Fig S1

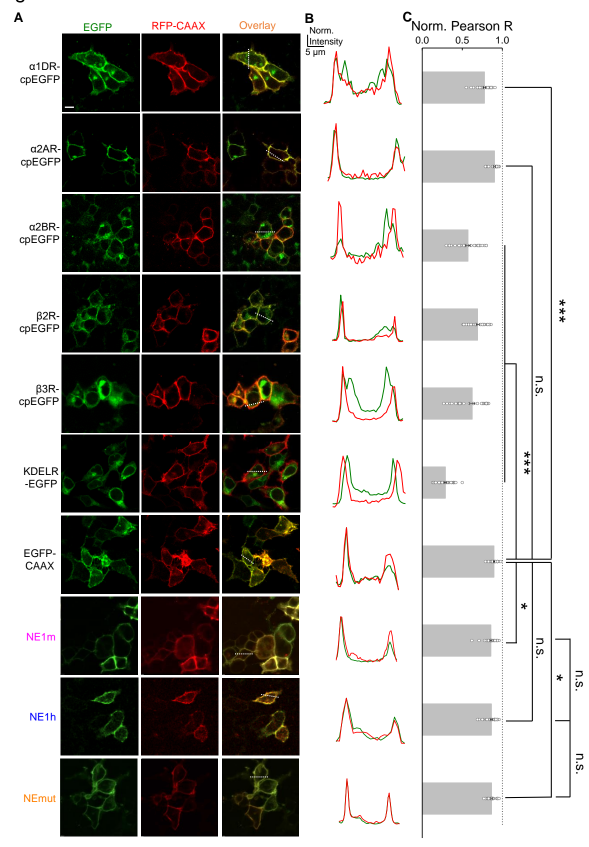


Fig S2

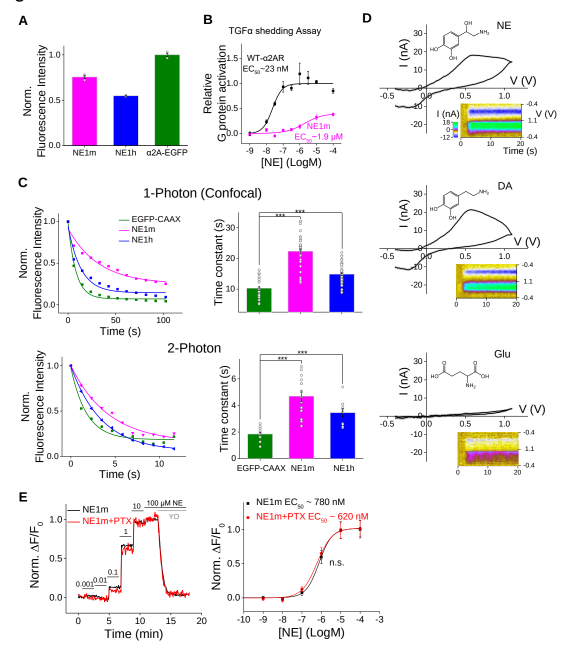


Fig S3

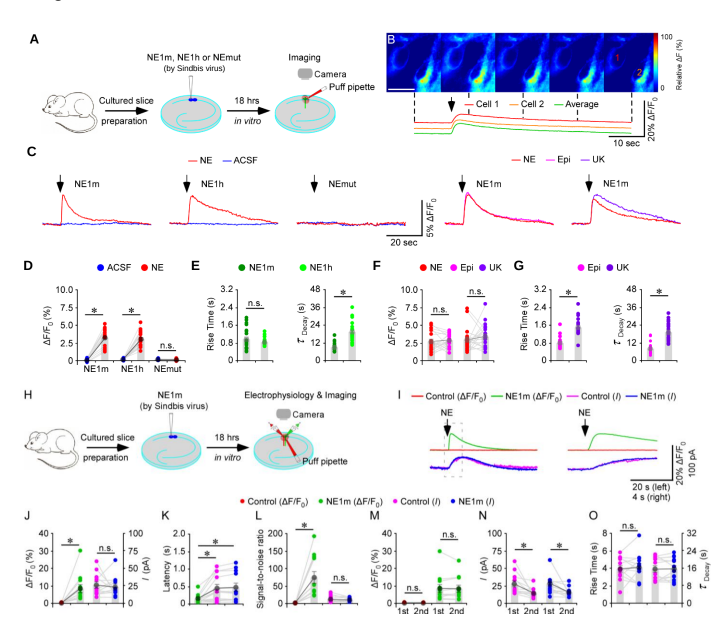
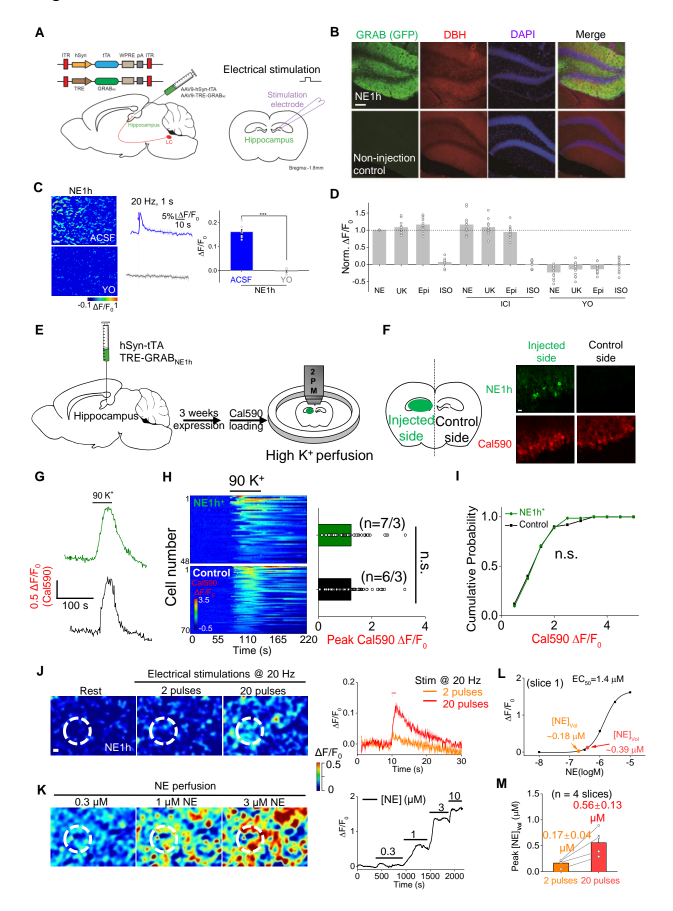


Fig S4



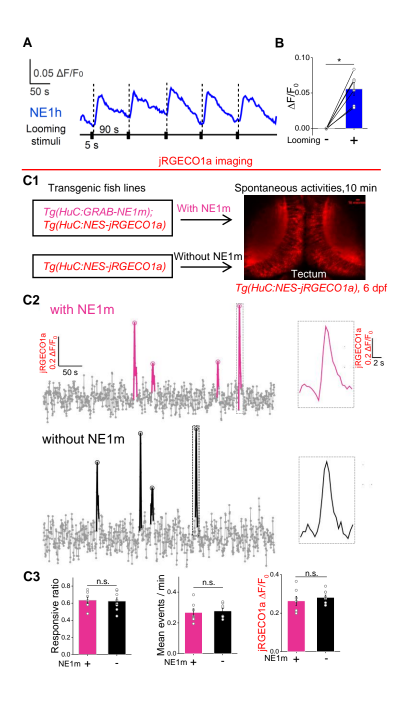
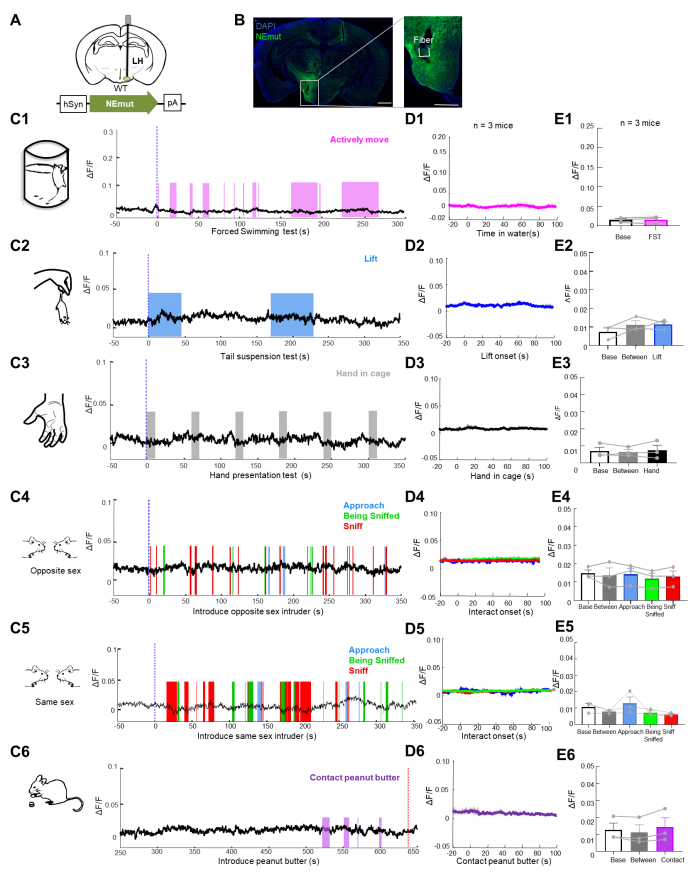


Fig S6



- 1 Figure S1. Characterization of the membrane trafficking of a panel of screening
- 2 candidates, related to Figure 1.
- 3 Representative images (A) of HEK293T cells co-transfected with the indicated screening
- 4 candidates (green) together with RFP-CAAX (red) to label the plasma membrane. KDELR-
- 5 EGFP was used as an ER marker. The dashed white lines indicate the line used for the
- 6 line-scanning data shown in (B) and summarized in
- 7 (C) n = 30 cells from 4-5 cultures.

- 8 The scale bars in (A) represent 10 μ m.
- 9 *p < 0.05 and ***p < 0.001; n.s., not significant (Student's t-test).

- 11 Figure S2. Further characterization of GRAB_{NE} sensors, related to Figure 2 and
- 12 **Figure 3.**
- 13 (A) Fluorescence intensity of GRAB_{NE1m} and GRAB_{NE1h} expressed relative to EGFP-α2AR.
- 14 $n \ge 2$ wells with 300-500 cells per well.
- 15 (**B**) G protein activation mediated by GRAB_{NE1m} and wild-type α2AR was measured using
- the TGFα shedding assay and is expressed relative to α2AR. n = 4 wells with $\ge 10^5$ cells
- per well.
- 18 (C) Exemplar (left) and summary data (right) showing the photostability of GRAB_{NE}
- sensors and EGFP-CAAX using confocal (top) and 2-photon (bottom) microscopy. n > 10
- 20 cells from at least 3 cultures.
- 21 (**D**) Exemplar cyclic voltammograms for 10 μM NE (**top**), 10 μM DA (**middle**), and 10 μM
- 22 Glu (bottom) measured using FSCV are shown. The traces were averaged from separate
- 23 **200 trials**.

- 24 (E) Disrupting of G protein activation with pertussis toxin does not affect the NE-induced
- 25 fluorescence change in GRAB_{NE1m}-expressing neurons. n = 27 neurons from 3 cultures.
- 26 ***p < 0.001 (Student's *t*-test).

Figure S3. GRAB_{NE} sensors respond selectively to noradrenergic agonists in brain

- 29 slices without affecting endogenous NE receptor functions, related to Figure 4.
- 30 (A) Schematic drawing showing the experimental design for measuring CA1 pyramidal
- 31 neurons in cultured rat hippocampal slices.
- 32 **(B)** Heat-map images of the change in fluorescence in GRAB_{NE1m}-expressing CA1 neurons
- in response to a 10-ms local application of NE (20 µM). The red and orange traces show
- 34 the fluorescence responses of two neurons, and the green trace shows the average
- 35 response of all neurons in the field.
- 36 (C) Fluorescence responses measured in GRAB_{NE1m}-, GRAB_{NE1h}-, and GRAB_{NEmut}-
- expressing CA1 neurons following a 10-ms puff (arrow) of ACSF, NE (20 μM), Epi (100
- μ M), or brimonidine (UK, 20 μ M).
- 39 (D) Maximum $\Delta F/F_0$ responses measured in GRAB_{NE1m}-, GRAB_{NE1m}-, and GRAB_{NEmut}-
- 40 expressing CA1 neurons following a 10-ms puff of ACSF or NE. n = 20-21 cells from 8
- 41 animals per group.
- 42 (E) Rise times and decay time constants measured in CA1 neurons expressing GRAB_{NE1m}-
- and GRAB_{NE1h}- expressing CA1 neurons in response to a puff of NE. n = 21 cells from 8
- 44 animals.

- Δ F/F₀ responses measured in GRAB_{NE1m}-expressing CA1 neurons following
- a puff of NE, Epi, or brimonidine (UK). n = 20-21 cells from 8 animals per group.
- 47 (G) Rise times and decay time constants measured in GRAB_{NE1m}-expressing CA1 neurons
- 48 following a puff of Epi or brimonidine (UK).
- 49 (H) Schematic drawing outlines the design of simultaneous imaging and
- 50 electrophysiological recording experiments in rat cultured hippocampal slices.
- 51 (I) Left, simultaneous fluorescence and current responses of a pair of GRAB_{NE1m}
- 52 expressing and neighboring control non-expressing CA1 neurons to a 10-ms puff
- application of 0.2 mM norepinephrine (NE). Right, the responses in the left rectangle box
- are shown again in an expanded time scale. Note the different latencies of fluorescence
- 55 and current responses.
- (J) Values for the amplitude of noradrenergic fluorescence (GRAB_{NE1m}: 8.52 ± 2.26%; Ctrl:
- 57 0.14 \pm 0.02%; Z = 3.059; p = 0.002; n = 12 from 12 animals) and current (GRAB_{NE1m}: 23.4
- \pm 4.2 pA; Ctrl: 26.4 \pm 4.7 pA; Z = 0.078; p = 0.937; n = 12 from 12 animals) responses of
- 59 GRAB_{NE1m} expressing CA1 neurons compared to non-expressing neurons.
- 60 (K) Values for the latency of noradrenergic current responses in GRAB_{NE1m} expressing
- (GRAB_{NE1m}: $462.0 \pm 124.2 \text{ ms}$; Z = 2.578; p = 0.01) and non-expressing CA1 neurons (Ctrl:
- 440.6 \pm 113.1 ms; Z = 2.432; p = 0.015) compared to those of fluorescence responses of

- GRAB_{NE1m} expressing neurons (GRAB_{NE1m}: 145.8 ± 36.4 ms; n = 12 from 12 animals).
- 64 (L) Values for the signal-to-noise ratio (SNR) of noradrenergic fluorescence responses of
- 65 GRAB_{NE1m} expressing CA1 neurons compared to non-expressing neurons (GRAB_{NE1m}:
- 75.9 \pm 17.1; Ctrl: 2.5 \pm 0.3; Z = 3.509; p = 0.002; n = 12 from 12 animals) and noradrenergic
- 67 current responses of GRAB_{NE1m} expressing CA1 neurons compared to non-expressing
- neurons (GRAB_{NE1m}: 9.6 ± 1.6 ; Ctrl: 11.3 ± 2.8 ; Z=-0.235; p=0.814; n=12 from 12 animals).
- 69 Note the larger SNR of noradrenergic fluorescence responses of GRAB_{NE1m} expressing
- 70 CA1 neurons compared to current responses of GRAB_{NE1m} expressing and non-expressing
- 71 CA1 neurons (GRAB_{NE1m}: Z = -3.509; p = 0.002; Ctrl: Z = -2.981; p = 0.002).
- 72 (M) Values for the two consecutive fluorescence responses of GRAB_{NE1m} expressing (1st:
- 73 8.56 \pm 0.02%; 2nd: 8.43 \pm 0.02%; Z = 0; p = 1; n = 12 from 12 animals) and control non-
- 74 expressing (1st: 0.14 \pm 0.02%; 2nd: 0.11 \pm 0.01%; Z = -1.832; p=0.067; n = 12 from 12
- 75 animals) CA1 neurons.
- 76 (N) Values for the two consecutive noradrenergic current responses in GRABNE1m
- expressing (1st: 28.6 ± 5.0 pA; 2^{nd} : 15.6 ± 2.0 pA; Z = -2.51; p=0.012; n = 12 from 12
- 78 animals) and control non-expressing (1st. 27.7 \pm 4.4 pA; 2nd: 14.9 \pm 2.0 pA; Z = -3.059;
- 79 p=0.02; n=12 from 12 animals) CA1 neurons.
- 80 (**O**) Values for the rise time (GRAB_{NE1m}: 4.14 ± 0.46 s; Ctrl: 3.98 ± 0.38 s; Z = 0.314; p =
- 81 0.754; n = 12 from 12 animals) and decay time constant (GRAB_{NE1m}: 15.81 ± 1.50 s; Ctrl:
- 82 15.70 \pm 1.53 s; Z = 0.784; p = 0.433; n = 12 from 12 animals) of noradrenergic current
- 83 responses in GRAB_{NE1m} expressing neurons compared to control non-expressing CA1
- neurons. Large gray dots indicate average responses and asterisks indicate *p*<0.05
- 85 (Wilcoxon tests).

- Figure S4. GRAB_{NE} sensors respond selectively to noradrenergic agonists in brain slices without disturbing neuronal activities, related to Figure 4.
- 90 **(A)** Schematic illustration depicting AAV-mediated delivery of GRAB_{NE1h} in the mouse 91 hippocampus and bath application of various agonists in the dentate gyrus.
- 92 (B) Example images showing GRAB_{NE1h} (green) expression and dopamine beta
- 93 hydroxylase (DBH) immunostaining (red) in the dentate gyrus of AAV-GRAB_{NE1h}- and
- 94 control-injected hippocampus. The nuclei were counterstained with DAPI. The scale bar
- 95 represents 100 μm.
- 96 **(C)** Electrical stimulation evokes NE release in the hippocampus measured as a change in
- 97 GRAB_{NE1h} fluorescence. The response was blocked by batch application of yohimbine
- 98 (YO). Exemplar images (left), representative traces (middle), and the summary data (right)
- 99 are shown.
- 100 (D) Normalized change in GRAB_{NE1h} fluorescence in response to bath application of the
- indicated noradrenergic agonists in the presence or absence of ICI 118,551 or yohimbine.
- 102 (E) Schematic illustration of the calcium imaging experiments in acute brain slices. The
- 103 AAVs expressing GRAB_{NE1h} were injected unilaterally into the dentate gyrus and acute
- brain slices were prepared after 3 weeks and loaded with Cal590 red calcium dye for
- 105 imaging.
- 106 (F) The fluorescent signal of GRAB_{NE1h} sensor (green) and calcium dye Cal590 (red) in
- 107 acute brain slices. Scale bar, 20 µm.
- 108 (G) Representative traces of the Cal590 fluorescent response in either a GRAB_{NE1h}-
- 109 expressing neuron (upper) or a non-expressing control neuron (lower) to the perfusion of
- 110 high potassium solution (90mM K⁺).
- 111 (H) The group data of the Cal590 fluorescence responses in GRAB_{NE1h}-expressing
- neurons or non-expressing control neurons to the perfusion of high potassium solution
- 113 (n=48 neurons from 7 slices of 3 mice for GRAB_{NE1h}, n=70 neurons from 6 slices of 3 mice
- for control, p=0.95, student-t test)
- (I) Cumulative plot of the Cal590 fluorescence response. (P=0.93, Kolmogorov-Smirnov
- 116 test)
- 117 (J,K) Fluorescence responses of GRAB_{NE1m}-expressing cells in an acute LC slice to
- electrical stimulation of different pulses at 20 Hz in (J), or to the exogenous perfusion of
- 119 different concentrations of NE in (K). Left, pseudocolor snapshots of GRAB_{NE1m}
- 120 fluorescence responses. The white dash circles indicate ROI (50 µm in diameter) used for
- the fluorescence analysis. Right, corresponding fluorescence responses of left.
- 122 **(L)** Dose-dependent curve of fluorescence response to different concentrations of NE.
- Response data were fitted by the Boltzmann equation, and the evoked volume-averaged

- NE concentration ([NE]vol) was estimated based on the calibration curve from the same
- 125 **slice**.
- (M) Group data of the evoked [NE]_{Vol} during electrical stimulations (n = 4 slices from 3
- mice). Error bars indicate S.E.M.
- 128 The scale bar shown in (**B**) represents 100 μm. The scale bar shown in (**C**) and (**J**)
- 129 represent 10 μm.
- 130 ***p < 0.001 (Student's t-test).

| 131 | Figure S5. GRAB _{NE1h} can sense endogenous NE release and optic tectal neurons |
|-----|--|
| 132 | with or without HuC:GRAB _{NE1m} overexpression show no difference in spontaneous |
| 133 | calcium responses, related to Figure 5. |
| 134 | (A,B) Detection of endogenous NE release in the midbrain of GRAB _{NE1h} zebrafish |
| 135 | evoked by visual looming stimuli. Quantification data is shown in (B). n = 6 fish. |
| 136 | (C) Spontaneous calcium activities of optic tectal neurons revealed by jREGCO1a |
| 137 | fluorescent signals show no difference with- or without HuC:GRAB _{NE1m} expression. |
| 138 | Experimental diagram is shown in (C1). Traces for representative calcium responses are |
| 139 | shown in (C2). Group data are shown in (C3). n = 7 for transgenic HuC:GRAB _{NE1m} |
| 140 | zebrafish, and $n = 8$ for fish without expressing GRAB _{NE1m} . |
| 141 | |
| 142 | *p< 0.05 and ***p< 0.001; n.s., not significant (Student's t-test, Wilcoxon test, or Mann- |
| 143 | Whitney rank sum test). |
| 144 | |
| | |

- Figure S6. No detectable changes in noradrenergic activity are observed in freely
- moving mice after expression of the GRAB_{NEmut} sensor during stress, social
- interactions and food related behaviors, related to Figure 7.
- 148 (A) Schematic diagrams depicting the AAV virus injection, and recording sites.
- (B) Histology showing the expression of GRAB_{NEmut} (green) and placement of the
- recording; the nuclei were counterstained with DAPI (blue). Scale bar: 1 mm (left), 500µm
- 151 (right).
- 152 (C1-E5) Representative traces (C1-C5), average per-stimulus histograms (D1-D5), and
- summary data (**E1-E5**) showing normalized GRAB_{NE1m} fluorescence (ΔF/F) before and
- during the forced swim test (1), and before, between and during the tail suspension test
- 155 (2), the hand presentation test (3), social interaction with an intruder of the opposite sex
- (4), an intruder of the same sex (5) and presentation of peanut butter (6). n = 3 animals
- 157 each.
- 158 The Shapiro-Wilk normality test was performed; if the test revealed it followed a normal
- distribution, a paired Student's *t*-test or one-way repeated measures ANOVA followed by
- Tukey's multiple comparisons was performed. If the values did not follow a normal
- distribution, a non-parametric ANOVA (Friedman's test) was performed followed by
- Dunn's multiple comparisons test. In (C) and (D), the blue dotted lines represent the start
- of the stimulus, and the red dotted lines represent the end of the trial.
- $^*p < 0.05$ and $^*p < 0.01$.