~"D>C",prop_c>prop_d~"C>D"))%>% ###set abundance based on
which proportion is dominant, create character factor
mutate(sd_warning=case_when((c_sd>1&d_sd>1)~"cd*",c_sd>1~"c*",d_sd>1~"d*"))%>% ###set warnings where
standard deviation of tech replicates is >1
mutate(ct_warning=case_when((c_mean>38&d_mean>38)~"cd*",c_mean>38~"c*",d_mean>38~"d*"))%>% ###set
warnings where ct values is later than 34
mutate(rep=case_when((is.na(d_sd)&d_mean>0)|(is.na(c_sd)&c_mean>0)~"1replicate"))%>% ###set warning if only
one replicate processed (no standard deviation but mean present)
separate(sample_name,into=c("Timepoint","Fragment"),sep="-")%>% ### customize this to fit your well names
mutate_at("Timepoint", as.factor)%>%
mutate_at("Fragment", as.factor)%>%
mutate(C=as.integer(prop_c*100))%>%
mutate(D=100-C)%>%
filter(is.na(rep))

# this is your final file with the result of all your qPCRs. You can now do stats and visualize your results!

```