

# Coupled biophysical modeling to understand the spread of a deadly coral disease in Florida

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## 2 ABSTRACT

3 For about six years, the Florida Reef Tract (FRT) has been experiencing an outbreak of the  
4 Stony Coral Tissue Loss Disease (SCTLD). First reported off the coast of Miami-Dade County  
5 in 2014, the SCTLD has since spread throughout the entire FRT with the exception of the Dry  
6 Tortugas. However, the causative agent for this outbreak is currently unknown. Here we show  
7 how a high-resolution bio-physical model coupled with a modified patch SIR epidemic model can  
8 inform on the potential characteristics of the causative agent of the disease and its vector. In this  
9 study, the agent is assumed to be transported within composite material (such as coral mucus,  
10 dying tissues and/or resuspended sediments) driven by currents and potentially persisting in the  
11 water column for extended periods of time. In this framework, our simulations suggest that the  
12 SCTLD is likely to be propagated within neutrally buoyant material driven by mean barotropic  
13 currents. Calibration of our model parameters with field data show that corals are then infected  
14 within a mean transmission time of 6.45 days, with a basic reproduction number slightly above  
15 1. Furthermore, the propagation speed of the disease through the FRT is shown to occur for  
16 a well defined range of values of an infection threshold, defined as the fraction of infectious  
17 corals that causes an exponential growth of the disease in the colony. Our results present a new  
18 connectivity-based approach to understand the spread of the SCTLD through the FRT. Such a  
19 method can provide a valuable complement to field observations and lab experiments to support  
20 the management of the epidemic as well as the identification of its causative agent.

21 **Keywords:** stony-coral-tissue-loss disease, biophysical modeling, Florida reef tract, spatial epidemiology, connectivity

## 1 INTRODUCTION

Coral diseases are a major threat to coral reef ecosystems and have led to significant declines in coral cover especially within the Caribbean region (Richardson et al., 1998; Sutherland et al., 2004; Aronson and Precht, 2001; Harvell et al., 2007; Miller et al., 2009; Brandt and McManus, 2009). Indeed, the Florida Reef Tract (FRT), which was dominated by *Acropora palmata* and *Acropora cervicornis*, and often had 30% coral cover until the 1970s/80s (Dustan and Halas, 1987; Porter and Meier, 1992), is now dominated by bare substrate, octocorals, and macroalgae with only approximately 5% stony coral cover remaining (Ruzicka et al., 2013). The loss of the branching Acroporid species was attributed primarily to a disease outbreak, termed white band disease (Aronson and Precht, 2001), but several other threats such as habitat reduction, eutrophication, overfishing, hurricanes, and bleaching likely all contributed to these species decline (Team, 2005). Subsequent losses of coral cover within the region were often linked to additional disease incidences and repeated regional coral bleaching events as a result of global climate change (Kuta and Richardson, 1996; Richardson et al., 1998; Sutherland et al., 2004; Gardner et al., 2003; Aronson and Precht, 2006; Kuffner et al., 2015; Manzello, 2015). A novel coral disease outbreak, termed Stony Coral Tissue Loss Disease (SCTLD), is now threatening the last vestiges of coral throughout the Florida Reef Tract (FRT).

SCTLD was first documented off the coast of Miami-Dade County in the summer of 2014 by Precht et al. (2016) and has since spread throughout the entire FRT with the exception of the Dry Tortugas. To date, SCTLD has been observed affecting over 20 different stony corals species. A case definition of SCTLD has been compiled to describe the visual appearance and ecology of SCTLD (NOAA, 2018). Briefly, the gross morphology of SCTLD is described as focal or multifocal, with locally extensive to diffuse areas of acute to subacute tissue loss distributed basally, peripherally, or both. In some cases, tissues bordering areas of chronic tissue loss show indistinct bands (1–5 cm) of pallor, progressing to normal pigmentation away from the denuded skeleton. There is also a range in coral susceptibility to SCTLD, with species categorized as highly susceptible (e.g., *Dendrogyra cylindrus*, *Dichocoenia stokesii*, *Meandrina meandrites*), moderately susceptible (e.g., *Orbicella* spp., *Montastraea cavernosa*, *Siderastrea siderea*, *Stephanocoenia intersepta*), or tolerant (e.g., *Porites* spp., *Acropora* spp.). Unfortunately, SCTLD has not remained isolated in the FRT and has now been recorded in Mexico (Alvarez-Filip et al., 2019), the US Virgin Islands (Blondeau et al., 2020) and several other locations around the Caribbean (Kramer et al., 2019). The continued persistence of the outbreak, the high number of species affected, and the large geographical range of reports consistent with the case definition suggests that SCTLD is the largest coral disease outbreak on record

Large-scale spatial epidemiologic analyses showed that the reefs in Florida with SCTLD are clustered, supporting a contagious mode of transmission (Muller et al., 2020). Similarly, aquaria-based experiments indicate SCTLD can be transmitted through direct contact or indirectly through the water column (Aeby et al., 2019) suggesting water can be a SCTLD vector, at least within a controlled setting. The initial exponential increase in spread among reefs from the disease epicenter (Precht et al., 2016) and the persistent subsequent linear rate of spread of SCTLD (Muller et al., 2020), north along South Florida reefs and south into the Florida Keys, indicates that water currents may play a role in disease transmission. Furthermore, the rate of spread, estimated at 100 m per day, suggests surface currents are likely too fast to have spread SCTLD within the region. These results imply that the bottom boundary layer, which is significantly slower than surface currents, may be the vertical location in which transmission occurs (Aeby et al., 2019). However, to date, there have been no attempt to link local hydrodynamic modeling efforts with the spatio-temporal dynamics of SCTLD in Florida.

64 Estimating the transport of the disease causative agent from reef to reef by currents cannot be performed  
65 empirically. However, experimentally-calibrated numerical models that simulate currents can provide a  
66 realistic picture of the dispersal of disease agents. Nonetheless, accurately modeling water circulation at  
67 the spatial scales that affect this dispersal remains a key challenge, as small-scale flow features such as  
68 recirculation eddies around reefs and islands strongly impact exchanges between reefs (Wolanski, 1994;  
69 Burgess et al., 2007; Figueiredo et al., 2013). In this context, models that can explicitly simulate flow  
70 features down to the reef scale are needed. This represents a spatial resolution of the order of 100-1,000 m  
71 in dense reef systems. As of today, most regional ocean models using traditional numerical methods cannot  
72 achieve such resolution because of the computational resources it requires. To our knowledge, the best  
73 resolution currently available among these models in the FRT is  $\sim 900$  m with the FKEYS-HYCOM model  
74 that has been developed for the Florida Keys region (Kourafalou and Kang, 2012; Sponaugle et al., 2012;  
75 Vaz et al., 2016). Unstructured-mesh ocean models offer a potential solution to this resolution issue by  
76 locally increasing the model resolution close to reefs and islands (Lambrechts et al., 2008; Thomas et al.,  
77 2014, 2015), in order to focus the computational resources where they are most needed. High resolution  
78 bio-physical dispersal models can be used to build the potential connectivity between reefs and therefore  
79 approximate exchanges between colonies in the complex topography of the coral reef systems (Frys et al.,  
80 2020).

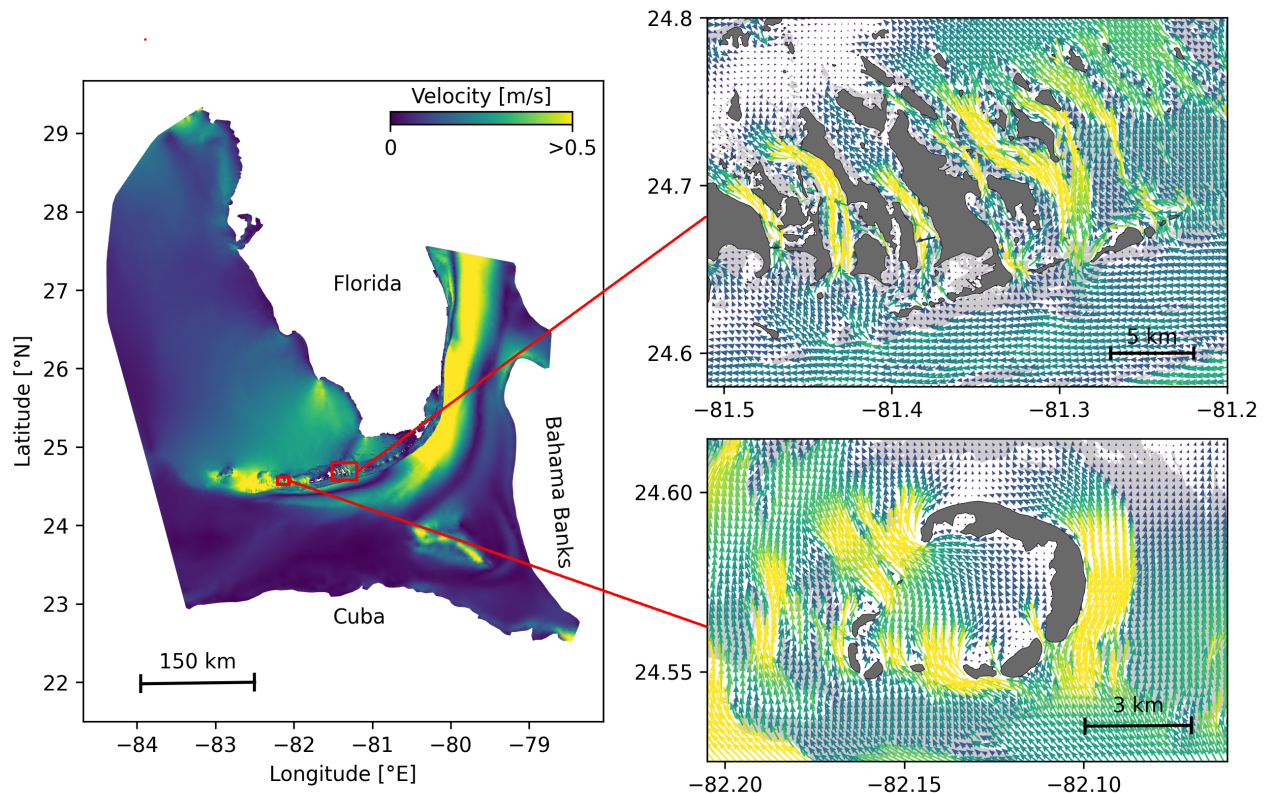
81 Marine diseases differ significantly from better studied terrestrial diseases, namely due to the potential  
82 for long environmental residence times, during which pathogens may survive and disperse through the  
83 water (Harvell et al., 2007; Sokolow et al., 2009). Several recent studies have attempted to adapt traditional  
84 epidemic models (Susceptible-Infectious-Recovered, or SIR models) to coral reef systems (Sokolow et al.,  
85 2009; Bidegain et al., 2016a,b). Novel approaches have included developing pathogen pools (Bidegain  
86 et al., 2016a,b), and to model at the metapopulation scale, rather than at the scale of coral holobionts  
87 (Sokolow et al., 2009). Both of these approaches are attempting to address the same issue: infection  
88 occurs between patches of entirely sessile animals, through the dispersal of pathogen(s). Thus, there  
89 are internal within-patch disease dynamics and metapopulation-scale between-patch dynamics occurring  
90 simultaneously. The epidemic model developed in this study utilizes the same basic architecture of previous  
91 coral reef SIR models, but rather than assume pathogen pools (e.g. Bidegain et al. (2016a,b)) or ignore  
92 internal patch dynamics (e.g. Sokolow et al. (2009)), we have modeled both within-patch disease dynamics  
93 and the dispersal of pathogens explicitly using potential connectivity networks.

94 The objective of this study is to deduce the probable propagation mechanism of the SCTLD throughout  
95 the FRT by developing an experimentally-calibrated epidemio-hydrodynamic model. With a resolution of  
96 about 100 m, this model can capture potential exchanges of disease-carrying material, further denominated  
97 as "infectious" in our modeling framework, between reefs that would be ignored by coarser models. By  
98 reproducing the observed spread of disease between 1st May 2018 and 1st April 2018, we provide insight  
99 on the characteristics of the disease agent and its vector. Ultimately, our model, coupled with lab and field  
100 works, would support the management of the epidemic and the identification of its causative agent.

## 2 METHODS

### 101 2.1 Modeling reef connectivity

102 In this study, we focus on the exchanges of infectious material between coral reefs driven by ocean  
103 currents, which therefore have to be accurately simulated. An ocean model should provide a realistic  
104 large-scale circulation while also resolving small-scale flow features down to the scale of individual reefs.



**Figure 1.** Model computational domain and close-up views of the mesh with snapshots of the currents on May, 25 2018 at 00:00, for the Marquesas Keys (bottom) and the Lower Keys (top). This illustrates the benefits of unstructured meshes to represent the fine-scale details of the topography and hence simulate currents down to the scale of individual reefs (shown in light grey) and islands (shown in darker grey).

In this context, we use the unstructured-mesh depth-integrated coastal ocean model SLIM<sup>1</sup> to simulate ocean currents over an area that includes the FRT but also the Florida Strait and part of the Gulf of Mexico (Fig. 1). By using an unstructured mesh, we can increase the model resolution only over the FRT and hence concentrate computational resources where they are most needed. SLIM has already been successfully applied in complex coastal systems such as the Great Barrier Reef (Lambrechts et al., 2008; Thomas et al., 2014) and is well suited to shallow-water flows. Details of the model formulation and validation are provided in Frys et al. (2020).

The mesh resolution depends only on the distance to the coast, but we distinguish between the coastlines along the FRT where we impose a maximum resolution of 100 m and the other coastlines along which the finest resolution is 2500 m. The mesh has been generated with the open-source mesh generator GMSH (Geuzaine and Remacle, 2009) and has about  $7 \times 10^5$  elements. The coarsest elements, far away from the FRT, have a characteristic length size of about 10 km. An illustration of ocean currents simulated on that mesh are shown in Fig. 1. It shows how a 100-m spatial resolution allows us to simulate fine-scale details of the flow, such as recirculation eddies and currents within the dense reef system in the Lower Keys that consist of many individual reefs with narrow passages in between.

The simulated currents can then be used to model dispersal of infectious material throughout the FRT. In this study, 3 types of potential vector carrying the disease causative agent were considered: positively

<sup>1</sup> <https://www.slim-ocean.be>



buoyant (e.g. mucus and surfactant), neutrally buoyant (e.g. fines, pelagic organisms) and negatively buoyant (e.g. sediments, composites, demersal organisms). As SLIM is a depth-averaged model, the mean currents it generates are well suited to model the dispersal of neutrally buoyant material remaining within the water column. However, these currents must be modified to correctly represent the dynamics of material evolving in the surface and bottom boundary layers. Therefore, surface current response to winds is estimated by adding 1.5% of the wind speed to SLIM currents with a stress-layer veering angle of 45° to the right for positively buoyant particles. Such parameterization is shown to be an accurate approximation of wave-induced Stokes drift and quasi-Eulerian surface currents by Ardhuin et al. (2009). For negatively buoyant material, on the other hand, bottom currents are obtained by taking 60% of SLIM currents velocity with a veering angle of 15° to the left. This is an approximation based on observations of bottom currents and whole water column current profiles in the shallow waters (<15 m) of Hawk Channel in the middle Florida Keys by Smith (2009), as well as observations obtained during the Atlantic Ocean Acidification Testbed project (Gramer, pers. comm.). It is also consistent with the theory of current veering in the bottom Ekman layer, albeit that was previously observed in deeper (30-90 m) coastal waters, e.g., by Perlin et al. (2007) and Kundu (1976).

Using these three velocity fields, virtual particles are then released on all the reefs composing the FRT to model the dispersal of infected material carrying the disease causative agent. The locations of the reefs of Florida are extracted from the "coral reefs and hardbottom" layer of the Unified Florida Reef Tract Map (FWC-FWRI, 2017). The polygons of this reef map are then further divided into 500 m × 500 m squares in order to track the propagation of the disease with a finer geographical resolution, generating a total of 16,823 polygons. At the beginning of each simulated month and for each type of current, a total of about  $1.5 \times 10^6$  particles are released over all the reef polygons. These particles have a state composed of their polygon of origin as well as their mass, that they loose at a constant rate  $\gamma$  as they are moved by surface, mean or bottom currents. In this study, the value of  $\gamma$  is chosen so that particles have a half life of 30 days. When the particles are brought over a reef polygon by currents, the amount of infected mass that lands on the polygon is recorded in monthly potential connectivity matrices, whose entries are denoted  $C_{ij}$ . The matrix rows correspond to the source reefs and the columns correspond to the destination reefs. Hence  $C_{ij}$  represents the mass of diseased material originating from sub-reef  $i$  that has settled on sub-reef  $j$ . This matrix is then normalized by dividing each of its rows  $i$  by the total mass of particles released on polygon  $i$  in order to obtain the normalized potential connectivity matrix  $\tilde{C}$ , whose entry  $\tilde{C}_{ij}$  gives the probability that disease agents produced on sub-reef  $i$  settle on sub-reef  $j$ . Connectivity matrices are computed for each type of current and for each month of the simulated period.

These connectivity matrices can be more easily handled by interpreting them as large graphs whose vertices are sub-reefs and whose edges represent connectivity pathways. They can then be analyzed using graph theory tools. In this study, four potential connectivity measures are used to interpret the monthly computed graphs. These indicators are described in Table 1. The first indicator is the weighted connectivity length (WCL), that gives the average dispersal distance from origin to destination for material produced on a given reef. The weighted connectivity of reef polygon  $i$  writes:

$$\text{WCL}_i = \frac{\sum_j \tilde{C}_{ij} L_{ij}}{\sum_j \tilde{C}_{ij}} \quad (1)$$

where  $L_{ij}$  is the distance between origin reef  $i$  and destination reef  $j$ . Another measure of the spreading potential of reef  $j$  is its out-degree, *i.e.* the product of the number of connections originating from reef  $j$  by the quantity of infectious material it sends to the network. This indicator is obtained by computing

Indicators	Description	What it shows
Weighted connectivity length (WCL)	Average dispersal distance from origin to destination reef for all disease agents released over a reef	Average distance at which a reef can send disease agents
Out-degree	Number of out-going connections originating from a given reef multiplied by the total mass exchanged	Potential for a reef to spread the disease
Fraction exchanged	Fraction of infectious material produced on a given reef that settles on other reefs	Success rate of potential disease spread
Self recruitment	Fraction of infectious material settling on a given reef that has been released on the same reef	Potential for disease to settle on reef

**Table 1.** Indicators used to analyze the modeled exchanges of infected material for each considered type of currents and for each simulated month

the number of non-zero entries of row  $i$  in the potential connectivity matrix  $C$  and multiplying it with  $\sum_j C_{ij}$ . The information given by the out-degree is complemented by the fraction of infectious material produced on reef  $i$  that successfully settles on a reef, called the fraction exchanged of reef  $i$ . This indicator is given by  $\sum_j \tilde{C}_{ij}$ . Finally, the isolation of reef  $i$  in the network is given by its self recruitment, *i.e.* the fraction of infectious material settling on reef  $i$  that originates from reef  $i$ , computed by  $C_{ii} / \sum_j C_{ji}$ . A large self-recruitment value indicates that little infectious material produced elsewhere settles on the reef and thus that it is isolated from the rest of the network.

## 2.2 Epidemiological modeling

### 2.2.1 Model equations

The spread of the SCTLD throughout the FRT is simulated using a connectivity-based Kermack and McKendrick (1927) SIR model. SIR models are among the most standard epidemiological models. They divide individuals into three compartments: susceptible (S), infectious (I) and removed (R). When affected by the disease, susceptible individuals become infectious and infect other susceptible individuals until they are removed from the system, either through recovery or death. Such models usually rely on the hypothesis of an homogeneous, well-mixed population. To account for the spatial heterogeneity of the FRT, the basic SIR formulation is here modified by considering the fractions of susceptible ( $S_j$ ), infectious ( $I_j$ ) and removed ( $R_j$ ) corals of each sub-reef  $j$ . In this epidemiological model, individual reefs interact through the exchange of infectious material as represented by the different connectivity matrices. For each sub-reef  $j$  and at any time, the following relations hold:  $0 \leq S_j, I_j, R_j \leq 1$  and  $S_j + I_j + R_j = 1$ . The evolution of these fractions through time is governed by the following equations:

$$\begin{aligned}
 \frac{dS_j}{dt} &= -\beta \sum_i \frac{A_i}{A_j} I_i \tilde{C}_{ij} S_j - \beta'(I_j) S_j I_j \\
 \frac{dI_j}{dt} &= \beta \sum_i \frac{A_i}{A_j} I_i \tilde{C}_{ij} S_j + \beta'(I_j) S_j I_j - \sigma I_j \\
 \frac{dR_j}{dt} &= \sigma I_j
 \end{aligned} \tag{2}$$

where  $\tilde{C}_{ij}$  is the entry corresponding to reef pair  $(i, j)$  of the normalized potential connectivity matrix [-],  $A_i$  is the area of reef polygon  $i$  [km<sup>2</sup>],  $\sigma$  is the mortality rate [day<sup>-1</sup>], and  $\beta$  and  $\beta'(I_j)$  are the inter- and intra-reef disease transmission rates [day<sup>-1</sup>], respectively. In this model, infectious corals of reef  $i$  can infect corals of reef  $j$  if there is non-zero probability of infectious material exchange from reef  $i$  to reef  $j$ , given by  $\tilde{C}_{ij}$ . Moreover, to account for coral resistance to the disease, the intra-reef transmission function  $\beta'(I_j)$  has the shape of a smooth step function of the fraction of infectious corals  $I_j$  and writes:

$$\beta'(I_j) = \frac{\beta'_0}{2}(1 + \tanh[(I_j - I_0)/\tau]), \quad (3)$$

where  $I_0$  is a threshold on the infection population above which intra-reef transmission becomes significant, and  $\tau$  is a measure of the interval over which the transition from low to high transmission occurs. As long as the fraction of infectious corals on reef  $j$  is below  $I_0$ , the only infection mechanism taking place is connectivity-driven transmission at rate  $\beta$ . Once the threshold is approached, intra-reef transmission with rate  $\beta'_0$  is activated. A larger value of threshold  $I_0$  corresponds to a greater resistance of corals to the disease, and therefore a slower spread of the disease within reef  $j$ . Coral birth and natural (*i.e.* non SCTL-related) death rates are not taken into account in this model, which amounts to assume that they balance each other. For this study the same values were used for  $\beta$  and  $\beta'_0$ .

## 2.2.2 Calibration

Transmission and removal parameters  $\sigma$  and  $\beta'_0$  are fitted to disease prevalence observations averaged over all colonies from 6 permanent monitoring sites in the Lower Keys in order to accurately simulate the temporal evolution of  $S_j$ ,  $I_j$ ,  $R_j$  on each infected reef polygon. Three focal reef sites were established in the lower Florida Keys, one offshore (Acer 17/18), one mid-channel (Wonderland), and one nearshore reef (N. Birthday). Sites were established in May 2018, when all colonies appeared healthy. Within each site, two 10 m × 10 m quadrats were established. They were generally set up from east to west although N. Birthday was established with one quadrat further north of the other two to better capture coral cover in the site. All coral colonies > 10 cm in size were mapped using SCUBA. Each coral was given an  $(x, y)$  coordinate, identified to species, and maximum diameter was noted. After the initial data collection surveys, each site was visited every two to three weeks for rapid assessments to determine whether SCTL was present. During these site visits, two divers conducted a visual assessment at each of the 6 quadrats. Disease was first observed in early October 2018. Detailed surveys were conducted every two to four weeks until December 2019. During the surveys, each individual coral was visually assessed for signs of SCTL, including discoloration and tissue loss. Prevalence of diseased, apparently healthy, and dead were assessed for each time period. To relate our model framework to the compiled data, Eqs. 2 are simplified to a single-reef SIR model:

$$\begin{aligned} \frac{dS}{dt} &= -\beta'_0 SI \\ \frac{dI}{dt} &= \beta'_0 SI - \sigma I \\ \frac{dR}{dt} &= \sigma I \end{aligned} \quad (4)$$

Due to the low values of the entries in the normalized connectivity matrix  $\tilde{C}_{ij}$ , intra-reef transmission, when activated, is the dominant infection mechanism of Eqs. 2. Consequently, Eqs. 4 give a reasonable approximation of the evolution of the disease on sub-reefs for which  $I_j > I_0$ . Using this approximation, the ratio  $\beta'_0/\sigma$  is imposed by matching the modeled fraction of susceptible corals remaining after the disease

has vanished ( $S_\infty$ ) with observations. A standard property of a SIR model solution is indeed that

$$S_\infty - \frac{\sigma}{\beta'_0} \log(S_\infty/S_0) = 1 \quad (5)$$

where the initial fraction of susceptible corals ( $S_0$ ) is taken equal to  $1 - I_0$  (see for instance Murray (2007)). In the framework of Eqs. 4, the ratio  $\beta'_0/\sigma$  gives the value of the basic reproduction number  $R_0$ , defined as the average number of secondary cases produced by one infected individual introduced into a population of susceptible individuals (Keeling and Rohani, 2007). This number is used in epidemiological models to determine whether an emerging infectious disease can spread in a population ( $R_0 > 1$ ) or not ( $R_0 < 1$ ). The obtained basic reproduction number is then used to express  $\sigma$  in terms of  $\beta'_0$  and calibrate its value in order to reproduce as well as possible the temporal evolution of the colonies-averaged susceptible population shown in Fig. 5.

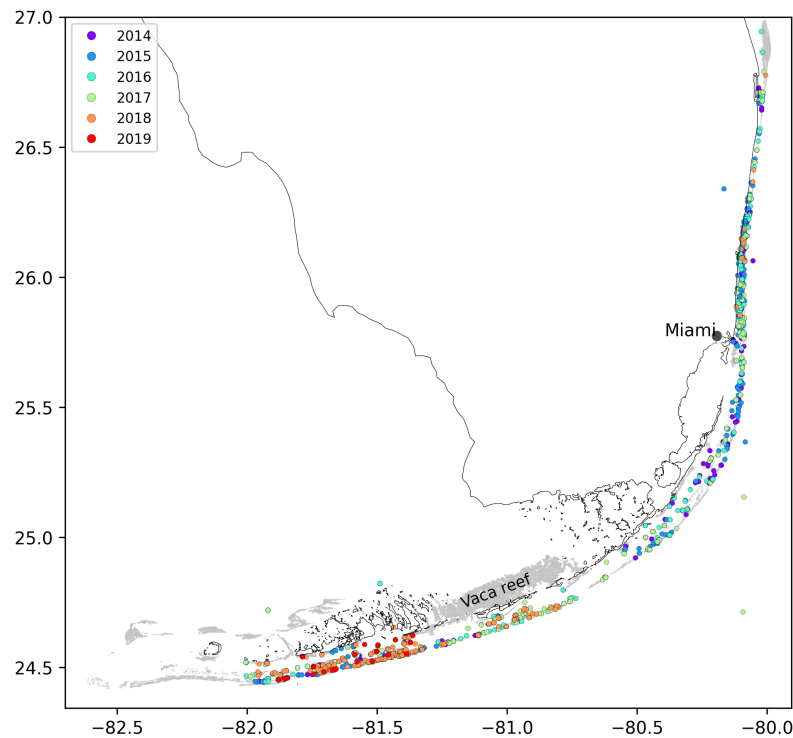
### 2.2.3 Initialization

In order to solve Eqs. 2, initial conditions are needed, *i.e.* fractions of susceptible, infectious and recovered corals at the beginning of the simulated period. This information is constructed based on 9 different field-collected datasets: (i) Coral Reef Evaluation and Monitoring Project (CREMP; 2014–2017), (ii) CREMP Presence/Absence Data (CREMP P\_A; 2016–2017), (iii) Southeast Florida Coral Reef Evaluation and Monitoring Project (SECREMP; 2014–2017), (iv) Florida Reef Resilience Program Disturbance Response Monitoring (FRRP; 2014–2017), (v) Hurricane Irma Rapid Reef Assessment (IRMA; 2017, Viehman et al. (2018)), (vi) the Southeast Florida Action Network citizen science program (SEAFAN; 2014–2017), and (vii) the Southern Coral Disease Margin field effort (2017 and 2018; Neely (2018)), (viii) Mote Marine Laboratory's Field operations data (2018-2019) and (ix) data compiled through Mote's citizen science BleachWatch program (2018). Every dataset provided data on the presence or absence of the SCTLD (or tissue loss consistent with the SCTLD case definition) within each survey. Some also provided detailed disease metrics such as the species affected and the disease prevalence, which was subsequently compiled into presence/absence of SCTLD data by surveyed site. The locations of these observations are shown in Fig. 2. Using this information, we first delineate an infected zone by constructing the concave hull of the points where the disease was observed before May 2018. The reefs infected prior to the beginning of our simulated period are then defined as the reefs located inside the constructed zone. The time of observed infection is then spatially interpolated on each reef of the infected zone by kriging with a Gaussian semivariogram using Python `pyKrig` module (Murphy, 2014). Assuming an initial state  $(S, I, R) = (1 - I_0, I_0, 0)$  when the disease was observed, the fractions of susceptible, infectious and removed corals on each reef of the infected zone on the 1st May 2018 is finally approximated using Eqs. 4. Reefs outside of the infected zone are initialized with an entirely susceptible population.

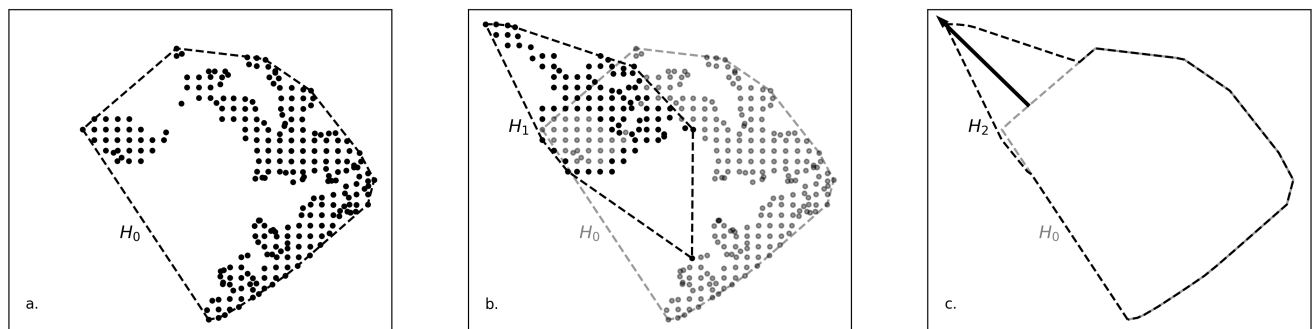
### 2.2.4 Computation of front speed

Muller et al. (2020) estimated the speed of the spreading STCLD epidemics at around 92 m/day in the southern section of the FRT. In order to assess our simulation results in regard to this value, we developed a methodology to compute the displacement of the disease front during a given time interval within our simulated period. First, the concave hull  $H_0$  of the infected polygons at the beginning of the time interval is delineated. Then the concave hull  $H_1$  of the polygons infected during the time interval is computed while the concave hull  $H_2$  is defined as the union of  $H_0$  and  $H_1$ . This methodology is illustrated in Fig. 3. The distance traveled by the disease front is then obtained by computing the maximum distance between





**Figure 2.** Locations of the disease observations between 2014 and 2019 recorded in the data sets used in this study



**Figure 3.** Method used to compute the disease front displacement during a simulated time interval. **a.** Concave hull  $H_0$  of the infected sub-reefs at the beginning of the simulated period. **b.** Concave hull  $H_1$  of the sub-reefs infected during the simulated time interval. **c.** Arrow showing the computed front displacement during simulated time interval between  $H_0$  and  $H_2$ , the union of  $H_0$  and  $H_1$ .

all pairs of points of  $H_0$  and  $H_2$ . The epidemics front speed is finally obtained by dividing the resulting distance by the number of days in the simulated time interval.

### 2.3 Transmission experiments

In parallel to this modeling study, laboratory-based transmission experiments of SCTLD were conducted by several independent groups for various end points including transmission dynamics and samples for molecular and histological analysis. Requests for transmission data were sent to members of the ‘Transmission’ sub-group of the Florida Disease Advisory Committee’s ‘Research’ working group as well as any other additional researchers that may have been conducting transmission studies on SCTLD. Data

that was requested and subsequently provided included the location, dates, and duration of the experiment, the species used as the diseased colony (donor of disease agents) and apparently healthy colony (exposed to disease agents), the number of successful transmissions as well as the incubation period following a contact with disease agents prior to disease signs. Additional information included the size of the colonies used in the experiment, the percent tissue loss of the diseased (donor) colony at beginning of the experiment, and whether the apparently healthy (exposed) fragment was touching the diseased colony or not.

The average probability of successful disease transmission was determined by taking the mean of the number of colonies exposed to the disease in each study divided by the total number of coral colonies exposed to diseased colonies. The ‘incubation’ period was identified as the average number of days after an apparently healthy coral colony was exposed to a diseased colony before visual disease signs occurred (i.e., active tissue loss). Only corals that eventually showed disease signs were integrated within the incubation period calculation.

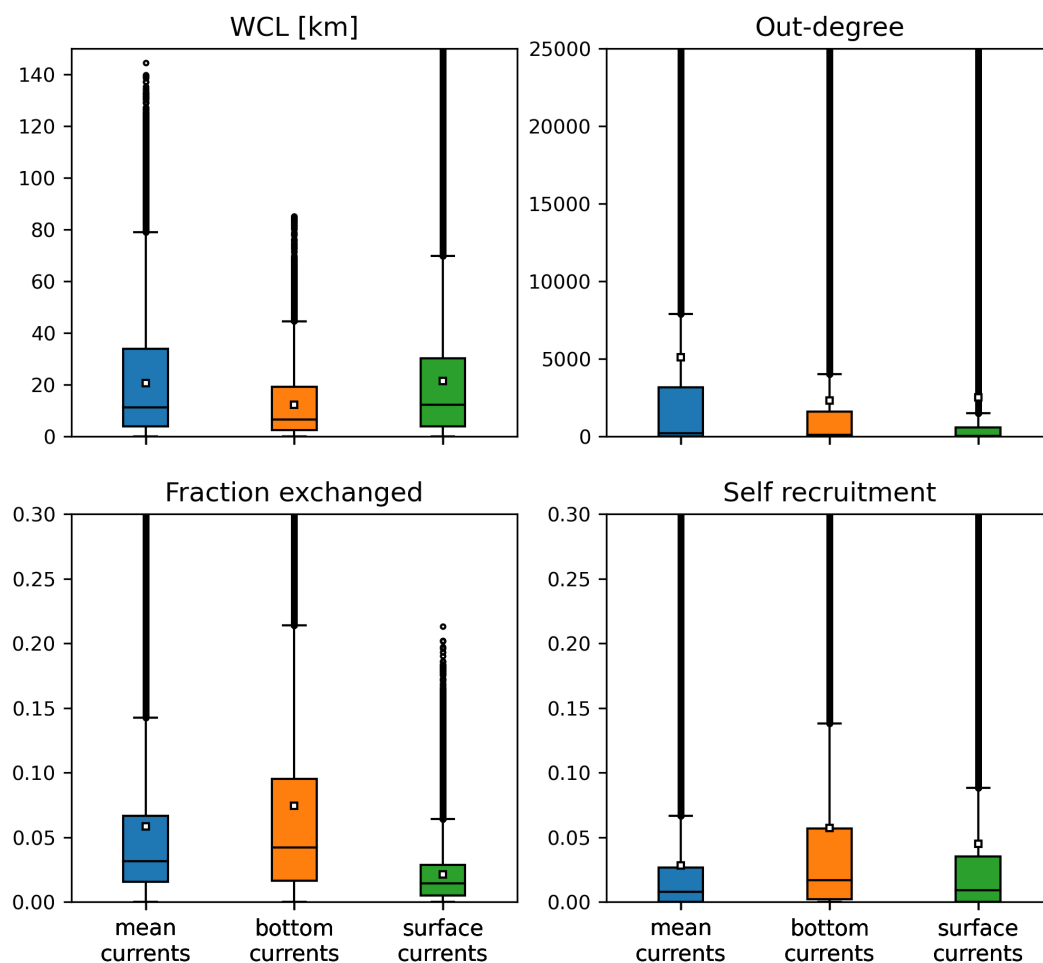
Data was provided from 8 different research groups representing 15 institutions and 19 total collaborators providing a total of 109 data points (see table 2 in appendix). After amalgamating the contributed data, the mean probability of transmission of SCTLD to an apparently healthy coral had a likelihood of approximately  $44.8 \pm 3.6$  %. The probability of transmission ranged from 0 to 100% depending on the experiment. Additionally, the time between exposure of an apparently healthy coral to a diseased coral and subsequently showing initial signs of tissue loss (i.e., incubation period) was  $9.7 \pm 1$  days.

### 3 RESULTS

#### 3.1 Exchanges of infected material

Among the three modes of transport, bottom currents exhibited the lowest propagation range as they generate the networks with the smallest weighted connectivity length (Fig. 4). However, infectious material transported by bottom currents has the largest settlement success rate as these currents generate the graphs with the largest fraction exchanged. Therefore, bottom currents tend to transport more infectious material on closer reefs compared to the two other modes of transport. Mean and surface currents, on the other hand showed similar spreading ranges with mean WCL of 20.63 km and 21.39 km respectively. However, the infectious material that surface currents transport have the weakest probability to successfully settle on reefs. Consequently, surface currents and bottom currents produce networks with similar mean out-degree, although surface currents have the potential to transport infectious material on larger distances. Nonetheless, networks have larger median out-degree with bottom currents than with surface currents, which suggests that surface currents have a lower spreading potential than bottom currents. As a result of their large WCL and fraction exchanged, barotropic currents on the other hand exhibit the largest mean out-degree, which indicates that they have the strongest dispersal potential.

Self recruitment gives the fraction of infectious material settling on a reef that was produced on the same reef. The greater its values, the more the reef is isolated from the rest of the network. Since infectious material is less likely to settle on isolated reefs, self recruitment measures the probability for the disease to reach a given reef, whereas all three other indicators inform on the reef spreading potential. Fig. 4 shows that the disease is more likely to settle on the reefs of networks generated by mean currents. This result is consistent with the values of the other connectivity measures, as reefs tend to be more strongly connected with mean currents. On the other hand, reefs are more isolated with bottom currents, as they produce the graphs with lowest WCL and out-degree. Finally, surface currents generate larger self recruitment values than mean currents as they exhibit the lowest fraction exchanged. Therefore, although bottom currents



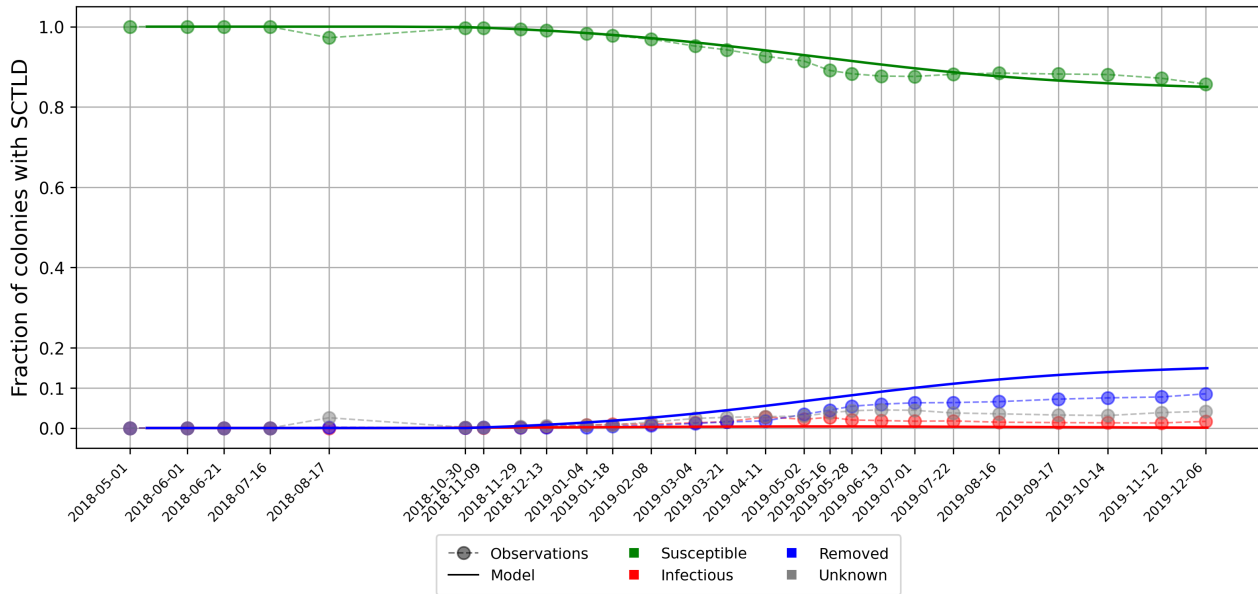
**Figure 4.** Distribution of the indicators derived from the monthly connectivity matrices computed for each type of current during our simulated period. Mean values are indicated by white squares

306 exhibit stronger spreading potential than surface currents, reefs are more sensitive to infection with surface  
 307 currents.

### 308 3.2 Epidemiological model results

309 As aggregated observations show a fraction of susceptible individuals of about 85% at the end of the  
 310 outbreak, a basic reproduction number  $R_0 = \beta'_0 / \sigma = 1.0345$  is found with Eq. 5. Using this ratio, best  
 311 fit to averaged disease prevalence observations is obtained with transmission rate  $\beta'_0 = \frac{1}{6.45} \text{ days}^{-1}$  and  
 312 mortality rate  $\sigma = \frac{1}{6.99} \text{ days}^{-1}$ . Comparison of the evolution of the state described by Eqs 4 results  
 313 with observations is shown in figure 5. Our model results accurately reproduce the observed fraction of  
 314 susceptible individuals on colonies through time. However, the modeled fraction of removed individuals  
 315 overestimates observations by about 5%. These discrepancies might be explained by the presence of  
 316 "Unknown" values in our data sets as well as the simplifying assumptions of SIR models. Since infection  
 317 and removal occur at very close rates, the instantaneous fraction of infectious individuals on the reefs  
 318 remains low through the outbreak, with a maximum value of about 0.4%.

319 Once the model calibrated, epidemio-hydrodynamic model simulations were performed from 1st May  
 320 2018 to 1st April 2019 for each type of currents and different values of the infection threshold  $I_0$ . Two

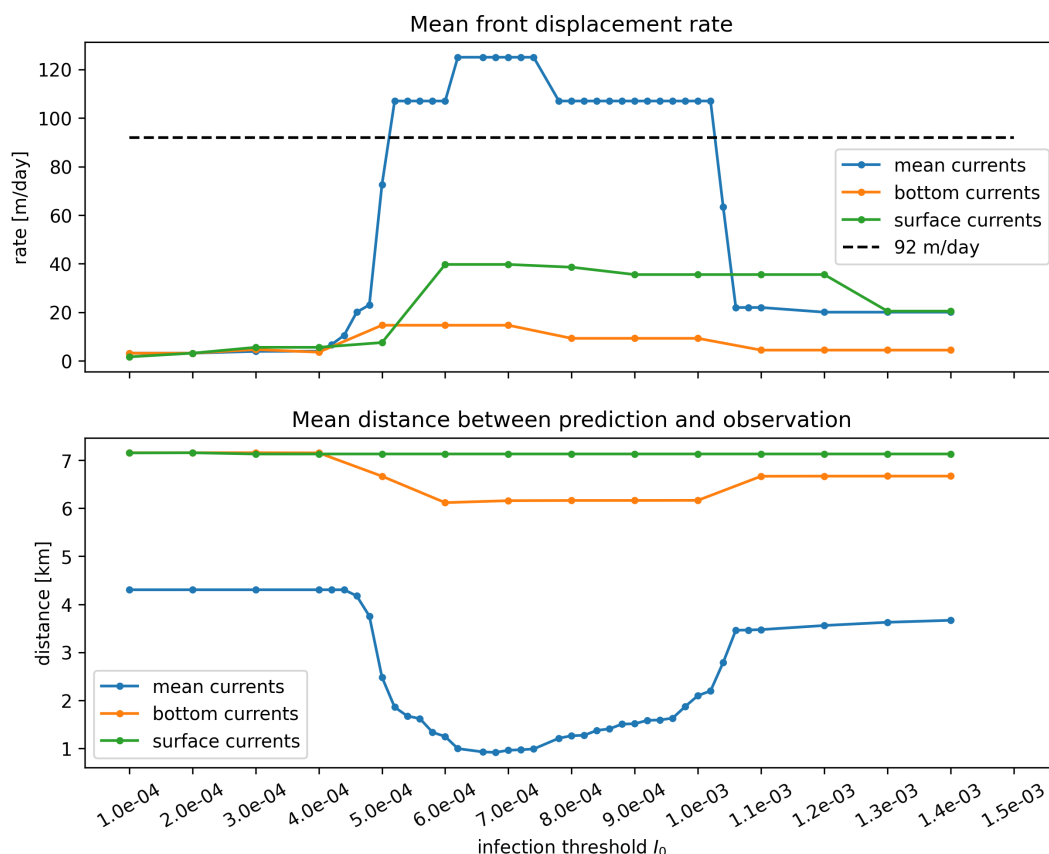


**Figure 5.** Disease prevalence averaged over all monitored sites over time as modeled by Eqs. 4 using calibrated transmission and removal parameters  $\beta'_0 = \frac{1}{6.45} \text{ days}^{-1}$  and  $\sigma = \frac{1}{6.99} \text{ days}^{-1}$ .

metrics were used to assess the accuracy of the model. First, the modeled front speed was compared to the reference rate of 92 m/day derived by Muller et al. (2020). Furthermore, we computed the mean of the distances between each point where SCTLD had been observed during our simulated period (extracted from the 2018-2019 data sets described in section 2.2.3) and the centroid of the closest reef polygon predicted to be infected by our model during the same period (Fig. 6). Bottom currents produced the slowest modeled disease propagation with a maximum front speed of  $\sim 20$  m/day, while simulations performed with surface currents spread the disease at a maximum speed of of about 60 m/day. However, surface currents tend to propagate the disease to the north, rather than westward, along the Florida Keys. This explains why bottom currents predict disease occurrence closer to field observations despite exhibiting slower front speed. Finally, mean barotropic currents outperform other types of current predictions regarding both criteria with a front speed of 107 m/day and a mean geographical accuracy of  $\sim 1.2$  km. This suggests that the disease agents of SCTLD may be transported within neutrally buoyant material driven from reef to reef inside the water column by mean currents.

Moreover, Fig. 6 shows a strong dependence of the model results to infection threshold  $I_0$ , that gives the fraction of infectious individual that colonies can withstand before exponential disease growth is triggered on the reef. Front speeds of both mean and bottom currents reach a plateau for values of infection threshold between  $I_0 = 0.05\%$  and  $I_0 = 0.1\%$ , while the minimal prediction error is reached around  $I_0 \approx 0.078\%$  with mean currents. For  $I_0 > 0.1\%$ , intra-reef infection is strongly impeded and populations of infectious individuals on infected reefs are not able to become sufficiently large to infect other colonies on reefs they are connected to. For values of  $I_0$  lower than  $0.05\%$  on the other hand, intra-reef infection dominates and coral population on infected reefs is removed too fast to efficiently spread the disease through the network. Since disease propagation throughout the FRT only occurs for fairly small values of  $I_0$  in our model, corals are expected to have low resistance to the causative agent of the SCTLD.

The results shown in Fig. 6 were obtained by removing the large reef located North to Vaca key, denoted Vaca reef in Frys et al. (2020), from our reef polygons. Preliminary simulations showed that this reef



**Figure 6.** Summary of epidemiological model simulations with calibrated transmission parameters. **Top:** Modeled disease front speed for each type of current with respect to intra-reef infection threshold  $I_0$ . **Bottom:** Mean distance between predicted infected reefs and observed disease points. These results show that mean barotropic currents outperform other modes of transport at reproducing the observed spread of the disease. The appearance of a plateau suggests that the modeled spread of the disease occurs for a well-define range of values of  $I_0$ .

has close to no impact on the modeled spread of the disease to the rest of the FRT, as it sends very little infectious material to southerly and easterly neighboring reefs. Moreover, Vaca reef has a low coral coverage (0 – 10%), which strongly impedes disease spread on the reef. However, as coral coverage is not taken into account in our epidemiological model, propagation of the disease on the reef was overestimated. This led to unrealistically strong modeled front speed variations due to the large size of the reef. Consequently, and in the absence of SCTLD observations on Vaca reef, it has been removed from our reef polygons in order to avoid overestimating the front speed.

## 4 DISCUSSION AND CONCLUSIONS

We have developed an epidemio-hydrodynamic model to simulate the spread of the SCTLD through the entire FRT. Calibrating our model with colonies-averaged prevalence observations, we estimated the species-averaged reproduction number  $R_0$  to be slightly larger than one. Our model simulations suggest that only the barotropic currents are able to reproduce the observed spread of the disease. Bottom current do not spread infectious material far enough while surface currents do not allow infectious material to spend enough time over the reefs to strongly affect them. The causative agent of the SCTLD is therefore expected to be transported within neutrally buoyant particles inside the water column. With this mode of



transport, the propagation of the disease from reef to reef only occurs for a well-defined range of values of the infection threshold  $I_0$ . This threshold is defined as the fraction of all reef colonies that have to be infected to trigger a rapid spread of the disease over the entire reef. Our results suggest that this occurs as soon as 0.05 – 0.1% of colonies are infected. On average, corals are thus expected to have a low resistance to the SCTLD.

After calibration, we estimated the species-averaged basic reproduction number  $\beta'_0/\sigma$  to be equal to 1.0835. This value being close to 1, modeled infectious individuals are removed from the system almost as fast as susceptible individuals get infected. This causes the fraction of infectious corals on the reefs to remain pretty low (*i.e.*  $\leq 0.4\%$ ) through time. This suggests that only a small fraction of the colony causes the disease to spread on the reef during the outbreak. The observation-based species-averaged transmission period of 6.45 days used in this model seems to be a reasonable estimation of the disease transmission dynamics as it is of the same order of magnitude as the experimentally-derived mean incubation period of 9.7 days. The difference between the two values can be explained by field measurement uncertainties as well as the inability to perfectly mimic field conditions in laboratory. In this study, the same values were used for inter- and intra-reef rates  $\beta$  and  $\beta'_0$ . This implies that the infectiousness of the causative agent is not reduced during its journey from reef to reef. However, to assess the impact of this assumption, epidemiological model simulations were performed with  $\beta = \beta'_0/2$ . The resulting disease front speeds did not exceed 20 m/day. This strong decrease can be explained by the interplay between inter- and intra-reef infection. Reducing inter-reef transmission rates decreases the fraction of infectious corals on reefs attained by infectious material, which in turn reduces the amount of infectious material sent to the rest of the network. This suggests that, to reproduce the observed spread, inter- and intra-reef transmission rates must have similar magnitude, *i.e.* that the causative agent is almost not degraded while traveling from reef to reef.

The fact that mean barotropic currents outperform the other modes of transport can be explained by considering the trajectories of the particles used to model the transport of the disease causative agent. Due to the impact of winds on positively buoyant material, particles driven by surface currents are likely to be blown away from the reefs. Moreover, even when winds are pushing particles along the reef line, these particles spend less time over the same region than particles driven by mean and bottom currents. Smaller amounts of particle mass will therefore settle on reef polygons, leading to lower entries of the potential connectivity matrix, *i.e.* lower exchange of infectious material between reefs. Hence, despite being able to transport the disease over greater distances, surface currents are less likely to drive the propagation of the disease. Particles driven by bottom currents, on the other hand, remain longer over the same region, producing larger entries of the potential connectivity matrix. Due to these large exchange probabilities between reefs, bottom currents are better at propagating the disease (Fig. 6). Nevertheless, bottom currents being slower, exchanges of infectious material occur on a limited geographical range. Mean barotropic currents, that carry particles on greater distances while allowing for sufficiently large amounts of infected mass to settle on reef polygons, are thus best suited to propagate the disease (Fig. 6).

Since mean currents are the only mode of transport that successfully reproduces the observed propagation speed of the disease in our model, the disease causative agent is expected to be transported within neutrally buoyant material inside the water column. Current-driven propagation seems reasonable as water-borne transmission is an important spreading mechanism for multiple coral diseases, including white band disease, white plague disease, white pox disease, white syndrome disease, *Porites* ulcerative white spots diseases, skeletal eroding band disease (Shore and Caldwell, 2019). The causative agent might for instance be transported within fine sediments such as silt, as suggested by Rosales et al. (2020). Such sediments are

easily eroded in shallow areas around coral reefs and would therefore be mostly transported inside the water column by mean barotropic currents. This hypothesis might be tested by adapting the deposition rate  $\gamma$  used in our experiments to be consistent with the sedimentation rate of silt. However, such modification of  $\gamma$  would alter the entries of our potential connectivity matrices. Nonetheless, the sensitivity of the connectivity matrices to the value of  $\gamma$  has been briefly assessed by generating new matrices using particles with a half-life of 15 days ( $\gamma$  increased by a factor two). Although these matrices exhibited stronger short-range connectivity, the impact on connectivity indicator values remained limited ( $< 10\%$ ). This suggests that the main results of this study would remain valid for larger deposition rates.

Coral resistance to the SCTLD is represented by parameter  $I_0$ , defined as the maximum fraction of the colony that can get infected without causing the disease to spread to the rest of the colony. The plateau shown in Fig. 6 highlights the impact of this parameter on the modeled propagation of the disease. On the one hand, when corals are strongly susceptible to the disease, infectious individuals are removed from the system too fast to become sustainable sources of infectious material in the network. On the other hand, if corals are weakly susceptible to the disease, very few corals get infected and the disease barely propagates. Our simulations suggest that this value must be fairly low (around 0.01%) in order to successfully spread the disease throughout the FRT. This seems to imply that susceptible coral species have very weak defense mechanisms against the causative agent of the disease.

As with any modeling study, it is important to understand the assumptions on which the model is based. Here, we have used a 2D barotropic ocean model forced by the 3D model HYCOM (Chassignet et al., 2007) in order to indirectly represent baroclinic phenomena. Such model is well suited to simulate the fate of neutrally-buoyant material in shallow regions. However, as depth-averaged currents do not accurately approximate the motion of particles in the bottom and surface layers, they have been modified to simulate the exchanges of negatively and positively buoyant material. Surface current response to wind parameterization is based on the results of Ardhuin et al. (2009), consistent with observations. In this study, measured surface currents are shown to be in the order of 1.0% – 1.8% of the wind speed, in a direction  $10^\circ - 40^\circ$  to the right of the wind. Moreover, the norm and veering angle used to parameterize bottom currents are expected to be reasonably accurate approximations as they are consistent with both Ekman theory and current observations in the Florida Keys (Perlin et al., 2007; Kundu, 1976; Smith, 2009). Although such estimation of surface and bottom currents is disputable, using a 2D model allows for reef-scale resolution throughout the whole FRT. Such high-resolution allows us to explicitly represent recirculation eddies around islands and reefs, that significantly impact the weighted connectivity length as well as the local retention on the reefs.

The appearance of an interval of optimal values of threshold  $I_0$  for the propagation of the disease in our results highlights the impact of coral resistance on the spread of SCTLD through the FRT. Therefore, a further step in our modeling approach would be further dividing coral populations of our polygons into highly susceptible (e.g. *Dichocoenia stokesii*, *Meandrina meandrites*), intermediately susceptible (e.g. *Orbicella faveolata*, *Montastrea cavernosa*), and weakly susceptible (e.g. *Acropora Palmata*, *Acropora cervicornis*) sub-populations. The fractions of susceptible, infectious and removed individuals within these sub-populations would then be modeled with specific transmission ( $\beta$ ,  $\beta'_0$ ) and removal ( $\sigma$ ) rates as well as specific infection thresholds  $I_0$ . Such approach would however require a fine knowledge of the distribution of the different coral species throughout the FRT. This knowledge about coral coverage could also be used to avoid overestimation of the front propagation, as in the case of Vaca reef.

Despite the limitations of its current formulation, we believe that our model brings unprecedented perspectives on the propagation mechanism of the SCTLD through the FRT. Using a reef-scale spatial

resolution, we determined the most probable mode of transport for the vector of the disease agent and deduced its species-averaged reproduction number based on prevalence observations. Besides, our model formulation provides a framework to quantify coral resistance to the disease. As our model results are continuous through time, they can exhibit the variability of the propagation of the SCTLD through time and therefore bring additional insight to observation data. This study therefore provides much-needed complementary insight on the identification of the causative agent of the SCTLD and the management of the crisis it generates. Furthermore, our modeling approach could be applied to other affected areas of the Caribbean, where there is still time to perform active management of the disease.

## APPENDIX

### Transmission data contributors

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Laura Mydlarz	University of Texas - Arlington
Dan Holstein	Louisiana State University
Amy Apprill	Woods Hole Oceanographic Institute
Tyler Smith	University of the Virgin Islands
Sonora Meiling*	University of the Virgin Islands

**Table 2.** Data contributors to the transmission experiments described in section 2.3, to which the calibrated model parameters were compared. Contributors highlighted with "\*" conducted the Data Sharing

## CONFLICT OF INTEREST STATEMENT

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

## AUTHOR CONTRIBUTIONS

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## REFERENCES

- Aeby, G., Ushijima, B., Campbell, J. E., Jones, S., Williams, G., Meyer, J. L., et al. (2019). Pathogenesis of a tissue loss disease affecting multiple species of corals along the Florida Reef Tract. *Frontiers in Marine Science* 6, 678
- Alvarez-Filip, L., Estrada-Saldívar, N., Pérez-Cervantes, E., Molina-Hernández, A., and González-Barrios, F. J. (2019). A rapid spread of the stony coral tissue loss disease outbreak in the Mexican Caribbean. *PeerJ* 7, e8069
- Ardhuin, F., Marié, L., Rasclé, N., Forget, P., and Roland, A. (2009). Observation and estimation of Lagrangian, Stokes, and Eulerian currents induced by wind and waves at the sea surface. *Journal of Physical Oceanography* 39, 2820–2838
- Aronson, R. B. and Precht, W. F. (2001). White-band disease and the changing face of Caribbean coral reefs. In *The ecology and etiology of newly emerging marine diseases* (Springer). 25–38
- Aronson, R. B. and Precht, W. F. (2006). Conservation, precaution, and Caribbean reefs. *Coral reefs* 25, 441–450
- Bidegain, G., Powell, E., Klinck, J., Ben-Horin, T., and Hofmann, E. (2016a). Microparasitic disease dynamics in benthic suspension feeders: infective dose, non-focal hosts, and particle diffusion. *Ecological Modelling* 328, 44–61
- Bidegain, G., Powell, E. N., Klinck, J. M., Ben-Horin, T., and Hofmann, E. E. (2016b). Marine infectious disease dynamics and outbreak thresholds: contact transmission, pandemic infection, and the potential role of filter feeders. *Ecosphere* 7, e01286
- Blondeau, J., Brandt, M., Donovan, C., Eakin, M., Edwards, K., Edwards, K., et al. (2020). Coral reef condition: A status report for the US Virgin Islands
- Brandt, M. E. and McManus, J. W. (2009). Dynamics and impact of the coral disease white plague: insights from a simulation model. *Diseases of aquatic organisms* 87, 117–133
- Burgess, S. C., Kingsford, M. J., and Black, K. P. (2007). Influence of tidal eddies and wind on the distribution of presettlement fishes around One Tree Island, Great Barrier Reef. *Marine Ecology Progress Series* 341, 233–242
- Chassignet, E. P., Hurlburt, H. E., Smedstad, O. M., Halliwell, G. R., Hogan, P. J., Wallcraft, A. J., et al. (2007). The HYCOM (hybrid coordinate ocean model) data assimilative system. *Journal of Marine Systems* 65, 60–83
- Dustan, P. and Halas, J. C. (1987). Changes in the reef-coral community of Carysfort Reef, Key Largo, Florida: 1974 to 1982. *Coral Reefs* 6, 91–106
- Figueiredo, J., Baird, A. H., and Connolly, S. R. (2013). Synthesizing larval competence dynamics and reef-scale retention reveals a high potential for self-recruitment in corals. *Ecology* 94, 650–659
- Frys, C., Saint-Amand, A., Le Hénaff, M., Figueiredo, J., Kuba, A., Walker, B., et al. (2020). Fine-scale coral connectivity pathways in the Florida Reef Tract: Implications for conservation and restoration. *Frontiers in Marine Science* 7, 312
- FWC-FWRI (2017). Unified Reef Map v2. 0. *FWC-FWRI (Florida Fish and Wildlife Conservation Commission-Fish and Wildlife Research Institute)*

- Gardner, T. A., Côté, I. M., Gill, J. A., Grant, A., and Watkinson, A. R. (2003). Long-term region-wide declines in Caribbean corals. *science* 301, 958–960
- Geuzaine, C. and Remacle, J.-F. (2009). Gmsh: A 3-d finite element mesh generator with built-in pre-and post-processing facilities. *International journal for numerical methods in engineering* 79, 1309–1331
- Harvell, D., Jordán-Dahlgren, E., Merkel, S., Rosenberg, E., Raymundo, L., Smith, G., et al. (2007). Coral disease, environmental drivers, and the balance between coral and microbial associates. *Oceanography* 20, 172–195
- Keeling, M. and Rohani, P. (2007). Stochastic dynamics. *Modeling Infectious Diseases in Humans and Animals*, 190–230
- Kermack, W. O. and McKendrick, A. G. (1927). A contribution to the mathematical theory of epidemics. *Proceedings of the Royal Society of London. Series A, Containing papers of a mathematical and physical character* 115, 700–721
- Kourafalou, V. H. and Kang, H. (2012). Florida current meandering and evolution of cyclonic eddies along the Florida Keys Reef Tract: Are they interconnected? *Journal of Geophysical Research: Oceans* 117 [Dataset] Kramer, P., Roth, L., and Lang, J. (2019). Map of stony coral tissue loss disease outbreak in the Caribbean. [www.agrra.org](http://www.agrra.org). ArcGis Online. (accessed June 12, 2020)
- Kuffner, I. B., Lidz, B. H., Hudson, J. H., and Anderson, J. S. (2015). A century of ocean warming on Florida Keys coral reefs: historic in situ observations. *Estuaries and Coasts* 38, 1085–1096
- Kundu, P. K. (1976). Ekman veering observed near the ocean bottom. *Journal of Physical Oceanography* 6, 238–242
- Kuta, K. and Richardson, L. (1996). Abundance and distribution of black band disease on coral reefs in the northern Florida Keys. *Coral reefs* 15, 219–223
- Lambrechts, J., Hanert, E., Deleersnijder, E., Bernard, P.-E., Legat, V., Remacle, J.-F., et al. (2008). A multi-scale model of the hydrodynamics of the whole Great Barrier Reef. *Estuarine, Coastal and Shelf Science* 79, 143–151
- Manzello, D. P. (2015). Rapid recent warming of coral reefs in the Florida Keys. *Scientific reports* 5, 16762
- Miller, J., Muller, E., Rogers, C., Waara, R., Atkinson, A., Whelan, K., et al. (2009). Coral disease following massive bleaching in 2005 causes 60% decline in coral cover on reefs in the US Virgin Islands. *Coral Reefs* 28, 925
- Muller, E. M., Sartor, C., Alcaraz, N. I., and van Woesik, R. (2020). Spatial epidemiology of the Stony-Coral-Tissue-Loss Disease in Florida. *Frontiers in Marine Science* 7, 163
- Murphy, B. S. (2014). Pykrige: development of a kriging toolkit for Python. *AGUFM* 2014, H51K–0753
- Murray, J. D. (2007). *Mathematical biology: I. An introduction*, vol. 17 (Springer Science & Business Media)
- Neely, K. (2018). Surveying the Florida Keys southern coral disease boundary. *Florida DEP. Miami, FL*, 1–15
- [Dataset] NOAA (2018). Stony Coral Tissue Loss Disease Case Definition. Available online at: <https://nmsfloridakeys.blob.core.windows.net/floridakeys-prod/media/docs/20181002-stony-coral-tissue-loss-disease-case-definition.pdf> (accessed June 4, 2020)
- Perlin, A., Moum, J., Klymak, J., Levine, M., Boyd, T., and Kosro, P. (2007). Organization of stratification, turbulence, and veering in bottom Ekman layers. *Journal of Geophysical Research: Oceans* 112
- Porter, J. W. and Meier, O. W. (1992). Quantification of loss and change in floridian reef coral populations. *American Zoologist* 32, 625–640



- 547 Precht, W. F., Gintert, B. E., Robbart, M. L., Fura, R., and Van Woesik, R. (2016). Unprecedented  
548 disease-related coral mortality in Southeastern Florida. *Scientific Reports* 6, 1–11
- 549 Richardson, L. L., Goldberg, W. M., Kuta, K. G., Aronson, R. B., Smith, G. W., Ritchie, K. B., et al. (1998).  
550 Florida's mystery coral-killer identified. *Nature* 392, 557–558
- 551 Rosales, S. M., Clark, A. S., Huebner, L. K., Ruzicka, R. R., and Muller, E. (2020). Rhodobacterales and  
552 Rhizobiales are associated with stony coral tissue loss disease and its suspected sources of transmission.  
553 *Frontiers in Microbiology* 11, 681
- 554 Ruzicka, R., Colella, M., Porter, J., Morrison, J., Kidney, J., Brinkhuis, V., et al. (2013). Temporal changes  
555 in benthic assemblages on Florida Keys reefs 11 years after the 1997/1998 El Niño. *Marine Ecology*  
556 *Progress Series* 489, 125–141
- 557 Shore, A. and Caldwell, J. M. (2019). Modes of coral disease transmission: How do diseases spread  
558 between individuals and among populations? *Marine biology* 166, 45
- 559 Smith, N. P. (2009). The influence of wind forcing on across-shelf transport in the Florida Keys. *Continental*  
560 *Shelf Research* 29, 362–370
- 561 Sokolow, S. H., Foley, P., Foley, J. E., Hastings, A., and Richardson, L. L. (2009). Editor's choice: Disease  
562 dynamics in marine metapopulations: modelling infectious diseases on coral reefs. *Journal of Applied*  
563 *Ecology* 46, 621–631
- 564 Sponaugle, S., Paris, C., Walter, K., Kourafalou, V., and Alessandro, E. (2012). Observed and modeled  
565 larval settlement of a reef fish to the Florida Keys. *Marine Ecology Progress Series* 453, 201–212
- 566 Sutherland, K. P., Porter, J. W., and Torres, C. (2004). Disease and immunity in Caribbean and Indo-Pacific  
567 zooxanthellate corals. *Marine Ecology Progress Series* 266, 273–302
- 568 Team, A. B. R. (2005). Atlantic Acropora status review document. *Report to National Marine Fisheries*  
569 *Service*, 152
- 570 Thomas, C. J., Bridge, T. C., Figueiredo, J., Deleersnijder, E., and Hanert, E. (2015). Connectivity between  
571 submerged and near-sea-surface coral reefs: Can submerged reef populations act as refuges? *Diversity*  
572 *and Distributions* 21, 1254–1266
- 573 Thomas, C. J., Lambrechts, J., Wolanski, E., Traag, V. A., Blondel, V. D., Deleersnijder, E., et al. (2014).  
574 Numerical modelling and graph theory tools to study ecological connectivity in the Great Barrier Reef.  
575 *Ecological Modelling* 272, 160–174
- 576 Vaz, A. C., Paris, C. B., Olascoaga, M. J., Kourafalou, V. H., Kang, H., and Reed, J. K. (2016). The perfect  
577 storm: match-mismatch of bio-physical events drives larval reef fish connectivity between Pulley Ridge  
578 mesophotic reef and the Florida Keys. *Continental Shelf Research* 125, 136–146
- 579 Viehman, S., Gittings, S., Groves, S., Moore, J., Moore, T., and Stein, J. (2018). NCCOS Assessment: Coral  
580 Disturbance Response Monitoring (DRM) Along the Florida Reef Tract Following Hurricane Irma From  
581 2017-10-09 to 2017- 10-18 (NCEI Accession 0179071). *NOAA National Centers for Environmental*  
582 *Information*. Silver Spring, MD: National Centers for Coastal Ocean Science. Available online at:  
583 <https://doi.org/10.25921/sscd-6h41> (accessed June 22, 2020)
- 584 Wolanski, E. (1994). *Physical oceanographic processes of the Great Barrier Reef* (CRC Press)