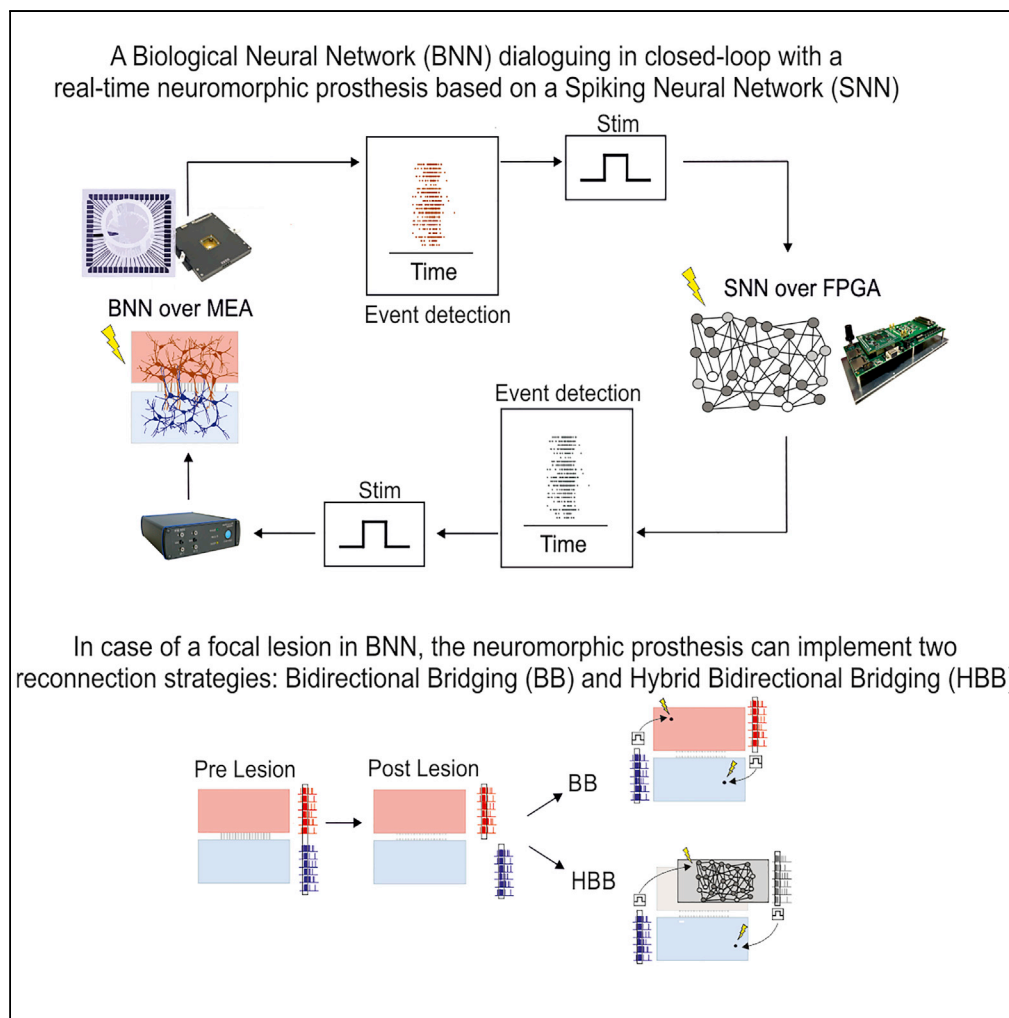


## Article

# A Neuromorphic Prosthesis to Restore Communication in Neuronal Networks



Stefano Buccelli,  
Yannick Bornat,  
Ilaria Colombi, ...,  
Paolo Massobrio,  
Timothée Levi,  
Michela  
Chiappalone

timothee.levi@u-bordeaux.fr  
(T.L.)  
michela.chiappalone@iit.it  
(M.C.)

## HIGHLIGHTS

We developed a novel real-time neuromorphic system acting as a neuroprosthesis

The neuroprosthesis can bidirectionally reconnect two disconnected networks

The neuroprosthesis implements a real-time spiking neural network

The spiking network can successfully replace the activity of a neural population

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## Article

# A Neuromorphic Prosthesis to Restore Communication in Neuronal Networks

Stefano Buccelli,<sup>1,2,3,13</sup> Yannick Bornat,<sup>4,13</sup> Ilaria Colombi,<sup>2,3,14</sup> Matthieu Ambroise,<sup>4,14</sup> Laura Martines,<sup>3,5,14</sup> Valentina Pasquale,<sup>3</sup> Marta Bisio,<sup>3,6</sup> Jacopo Tessadori,<sup>3</sup> Przemysław Nowak,<sup>5,7</sup> Filippo Grassia,<sup>4,8</sup> Alberto Avena,<sup>1,2,3</sup> Mariateresa Tedesco,<sup>5</sup> Paolo Bonifazi,<sup>9,10,11</sup> Francesco Difato,<sup>3</sup> Paolo Massobrio,<sup>5</sup> Timothée Levi,<sup>4,12,\*</sup> and Michela Chiappalone<sup>1,3,15,\*</sup>

## SUMMARY

**Recent advances in bioelectronics and neural engineering allowed the development of brain machine interfaces and neuroprostheses, capable of facilitating or recovering functionality in people with neurological disability. To realize energy-efficient and real-time capable devices, neuromorphic computing systems are envisaged as the core of next-generation systems for brain repair. We demonstrate here a real-time hardware neuromorphic prosthesis to restore bidirectional interactions between two neuronal populations, even when one is damaged or missing. We used *in vitro* modular cell cultures to mimic the mutual interaction between neuronal assemblies and created a focal lesion to functionally disconnect the two populations. Then, we employed our neuromorphic prosthesis for *bidirectional bridging* to artificially reconnect two disconnected neuronal modules and for *hybrid bidirectional bridging* to replace the activity of one module with a real-time hardware neuromorphic Spiking Neural Network. Our neuroprosthetic system opens avenues for the exploitation of neuromorphic-based devices in bioelectrical therapeutics for health care.**

## INTRODUCTION

One of the greatest challenges of modern neuroscience is to find reliable and sustainable treatments for the disabling effects caused by many chronic and incurable brain conditions. With the greatest impact carried by stroke (Feigin et al., 2017) and traumatic brain injury (Maas et al., 2017), brain disorders are among the leading causes of disabilities worldwide. Owing to recent advances in bioelectronics and in neural and neuromorphic engineering, direct interfacing of artificial circuits with large neuronal networks is possible to develop novel “neurobiohybrid” systems (such as neuroprostheses Vassanelli and Mahmud, 2016), which are envisaged as potentially interesting clinical applications for brain lesions (Broccard et al., 2017). In this paper, we introduce an innovative bioelectronic system acting as a neuroprosthesis, which, thanks to a neuromorphic real-time hardware interface, can re-establish the communication between two disconnected neuronal populations.

Neural interfaces are promising solutions for brain repair (Soekadar et al., 2015). Modern neural interfaces are mainly designed to restore lost motor functions in only one direction, i.e., from the brain to the body (Abdulkader et al., 2015) or from the body to the brain (Flesher et al., 2016). Additionally, recent neuroprosthetic developments have shown the enormous potential of neural interfaces to aid and accelerate functional recovery (Bouton et al., 2016; Rosin et al., 2011). However, a major obstacle in developing novel neuroprostheses for bidirectional communication with and within the brain is the complex nature of interactions among different brain areas, which in turn presents a challenge for the development of appropriate stimulation protocols as well as for testing such devices using *in vivo* models (Kohler et al., 2017).

Despite very recent technological progress (Jun et al., 2017; Steinmetz et al., 2018), *in vivo* models still have two main bottlenecks. The first bottleneck is the technical challenge to faithfully reproduce specific/focal network lesions (mainly due to their complexity) that the neuroprosthesis aims to treat, whereas the second is the difficulty in disentangling the actual effect of the adopted electrical therapy from the complex activity of a brain constantly processing sensory inputs and producing behavior. Therefore, since *in vivo* models exhibit inherent complexity and low controllability, using *in vitro* reduced neuronal systems to model

<sup>1</sup>Rehab Technologies IIT-INAIL Lab, Istituto Italiano di Tecnologia, Via Morego 30, 16163 Genova, Italy

<sup>2</sup>Department of Neuroscience, Rehabilitation, Ophthalmology, Genetics and Maternal and Child science (DINO GMI), University of Genova, L.go P. Daneo 3, 16132 Genova, Italy

<sup>3</sup>Department of Neuroscience and Brain Technologies, Istituto Italiano di Tecnologia, Via Morego 30, 16163 Genova, Italy

<sup>4</sup>Laboratoire de l'Intégration du Matériau au Système (IMS), University of Bordeaux, Bordeaux INP, CNRS UMR 5218, 351 Cours de la Libération, 33405 Talence Cedex, France

<sup>5</sup>Department of Informatics, Bioengineering, Robotics, System Engineering (DIBRIS), University of Genova, Via all'Opera Pia 13, 16145 Genova, Italy

<sup>6</sup>Department of Neurosciences, University of Padova, Via Nicolò Giustiniani 5, 35128 Padova, Italy

<sup>7</sup>Institute of Information Technology, Lodz University of Technology, ul. Wolczanska 215, 90-924 Lodz, Poland

<sup>8</sup>University of Picardie Jules Verne, Laboratory of Innovative Technologies (LTI, EA 3899), Avenue des Facultés, Le Bailly, 80025 Amiens, France

<sup>9</sup>School of Physics and Astronomy, Tel Aviv University, 69978 Tel Aviv, Israel

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precise and reproducible neuronal network lesions and test neuroprosthetic devices for their treatment may be advantageous. This approach is also justified by a growing recognition that *in vitro* testing of both research and medical devices can be more effective in terms of cost, time consumption, and ethical issues and much more reliable than *in vivo* testing (Myers et al., 2017).

In this work, we bidirectionally interfaced in real-time a neuroprosthetic system with an *in vitro* culture constituted by interconnected “neuronal assemblies” (Hebb, 1949). Therefore, our first objective was to create a simplified yet plausible *in vitro* model of a focal brain lesion by using bimodular cultures grown onto microelectrode arrays (MEAs) (Bisio et al., 2014; Bonifazi et al., 2013; Shein-Idelson et al., 2011) with reciprocal connections cut by a custom-made laser setup (Difato et al., 2011) to mimic the pathological effect of a traumatic brain injury (Hayes et al., 2016). We created a neurobiohybrid system connecting the biological element (the bimodular culture) following the lesion with a neuroprosthetic prototype. Our hardware neuroprosthesis could perform low-power computations in hard real-time (Pirog et al., 2018), collecting the inputs coming from neural recordings, processing those signals, and generating suitable electrical stimulation triggers as an output. With this experimental setup, we tested two specific applications, namely, *bidirectional bridging* (BB) to artificially reconnect two disconnected neuronal modules and *hybrid bidirectional bridging* (HBB) in which a real-time spiking neural network (SNN) replaced the activity of one of the two modules in real-time while implementing bidirectional connectivity with the remaining neuronal module.

The motivation of our research is to provide a new technological instrument as a novel form of neuroprosthesis aimed at treating disabling brain pathologies. The hardware choice (field-programmable gate array, FPGA) maximizes the real-time performances of the system and allows for a faster development of a future implantable bioelectronic device for biomedical applications. The adoption of bidirectional communication allows the development of a generalized non-specific approach that is applicable to the central nervous system (CNS) or peripheral nervous system. In particular, prostheses for the CNS should restore the communication between two or more neuronal assemblies whose functional and anatomical path could be distributed and sparse and not necessarily known *a priori*.

Indeed, our idea to develop a generalized approach comes from the future perspective of creating a cerebral neuroprosthesis for direct implantation in the brain that could be used by patients affected by stroke or brain injury. Our proof-of-principle results are the first for a next-generation neurobiohybrid system to restore brain functions (Broccard et al., 2017; Vassanelli and Mahmud, 2016).

## RESULTS

### Neuroprosthetic Architecture

To create bimodular *in vitro* systems, we developed Polydimethylsiloxane (PDMS) masks with two connecting compartments that constrained the growth of neuronal cells in two precise areas over 60-electrode MEAs (Figure S1). The obtained bimodular neuronal culture constitutes the biological neuronal network (BNN) of our system (Figure 1A). The signal from the BNN was amplified by a commercial system and acquired by a custom-developed FPGA-based neuromorphic board (Figure S2) previously configured by a custom-made MATLAB code (MathWorks, Natick, MA, USA) running on a general-purpose personal computer. The neuromorphic board triggered a commercial stimulator to close the loop with the BNN. The general protocol designed for this study involved three steps. First, spontaneous activity in both neuronal modules was recorded (“pre-lesion condition”). Then, laser ablation of the biological connections between the two modules (see Video S1) was performed (see Transparent Methods), followed by recording of spontaneous activity in both modules (“post-lesion condition”) to assess the viability of the networks. Finally, we tested our neuroprosthetic device using two experimental frameworks. In the first case, we applied a reconnection strategy using a bidirectional activity-dependent stimulation (i.e., BB paradigm), whereas in the second case, we interfaced a hardware-implemented biomimetic SNN with one of the two neuronal modules (i.e., HBB paradigm) to simulate a “replacement” strategy that utilizes the bidirectional interaction between the biological system and its artificial counterpart (Figure 1B). The sequence of algorithms (e.g., spike detection, network burst detection) implemented on the board to realize both experimental approaches (BB or HBB) is schematically depicted in Figure 1C.

### Control Experiments

To evaluate the stability and effect of the focal lesion on bimodular BNN activity, we first performed two sets of control experiments. In the first set defined as “experiments with no lesion” (Figure 2A1), we

<sup>10</sup>Computational Neuroimaging Laboratory, Biocruces Health Research Institute, Hospital Universitario Cruces, Baracaldo, Vizcaya 48903, Spain

<sup>11</sup>Ikerbasque: The Basque Foundation for Science, Bilbao, Bizkaia 48013, Spain

<sup>12</sup>LIMMS CNRS-IIS, The University of Tokyo, 153-8505 Tokyo, Japan

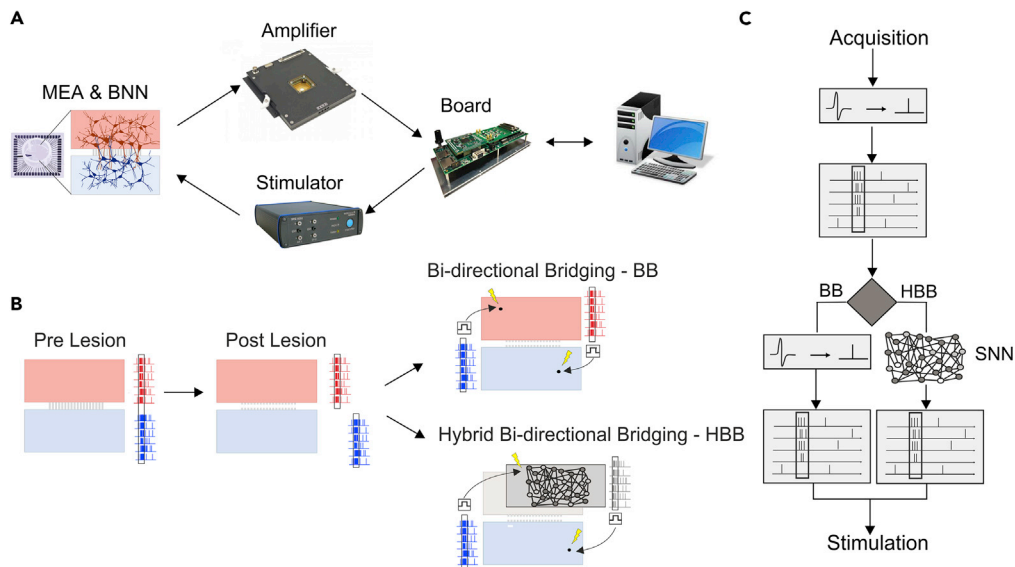
<sup>13</sup>These authors contributed equally

<sup>14</sup>These authors contributed equally

<sup>15</sup>Lead Contact

\*Correspondence: timothee.levi@u-bordeaux.fr (T.L.), michela.chiappalone@iit.it (M.C.)

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**Figure 1. Interfacing a Biological Neural Network and Neuromorphic Neuroprosthesis**

(A) Schematic representation of the main elements of the setup: cartoon of an MEA coupled with a BNN; picture of the amplification system; picture of the custom FPGA board; picture of the stimulus generator. Out of the loop, we used a PC to configure the board.

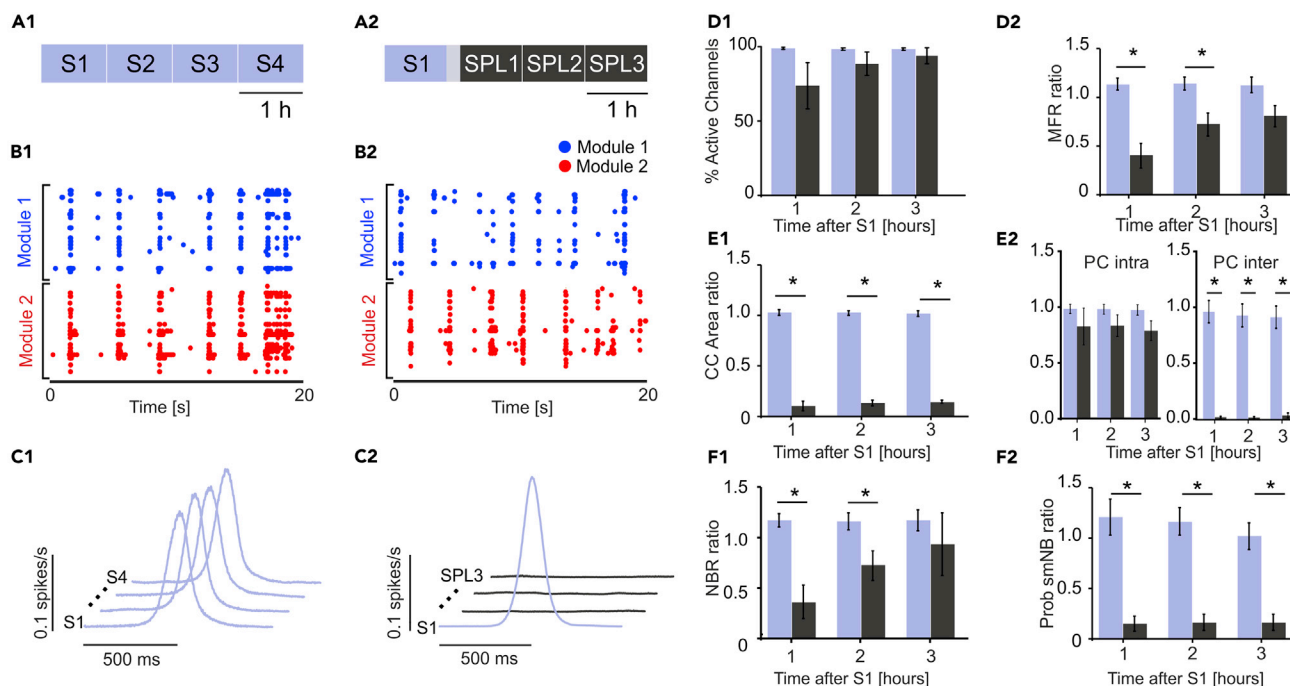
(B) Schematic representation of the different phases of two experimental approaches that share a pre-lesion, a lesion (performed through laser ablation, not shown), and a post-lesion phase. The final experimental phase can be either bidirectional bridging (BB) or hybrid bidirectional bridging (HBB).

(C) Schematic of real-time data processing performed by the board: the first step is spike detection followed by network burst (NB) detection monitoring module 1. After NB detection, delivering stimulation to module 2 of the BNN (BB approach) or to the SNN is possible. In the second modality, there is also NB detection of the SNN, which can result in stimulation delivered to the BNN (HBB approach).

recorded four consecutive hours of spontaneous activity (S1-S4,  $n = 9$  cultures). In the second set defined as “experiments with a lesion” (Figure 2A2), we recorded 1 h of spontaneous activity (S1), followed by laser ablation of the connections between the two modules, which usually took less than 20 min. Next, we recorded 3 h of spontaneous activity post lesion (SPL1-SPL3) to quantify the effects of laser ablation ( $n = 4$  cultures). As depicted in the raster plot of one representative experiment (Figure 2B1), bimodular neuronal networks exhibited spontaneous, synchronized, multi-unit activity composed of network-wide bursts (NBs) spreading over the two modules. Following laser ablation, the propagation between compartments was disrupted, as shown in Figure 2B2.

With no lesion, the percentage of active channels with respect to the first hour of recording (S1) was higher than 98% and was maintained for the entire duration of the experiment (Figure 2D1, light blue bars). The mean firing rate (MFR) was stable for all control experiments with no lesion (from S1 to S4, Figure S3A). Alternatively, control experiments with lesions showed a reduced number of active channels (close to 73%) during the first hour after ablation (SPL1). During the following 2 h (SPL2-SPL3), this value increased and reached 93% at the end of the recording (Figure 2C1, dark gray bars). No significant differences were found. The MFR was quite stable for all control experiments with lesions except between S1 and SPL1 (Figure S4A). The activity level with respect to the S1 phase, expressed by the MFR ratio with respect to S1, was stable during the control experiments without lesions (Figure 2D2, light blue bars). The lesion produced a clear decrease in activity in most cultures, especially during the first 2 h (SPL1-SPL2, Figure 2D2). We found a significant difference between the two experimental sets during the first two hours after S1 but not during the last hour. This result suggests that 2 h after a lesion, almost complete spontaneous recovery occurred in terms of the firing rate for the two neuronal modules.

To evaluate changes in the synchronicity between the two modules, we performed cross-correlation (CC) analysis between the collapsed spike trains of each module. The shape of the CC function was stable throughout the entire recordings in experiments without lesions, as reported in Figure 2C1 for a



**Figure 2. A Laser Ablation Induced Lesion Can Disconnect Two Neuronal Modules**

(A–F) (A1) Schematic of the first experimental protocol. Experiments with no lesion: we recorded four consecutive hours of spontaneous activity (S1–S4). (A2) Schematic of the second experimental protocol. Experiments with lesion: we recorded 1 h of spontaneous activity (S1) followed by laser ablation and three consecutive hours of spontaneous activity post lesion (SPL1–SPL3). The gray-shaded area indicates 20 min of no recording due to the execution of the lesion. (B1) A 20-s raster plot of the network bursting activity of one representative experiment during the S1 phase. (B2) A 20-s raster plot of the network bursting activity of one representative experiment during SPL3. (C1) Cross-correlation (CC) function for one representative experiment during the S1–S4 phases. The CC profiles between the spike trains of each module (light blue) in the four phases of the experiment were high and stable (lines shifted for the sake of clarity). Time axis [–500, +500] ms. (C2) CC profile between the spike trains of each module for one representative experiment with lesion. Before the lesion (light blue profile), CC was high; following the lesion (dark gray), CC collapsed to zero (lines shifted for the sake of clarity). Time axis [–500, +500] ms. (D1) Percentage of active channels with respect to S1 for the experiments with no lesions (light blue columns,  $n = 9$ ) and with lesions ( $n = 4$ , dark gray columns). No significant difference was found using the Mann-Whitney test (S2 VS SPL1:  $p = 0.2042$ ; S3 VS SPL2:  $p = 0.31608$ ; S4 VS SPL3:  $p = 0.70769$ ). (D2) Mean firing rate (MFR) ratio with respect to S1 for experiments without (light blue bars) and with lesions (dark gray bars). No significant difference was found during the last hour using the Mann-Whitney test (S2 vs SPL1:  $p = 0.0028$ ; S3 vs SPL2:  $p = 0.01119$ ; S4 vs SPL3:  $p = 0.10629$ ). (E1) Comparison of the CC area ratio with respect to S1 for the experiments without (light blue bars) and with lesions (dark gray bars) (Mann-Whitney test; S2 vs SPL1:  $p = 0.0028$ ; S3 vs SPL2:  $p = 0.0028$ ; S4 vs SPL3:  $p = 0.0028$ ). (E2) On the left, comparison of the intra-module correlation coefficient (i.e., Pearson Correlation, PC) ratio with respect to S1 for the experiments without (light blue bars) and with lesions (dark gray bars). No significant difference was found using the Mann-Whitney test (S2 vs SPL1:  $p = 0.71049$ ; S3 vs SPL2:  $p = 0.14825$ ; S4 vs SPL3:  $p = 0.07552$ ). On the right, the same comparison regarding inter-module PC that showed clear differences between experiments without and with lesion (Mann-Whitney test; S2 vs SPL1:  $p = 0.0028$ ; S3 vs SPL2:  $p = 0.0028$ ; S4 vs SPL3:  $p = 0.0028$ ). (F1) Network burst rate (NBR) ratio with respect to S1 showing significant differences during the first and second hour after the lesion (Mann-Whitney test; S2 vs SPL1:  $p = 0.0028$ ; S3 vs SPL2:  $p = 0.01119$ ; S4 vs SPL3:  $p = 0.26014$ ). (F2) Probability of single-module NB (Prob smNB). The ratio with respect to S1 shows stability for experiments without (light blue bars) and with lesions (dark gray bars) (Mann-Whitney test; S2 vs SPL1:  $p = 0.0028$ ; S3 vs SPL2:  $p = 0.0028$ ; S4 vs SPL3:  $p = 0.0028$ ). Data in the bar graphs is reported as mean  $\pm$  standard error of the mean.

representative experiment. After a lesion, the CC function collapsed to zero and did not recover during the experiment (Figure 2C2: representative experiment). To quantify this difference, we integrated the CC function in a range of  $\pm 500$  ms to obtain the CC area. We did not find any significant change in the CC area values for all experiments with no lesion (Figure S3B). By contrast, the CC area values showed a marked decrease following the lesion (SPL1). This decrease was due to the lack of anatomical connections between the compartments and did not recover by itself (Figure S4B). Comparing the CC area ratio between later phases and S1 resulted in a significant difference between matching periods in “lesion” and “no lesion” experiments (Figure 2E1). We also computed the correlation coefficient (i.e., Pearson Correlation, PC) among all the active channels both intra module and inter module (Figure 2E2). The intra-module PC was constant across experimental phases for controls with no lesion (light blue bars). On the other hand, for controls with lesion (dark gray bars) there was a drop in the intra-module PC, related to the reduced firing rate following the lesion, but no statistical difference was found (Figure 2E2, left panel). The inter-module CC was stable for controls with no lesion. On the other hand, following the lesion the inter-module

PC collapsed and never recovered by itself (Figure 2E2, right panel) as already demonstrated with the previous analysis.

The network bursting rate (NBR) was stable during all experiments with no lesions (Figure S3C). For the experiments with lesions, this parameter was less stable but with no significant differences between phases (Figure S4C). When comparing the two experimental protocols with the NBR ratio with respect to S1, we found significant differences during the first and the second hour post lesion (Figure 2F1). The mean probability to have NBs composed of spikes belonging to a single module (i.e., Prob smNB, see [Transparent Methods](#)) was close to 0.2 in the experiments without lesions (Figure S3D), meaning that the majority of NBs in an intact bimodular network involved both modules. Alternatively, following the lesion, the probability became close to 1 (Figure S4D), meaning a total loss of functional communication between the two compartments. Using the Prob smNB ratio with respect to S1 (Figure 2F2), we found significant differences between the two experimental groups during all phases post lesion (Mann-Whitney test;  $p < 0.05$ ). Thus, for the no lesion experiment, the Prob smNB remained very similar to the initial values, whereas for the lesion experiments, it changed abruptly due to the lesion. This result further confirmed that the lesion was effective in functionally disconnecting the two modules.

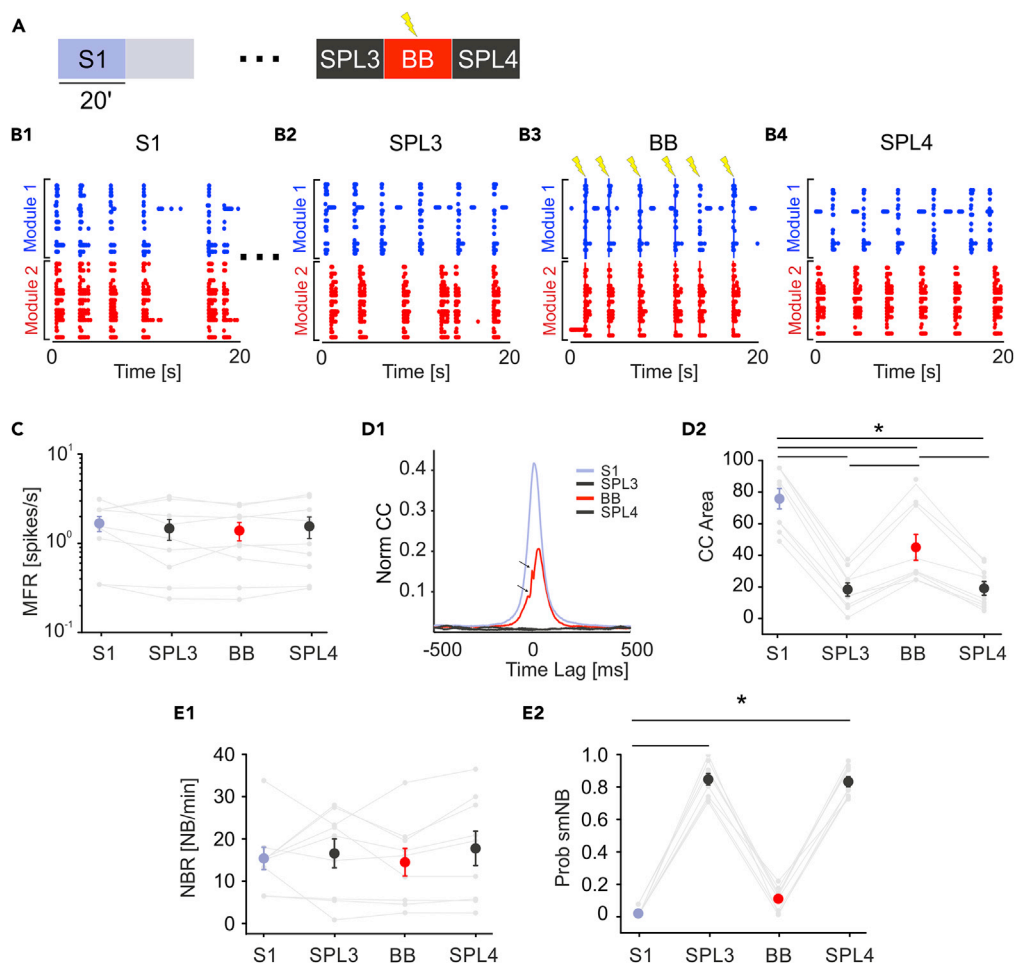
### Experiment 1: Bidirectional Bridging

The goal of this experiment is to restore communication between two neuronal assemblies after lesion-induced separation. To achieve this goal, we designed and implemented a stimulation reactive paradigm inspired by the “activity-dependent stimulation” (ADS) described in [Guggenmos et al. \(2013\)](#) in our neuro-morphic board. In contrast to the control experiments, the general protocol (Figure 3A) included 20-min recordings of spontaneous activity before the lesion (S1). Upon the lesion execution, we waited for 2 h to reach stable activity in both modules, as shown by the results of control experiments (see Figure 2). Then, we recorded 20 min of spontaneous activity (SPL3). The raster plot of a representative experiment is reported in Figure 3B. Before the lesion (S1), the bursting activity involved both modules (Figure 3B1), whereas after the lesion (SPL3), the activity was characterized by single-module NBs (Figure 3B2). To choose the best parameters (threshold and window time, see [Transparent Methods](#)) that allowed us to reliably detect NBs in both modules, we performed offline NB detection. After the FPGA was updated with these parameters, a 20-min session of BB was conducted (see Figures 1B and 1C for the description of the BB protocol). The BB approach implemented a reactive paradigm: every time an NB was detected in one module, a stimulation pulse was delivered to an electrode in the other module (see [Transparent Methods](#)) in both directions. During the BB phase, the bursting activity involved both modules similar to the intact condition due to the bidirectional stimulation pulses (Figure 3B3, blue and red lines represent electrical stimulation pulses delivered from module 1 to module 2 and vice versa). The last phase of the protocol involved 20 additional minutes of spontaneous activity (SPL4), which showed the same activity as SPL3 (Figure 3B4). We did not observe significant changes in spiking activity (i.e., MFR) throughout the recordings (Figure 3C).

Next, we evaluated the effect of this configuration in terms of CC (Figures 3D1 and 2). During spontaneous activity before the lesion (S1), the CC peak was high and stable due to the functional and anatomical connections between the two modules, which was also reported for the control experiments. After the lesion (SPL3), there was a decrease in CC that was not expected to recover without external intervention, as we demonstrated earlier (see Figure 2D). The bidirectional stimulation at least partially recovered the CC area and consequently the communication between modules (Figure 3D2), as demonstrated by statistical analysis. Regarding the number of NBs, we did not find any significant difference between the experimental phases (Figure 3E1). However, the probability of isolated NBs was not uniform; it reached the maximum value after the lesion, as we previously observed in the control experiments with lesions (see Figure 2D2). During bidirectional stimulation, these values became closer to the spontaneous recording (Figure 3E2), meaning that NBs mainly involved both modules. This finding further confirmed that the BB protocol could reconnect two disconnected modules through a real-time ADS acting in both directions (from module 1 to module 2 and from module 2 to module 1).

### Experiment 2: Hybrid Bidirectional Bridging

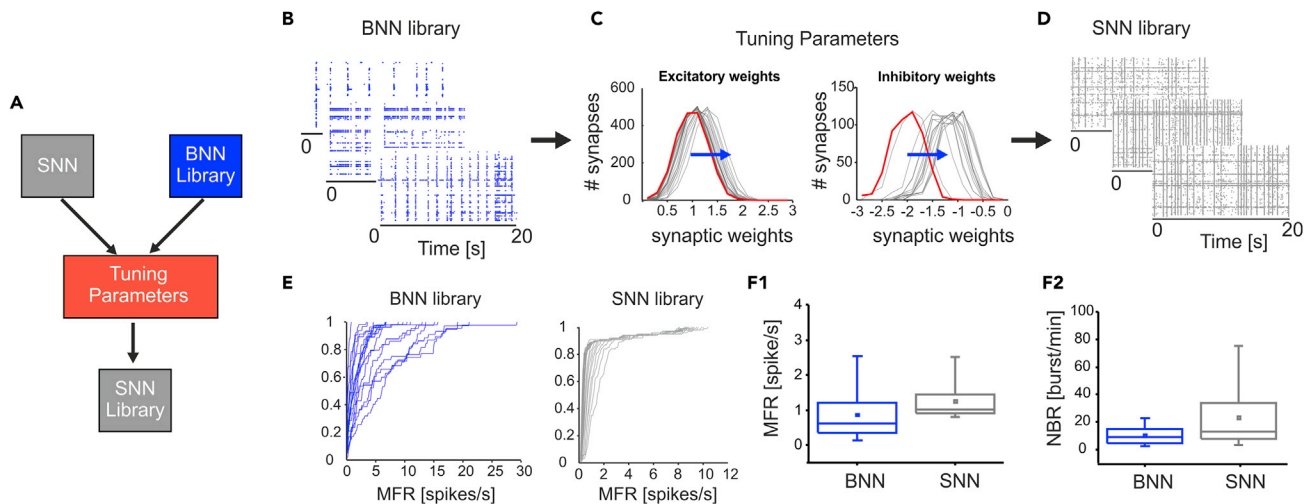
With an injury causing damage to an entire neuronal subnetwork, a reconnection strategy such as the BB illustrated earlier would not be feasible. For this reason, we developed a second reconnection strategy based on the use of a hardware SNN that can interact in real-time with its biological counterpart, HBB (see Figure 1). We created a set of SNNs (i.e., SNN library) by tuning the mean value of the synaptic weight



**Figure 3. Bidirectional Bridging Is Effective in Reconnecting Functionally and Anatomically Disconnected Neuronal Modules**

(A and B) (A) Schematic of the experimental protocol. We recorded 20 min of spontaneous activity (S1) followed by laser ablation. The gray-shaded area indicates 20 min of no recording during ablation. Dots represent 2 h of no recording after the lesion to maintain a stable activity in both modules. Then, we recorded 20 min of SPL activity (SPL3) followed by 20 min of the bidirectional bridging (BB) protocol and another 20 min of spontaneous activity (SPL4). (B1–B4) The 20-s-long raster plots of representative experiments (respectively, from phases S1, SPL1, BB, and SPL4). In (B3) blue and red lines represent electrical stimulation pulses delivered from module 1 to module 2 and vice versa, respectively. (C–E) (C) MFR during the four experimental phases was stable (color code as in panel A: S1: light blue dot; SPL3, SPL4: dark gray dots; BB: red dot). No significant difference was found (one-way RM ANOVA,  $p = 0.469$ ,  $DF = 3$ ,  $F = 0.872$ ). (D1) CC function during the four experimental phases. Small arrows indicate the blanking period of 8 ms following each stimulation. Color code, the same as that in panel A. Note that during BB, the cross-correlation function (red) recovers even if not completely with respect to the initial profile (light blue), whereas it stays at zero during the spontaneous activity phases post lesion (SPL3 and SPL4, dark gray profiles). (D2) CC area was highly reduced during the post-lesion phases. The CC area partially recovered during the BB protocol and collapsed again when stimulation was switched off (one-way repeated measures analysis of variance; degrees of freedom = 3;  $F = 101,832$ . S1 vs SPL3  $p = 5.67 \times 10^{-13}$ ; S1 vs SPL4  $p = 7.54 \times 10^{-13}$ ; S1 vs BB  $p = 1.60 \times 10^{-7}$ ; BB vs SPL3  $p = 1.77 \times 10^{-6}$ ; BB vs SPL4  $p = 2.81 \times 10^{-6}$ ; SPL4 vs SPL3  $p = 1$ ). (E1) NBR remained stable during the experiments. No significant difference was found (one-way repeated measures analysis of variance:  $p = 0.501$ ,  $DF = 3$ ;  $F = 0.810$ ). (E2) Probability of the single-module NB (Prob smNB) was close to one after the lesion. During the BB protocol, the probability was similar to the pre-lesion condition (Friedman’s repeated measures analysis of variance;  $p < 0.001$ ,  $DF = 3$ , Chi-square = 24.3. SPL4 vs S1 and SPL3 vs S1:  $p < 0.001$ ). Data in the plots is reported as mean  $\pm$  standard error of the mean.

distributions of our models to cover the variability of the BNNs (i.e., BNN library, Figure 4A). The biomimetic SNN (see Figure S5), working in hardware real-time to allow bidirectional communication with living neurons, was modeled as a network of 100 Izhikevich (IZH) neurons (Izhikevich, 2003), with 80 excitatory and 20 inhibitory neurons (see Transparent Methods), according to the biological composition of dissociated



**Figure 4. Spiking Neural Network (SNN) Design and Characterization**

(A) Schematic of the procedure used to create a library of SNNs. Starting from the Izhikevich model implemented on the FPGA and a library of 34 BNNs with a large spectrum of activity, we tuned the mean value of the synaptic weight distribution to obtain and select from a collection of SNNs (SNN library, comprising 27 different configurations).

(B) Representative 20-s-long raster plots of different BNNs showing different NB rates.

(C) Left, distribution of excitatory synaptic weights from the 27 SNNs. In red, the slower SNN of the library (SNN 1). The blue arrow indicates the shift of the mean value (from 0.99 to 1.34) of the normal distribution with standard deviation = 0.3 to obtain increasing NBR values. Right, distribution of inhibitory synaptic weights from the 27 SNNs. In red, the slower SNN of the library (SNN 1). The blue arrow indicates the shift of the mean value (from  $-2.02$  to  $-1.02$ ) of the normal distribution with standard deviation = 0.3 to obtain increasing NBR values.

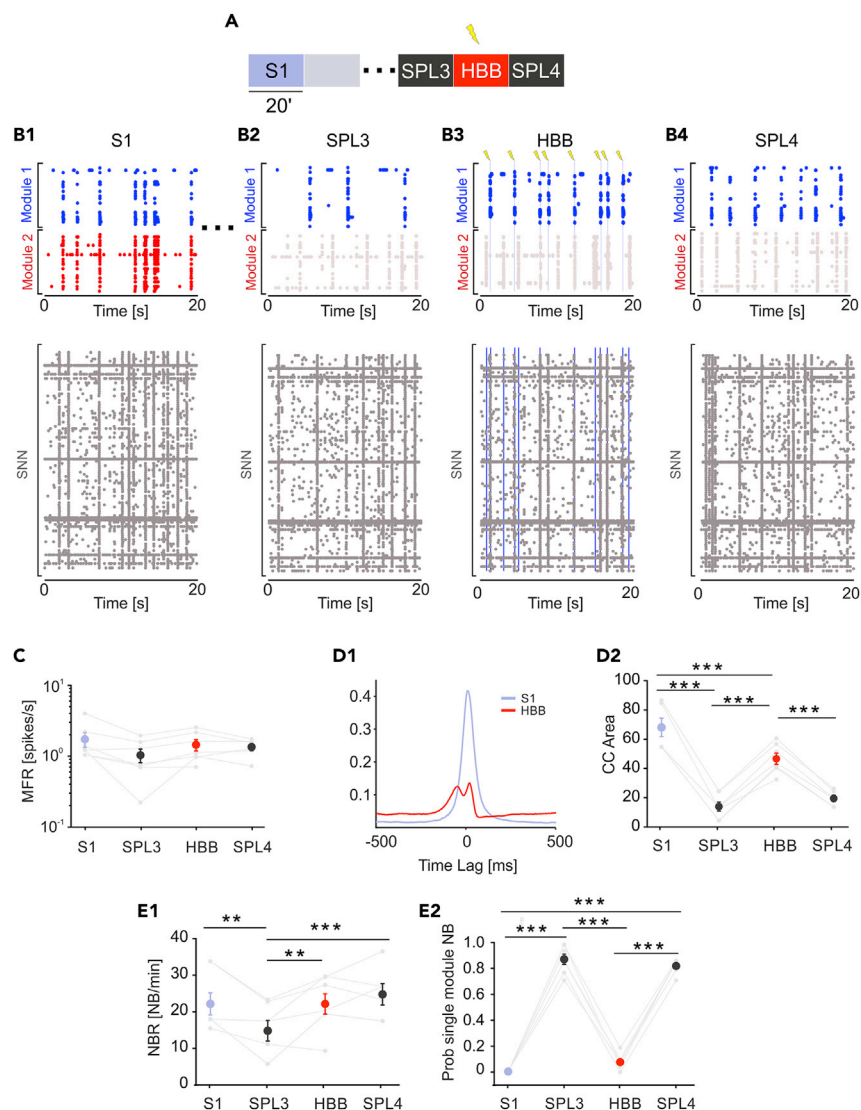
(D) Representative 20-s-long raster plots of different SNNs showing different NB rates.

(E and F) (E) Left, cumulative MFR profile for the BNN library. Right, cumulative MFR profile for the SNN library. (F1) Box plot showing the comparison between BNN and SNN libraries in terms of network MFR (i.e., the mean value of all active electrodes for BNN and neurons for SNN). (F2) Box plot showing the comparison between BNN and SNN NBR, showing that the SNN library covers the BNN variability and contains networks with a higher NBR. For each box plot (F1- F2), the small square indicates the mean, the central line illustrates the median and the box limits indicate the 25th and 75th percentiles. Whiskers represent the 5th and the 95th percentiles.

cultures (Bonifazi et al., 2005; Hayashi et al., 2003). Synaptic noise (Grassia et al., 2016), inhibitory and excitatory synapses (Izhikevich, 2004), short-term plasticity (Izhikevich and Edelman, 2008), and axonal delays were included in the model to recreate the network dynamics (see [Transparent Methods](#) and [Figures S6A1](#) and [S6A2](#)). Regarding the connectivity rules, we set the outdegree (i.e., the number of post-synaptic neurons) to 25 for all neurons in the network, whereas the indegree (i.e., the number of pre-synaptic neurons) followed a normal distribution with a mean value of 25 and a standard deviation of 4.3 ([Figure S6](#), R-Square = 0.806). The goal of creating an SNN library was to cover a wide range of NBRs because NB was chosen as the triggering event for our reconnection paradigm, as explained earlier. To this end, we tuned only the mean value of the normal distribution of synaptic weights (the standard deviation was kept constant at the value of 0.3). By increasing or decreasing the mean synaptic weights, we tuned the NBR. For excitatory synapses, the mean value ranged from 0.99 to 1.34 ([Figure 4C](#) left), whereas that for inhibitory synapses ranged from  $-2.02$  to  $-1.02$  ([Figure 4C](#) right and [Table S1](#)). As previously stated, our goal was to cover the NBR variability and not the MFR. The MFR variability in our BNN library was higher than that obtained with our SNN library ([Figures 4E](#) and [4F1](#)). Nevertheless, the BNN variability in terms of NBR was completely covered by our SNN library, which also contains networks with a much higher NBR than that in the BNN library ([Figures 4F2](#)).

The general HBB protocol ([Figure 5A](#)) is similar to the BB protocol. The HBB procedure included a 20-min recording of spontaneous activity before the lesion. This recording was used to quantify activity in terms of the NB rate of the network. This feature was used to choose one SNN from the SNN library that had an NB rate closer to its biological counterpart. We waited 2 h after the lesion to allow activity in both modules to stabilize, as shown by the results of control experiments (e.g., [Figure 2](#)). Then, we recorded 20 min of spontaneous activity. After setting FPGA detection parameters, we performed a 20-min HBB session (see [Figures 1B](#) and [1C](#) for the description of the HBB protocol). As anticipated, the HBB approach also implemented an ADS paradigm; every time an NB was detected on the “surviving” module (i.e., when one of the





**Figure 5. The Hybrid Bidirectional Bridging Approach Is Effective When a Neuronal Assembly Must Be Replaced**

(A and B) (A) Schematic of the experimental protocols. We recorded 20 min of spontaneous activity (S1) followed by laser ablation. The gray-shaded area indicates 20 min of no recording during ablation. Dots represent 2 h of no recording after the lesion to obtain stable activity in both modules and to test different stimulation channels. Then, we recorded 20 min of SPL activity (SPL3) followed by 20 min of a hybrid bidirectional bridging (HBB) protocol and another 20 min of spontaneous activity (SPL4). (B1) Top, 20-s-long raster plot depicting the BNN bursting activity involving both modules before lesion. Bottom, activity of SNN uncorrelated with the BNN. The networks are not linked. (B2) Top, 20-s-long raster plot after lesion showing uncorrelated bursting activity on BNN modules 1 and 2. Bottom, same as that in B1. (B3) Twenty-second-long raster plot during HBB depicting two hybrid events. The first event on the left was an NB detected on module 1 of the BNN. The detection resulted in a stimulation pulse delivered to 10 excitatory neurons of the SNN (blue line, bottom). An NB on the SNN was detected 18 ms after the stimulation and triggered the delivery of a stimulation pulse to module 1 of the BNN (gray line, top). (B4) Twenty-second-long raster plot depicting the uncorrelated activity of BNN modules (top) and SNN network (bottom). (C–E) (C) MFR during the four experimental phases was stable (color code as in panel A: S1: light blue dot; SPL3, SPL4: dark gray dots; HBB: red dot). No significant difference was found (one-way repeated measures analysis of variance.  $p < 0.001$ ,  $DF = 3$ ,  $F = 3.16$ ; Bonferroni test: all comparisons with  $p > 0.05$ ). (D1) CC function during the four experimental phases. Color code the same as that in panel A. Note that during BB, the cross-correlation function (red) recovers even if not completely with respect to the initial profile (light blue). (D2) CC area was highly reduced during the post-lesion phases. The CC area partially recovered during the BB protocol and collapsed again when stimulation was switched off (one-way repeated measures analysis of variance.  $p < 0.001$ ,  $DF = 3$ ;  $F = 70,448$ ; S1 vs SPL3:  $p = 5.80E-10$ ; S1 vs SPL4  $p = 2.72 \times 10^{-9}$ ; S1 vs HBB:  $p = 9.16 \times 10^{-4}$ ; HBB vs SPL3  $p = 1.16 \times 10^{-7}$ ; HBB vs SPL4  $p = 1.02 \times 10^{-6}$ ; SPL4 vs SPL3  $p = 0.73643$ ). (E1) NBR did not change during HBB with respect to the S1 phase (one-way repeated measures analysis of variance.  $p = 0.005$ ,  $DF = 3$ ;  $F = 6,069$ ; S1 vs SPL3  $p = 0.02482$ ; S1 vs

**Figure 5. Continued**

SPL4  $p = 1$ ; S1 vs HBB  $p = 1$ ; HBB vs SPL3  $p = 0.01022$ ; HBB vs SPL4  $p = 1$ ; SPL4 vs SPL3  $p = 0.00674$ ). (E2) Probability of a single-module NB (Prob smNB) was close to one after the lesion. During the HBB protocol, the probability was similar to that in the pre-lesion condition (one-way repeated measures analysis of variance.  $p < 0.001$ ,  $DF = 3$ ;  $F = 453,439$ ; S1 vs SPL3  $p = 4.96 \times 10^{-13}$ ; S1 vs SPL4  $p = 1.03 \times 10^{-12}$ ; S1 vs HBB  $p = 0.22606$ ; HBB vs SPL3  $p = 1.94 \times 10^{-12}$ ; HBB vs SPL4  $p = 4.30 \times 10^{-12}$ ; SPL4 vs SPL3  $p = 1$ ). Data in the plots is reported as mean  $\pm$  standard error of the mean.

two modules was completely damaged), a stimulation pulse was delivered to the SNN. The board implemented the corresponding paradigm in the opposite direction. Detection of NBs occurred in the SNN, whereas stimulation was delivered to the “surviving” module, thus avoiding the imposition of any predefined unidirectional communication. Next, we recorded 20 additional minutes of spontaneous activity.

We did not observe significant changes in terms of spiking activity (i.e., MFR) throughout the recordings (Figure 5C). Then, we evaluated the effect of this configuration in terms of CC (Figures 5D1 and 2). During spontaneous activity before the lesion (S1), the CC peak was high and stable due to the functional and anatomical connections between the two modules, which was also reported for the control experiments. As expected, with no external intervention, CC decreased sharply after the lesion (SPL3), as we previously observed. One of the two modules was damaged, whereas the correlation was evaluated between the SNN and the surviving module during the HBB phase. The bidirectional stimulation created a relevant correlation area between SNN and the surviving module, as demonstrated by statistical analysis. Regarding the number of NBs, we did not find a significant difference between the S1 and HBB phases (Figure 5E1). However, the probability of isolated NBs was not uniform; it reached the maximum value after the lesion, as we previously observed in the control experiments with lesions (see Figure 3E2). During the hybrid bidirectional stimulation, these values became closer to the spontaneous recording (Figure 5E2), meaning that NBs mainly involved both modules. This finding further confirmed that the HBB protocol created a hybrid system with the surviving biological module though real-time ADS acting in both directions (from BNN to SNN and from SNN to BNN).

**DISCUSSION**

We presented an innovative neuromorphic prosthesis based on an FPGA board and demonstrated two successful reconnection paradigms for a lesion interrupting the communication between two neuronal populations *in vitro*.

According to previous reports, *in vitro* systems constitute a successful experimental model of neuronal dynamics (Javier et al., 2013; Johnson et al., 2010), thus providing an excellent test bed for adaptive closed-loop neural interfaces (Potter, 2010). Starting from our recently developed methodology (Kanner et al., 2015), we created custom bimodular cultures with the goal of reproducing two interacting neuronal populations, thus mimicking the intrinsic modularity of the brain (Bonifazi et al., 2013). Our bimodular cultures were highly temporally stable in terms of firing properties at the whole network level as the activity between the two populations remained highly correlated for the entire duration of the recording. A lesion produced via laser ablation was employed to physically cut the connections between the two modules. This methodology was proven to be safe because it produced localized damage by selectively ablating subcellular compartments without damaging adjacent structures (Difato et al., 2011; Habibey et al., 2015; Soloperto et al., 2016). We assume that such a focal damage, allowing to specifically cut few connections among those available in the network, together with possible intrinsic compensatory mechanisms of synaptic scaling (le Feber et al., 2017; Turrigiano, 2008), was responsible for the spontaneous recovery of the firing rate on a timescale of 2–3 h. This demonstrates the effectiveness of our technique in preserving the functionality of the two modules while decoupling their activity, as proven by the loss of correlation of bursting behavior.

Two different applications of our neuroprosthesis, BB and HBB, were tested. Our neuromorphic prosthesis, independently of the stimulation paradigm, works according to a closed-loop reactive policy as follows: each time a condition is met (i.e., an “event” is detected), a stimulus is delivered. The hardware architecture was designed to be flexible enough to allow the implementation of different experimental paradigms and the definition of different triggering events. In our study, we chose NBs as trigger events (see Transparent Methods). The choice to deliver a stimulation depending on a network-wide event has two main advantages as follows: first, NB frequency is low enough to avoid inducing plasticity phenomena by electrical stimulation in our cultures (Wagenaar et al., 2006), which could confound the final results and effectiveness of neuroprosthetic reconnection. The second point is anticipation of the following major issue that will

emerge during *in vivo* recordings: monitoring single neurons presents problems at both theoretical (Guggenmos et al., 2013) and practical levels. Namely, how much information on complex functions can be obtained by single-neuron observation remains unclear (Luczak et al., 2015; Panzeri et al., 2017), whereas tracking the activity of the same neuron for extended periods of time is problematic (Kozai et al., 2015). Taking multiple input sources into account was also used in the work of Berger (Berger et al., 2012), but they employed a neuroprosthetic strategy different from ours. In particular, these authors used a generalized linear model to predict the CA1 activity from spikes in CA3 of the hippocampal circuit. Our system is considered more flexible and adaptable to networks with different connectivity, not just feedforward similar to that in the hippocampus. Moreover, we were interested in mimicking the overall spiking activity of the network and not mapping an input-output transformation only.

Another important novelty of our system regards directionality. To the best of our knowledge, this neuroprosthetic system is the first to implement a truly bidirectional interaction with an SNN through a hard real-time interface. We recorded activity from the first module (via multiple sources); when a criterion was met, the device stimulated the second module (this is how a “typical” closed-loop in neuroscience works; for a review see Greenwald et al., 2016). The novelty is simultaneous monitoring of multiple sources from another module and delivery of the stimulation when the triggering event is detected. To date and as far as we are aware of, only Jung and colleagues (Jung et al., 2001) have performed a bidirectional interface to a neuromorphic device, but their models were not precise at the spike level (modeling neuron populations) and they used non-configurable analogue electronics, which resulted in an experiment-specific setup. The other neuroprosthetic devices that have been proposed in the literature can implement a “unidirectional” artificial link only from one area to another (or maybe the same) but not double it. Here, we are not imposing any preferred directionality to the communication; networks are self-organizing on the basis of their intrinsic natural relationship (we are not imposing who is driving whom). This approach has the main advantage of informing both brain regions (i.e., neuronal modules, in our case) that an event occurred in the other region, given that interaction in the brain is intrinsically bidirectional (Roelfsema and Holtmaat, 2018). For example, in the sensorimotor system, sensory stimulation can help motor recovery (Cuppone et al., 2018) and motor learning can enhance sensory functions (Ostry et al., 2010; Takeuchi and Izumi, 2013). Applications of our neuroprosthetic systems to conditions where the sensorimotor interaction is impaired would allow restoration of both communication channels, suggesting improvements in current rehabilitation therapies. Moreover, although tested on a bimodular system, the neuromorphic FPGA board can be easily upgraded to play the bridging role on an arbitrary number (within reason) of different neuronal circuits. A recent work (Forró et al., 2018) developed directional networks of primary hippocampal neurons on MEA and compared the information flow of these networks with respect to bidirectional networks (similar to our bimodular preparations). They found that, without physically imposing a unidirectional configuration, there is a continuous back and forth communication between nodes thus suggesting the importance of a bidirectional communication in a healthy network.

The second paradigm we tested was based on the use of a biomimetic SNN to “substitute” a missing/damaged neuronal population. Currently, SNN applications span different fields, including computational neuroscience (Markram, 2012; Melozzi et al., 2017), and very recently, they were used for sensory encoding in hand prosthesis for amputees (Osborn et al., 2018; Valle et al., 2018). SNNs can be simulated in software (Goodman and Brette, 2009) and/or neuromorphic hardware (Thakur et al., 2018). As time and energy consumption are fundamental in neuroprosthetic applications for translational purposes, the use of hardware-based computing systems becomes mandatory.

In general, hybrid systems composed of *in vitro* BNNs coupled to SNNs are rare. In one approach, the SNN served as a self-organizing classifier of activity patterns exhibited by the BNN, with output of the SNN being subsequently used to control the behavior of a robot (Pizzi et al., 2009). Other studies focused on the unidirectional or bidirectional influence of the two networks, investigating the dynamics of the interaction between the BNN and SNN in which the SNN played a role of an artificial counterpart of its biological original (Bruzzone et al., 2015; Chou et al., 2015). However, closed-loop effects in those hybrid networks were not thoroughly determined. In one of these studies, only unidirectional connectivity was considered with input from the SNN to the BNN, which was also simulated beforehand (Bruzzone et al., 2015). In this study, we established hybrid communication in the case of an entire neuronal population that needed substitution.

A study by Chou et al. (2015) implemented a bidirectional interface between an SNN and a retinal slice obtained from an adult rat and recorded by an MEA. This system is quite interesting, but there is a 1-s delay

between the BNN and SNN interactions. Therefore, this delay in Chou's setup is three orders of magnitude larger than that in our work in which the sampling of biological activity is never interrupted and the step size of the SNN is 1 ms. The difference between the two systems is radical; bidirectional communication in real-time allows actual clinical application, whereas delays in the range of seconds prevent (or at least seriously reduce) the possibility of meaningful control of a biological system.

A recent study, inspired by a previous work (Hogri et al., 2015), implemented a hybrid interaction (Xu et al., 2018) between the cerebellum of a rat and an SNN implemented on FPGA. Their model involved 10k leaky integrate and fire (LIF) neurons and did not integrate other biomimetic behaviors, such as axonal delay, short-term plasticity, and synaptic noise, unlike the IZH neurons implemented in our system. Both the hard real-time processing and simplified neuronal model (which allow mimicking the richness of the electrophysiological patterns *in vivo*) are mandatory for reproducing the biological dynamics of living neural networks and for performing useful real-time hybrid experiments.

In this work, we demonstrated the possibility of designing a neuromorphic all-hardware prosthesis capable of artificially reconnecting two disconnected neuronal networks or artificially replacing one entire neuronal sub-network.

The use of a fully integrated hardware computing system allowed hard real-time performances and low power consumption, which are crucial for translational purposes related to therapeutic applications in humans (Kipke et al., 2008; Wang et al., 2010).

### Limitations of the Study

A limitation of the present work is that we deliberately chose to downsample both the number of biological neurons recorded through a low-density MEA and the number of artificial neurons implemented on the FPGA. For the purpose of detecting network-wide activity, this oversimplification of biological complexity can be acceptable to test the functionality of the device and the feasibility of the approach, but if the goal is to functionally replace a biological network, a higher resolution would be preferable. It is worth underlining that this hardware implementation is not exploiting the full resources of the FPGA, thus in follow-up studies, also thanks to the flexibility of our system, we foresee to scale up the number of neurons and synapses and to upgrade the computational algorithms to deal with more complex experimental designs. From a technological point of view, the current state of the art makes possible the use of devices with a large number (thousands) of recording electrodes (Berdoncini et al., 2009; Frey et al., 2009). Such an improvement would also allow one to have more information about functional connectivity of the biological network (Pastore et al., 2018) and thus develop more realistic, in terms of topology, artificial models. In this work, we arbitrarily modeled the network connectivity with a random adjacency matrix since the use of MEAs with 60 electrodes made impossible to correctly identify the topological properties (e.g., hubs, recurring connections, modules) of the network under investigation.

We are aware that the road is still long to target human applications. Despite this, we think that the extensive work performed represent an important milestone to start from. The next fundamental (and critical) step would be to test the neuromorphic prosthesis *in vivo*, for example, on animal models affected by ischemic or traumatic lesions (Guggenmos et al., 2013). Even if the adaptation to the new experimental setups will require time, we believe these are necessary steps to further push the translational potential of our system, which will be able to create real innovation in the clinical therapeutics.

### METHODS

All methods can be found in the accompanying [Transparent Methods supplemental file](#).

### SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.isci.2019.07.046>.

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## AUTHOR CONTRIBUTIONS

T.L., Y.B., P.M., P.B., and M. C. designed the study. Y.B. and T.L. designed and fabricated the hardware board. S.B., V.P., F.D., and M.C. designed the experiments. M.A., P.M., P.N., F.G., and T.L. designed and worked on SNN. I.C., M.B., and M.T. prepared the bimodular cultures. S.B., L.M., A.A., and F.D. performed the experiments. S.B., L.M., J.T., V.P., and M.C. designed the analyses. S.B. and L.M. performed the analyses. S.B., I.C., and V.P. performed the statistical analyses. S.B. and I.C. prepared the original figures. S.B., I.C., T.L., and M.C. wrote the manuscript. T.L., P.M., P.B., and M.C. acquired funding to conduct the research. All authors have read, corrected, and approved the final version of the manuscript.

## DECLARATION OF INTERESTS

The authors report no competing interests.

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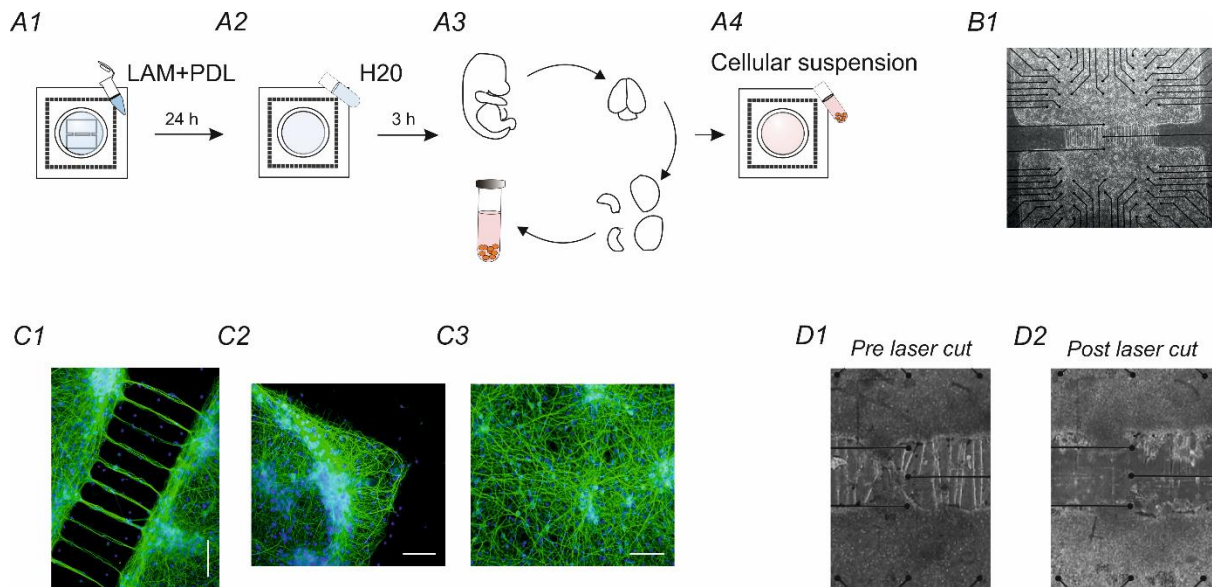
## **Supplemental Information**

### **A Neuromorphic Prosthesis to Restore Communication in Neuronal Networks**

**Stefano Bucci, Yannick Bornat, Ilaria Colombi, Matthieu Ambroise, Laura Martines, Valentina Pasquale, Marta Bisio, Jacopo Tessadori, Przemysław Nowak, Filippo Grassia, Alberto Averna, Mariateresa Tedesco, Paolo Bonifazi, Francesco Difato, Paolo Massobrio, Timothée Levi, and Michela Chiappalone**

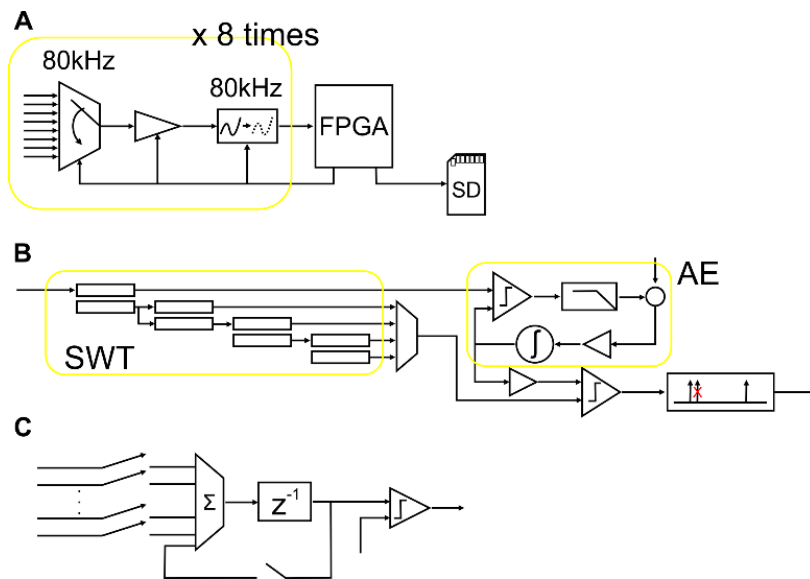
## Supplemental Information

### Supplemental Figures

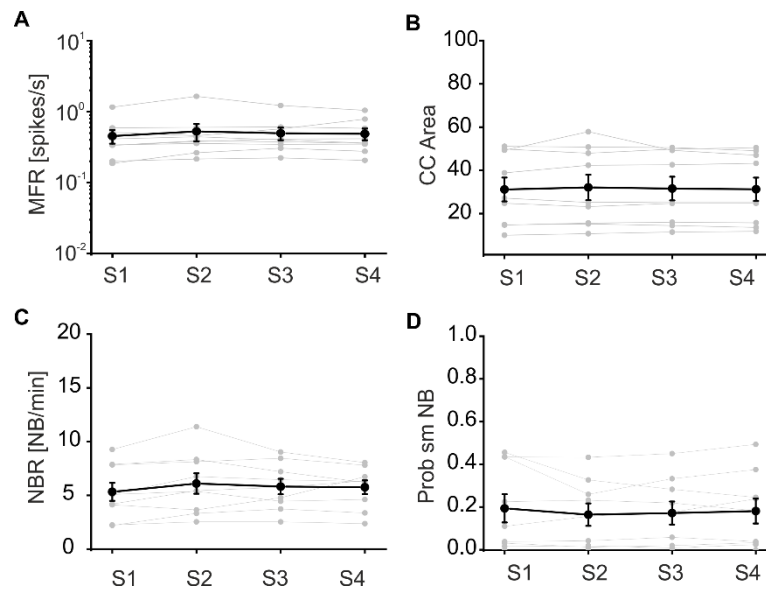


**Figure S1. Bimodular neuronal network preparation steps and morphological characterization, Related to Figure 2.** **A1** MEA surface coating using poly-D-lysine (PDL) and laminin (LAM) to promote cell adhesion. **A2** Three consecutive surface rinsing steps using sterilized water and 3 hours of waiting before plating cells. **A3** Dissection and dissociation procedures of E18 rat neocortex followed by resuspension of the resulting tissue in Neurobasal medium (see 'Methods'). **A4** Cellular solution plating onto a 60-channel MEA surface. **B1** A 60-channel MEA (Multichannel Systems MCS, Reutlingen, Germany) with square layout (4Q) plated with a bimodular pattern. **C1** Immunofluorescence micrograph of MAP2 of a representative bimodular culture on a coverslip at DIV 25 with magnification of the connections between modules. **C2**, **C3** Immunofluorescence micrographs of MAP2 of a representative uniform culture and of one corner on a coverslip at DIV 25. Scale bar = 200  $\mu\text{m}$ . **D1** Magnification of the connections between the two modules before performing the laser cut. **D2** Magnification of the cut connections between the two modules.

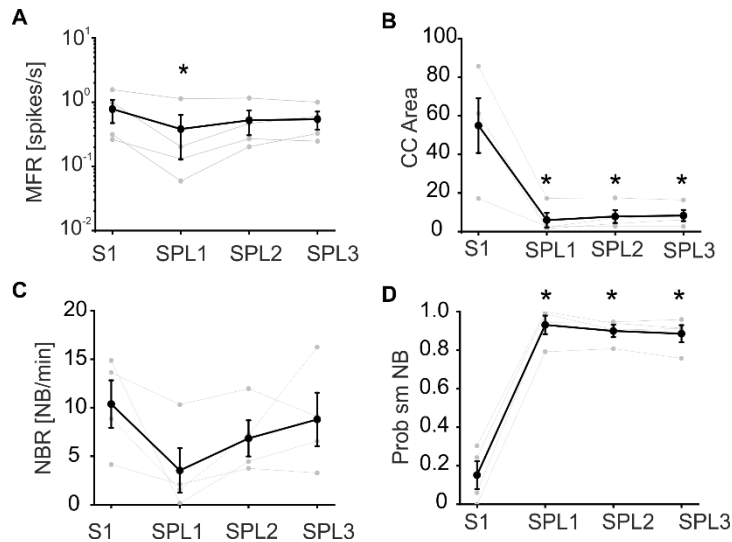




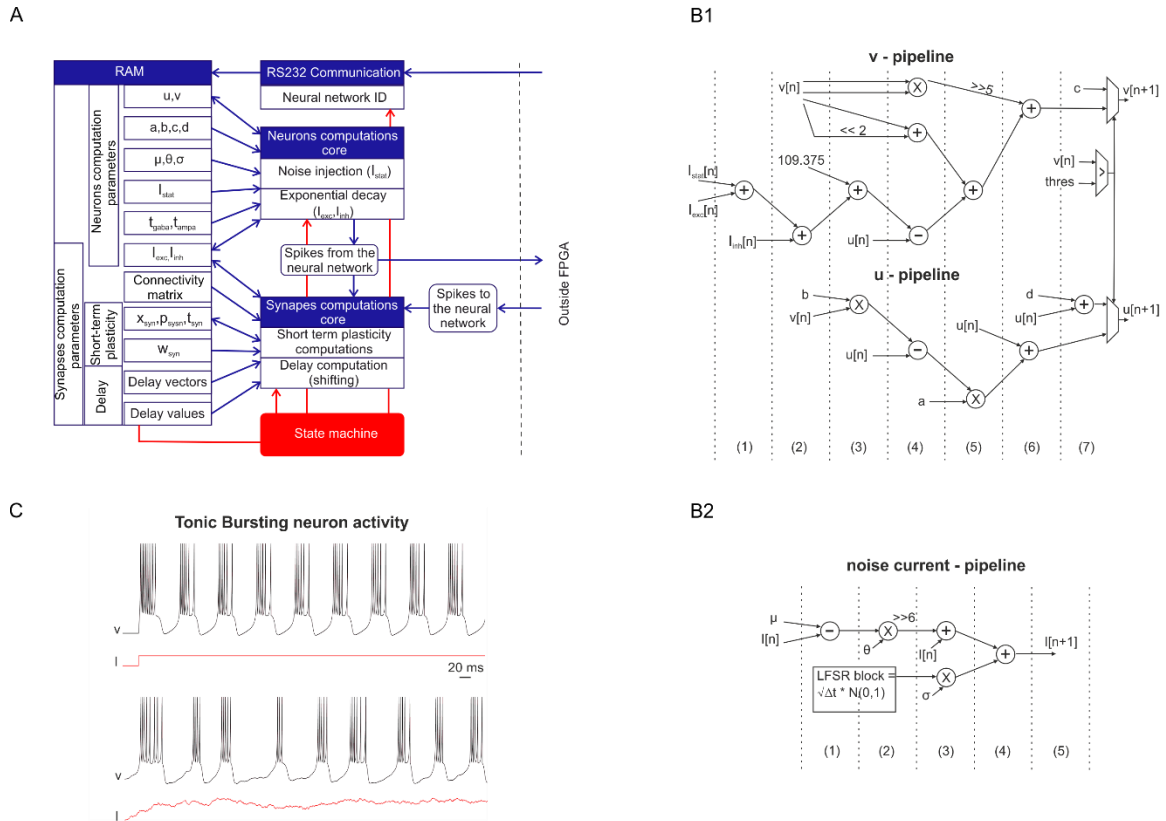
**Figure S2. Schematic view of the hardware computation architecture, Related to Figure 1. A,** Integrated circuit data flow on the device (Multiplexer: ADG1408, amplifier: LTC6912, analogue to digital converter: AD7686, FPGA: XC6SLX150). Each flow makes it possible to sample 8 channels at 10 kHz (80 kHz multiplexing), eight equivalent structures are connected to the FPGA to sample 64 inputs, and 60 of these inputs are connected to the MEA. **B,** Hardware spike detection architecture embedded in the FPGA. SWT represents the wavelet-based signal enhancement (stationary wavelet transform). The SWT consists of a series of FIR filters. The first two filters (on the left) are on the 8th order, and the following filters are on the 16th order with 8 null coefficients. AE represents the signal amplitude estimator used to set the final threshold. The loop estimates the standard deviation as the value above 15.9% of samples. The comparator and lowpass filter measure the ratio of samples above the estimated standard deviation; then, the target value is subtracted to identify the relative error. The regulator is composed of an amplifier and integrator to update the estimation of the standard deviation. After the final threshold, an event filter drops each second event too close to the previous one. **C,** Hardware network burst detector, incoming sources are events produced by the spike detector or the ISI-based burst detector, and the event count is stored in the accumulator ( $z^{-1}$ ) and cleared according a timer (the accumulator is not actually cleared, but its value is updated without considering the number of accumulated events thus far, which results in data clearance). The number of events counted since the beginning of the time window is compared with the user-defined threshold to decide whether the network is in a burst state.



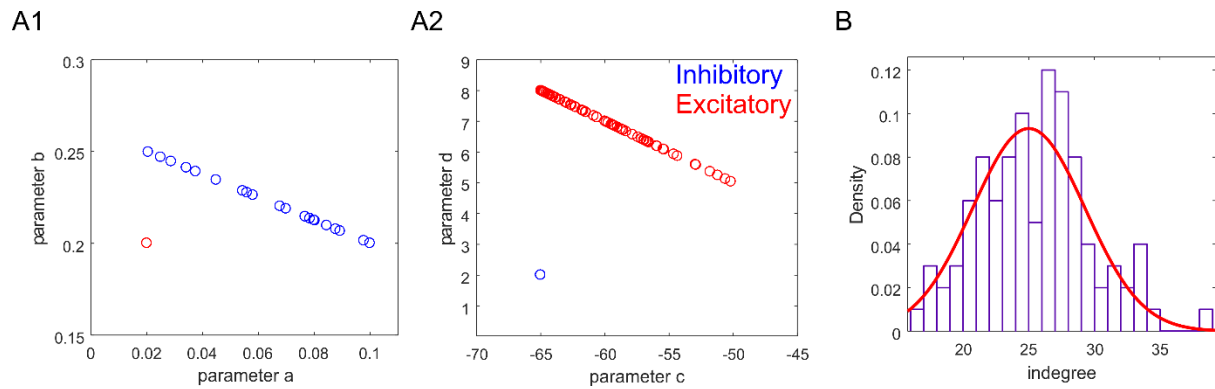
**Figure S3. Control no lesion, Related to Figure 2.** **A**, Mean firing rate (MFR) was stable for all control with no lesion experiments for all experimental phases (from S1 to S4). No significant difference was found (Friedman's repeated measures analysis of variance on ranks;  $n=9$ ;  $p=0.04$ ,  $DF=3$ , Chi-square= 8,333 but no significant difference between rank sums was found by the Tukey test). **B**, Cross-correlation (CC) area (obtained integrating the CC function of the collapsed spike trains from module 1 and 2 in a range of  $\pm 500$  ms) was stable for all control with no lesion experiments for all experimental phases (from S1 to S4). No significant difference was found (Friedman's repeated measures analysis of variance on ranks;  $n=9$ ;  $p=0.833$ ,  $DF=3$ , Chi-square= 0.867). **C**, Network burst rate (NBR) was stable for all control with no lesion experiments for all experimental phases (from S1 to S4). No significant difference was found (one-way repeated measures analysis of variance.  $n=9$ ;  $p=0.308$ ,  $DF=3$ ,  $F= 1.267$ ). **D**, Probability of single-module network burst (Prob smNB) was stable for all control with no lesion experiments for all experimental phases (from S1 to S4). No significant difference was found (Friedman's repeated measures analysis of variance on ranks;  $n=9$ ;  $p=0.789$ ,  $DF=3$ , Chi-square= 1.050).



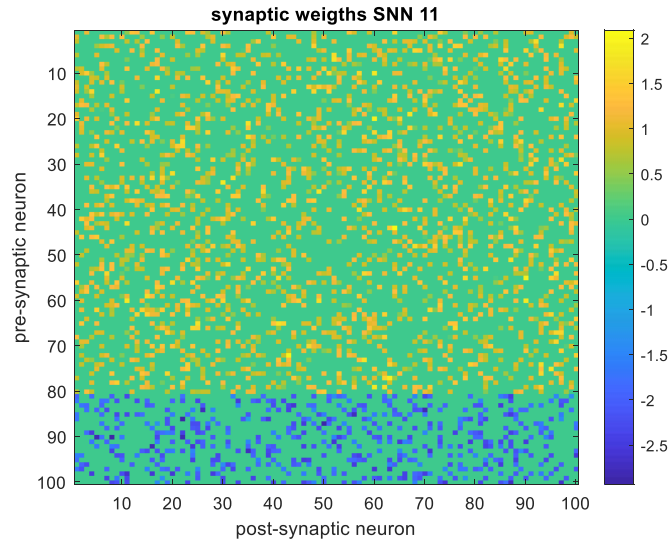
**Figure S4. Control with lesion, Related to Figure 2.** **A**, Mean firing rate (MFR) was almost stable for all control with lesion experiments. The only significant difference was found between the first spontaneous phase (S1) before the lesion and the first spontaneous phase post lesion (SPL1). Starting from the second hour post lesion, no significant difference was found between SPL2 and S1 and SPL3 and S1 (one-way repeated measures analysis of variance.  $n=4$ ;  $p=0.037$ ,  $DF=3$ ,  $F= 4,386$ ; all pairwise multiple comparison procedures (Bonferroni t-test): S1 vs. SPL1:  $p=0.036$ ; S1 vs. SPL2:  $p=0,282$ ; S1 vs. SPL3:  $p=0.376$ ; SPL3 vs. SPL1:  $p=1$ ; SPL3 vs. SPL2:  $p=1$ ; SPL2 vs. SPL1:  $p=1$ ). **B**, Cross-correlation (CC) area (obtained integrating the CC function of the collapsed spike trains from modules 1 and 2 in a range of  $\pm 500$  ms) collapsed following laser ablation. We found significant differences between all phases post lesion and the pre-lesion phase (one-way repeated measures analysis of variance.  $n=4$ ;  $p<0.001$ ,  $DF=3$ ,  $F= 16,555$ ; All Pairwise Multiple Comparison Procedures (Bonferroni t-test): S1 vs. SPL1:  $p=0.001$ ; S1 vs. SPL2:  $p=0.002$ ; S1 vs. SPL3:  $p=0.002$ ; SPL3 vs. SPL1:  $p=1$ ; SPL3 vs. SPL2:  $p=1$ ; SPL2 vs. SPL1:  $p=1$ ). **C**, Network burst rate (NBR) was stable for all control with lesion experiments (from S1 to SPL3). No significant difference was found (one-way repeated measures analysis of variance.  $n=4$ ;  $p=0.142$ ,  $DF=3$ ,  $F= 2.331$ ). **d**, Probability of single-module network burst (Prob smNB) before the lesion (S1) was on average close to 0.2, meaning that the majority of NBs involved both modules. Following the lesion, the probability increased to an average value higher than 0.85, meaning that the large majority of NBs involved only one module or the other. We found significant differences between SPL1 and S1 (one-way repeated measures analysis of variance.  $n=4$ ;  $p<0.001$ ,  $DF=3$ ,  $F= 109.911$ ; all pairwise multiple comparison procedures (Bonferroni t-test): SPL1 vs. S1:  $p<0.001$ ; SPL1 vs. SPL3:  $p=1$ ; SPL1 vs. SPL2:  $p=1$ ; SPL2 vs. S1:  $p<0.001$ ; SPL2 vs. SPL3:  $p=1$ ; SPL3 vs. S1:  $p<0.001$ ).



**Figure S5. Hardware system organization, pipeline implementation and synaptic noise results, Related to Figure 4.** **A**, Organization of the SNN system with two computation cores, the neuron and synapse. A state machine allows communication between the different blocks. The RAM stacks all the parameters and updates all values every 1 ms. RS232 communication allows communication with a computer to send the initial neural network topology. Spikes to the neural network block are for hybrid experiments where all detected spikes from the neuron culture are sent to the SNN via external synapses. **B1**, **B2**, Architecture of 'v', and 'w' and 'I' pipelines for digital implementation. The computation cycles are separated by dotted lines. Each of these dotted lines represents a rising clock edge on which the result of each operation is saved. For the sake of clarity, we have not shown the sequences of the flip-flops. 'u' and 'v' need 7 clock cycles (140 ns), and 'I' needs 5 clock cycles (100 ns). The synaptic noise current 'I' is modelled using an Ornstein-Uhlenbeck process, which makes the neuron implementation more biologically plausible, as shown in C. The parameter  $\mu$  represents the equilibrium or mean value for the process. We set this value equal to 0 with a bias current value necessary for tonic bursting activity without noise. For the other parameters,  $\sigma$  represents the degree of volatility around the mean value caused by shocks, and  $\theta$  represents the rate by which these shocks dissipate and the variable reverts towards the mean. The stationary variance depends on that parameter. Therefore, in our case, we can set the stationary (long-term) variance with  $\sigma$  and  $\theta$  parameters (neuron bursting activity) of 35 and 1, respectively. **C**, SNN output of the tonic bursting neuron in which the parameters are set to obtain tonic bursting activity with and without synaptic noise. The synaptic noise allows for better biomimetic dynamics.



**Figure S6. SNN model parameters, Related to Figure 4.** **A1**, Parameters 'a' and 'b' of the Izhikevich model (see Methods) for excitatory (red circle) and inhibitory (blue circles) neurons. All excitatory neurons have the same parameter  $a=0.02$  and  $b=0.2$ . All inhibitory neurons have different parameters  $a$  (from 0.02 to 0.1) and  $b$  (from 0.2 to 0.25). **A2**, Parameters 'c' and 'd' of the Izhikevich model for excitatory (red circles) and inhibitory (blue circles) neurons. All inhibitory neurons have the same parameter  $c=-65$  and  $d=2$ . All excitatory neurons have different parameters  $c$  (from -64.97 to -50.17) and  $d$  (from 5.04 to 7.99). **B**, Distribution of the indegree (i.e., the number of pre-synaptic neurons) with a Gaussian distribution superimposed (red curve). Mean value = 25, standard deviation = 4.3, R-Square = 0.806 (fit converged, Chi-square tolerance value of  $1e-9$  was reached).



**Figure S7. SNN connectivity matrix, Related to Figure 4.** The matrix represents the connectivity matrix of one of the 27 SNNs (SNN 11) belonging to the SNN library. Pre-synaptic neurons are on the y-axis and post-synaptic neurons are on the x-axis. Each element of the matrix represents the synaptic weight between the pre- and post-synaptic neuron. If the pre-synaptic neuron is excitatory (i.e. one of the first 80 neurons), the synaptic weight will be positive. Conversely, if the pre-synaptic neuron is inhibitory (i.e. one of the last 20 neurons), the synaptic weight will be negative. We set the outdegree (i.e., the number of post-synaptic neurons) to 25 for all neurons in the network, while the indegree (i.e., the number of pre-synaptic neurons) followed a normal distribution with a mean value of 25 and a standard deviation of 4.3 (Figure S6, R-Square=0.806).

## Supplemental Tables

	# NB	NB/min	Mean exc	Mean inh	Add exc	Add inh
SNN_1	39	1,95	0,99	-2,02	0,00	0,00
SNN_2	55	2,75	1,00	-2,02	0,01	0,00
SNN_3	73	3,65	1,02	-2,02	0,03	0,00
SNN_4	90	4,50	1,03	-2,02	0,04	0,00
SNN_5	111	5,55	1,04	-2,02	0,05	0,00
SNN_6	128	6,40	1,01	-1,22	0,02	0,80
SNN_7	136	6,80	1,00	-1,02	0,01	1,00
SNN_8	159	7,95	1,01	-1,03	0,02	0,99
SNN_9	164	8,20	1,02	-1,22	0,03	0,80
SNN_10	186	9,30	1,02	-1,04	0,03	0,98
SNN_11	208	10,40	1,08	-2,02	0,09	0,00
SNN_12	227	11,35	1,03	-1,05	0,04	0,97
SNN_13	239	11,95	1,08	-1,92	0,09	0,10
SNN_14	265	13,25	1,06	-1,22	0,07	0,80
SNN_15	295	14,75	1,07	-1,37	0,08	0,65
SNN_16	303	15,15	1,08	-1,52	0,09	0,50
SNN_17	329	16,45	1,07	-1,22	0,08	0,80
SNN_18	384	19,20	1,08	-1,22	0,09	0,80
SNN_19	420	21,00	1,09	-1,22	0,10	0,80
SNN_20	597	29,85	1,12	-1,14	0,13	0,88
SNN_21	743	37,15	1,15	-1,22	0,16	0,80
SNN_22	842	42,10	1,16	-1,18	0,17	0,84
SNN_23	968	48,40	1,18	-1,20	0,19	0,82
SNN_24	1080	54,00	1,20	-1,22	0,21	0,80
SNN_25	1303	65,15	1,24	-1,26	0,25	0,76
SNN_26	1510	75,50	1,27	-1,29	0,28	0,73
SNN_27	1882	94,10	1,34	-1,36	0,35	0,66

**Table S1. Library of SNN, Related to Figure 4.** SNN database was created to cover a wide range of NBRs from 1.95 NB/minute (SNN 1) to 94.1 NB/minute (SNN 27). To obtain such variability, we kept the same connectivity matrix with an average degree of 25% for all networks. The main parameters that we tuned to obtain such variability were the mean synaptic excitatory and inhibitory weights. Starting with SNN 1, we added a variable number to each synapse, maintaining separated excitatory (add exc) and inhibitory synapses (add inh). This SNN database can provide a wide range of activities and allows for a better choice of SNN for HBB experiments.

	bit registers			LUTs			RAM blocks			multipliers		
	numb.	Project	Device	numb.	Project	Device	numb.	Project	Device	numb.	Project	Device
Signal processing	4753	31%	3%	6274	31%	7%	11	5%	4%	3	7%	2%
SNN	3433	23%	2%	5747	29%	6%	82	39%	31%	42	93%	23%
Environment	6993	46%	4%	7996	40%	9%	118	56%	44%	0	0%	0%
Total	15179	100%	8%	20017	100%	22%	211	100%	79%	45	100%	25%

**Table S2. Hardware resource usage for the different functions of the hardware board, Related to Figure 4.** The signal processing part includes filters, spike detection, and burst detection (for events from electrodes and from the SNN). The SNN is considered a part by itself, and the environment part includes the necessary engineering to interface the system to the real world. This part includes analogue front-end management, computer interface, VGA display management and SD-card storage management. Bit registers (referenced as flip-flops in digital computing) are used to synchronously store local values or module states. Look-Up tables (LUTs) are asynchronous devices used to implement arbitrary 6-input logic gates. Random Access Memory (RAM) blocks are mostly used to store indexed values (e.g., channel samples for filters, and waiting queues). Multipliers are hard-coded embedded multiplier circuits that are available to increase both performance and density since multiplications are resource-demanding functions in hardware. The high majority of RAM blocks are required by the environment for VGA display and SD-card data buffer. The SNN part uses RAM blocks extensively to address the individual parameters of neurons and synapses. The SNN also requires many hardware multipliers because of the original mathematical formalism. Resources for signal processing are more focused on bit registers and LUTs because of pipeline structures that implement computations structures close to the device that stores data. The environmental cost in terms of resources is heavy, but this cost is constant and should not increase if implementing new signal processing features.



## Transparent Methods

### ***Bimodular confinement***

The polymeric structure for the physical confinement of neuronal cultures on conventional MEAs has been realized in polydimethylsiloxane (PDMS) by soft lithography and by using a photolithographically defined EPON SU-8 master on a silicon substrate. PDMS is an elastomer widely used for biomedical applications because of its high temperature, chemical and oxidation resistance; biocompatibility; transparency; and permeability to gases (Mata et al. 2005). Additionally, PDMS is an electrical insulator and nontoxic. These features also render PDMS particularly suited to the culture of primary neurons (Taylor and Jeon 2010). In addition, PDMS can be easily micro-structured by soft lithography, thus obtaining several low-cost replicas from a single master (Habibey et al. 2015; Weibel et al. 2007; Whitesides et al. 2001). In our case, replicas were produced for single use even if they could in principle be sterilized and re-used. In particular, PDMS stencil fabrication consisted of the following consecutive steps: i) structure design; ii) in silico wafer fabrication through a photolithographic technique; and iii) fabrication of PDMS replicas (for details, see (Bisio et al. 2014)). The developed modular neuronal networks were composed of two 3x2 mm modules that were interconnected by 25 channels 390  $\mu\text{m}$  long and 10  $\mu\text{m}$  wide with an inter-channel distance of 50  $\mu\text{m}$ . Once the PDMS masks were ready, they were positioned on the MEA substrates to include all the electrodes within the two modules. Next, the MEAs underwent the coating procedure for promoting cell adhesion, and the following day, the PDMS mask was removed immediately before the plating procedure, thus allowing neurons to slowly move only towards the promoting adhesion areas.

### ***Neuronal preparation***

Dissociated neuronal cultures were prepared from the neocortex of 18-day-old embryonic rats (pregnant Sprague-Dawley female rats were obtained from Charles River Laboratories Italia, Milano, Italy). All experimental procedures and animal care were conducted in conformity with institutional guidelines in accordance with the European legislation (European Communities Directive of 24 November 1986, 86/609/EEC) and with the NIH Guide for the Care and Use of Laboratory Animals. For our experiments, we used dissociated neurons arranged on a bimodular layout (cf. 'Bimodular confinement' paragraph for details) grown on MEAs. Culture preparation was performed as in (Bisio et al. 2014; Colombi et al. 2016). First, we coated MEAs overnight with poly-D-lysine and laminin to promote cell adhesion (Figure S1 A1). We washed the MEA devices at least 3 times with sterilized water before plating (Figure S1 A2). The neocortex of 4–5 embryos was dissected from the brain and dissociated first by enzymatic digestion in trypsin solution 0.125% (25–30 minutes at 37°C) and subsequently by mechanical dissociation with a fire-polished pipette. The resulting tissue was resuspended in Neurobasal medium supplemented with 2% B-27, 1% Glutamax-I, 1% Pen-Strep solution and 10% Foetal Bovine Serum (Invitrogen, Carlsbad, CA) at a final concentration of 500 cells/ul (Figure S1 A3). Cells were then plated onto 60-channel MEAs and maintained with 1 ml nutrient medium (i.e., serum-free Neurobasal medium supplemented with B27 and Glutamax-I). Then, cells were placed in a humidified incubator with an atmosphere of 5% CO<sub>2</sub>–95% air at 37°C. Half of the medium was changed weekly.

### ***Immunofluorescence staining and image analysis***

The cultures on coverslips were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) for 30 minutes. After permeabilization with 0.1% Triton X-100 in PBS for 10 minutes four times, the cultures were incubated with PBS containing 5% goat serum and 0.1% Triton X-100 for 1 hour. The permeabilized cultures were incubated with primary antibodies (anti-microtubule associated protein 2 [MAP2] mouse IgG; 1:100; Sigma-Aldrich) in PBS containing 5% goat serum overnight at 4°C and were rinsed with PBS for 10 minutes four times. Then, the cultures were incubated with a secondary antibody (Alexa Fluor 488-labelled anti-mouse IgG; Molecular Probes) in PBS containing 5% goat serum for 2 hours at room temperature and rinsed four times. The coverslips were removed from 12-well plates and mounted on glass slides with mounting media containing DAPI for nuclear staining. Fluorescence images were captured using a fluorescence microscope (Nikon eclips80). More details on the procedure can be found in (Ito et al. 2010).

### ***Focal lesion procedure and laser setup***

The laser dissection system (Difato et al. 2011) allowed for focal ablation of the sample in a three-dimensional confined volume due to the sub-nanosecond pulsed UVA laser source, which required delivering very low average power, thus confining the material breakdown to the focus spot. The setup was configured in an upright optical layout to allow optical surgery of neuronal networks plated on thick and non-transparent support such as the MEA chip. Therefore, simultaneously monitoring network activity using the MEA device and fluorescence calcium imaging during optical dissection of the connection of neuronal assembly was possible. The system was equipped with a custom micro-incubator (Aviv et al. 2013) maintaining the physiological parameters of the neuronal cultures (pH, osmolarity and temperature) to carry out long-term network activity recording before and after laser injury (Soloperto et al. 2016).

### ***Micro-electrode array recordings***

MEAs (Multi Channel Systems, MCS, Reutlingen, Germany) consisted of 60 TiN/SiN planar round electrodes (electrode diameter: 30  $\mu\text{m}$ ; inter-electrode distance: 200–500  $\mu\text{m}$ ). One recording electrode was replaced by a larger ground electrode. Each electrode provided information on the activity of the neural network in its immediate area. The array design using our bimodular mask allowed us to confine the growth of cells in two macro areas hosting 28 electrodes each (i.e., the ground electrode and the three central electrodes were excluded). A microwire connected each microelectrode of the MEA to a different channel of a dedicated amplifying system with a gain of 1100. The signals from the biological neural network (BNN) were amplified by a commercial system (MEA1060-Inv-BC amplification system, Multi Channel Systems, MCS, Reutlingen, Germany).

To reduce the thermal stress of the cells during the experiment, we maintained MEAs at 37°C with a controlled thermostat (MCS) and covered them with a custom PDMS cap to avoid evaporation and prevent changes in osmolarity (Blau et al. 2009). Additionally, we used a custom chamber to maintain a controlled atmosphere (i.e., gas flow of 5% CO<sub>2</sub> and 95% O<sub>2</sub> + N<sub>2</sub>) during the entire recording time (Frega et al. 2012).

### ***Electrical stimulation***

All electrical stimuli to BNN were triggered by our FPGA-based board (by a commercial stimulator, STG 4002, Multi Channel Systems, MCS, Reutlingen, Germany). The basic electrical stimulus was a biphasic voltage pulse, 300  $\mu$ s in the half-length phase and 750 mV of half-amplitude (in accordance with the literature (Wagenaar et al. 2004)). The necessary condition to choose the stimulating electrode was its capability to evoke a clear response on that module, meaning a network burst (NB) involving the entire module. In the pre-lesion condition, the stimulation evoked a response in both modules. In the post-lesion condition, the stimulation evoked a network burst confined to a single module.

### ***Experimental protocols and databases***

The general protocol included at least 30 minutes of no recording immediately after the culture was moved from the incubator to the amplifier to ensure the stability of the network in recording conditions. For control experiments without lesions, the four hours were recorded continuously. In controls with lesions, after the first hour of recording, we took the culture from the amplifier system, and we moved it to a separate room for laser ablation. The entire procedure usually took 20 minutes. After the lesion, the culture was moved back to the amplifier system where we immediately started the recording. The total number of experiments performed with these protocols was 9 for controls without lesions ( $26.8 \pm 0.8$  days *in vitro*, DIV) and 4 for controls with lesions ( $26 \pm 0.6$  DIV). For the bidirectional bridging (BB) protocol and hybrid bidirectional bridging (HBB), we performed similar protocols. We recorded 20 minutes of spontaneous activity before the lesion. Next, we performed laser ablation as described above and then waited for two hours to achieve stable activity in both modules, as shown by the results of control experiments (Figure 2). Then, we recorded 20 minutes of spontaneous activity after the lesion. To choose the best parameters that allowed us to reliably detect NBs in both modules, we performed offline NB detection. Next, we set these detection parameters on the FPGA with a custom-made MATLAB code and performed a 20 minute session of BB or HBB. The final step consisted of recording 20 minutes of spontaneous activity. The total number of experiments performed with these protocols was 9 for BB ( $26 \pm 0.8$  Days *in vitro*, DIV) and 7 for HBB ( $26 \pm 1.5$  DIV). In one HBB experiment, we did not record the last spontaneous phase because of a technical problem.

### ***Online processing***

#### ***Real-time spike detection***

The custom experimental system retrieved analogue data from MEA1060-Inv-BC pre-amplifiers through a standard MCS connector. Analogue input signals were packed in subgroups of 8 signals. For each subgroup, the signals were connected to the input of an analogue multiplexer that switched from one signal to the next one at a frequency of 80 kHz (each 12.5  $\mu$ s) as shown in Figure S2 A. Then, the output signals were amplified. The amplification gain ranged from 1 (no amplification) to 100. Multiplexed signals were finally converted to digital with 16-bit accuracy at a frequency of 80 kHz to provide one sample each time the multiplexer switched to a new input. Then, each of the 8 inputs of each multiplexer was sampled at a frequency of 10 kHz. The system relied on the antialiasing filters embedded in the pre-amplifier.

The spike detection module of the neuroprosthetic device was based on a threshold principle. A detection system was available for each input channel of the system. The first step of the detection was to increase the signal to noise ratio (SNR) of the input signal. The second step was to determine the appropriate threshold to perform the detection; this part of the processing is illustrated in Figure S2 B. To emphasize the amplitude of spike shapes and improve SNR, we used the wavelet decomposition technique – SWT (Quotb et al. 2012). This technique consisted of applying two orthogonal filters (one highpass and one lowpass) tailored to the characteristic shape of a spike.

Because of the impedance variation phenomenon, the input gain was expected to be different from one electrode to another and from one experiment to another. Therefore, we needed automatically set thresholds to avoid setting this parameter for each input channel individually. We determined these thresholds as a multiple of the standard deviation ( $\sigma$ ) of the first detail level from the SWT outputs. The module that computed  $\sigma$  was referenced as an amplitude estimator (AE) in Figure S2. The correspondence factor to determine the threshold was set system-wide by the experimenter depending on the experimental conditions. To compute  $\sigma$ , we used a continuous evaluation to avoid characteristic steps that occurred when performing computation on windows. To reduce computation resources, we digitized a method from analogue computing (Harrison 2003). With the hypothesis that the signal distribution was constant, we knew the proportion of samples below and above the standard deviation, and the system used this property to determine  $\sigma$  with a regulation loop. The signal distribution may change from experiment to experiment, but the estimated  $\sigma$  will always be relevant to the signal amplitude. Then, variations are compensated by adjusting the multiplier that determines the final threshold. Harrison (Harrison 2003) showed that in addition to its computation efficiency, this method was much more immune to accidental phenomena such as spike residues or stimulation artefacts.

As previously stated, the threshold was applied to the chosen output detail level of the SWT. The final step was to filter glitches that may come from noise around the threshold value and to change a spike-length pulse to a single event. This step was performed by a state machine that as far as the threshold had been crossed, provided a detected spike event and remained silent during a user-configured refractory period. Then, the output event was available as a spike event for the remaining computation modules of the system.

#### *Addressing the signal to noise ratio (SNR) with the stationary wavelet transform (SWT)*

As stated in the main text, the SWT technique we used is composed of a set of orthogonal filters (one highpass, one lowpass). The output of the highpass filter (detail level) was used as an output of the decomposition. The output of the lowpass filter (approximation level) was downsampled, and the filters were applied to provide the next detail and approximation outputs. The process is repeated as long as necessary. From the theoretical point of view, this method split the noise among the different detail outputs but restricted the spike contribution to one or two outputs, which improved the SNR for these specific outputs. The drawback is that the downsampling generated non-stationary behaviours and resulted in sub-optimal detection when the spike time corresponded to dropped samples. To avoid this phenomenon, we chose to replace signal downsampling by filter upscaling (transforming an 8<sup>th</sup> order filter to a 16<sup>th</sup> order filter containing 8 interleaved null coefficients). With this technique, stationary wavelet transform (SWT)(Pesquet et al. 1996), we kept the best performance for all samples with a

price of computational requirements twice as high, and memory requirements 4 times higher. This change in requirements was not an issue since we used hardware computing, and we added the computing resources according to our needs. Although our hardware was able to compute 8<sup>th</sup> order mother wavelets (highpass/lowpass filter couples), experience showed that the Haar mother wavelet (2<sup>nd</sup> order) was sufficient (Quotb et al. 2012). An empirical study showed that the third detail level provides the best performance for detection. As the first detailed output was expected to receive very few contributions from the spikes, it was used to determine the comparison threshold for spike detection.

#### *Real-time network burst detection*

The network burst detector (NBD) was based on the number of action potentials. Although this module was usable on a single channel, it was intended to gather events from multiple channels at the network scale as the preliminary experiments raised the necessity. Therefore, this module was not part of the computation flow build for each channel. There were 16 NBDs. Each NBD received events produced by all the incoming channels. Whether the input channels contributed to any of the 16 NBDs was user-defined. During a predefined time window, events on the selected channels were counted. At the end of the time window, the number of events was compared to a user-defined threshold to determine if an activity burst was occurring or not, and the counter was reset. The detector constantly kept track whether a burst was pending or not. Depending on user choice, each NBD could be independently configured to produce a single event when a burst started, a single event when a burst stopped, one event at each end of computing window detecting a burst, or continuously while the detector was in burst mode. The NBD could receive any event coming from the inputs on the system, including ISI-based burst detection events. It was possible to weight the event contributions such that an event coming from an ISI-based detected burst might have a much higher weight than an action potential on a single channel. The 16 NBDs worked on 1 kHz sampled signals because the application they fed was sampled at that frequency. Input events (10 kHz) were combined with a logical OR by group of 10 to produce the downsampling. There was no risk of losing an event because of the behaviour of neural cells that were unable to produce two action potentials within 1 ms.

An offline version of the NBD module was written in MATLAB to expedite the choice of parameters for reliable NB detection during experiments. By running the offline NBD with a combination of different windows in the range 1-50 ms and different thresholds in the range 1-250, we looked for a combination of parameters that could reliably detect NBs.

#### *Closed-loop implementation*

To close the loop, we chose to deliver a stimulation either to the BNN or the SNN based on the detection of a network burst (NB). The time to propagate the NB event depended on the source on which it is computed (BNN or SNN) and the stimulation destination (BNN or SNN). To follow the computation rhythm of the SNN, all events were processed at a frequency of 1kHz (so an event might have to wait for 900  $\mu$ s until the next step occurs).

NBs were computed within 37.4  $\mu$ s (including spike detection) and were available for processing 1.3  $\mu$ s later. Therefore, the BNN to SNN latency was 38.7  $\mu$ s. After a computation step of the SNN, the

stimulator was eventually activated after a delay of 3.9  $\mu\text{s}$ . As we measured the stimulator delay to be 85  $\mu\text{s}$ , the SNN to BNN latency was 88.9  $\mu\text{s}$ .

The BNN to BNN latency was composed of the BNN to SNN latency, the time to wait for the next computation step ( $T_w$ ), and the BNN to SNN latency.  $T_w$  was time dependant, values were equally distributed between 61.3  $\mu\text{s}$  and 961.3  $\mu\text{s}$  with a mean value of 511.3  $\mu\text{s}$ .

### **Spiking neural network**

The biomimetic SNN is a neuromorphic system with a most detailed level of analogy with the nervous system. The SNN is a network of silicon neurons connected via excitatory and inhibitory silicon synapses and plasticity rules. The characteristics of the systems for bio-hybrid experiments are compatible with the time constants of the real biological systems, complex neuronal models and plasticity that reproduce spatio-temporal patterns of activation. The hardware real-time implementation needs a low-resource neuron model.

#### *Neuron model*

The (Izhikevich 2003) IZH model is composed of two equations (1) and (2) in which the state variables 'v' represents the membrane potential of a neuron and 'u' represents the regeneration of the membrane potential, which takes into account the activation of ionic currents  $\text{K}^+$  and the inactivation of ionic currents  $\text{Na}^+$ .  $I_{Izh}$  describes the input current from other neurons. The integration time stamp  $dt$  of the IZH model is 1 ms.

$$\frac{dv}{dt} = 0.04v^2 + 5v + 140 - u + I_{Izh} \quad (1)$$

$$\frac{du}{dt} = a(bv - u) \quad (2)$$

with the after-spike resetting conditions:

$$\text{if } v \geq 30\text{mV} \rightarrow \begin{cases} v \leftarrow c \\ u \leftarrow u + d \end{cases} \quad (3)$$

In this model, parameter  $a$  describes the time scale of the regeneration variable  $u(t)$ . Lower values of  $a$  indicate slower regeneration. Parameter  $b$  describes the sensitivity of the regeneration variable influencing the under-threshold fluctuations of the membrane potential. Parameters  $c$  and  $d$  are the reset values of the membrane voltage and regeneration variable after an action potential, respectively. The IZH model can reproduce the behaviour of all known cortical neurons by changing only 4 parameters ( $a$ ,  $b$ ,  $c$  and  $d$ ), see Figure S6.

The detailed model and its implementation in digital hardware (Ambroise et al. 2013) are described below (see SNN implementation on FPGA).

#### *Synapse*

Synaptic transmission includes both excitatory and inhibitory models. In particular, the proposed model takes into account AMPA (i.e., an excitatory neurotransmitter) and GABA (i.e., an inhibitory neurotransmitter). Depolarization or hyperpolarization are represented by a positive or negative contribution to synaptic currents  $I_{syn}$ . Following the effects of AMPA and GABA, all excitatory or inhibitory synaptic currents tend to zero out, exponentially decreasing (Ben-Ari et al. 1997). These two synaptic

currents obey the same law of exponential decay  $\tau_{syn}$  (3 ms for excitatory decay and 10 ms for inhibitory decay). According to the literature (Izhikevich 2004), whenever a pre-synaptic neuron emits a peak, a synaptic weight ( $W_{syn}$ ) is added to the synaptic current of the post-synaptic neuron.

To connect the BNN to the SNN, we added external synapses to the SNN. Each neuron of the SNN was connected to one external synapse which can be tuned to be excitatory or inhibitory. Each event detected in the BNN could be used to stimulate the SNN through these external synapses. The SNN stimulation was performed through current stimulation via the external synapses. The stimulation duration depended on the type of synapses (excitatory or inhibitory) and the amplitude depended on the weight applied to the synapses. In our case study, excitatory synapses with 3ms exponential decay and with a weight of 9 are used for connection with BNN. We stimulated a maximum of 20 neurons to avoid over stimulation of the SNN.

#### *Short-term synaptic plasticity model*

To improve the biological behaviour of the synaptic model, we added short-term synaptic plasticity, which modified the synaptic weight according to the activity of pre-synaptic neurons.

Short-term plasticity is a biological phenomenon that modifies and regulates connection weights as a function of network activity. The model used in this work was proposed by (Izhikevich and Edelman 2008). This model is called "short-term plasticity" because the facilitation as well as the depression of a synapse is reabsorbed when no action potential has been emitted during a time constant. The implementation of the short-term plasticity and synapses are described below (parameters:  $x_{syn}$ ,  $P$  and  $t_{syn}$ , see *SNN implementation on FPGA*).

#### *Synaptic noise and axonal delay*

To enable spontaneous activity and the activity of our network, we added stochasticity by a source of current noise for each neuron in the neuron model. We used the Ornstein-Uhlenbeck (OU) process (with parameters: mean value = 0; degree of volatility = 35; dissipation rate = 1), which is a suitable model for modelling synaptic noise in a neural network (Rudolph and Destexhe 2005). The OU process  $X_t$  is a prototype of a noisy relaxation process and an example of a Gaussian process that has bounded variance and a stationary probability distribution. The process is stationary, Gaussian and Markovian. It satisfies the following stochastic differential equation:

$$dX_t = \theta(\mu - X_t)dt + \sigma dW_t \quad (4)$$

Where  $\theta > 0$ ,  $\mu$  and  $\sigma > 0$  are the parameters, and  $W_t$  is the Wiener process.

The parameter  $\mu$  represents the equilibrium or the mean value of the process. The stationary variance is given by:

$$var(X_t) = \frac{\sigma^2}{2\theta} \quad (5)$$

This form of current may represent an approximation to that resulting from the random opening and closing of ion channels on a neuron's surface or to randomly occurring synaptic input currents with exponential decay (Tuckwell et al. 2002).

Figure S5 C represents the dynamic behaviour of the neuron model (tonic bursting neuron) family in which the parameters are set for tonic bursting activity. This figure represents the same neuron in which we add the stochastic input currents. We noticed that the current noise source caused variability in the timing of action potentials. The digital implementation of this OU process is described in (Grassia et al. 2016) (see also SNN implementation on FPGA).

The synapse computation core also manages the axonal delay. The axonal delay is the phenomenon according to which an action potential emitted at time  $t$  arrives at time  $t + t_1$  to the post-synaptic neuron. In our implementation, the axonal delay was implemented by a 50-bit delay (shift register) vector, a  $D$  value and a multiplication factor. We stored the action potential (1 bit) in this vector delay at the position indicated by the value  $D$ . We can have a delay from 0 to 49 ms if the multiplication factor is equal to 1. We can increase this time by tuning this factor. The axonal delay makes it possible to describe a superposition of a network of neurons in 2D and consequently design pyramidal or 3D networks.

#### *Neural network architecture*

Our architecture of the neural network was based on blocks of RAM (storage of the parameters necessary for the definition of a network), two computing cores (a neuron and synapse), a block to manage the state machine and addresses and an "RS232 Communication" block (which will allow us to configure/modify the parameters of our system from a configuration file). Figure S5 A shows the interaction between these blocks. To simplify the figure, we did not represent the clock signal, but the entire architecture is synchronous. The limitation of the number of neurons is due to the following factors: 1) the clock frequency used to maximize the number of neurons in parallel and 2) the size of the RAM, which allows the storage of the parameters, especially the synapses. The resources used are presented in Table S2.

One of the advantages of our architecture was that each neuron and synapse were independent. A network configuration file was sent to the FPGA to select the connections and set the number of neurons, synapses, and used options. We could also define multiple networks with different connectivity between networks and within each network.

#### *Network configuration*

To add flexibility to our system, we added the ability to communicate with the FPGA via serial links to send the configuration, topology, and neural network options. Several options were possible depending on the desired application as follows: axonal delay, short-term plasticity, and synaptic noise. Then, we chose the family of neurons (excitatory, inhibitory), the percentage of connectivity between neurons, the inhibitory/excitatory ratio  $k$ , synaptic weight distributions, and the amplitude of the OU noise. From a library of configuration files, we could select the desired network for the given application and particularly in terms of frequency of NB. Figure 4 describes SNN library processing. Table S1 shows some metrics relative to the bursting features of the implemented artificial networks as well as the mean values of the excitatory and inhibitory weights.



The SNN library has been populated with network implementations sharing a fixed number of neurons (i.e. 100), the same topology (i.e. random connectivity) but with different synaptic weights (cf. Figure S7), necessary to reproduce different bursting behaviours.

Depending on the FPGA board, the maximum number of neurons varies. For instance, in a Spartan 6 FPGA board, we implemented 512 neurons, 66048 synapses with synaptic noise, axonal delay and synaptic plasticity. Table S2 describes the resources used. This biomimetic digital SNN worked in real-time, was tuneable and reproduced complex biological neural networks due to the biophysically neuron model, synapses, plasticity, axonal delay and synaptic noise.

### **SNN implementation on FPGA**

#### *Digital implementation of the Izhikevich model*

The mathematical equations must be modified to implement the IZH model in digital hardware. To make the IZH neural network more biorealistic, we split the current  $I_{Izh}$  into the following currents:  $I_{stat}$ ,  $I_{exc}$ , and  $I_{inh}$ .  $I_{stat}$  is the biasing current,  $I_{exc}$  is the positive contribution due to excitatory synapses and  $I_{inh}$  is the negative contribution of inhibitory synapses.

We also used the methodology developed at the University of Hopkins in the United States (Cassidy and Andreou 2008) by multiplying the equation of the membrane voltage by 0.78125 such that all parameters are decomposable with powers of 2, which allows easier digital implementation.

$$\begin{aligned} \frac{dv}{dt} &= \frac{1}{32}v^2 + 4v + 109.375 - u + I_{bias} + I_{exc} + I_{inh} \\ \frac{du}{dt} &= a.(bv - u) \end{aligned} \quad (6)$$

And  $\frac{dv}{dt} = \frac{v[n+1] - v[n]}{\Delta t}$  knowing that the computation time of the IZH model is 1 ms ( $\Delta t = 1$ ):

$$\begin{aligned} v[n+1] &= \frac{1}{32}v[n]^2 + 5v[n] + 109.375 - u[n] + I_{bias}[n] + I_{exc}[n] + I_{inh}[n] \\ u[n+1] &= u[n] + a.(b.v[n] - u[n]) \end{aligned} \quad (7)$$

Using a calculation pipeline, we can compute different neurons in parallel. All model parameters are stored in RAM. Figure S5 B1 describes the calculation pipelines for  $v$  and  $u$ .

#### *Digital implementation of synapse and synaptic plasticity*

*Synapse model:* When  $W_{syn}$  is positive, the synapse is excitatory, and when  $W_{syn}$  is negative, the synapse is inhibitory. The current  $I_{exc}$  is always positive, and the current  $I_{inh}$  is always negative. However, these two synaptic currents obey the same law of exponential decay  $\tau_{syn}$ .  $I_{syn}$ , the synaptic current, is written:

$$I_{syn}(t) = -\tau_{syn} \cdot I'_{syn}(t) = -\tau_{syn} \cdot \frac{I_{syn}(t+T) - I_{syn}(t)}{T} \quad (8)$$

$$I_{syn}(t+T) = \left(1 - \frac{T}{\tau_{syn}}\right) \cdot I_{syn}(t) \quad (9)$$

Where computation time  $T$  is 1 ms:

$$I_{syn}(t + 1) = \left(1 - \frac{1}{\tau_{syn}}\right) \cdot I_{syn}(t) \quad (10)$$

In digital implementation:

$$I_{syn}[n + 1] = I_{syn}[n] - \frac{1}{\tau_{syn}} \cdot I_{syn}[n] \quad (11)$$

*Short-term synaptic plasticity model:* This model is defined by 3 parameters:

- a scalar factor  $x_{syn}$ , which indicates the state of the synapse (depression or facilitation) and computes the synaptic weight.
- a percentage  $P$ , which will be multiplied by the factor  $x_{syn}$  after each emission of a pre-synaptic action potential. If this percentage is larger than 1, this synapse will describe a short-term facilitation. Otherwise, if this percentage is less than 1, this synapse will describe a short-term depression. In facilitation (or depression), the value added to the stimulation current of a post-synaptic neuron will increase (or decrease) with each emission of action potential.
- a time constant  $t_{syn}$  of exponential decay (or growth) in facilitation (or depression).

When a pre-synaptic spike occurs:

$$I_{syn}[n + 1] = I_{syn}[n] - \frac{1}{\tau_{syn}} \cdot I_{syn}[n] + W_{syn}[n] \quad (12)$$

Where  $W_{syn}$ , the synaptic weight, is defined by:

$$W_s[n] = x_{syn}[n] W_{syn} \quad (13)$$

$$x_{syn}[n+1] = P \cdot x_{syn}[n] \quad (14)$$

Parameters  $W_{syn}$ ,  $x_{syn}$ ,  $P$ ,  $\tau_{syn}$  are stacked into RAM.

*Synaptic noise implementation:*

The digital implementation of this noise (equation (4)) in the stimulation current of the neuron is given by:

$$I[n + 1] = I[n] + \theta(\mu - I[n])\Delta t + \sigma\Delta W[n] \quad (15)$$

Where  $n$  is the iteration step, and  $\Delta t = T/N$  is the time step after the partition of the interval  $[0, T]$  into  $N$  equal subintervals of width. Note that the random variables  $\Delta W[n]$  are independent and identically distributed normal random variables with expected value zero and variance  $\Delta t$ , thus  $\Delta W[n] \sim N(0, \Delta t) = \sqrt{\Delta t} * N(0, 1)$ .

Figure S5C represents the dynamic behaviour of the neuron model in which the parameters are set for tonic bursting activity.

### **Offline data analysis**

Offline data analysis was performed by custom scripts developed in MATLAB (MathWorks, Natick, MA, USA).

#### *Preprocessing*

To characterize the activity level of BNNs, we used the percentage of active channels and the mean firing rate (MFR). To compute the former, we considered active electrodes only those presenting a firing rate higher than 0.01 spikes per second (spikes/s), while the latter was defined as the mean number of spikes per second, computed over the total recording time, of the active channels. The low threshold guaranteed excluding only those electrodes that were not covered by cells or with very few spikes, keeping all the others (Bisio et al. 2014). In stimulation phases, the activity of stimulated channels was deleted.

#### *Cross-correlation*

To quantify the level of synchronization among multi-unit recordings, we applied correlation analysis (Chiappalone et al. 2007) to the spike trains. Cross-correlograms were built according to the method of the activity pairs described by (Eytan et al. 2004) as follows: given two trains (i.e.,  $X$  and  $Y$ ), we counted the number of events in the  $Y$  train within a time frame around the  $X$  event of  $\pm T$  ( $T$  set at 500 ms), using bins of amplitude  $\Delta t$  (set at 1 ms). The CC function  $C_{xy}(t)$  was obtained by a normalization procedure, according to the following formula (Pasquale et al. 2008):

$$C_{xy}(T) = \frac{1}{\sqrt{N_x N_y}} \sum_{t_i=(T-\frac{\Delta T}{2})}^{(T+\frac{\Delta T}{2})} X(t_s - t_i) \quad (16)$$

where  $t_s$  indicates the timing of an event in the  $X$  train,  $N_x$  and  $N_y$  are the total number of events in the  $X$  and  $Y$  trains, respectively, and  $\Delta T$  is the bin size. Equation (1) yields the symmetry between  $C_{xy}(t)$  and  $C_{yx}(t)$  (i.e.,  $C_{xy}(t)=C_{yx}(-t)$ ) (Eytan et al. 2004).

To evaluate the amount of correlation between two modules, we performed the computation between the collapsed spike trains for each module. A collapsed spike train is the result of a logic OR operation between the spike trains recorded on all the electrodes belonging to the same module. For control experiments and BB experiments, the modules were modules of the BNN. In the HBB protocol, we collapsed the activity of the entire SNN to a single spike train. We applied a locally weighted linear regression to smooth the curve. The area of the CC was the integral of the curve in the range from -500 to 500 ms.

#### *Offline network burst detection*

The offline NBD was performed using a software implementation of the NBD algorithm running online on the neuromorphic device, called the “fixed window accumulator”. Similar to the online version, the algorithm summed all the spikes fired from the neurons of the networks during a given time interval, and it detected an NB if this sum overcame a threshold. Each NB was highlighted by the generation of an event. The input parameters were the window time (i.e., time interval within which the spikes were summed), the threshold (i.e., the number of spikes to overcome to identify an NB event) and the source (i.e., the channels on which the analysis was performed).

Depending on the specific protocol we wanted to analyse, the source could be the whole BNN (in controls without lesions or pre-lesion spontaneous activity), a part of it (for example, to compute the NBR in only one module or to exclude stimulated electrodes), the SNN (for example, to characterize the SNN) or a hybrid network formed by both the SNN and whole BNN or part of it (in HBB). The offline version was also useful during the experimental protocols to set the best parameters (window time and threshold) to obtain a reliable NB detection and to choose the most suitable SNN, referring to NBR, for the HBB experiment.

#### *Probability single-module network burst*

To quantify the confinement of the electrophysiological activity in a single module, we used the single-module NB probability parameter. This parameter represented the probability that NBs were composed of spikes belonging to a single module (i.e., one of the modules of the BNN or the SNN). This parameter was computed by counting the number of spikes that belonged to each burst, followed by the percentage of spikes that belonged to each module. If this percentage was higher than 85%, we assumed that the burst was generated in a single module. Setting 85% as the threshold percentage allowed us to consider the possibility of having random spikes in the other module at the same time as the NB. To perform this analysis, we modified the NBD by adding start and stop thresholds. For each NB identified, the algorithm looked for an empty time window (i.e., a time window in which the number of spikes was equal or lower than the threshold) before and after the window in which an NB was detected to identify the first and last spikes of that event as the first spike in the first window and the last spike in the last window belonging to the NB, respectively. We set the start and stop thresholds to zero and 5, respectively, for BNN and SNN (the SNN threshold was higher because of the presence of fast spiking neurons).

#### *Statistics*

Data within the text is expressed as the mean  $\pm$  standard error of the mean (SEM), unless otherwise specified. Statistical tests were employed to assess significant differences among different experimental conditions. The normal distribution of experimental data was tested using the Shapiro-Wilk normality test. We performed one-way repeated measures ANOVA to compare data from the same group at different time points. When ANOVA gave a significant ( $p < 0.05$ ) result, the post hoc Bonferroni test was employed to assess differences between all phases. When normality was rejected, we used Friedman's repeated measures ANOVA on the ranks test to compare data from the same group at different time points. When ANOVA gave a significant ( $p < 0.05$ ) result, the post hoc Tukey test was employed to assess differences between all phases. To compare data from two different populations (Figure 2), we performed the Mann-Whitney test. Statistical analysis was carried out by using OriginPro (OriginLab Corporation, Northampton, MA, USA) and Sigma Stat (Systat Software Inc., San Jose, CA, USA).

### ***Data and Software Availability***

The data sets of the experiments presented in our manuscript are available in the Mendeley Data repository. The accession number for the experimental data reported in this paper is the following:

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The FPGA SNN is available upon request to Timotheé Levi, and it will be available in the website of the authors (Timothée Levi, <http://timothee.levi.free.fr>). Spike computation related modules are publicly available at <http://yannick.bornat.fr>.

The above accession numbers and links will be accessible to readers upon publication of our manuscript.

Upon reasonable request, data and information on the reported results can be asked to the corresponding authors.

## Supplemental References

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