Neuroactive compounds induce larval settlement in the scleractinian coral *Leptastrea purpurea*

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Settlement of pelagic coral larvae is commonly induced by chemical cues that originate from biofilms and coralline algae. These natural settlement cues initiate signal pathways leading to attachment and metamorphosis of the coral larva. In order to investigate the settlement process and its natural inducers, it is necessary to gain a better understanding of these signal pathways. At present, the pathways and neurotransmitters involved in this signal transduction are still widely unknown. In this study, we exposed larvae of the brooding coral *Leptastrea purpurea* to five neuroactive compounds known to be present in cnidarians, and K$^+$ ions. All compounds were applied at different dilutions and settlement behavior of the larvae was documented over 48 h. Dopamine, glutamic acid and epinephrine significantly induced settlement in the coral larvae. The highest observed metamorphosis response was 54% in $10^{-5}$ M dopamine. Serotonin, L-DOPA and K$^+$ ions did not have an influence on settlement behavior in our experiments. Exposing larvae to settlement-inducing neurotransmitters and thus bypassing the initial induction could be utilized in coral aquaculture. The active neurotransmitters should be used to further study the settlement process in *L. purpurea* in greater detail. Their role and relevance should also be assessed for other coral species as they may represent or reveal a universal inducer for coral settlement.

Coral reefs belong to the biologically most diverse ecosystems of our planet. By providing a multitude of ecosystem services, coral reefs generate significant revenue to economies of tropical coastal regions. Despite their significant role for sustaining the livelihoods of several hundred million people, coral reefs are threatened in many ways and continue to decline and disappear at a historically unprecedented rate. Rising sea surface temperatures caused by the ongoing climate change represent the most severe stressor. Like many sessile marine animals, scleractinian corals disperse through a larval stage before settling on a suitable substrate and metamorphosing into a primary polyp. The planula larvae have sensory systems consisting of chemoreceptors which allow them to locate suitable settlement substrates. Known settlement cues include light, reef sound, surface structure, and chemical cues. The presence of settlement-inducing cues triggers a sequence of processes leading to attachment and subsequent metamorphosis. The apical organ, which is a sensory structure located at the aboral pole of the planula larva, is most likely involved in the metamorphic process. Crustose coralline algae (CCA) and their associated bacteria are well-recognized substrates for settlement that are favored by many coral species. Settlement-inducing CCA usually grow under conditions that are beneficial to scleractinians, i.e. exposure to high light intensity, adequate water quality and sufficient water movement. Many scientists have been investigating the origin of settlement cues and, while it could be demonstrated that certain bacteria are an important source of settlement-inducing compounds, some settlement cues derive from CCA itself as well. Some of the compounds produced by CCA or their associated biofilms induce settlement in very few species of coral, others seem to have an effect on a larger number of coral species. The compound 11-deoxyfistularin-3 is a species-specific inducer which triggers settlement in larvae of the coral *Pseudosiderastrea tayamii*. In contrast, tetrabromopyrrol (TBP), which is produced by certain *Pseudoalteromonas* strains, induces metamorphosis in a number of coral species of different families such as *Acropora millepora*, *Porites astreoides*, *Orbicella franksi* and...
Acropora palmata\textsuperscript{11,13}. However, the ecological role that TBP plays is unclear because of its low natural concentration in the environment and the lack of larval attachment to the substrate in many corals\textsuperscript{11,13}. Another naturally occurring compound with an impact on settlement behavior is Luminaolide, which enhances settlement rates in larvae of Leptastrea purpurea and several Acroporidae species\textsuperscript{21,23}.

Although a variety of natural settlement-inducing agents have been isolated and identified\textsuperscript{21,22,23}, and gene expression during the settlement process has been described\textsuperscript{24}, the signal pathways and neurotransmitters involved in the settlement process are still widely unknown. GLW-amide is a neuropeptide that is known to induce metamorphosis, often without attachment in Acropora tenuis and A. palmata\textsuperscript{25–27}. It presumably “short-cuts” the normal developmental process and some of the behavioral and physiological changes normally associated with settlement\textsuperscript{27,28}. These findings suggest that other neurotransmitters are also involved in the regulation of natural settlement behavior.

Deciphering which transmitters regulate the processes leading to attachment and metamorphosis would be a vital step to a more complete understanding of the scleractinian life cycle. By exposing larvae to neurotransmitters, the individual steps of signaling pathways leading to settlement and the involved transmitters could be identified. This has been conducted with a number of marine invertebrates like polychaetes\textsuperscript{29–31}, gastropods\textsuperscript{32–36}, bivalves\textsuperscript{37–44} and holothuroids\textsuperscript{45,46}. The obtained knowledge found application in the commercial oyster culture where ammonia and the neurotransmitters L-DOPA, epinephrine, norepinephrine and yohimbine have been successfully used as settlement agents\textsuperscript{47,48}.

A number of neurotransmitters and artificial compounds have been discovered to induce settlement in marine cnidarians. Larvae of the hydrozoan Eudendrium racemosum exhibited settlement behavior in response to serotonin\textsuperscript{49}. Larvae of another hydrozoan, Halocordyle disticha settled when exposed to L-DOPA, dopamine and norepinephrine\textsuperscript{50}. As reviewed in Kass-Simon & Pierobon\textsuperscript{52}, both classical neurotransmitters, the fast-acting (acetylcholine\textsuperscript{53,54}, GABA\textsuperscript{55–59}, glutamate\textsuperscript{55,56,60,61}, glycine\textsuperscript{62–65}) and the slow-acting (catecholamines\textsuperscript{66,67} and serotonin\textsuperscript{68–72}) ones, as well as neuropeptides 73–85 play a role in the neurotransmission of cnidarians. Although many neurophysiologically active compounds have been tested on cnidarians and the corresponding receptors detected, there still is a large number of neurotransmitters whose effects on scleractinians and their role in settlement-related neurotransmission processes remain unknown.

A better understanding of the molecular basics of the settlement process of coral larvae is necessary to identify potential risks for that crucial life stage. Neuroactive environmental pollutants can interfere with the settlement of coral larvae like endocrine disruptors from sunscreen can induce calcification in larvae and cause bleaching in adults\textsuperscript{86,87}. More knowledge about the molecules involved in this critical step in a coral’s life can enable researchers to investigate potentially problematic compounds. Since most corals only reproduce once a year an interference with the larval settlement could significantly reduce an entire cohort. A better understanding of the settlement process could also find application in coral aquaculture as well as reef restoration purposes: large numbers of coral larvae could be settled on a substrate suitable for subsequent installation in the reef.

In this study, we conducted a series of experiments with larvae of the hermatypic coral species Leptastrea purpurea by exposing them to several neurotransmitters. We differentiated between attachment of larvae, metamorphosis and full settlement (attachment + metamorphosis) since metamorphosis can occur without attachment of the larva. The aim of the experiments conducted for this study was to widen the knowledge about the effects of commonly occurring neurotransmitters on the settlement of scleractinian coral larvae.

Materials and Methods

**Acquisition of coral larvae.** Larvae of the brooding coral Leptastrea purpurea were used for the experiments. L. purpurea is a brooding faviid coral which is common in the waters of Guam. Its unique mode of reproduction allows constant access to larvae. In contrast to other brooding species, colonies of the population in Guam, USA, release larvae on a daily basis. L. purpurea can be found in shallow water and easily be collected, e.g. at Luminao reef (13°27′25.66″N 144°37′31.55″E). The colonies used to obtain larvae for this study were detached from their substrate using hammer and chisel during free-diving. Upon collection, colonies were transported to the University of Guam Marine Laboratory (UOGML) and placed in large flow-through tanks. Larval collection started after three days of acclimatization of the 150 collected colonies. Larval collection was conducted as described in Moeller et al.\textsuperscript{88} and Nietzer et al.\textsuperscript{89}. In short: The corals were placed in 30 L plastic containers (30 colonies per container) with aeration and flow but no light throughout the night. The previous morning, colonies were placed back into the flow-through tanks where they stayed during the day. The water from the collection containers was carefully filtered through a 30 μm mesh to reduce their content to 2–3 L. Larvae of L. purpurea contain a green fluorescent protein (GFP) and can be easily detected in the collected material using a blue fluorescent lamp and yellow barrier filters (BlueStar\textsuperscript{90}, NIGHTSEA, Lexington, Massachusetts, USA). Larvae are 0.2–0.9 mm in length and can be collected with a plastic pipette without any optical magnification.

**Settlement assays.** The following compounds were tested: serotonin hydrochloride, (−)-epinephrine, dopamine hydrochloride, 3,4-dihydroxy-L-phenylalanine (L-DOPA), L-glutamic acid, KCl. All pharmacological agents were obtained from Sigma-Aldrich Chemie GmbH (Munich, Germany). Except for epinephrine, all compounds were dissolved at 1 M in MilliQ H₂O. Epinephrine is poorly soluble in water and was dissolved in 0.005 M HCl. Serial dilutions were compiled in order to achieve the desired end concentrations by adding 50 μl of the respective solution to 5 ml of filter-sterilized seawater (FSW).

Assays were conducted in 12-well plates. Test solutions were dissolved in 5 ml FSW and replicated five times. FSW was used as negative control and small pieces of live CCA (Hydrolithon reinboldii) served as positive control. Due to the use of HCl solution for the epinephrine experiment, controls were conducted with both FSW as well as 0.005 M HCl. All chemicals were tested at 10⁻³ M, 10⁻⁴ M, 10⁻⁵ M, 10⁻⁶ M, 10⁻⁷ M. These concentrations have been used in several other studies investigating neurotransmitters in marine invertebrates\textsuperscript{99,80,90,91}. In case
the highest concentration did not lead to 100% mortality of the exposed larvae, chemicals were tested in higher concentrations ($10^{-3}$ M, $10^{-2}$ M). The aim of this approach was to avoid missing potentially active concentrations.

Per replicate, 10 larvae were added. Larvae were examined under a dissecting microscope after 24 h and 48 h for metamorphosis, attachment and mortality. Effects are usually clearly visible after 24 h but more pronounced after 48 h; mesenterys are clearly visible in metamorphosed primary recruits whereas dead larvae start to disintegrate. The status after 48 h was used for statistical analysis and Fig. 1.

We defined the documented categories as: normal swimming (no effect, Fig. 2a), metamorphosis without attachment, metamorphosis with attachment (full settlement, Fig. 2b) and mortality.

**Statistical analysis.** Statistical analysis was conducted with SPSS 24 statistics. Metamorphosis data were not normally distributed, therefore a Generalized Linear Model (GLM) with Poisson variance was applied. All data points $>0$ were tested against negative control in a pairwise GLM with Wald Chi-Square Test.

**Results**

Positive controls (CCA pieces) yielded 98.0 ± 4.2% full settlement (metamorphosis and attachment) (Fig. 1g). In the negative controls (FSW), 2.0 ± 4.2% full settlement was documented. Metamorphosis without attachment was not observed in the controls (Fig. 1h).

Significant effects that lead to settlement behavior ($p < 0.05$) were detected for dopamine, epinephrine and glutamic acid (Fig. 1a–f).

The highest metamorphosis response of 54 ± 10.4% was documented with dopamine ($10^{-5}$ M) ($p < 0.001$). Stronger dilutions ($10^{-6}$ M, $10^{-7}$ M) had only marginally lower effects ($10^{-6}$ M: 42.0 ± 15.2%, $p = 0.001$; $10^{-7}$ M: 50.0 ± 19.6 $p < 0.001$) whereas higher concentrations ($10^{-3}$ M, $10^{-2}$ M) were lethal. Glutamic acid showed low but significant metamorphosis rates at $10^{-4}$ M (14.0 ± 5.1%, $p = 0.034$) and $10^{-3}$ M (14.0 ± 17.3%, $p = 0.034$). Epinephrine had a significant effect on metamorphosis at $10^{-4}$ M (14.0 ± 11.4%, $p = 0.034$). However, the same concentration was lethal for 42.0 ± 13.0% percent of the larvae. Higher dilutions showed significant metamorphosis rates peaking at $10^{-5}$ M (46.0 ± 21.8%, $p < 0.001$) and lower rates at $10^{-4}$ M (34.0 ± 27.4%, $p < 0.001$) and $10^{-3}$ M (28.0 ± 22.0%, $p < 0.001$). Phenylalanine, serotonin and K+ did not have significant effects on the settlement behavior of the larvae. Lethal concentrations were $10^{-3}$ M for phenylalanine, $10^{-3}$ M for serotonin and 1 M for K+.

**Discussion**

The conducted experiments revealed that dopamine, glutamic acid and epinephrine significantly induced settlement behavior in larvae of *Leptastrea purpurea*. All three compounds mostly induced full settlement (attachment and metamorphosis). A smaller proportion of the larvae underwent metamorphosis without attachment. As described in the introduction, GLW-amide has been documented to induce metamorphosis without attachment in larvae of *Acropora* sp., presumably by shortcircuiting the natural sequences and directly inducing metamorphosis. This suggests that GLW-amide could act as a transmitter in the metamorphic process. The results obtained in our experiments indicate that the three active compounds could also be relevant for the natural settlement process: i.e. the presence of specific settlement cues could trigger the release of these neurotransmitters and induce an upregulation of the expression of transmitter-associated genes. All tested neurotransmitters have previously been described as present in various cnidarian species (reviewed by Kass-Simon and Pierobon, 2007).

Dopamine yielded the strongest effect on settlement behavior at a concentration of $10^{-5}$ M: 54% of larvae underwent metamorphosis. Dopamine has been identified in both neurons and myoepithelial cells by Carlberg in *Hydra thomseni*. Westfall et al. found dopamine and serotonin in the epidermal synapses of tentacles of the sea anemone *Exaiptasia pallida*. Dopamine has also been found to play an inhibitory role in the spawning process of *Acropora* species. Apart from the mentioned examples, dopamine has been found in many members of the cnidarian phylum as reviewed by Kass-Simon and Pierobon. It is known to be a neurotransmitter in other marine invertebrates as well, and induces larval settlement in a number of invertebrates like mussels, sea cucumbers and barnacles. Dopamine and epinephrine are catecholamines that have been proposed to play an important role in the settlement process of other marine invertebrates and larvae. However, given the wide range of processes that dopamine could potentially be involved in, it is possible that an exposure to high concentrations could cause off-target effects like the observed metamorphosis without attachment.

Epinephrine also induced a significant settlement response in several tested concentrations. Highest settlement response was observed at $10^{-5}$ M. Epinephrine was localized in the octocorallian sea pansy, *Renilla koellikeri*. Norepinephrine-reactive neurons were found in the neurites, mesoglea and at the bases of its ectoderm and endoderm, and epinephrine, norepinephrine and dopamine could be identified in both neurons and myoepithelial cells. Based on its widespread occurrence in cnidarian organisms and its strong settlement response, epinephrine may be involved in the settlement signal transduction pathway. L-DOPA and dopamine are catecholamines that have been proposed to play an important role in the settlement process of other marine invertebrates and larvae. Our results indicate that *L. purpurea* larval settlement is also under endogenous regulation by a catecholaminergic mechanism.
and epinephrine in bivalve larvae. Our results suggest that this mechanism does not take place in larvae of *L. purpurea*.

High concentrations of glutamate have been documented in cnidarian nematocysts. In our assays, glutamic acid significantly induced settlement at $10^{-4}$ M. At a concentration of $10^{-5}$ M, glutamic acid induced full settlement but also metamorphosis without attachment. Apart from their role in the nematocysts, glutamic acids are also relevant components of immune reactions as well as feeding behavior of *Hydra* sp. Furthermore, glutamate has been shown to increase the output of both ectodermal and endodermal impulse generating systems.
The wide spectrum of activity in cnidarian physiology makes glutamic acids a potential signaling compound in the settlement process of coral larvae.

Despite serotonin being a known settlement inducer in different hydrozoan and scyphozoan species, it did not have any significant effects on the settlement behavior of the tested anthozoan larvae in our experiment. This suggests a different signal pathway. Experiments with Acropora millepora exposed to CCA revealed no effect on the expression of the serotonin receptor 5htr1. K+ ions are known to induce metamorphosis in marine invertebrates either by depolarization of cell membranes or by acting in the metamorphic signal-transduction pathway. The three highest concentrations of K+ ions tested led to a distinct ‘flower’ shape of the larvae within the first minutes of exposure. However, larvae exposed to these concentrations were either dead or swimming normally after 24 h. The same was observed after 48 h. Although the tested concentrations yielded consistently low and insignificant settlement rates, we cannot rule out that intermediate concentrations might lead to higher rates of settlement or metamorphosis without attachment. A mere depolarization by ions might not lead to settlement as well as the natural cue. Two or more compounds could have synergistic effects leading to a much stronger settlement response.

None of the tested compounds induced settlement as effective as the positive control which can have several reasons. It is likely that the tested concentrations were not the most effectives ones since we used a predefined dilution series. Intermediate or lower concentrations, especially in the case of dopamine, might yield settlement responses that are stronger than the documented ones. It is also possible that one single compound cannot induce settlement as well as the natural cue. Two or more compounds could have synergistic effects leading to a much stronger settlement response.

A broader understanding about coral settlement could make conservation efforts more effective. A detailed knowledge about the molecules involved in the settlement process could help to identify environmental pollutants that might be interfering. For example, oxybenzone, a compound in sunscreens has recently been shown to act as an endocrine disruptor in the calcification process leading to calcified and thus unviable coral larvae. Since reefs are exposed to a growing number of pollutants, knowledge about the endocrine system and signaling pathways in scleractinians is increasingly important.

Neurotransmitters could be used to induce settlement of coral larvae ex-situ on a large scale. If the tested compounds are also active in other scleractinian corals, mass production techniques of corals for the aquarium trade or reef restauration purposes could advance by using adequate settlement inducers such as neurotransmitters, as is common practice in the aquaculture of bivalves. Using CCA as a settlement substrate is highly effective but comes with a variety of negative aspects, especially when applied on a large scale. Finding adequate amounts of easily accessible CCA-covered rock or dead coral branches can be challenging. If large numbers (e.g., several tens of thousands) of coral recruits are produced, several kg of CCA-covered substrate have to be removed from the environment which might impact the habitat negatively. Since natural CCA-covered substrates are usually very porous, unwanted organisms can be introduced into the rearing facilities. Aside from various algae, Aiptasia anemones, benthic ctenophores, ciliates as well as corallivorous snails or crabs can find their way into the tanks and reduce the survival of the coral recruits, particularly in a long-term rearing operation. Live CCA also are known competitors to juvenile corals with the ability to shed their cuticula where a coral larva or any other another epibiont has settled. We observed this phenomenon on a regular basis. Additional to surface sloughing, an increased mucus production has been described as a protective measure from CCA against biofouling organisms. These defense mechanisms of live CCA can reduce the early survival of coral recruits.
Another problem could be the heterogeneity of CCA. If the corals need to settle on standardized substrates for experimental purposes, CCA chips can be a rather improper substrate. If artificial substrates such as ceramics are used for settlement, they have to be conditioned in natural seawater for several weeks in order to develop a settlement-inducing biofilm or CCA cover. In contrast, neuroactive compounds could be applied instantaneously and would likely be a cost-efficient alternative: a hypothetical application of 10−6 M dopamine-HCl as an inductive agent would amount to roughly $5 for 10,000 L of settlement medium. However, further studies should determine the most efficient concentrations and their long-term effects on settled recruits in order to assess whether neurotransmitters are a suitable way to settle large numbers of coral larvae. Although the experiments for this study were conducted with larvae obtained from 150 individual parent colonies which should eliminate phenotypic response biases, further studies with this species and a wider spectrum of compound concentrations should be conducted to corroborate the findings of this study.

The results obtained from the conducted experiments suggest that a number of neurotransmitters known to be common in cnidarians could play a role in coral larval settlement by regulating relevant biochemical signal pathways. The tested concentrations yielded a range of responses and were chosen to determine active and toxic concentrations for each of the tested neurotransmitters. Despite a moderate metamorphosis response of a maximum of 54% in dopamine, a full settlement consisting of both attachment and metamorphosis could be observed. This indicates that certain neurotransmitters could be harnessed to induce settlement of coral larvae in order to generate juvenile corals e.g. for reef restoration purposes. The identification of the settlement-inducing traits of three common neurotransmitters could be used as groundwork to further study the exact pathways involved in coral larval settlement on a molecular level. Although it is likely that the underlying pathways are of a universal nature in scleractinians, further research into the involved mechanisms with a range of other coral species from different families is indispensable.

Data Availability
Data is available on figshare.com: https://figshare.com/s/f808ab6db9236c3b0d75.

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Acknowledgements

We thank staff and students of the Marine Laboratory of the University of Guam for assistance during fieldwork and laboratory work. We thank the anonymous reviewers for their valuable comments. This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors.
Author Contributions
M.M. and S.N. executed the experiments and wrote the manuscript. M.M. prepared the figures. M.M., S.N. and P.S. planned the experiments, reviewed and revised the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

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