Learning Multi-Sensory Integration with Self-Organization and Statistics

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Abstract

Recently, we presented a self-organized artificial neural network algorithm capable of learning a latent variable model of its high-dimensional input and to optimally integrate that input to compute and population-code a probability density function over the values of the latent variables of that model. We did take our motivation from natural neural networks and reported on a simple experiment with simulated multi-sensory data. However, we focused on presenting the algorithm and evaluating its performance, leaving a comparison with natural cognition for future work. In this paper, we show that our algorithm behaves similar, in important behavioral and neural aspects, to a prime example of natural multi-sensory integration: audio-visual object localization.

1 Introduction

Imagine you are given a sheet of paper with unlabeled numbers, told that these numbers contain information about the value of some quantity, and asked what you think the value of that quantity is. This is clearly an impossible task. What if you are repeatedly shown such numbers, for different values of the quantity in question, but never given the right answer? Biological neurons face a similar situation. It is their function to produce activity corresponding to some quantity in the outside world. And all they have, to estimate that quantity, is the activity at their incoming synapses, which carries no information about its origin. A comparable situation also exists in unsupervised machine learning. Models of unsupervised neural learning have therefore found applications in general machine learning.

We have recently presented an artificial neural network (ANN) algorithm with possible applications in general machine learning based on the self-organizing map (SOM) [Bauer and Wermter, 2013]. This algorithm was inspired by the apparent ability of humans to utilize the sensory information they get in a statistically optimal fashion in many situations [Ernst and Banks, 2002; Landy *et al.*, 2011]. In particular, it aims to model how neural populations like the superior colliculus (SC) learn to make sense of uni- and crosssensory stimuli. The result is an algorithm which takes highdimensional data as input, learns a low-dimensional latentvariable model and computes for a given input a probability density function (PDF) over the values of the latent variables.

We have shown that our algorithm can perform nearoptimally on uni-sensory input and we have shown that it can handle multiple sensory modalities with different response and noise characteristics. However, we have not fully closed the loop in 1) trying to understand a problem faced by a natural system, 2) trying to find a solution 3) comparing that solution to the one found, tried, and tested by nature [Jacobs and Kruschke, 2011; Landy *et al.*, 2011]. In this paper, we compare the response properties of our algorithm to the wellestablished principles of multi-sensory integration in the SC.

In the next section, we will first briefly review our algorithm, focussing on giving a good intuition of the main principles behind it, and referring to the original paper for details. We will then report on experiments we carried out in which we simulated various conditions of multi-sensory integration. We will review the known neurophysiological and psychophysical effects and compare them to our network's responses. Finally, we will interpret our results in the broader scope of computational neuroscience and its interaction with general artificial intelligence.

2 The Model

Before we start describing our solution, let us again look at the problem we are trying to solve. A population of neurons is to collaborate in learning to compute a PDF for the latent variables behind patterns of neural activity.¹ This input activity can be uni- or cross-sensory; conceptually, both can be treated the same by introducing a logical population which simply concatenates the separate input populations' activity vectors (see Fig. 1).

Our approach is to have the network learn to represent the PDF in a population code, where each neuron codes for the probability of a different value being the true value of the latent variable [Pouget *et al.*, 2003]. ANN models working with such population-coded PDFs have been proposed eg. by Cuijpers and Erlhagen [2008] and Beck *et al.* [2008]. Our model

¹From now on, we will assume wlog. a single latent variable. Note that strictly speaking a combination of latent variables can be seen as a single complex latent variable.



Figure 1: Population-coded PDF Computed from Multiple Physical Input Populations

focuses on how computing such a PDF from arbitrary neural responses can be learned unsupervised and without heavy assumptions on the noise properties. The population code realized by our network is to be spatially organized, i.e. close-by neurons code for similar values of the latent variable. This restriction on the more general definition of a population code seems natural and it also reflects biological evidence of topographic maps in various sensory brain areas [Stone, 2012; Hyde and Knudsen, 2000; Stein and Stanford, 2013; Kaas, 1997].

Kohonen's SOM [Kohonen, 1982] was inspired by the formation of such topographic maps in the brain. It is a selforganizing ANN algorithm which has been shown to be able to learn latent-variable models [Yin, 2007; Klanke, 2007]: It learns topography-preserving mappings from points in a data space to its spatially ordered units, or neurons. Since a SOM models a population of neurons and each neuron has a response to a stimulus, it is possible to read out a population code from a SOM [Zhou *et al.*, 2011].

In a standard SOM, the response is just the Euclidean distance of the stimulus as a vector from the preferred stimulus of each neuron. This response is used to find the bestmatching unit (BMU), the neuron with the strongest response and the neuron the stimulus is mapped to. In our algorithm, the network simultaneously learns the latent-variable model and the noise properties of the input. The response of each neuron then is an estimate of the probability of the neuron's preferred value being the actual value of the latent variable, given what the network knows about the noise. This is done basically by keeping weighted counts of previous activities at each input connection of each neuron. Input activities at the neurons' synapses are discrete and treated nonmetrically. Therefore, the algorithm lends itself to learning problems where data points have nominal dimensions. The main benefit over previous approaches [Zhou et al., 2011; Bauer et al., 2012b] is that our approach does not assume any specific noise model. We refer to our original paper [Bauer and Wermter, 2013] for details on the learning algorithm and its motivation and derivation.

3 Experiments

In the experiments described below, we will compare our network's performance and response properties to those found in psychophysical and neurophysiological studies. Specifically, we will examine the responses of our network in light of biological data about the SC. The SC, or optic tectum (OT), as it is called in non-mammalian vertebrates, is an evolutionarily stable midbrain structure concerned with localizing objects in space on the basis of visual, auditory, and somatosensory stimuli. It is involved in generating orienting movements on the basis of uni- and cross-sensory stimuli [Stein and Stanford, 2013]. We choose it as a standard to which to compare our model because its input-output behavior is relatively wellunderstood and because it is likely that the general principles of SC functioning are realized similarly in other brain regions with comparable tasks [Stein, 2012].

For our simulations, we used a network of 350 output neurons connected to two populations $\mathbf{i}_{a,1}, \mathbf{i}_{a,2}, \ldots, \mathbf{i}_{a,40}$ and $\mathbf{i}_{v,1}, \mathbf{i}_{v,2}, \ldots, \mathbf{i}_{v,40}$ of input neurons. The two input populations each represented one sensory modality. Each input neuron $\mathbf{i}_{m,k}$ had a preferred value of $\mathfrak{p}_k = \frac{k-1}{39}$, for $m \in \{a, v\}$, and responded to a stimulus $\mathfrak{p} \in [0, 1]$ according to a poisson-noisy Gaussian:

$$\mathbf{a}_{m,k}(\mathbf{p}) \sim \Pr(a_m \exp(-(\mathbf{p}_k - \mathbf{p})^2 / \sigma_m^2)), \qquad (1)$$

for $a_a = 4$, $\sigma_a^2 = 0.01$ and $a_v = 6$, $\sigma_v^2 = 0.005$. In the following, the neurons $\mathbf{i}_{a,k}$ and $\mathbf{i}_{v,k}$, $k = 1, 2, \ldots, 40$ will be referred to as 'auditory' and 'visual', respectively, to make presentation more intuitive.

We trained the network with congruent stimuli until it had developed spatial organization and learned the simulated input noise statistics. We then started simulating our chosen psychophysical and neurophysiological experiments. For these experiments, we changed the simulated stimulus conditions as will be described below.

Enhancement/Depression. It is a well-established fact that multi-sensory SC neurons tend to react more strongly to cross-sensory stimuli in their receptive fields than to unisensory stimuli [Stein and Meredith, 1993]. This effect is called enhancement. Depression, on the other hand, occurs when stimuli are temporally or spatially incongruent: In the spatial case, this means that the reaction of a multi-sensory neuron to a visual stimulus in its receptive field will actually be weaker if that stimulus is accompanied by a sound coming from a different point in space (and vice-versa).

We simulated this condition by presenting, in each trial, one random stimulus $\mathfrak{p}_{a, \in} [0, 1]$ to neurons $\mathbf{i}_{a,k}$ and a different stimulus $\mathfrak{p}_v \in [0, 1]$ to neurons $\mathbf{i}_{v,k}$. We recorded the network's response to the combined, incongruent, crosssensory input population response. Fig. 2 shows the mean response over all trials of the output neuron at whose center was the visual stimulus depending on the absolute distance of the incongruent auditory stimulus. Although somewhat noisy, the graph clearly shows that congruent stimuli elicit much stronger responses than incongruent responses, which is in accordance with the phenomena of enhancement and depression explained above.



Figure 2: Response of an Output Neuron to a Visual Stimulus in its Receptive Field and an Auditory Stimulus at Various Distances from the Visual Stimulus

MLE. The effects discussed so far are on the level of single multi-sensory neurons. Since these neurons are part of a sensory-motor processing circuit, it is to be expected that they manifest themselves in observable behavior. Alais and Burr [2004] found that their test subjects's performance in an audio-visual localization task was consistent with a maximum likelihood estimator (MLE) model of multi-sensory integration [2004]. Other authors have found similar results for different combinations of sensory cues [Ernst and Banks, 2002; Hillis *et al.*, 2004].

The model used by Alais and Burr assumes Gaussian noise in sensory localizations. Under this assumption, MLE integrates two uni-sensory localizations l_a , l_v optimally according to a linear combination:

$$l_{MLE} = \frac{r_a}{r_a + r_v} l_a + \frac{r_v}{r_a + r_v} l_v,$$
 (2)

where $r_m = \frac{1}{\sigma^2}$ is the reliability of a modality m, if σ is the standard error of localizations by that model, that is, the mean absolute error between the localization and the actual location of the target.

The expected reliability r_{MLE} of the combined result is given by:

$$\frac{1}{r_{MLE}} = \frac{1}{r_a} + \frac{1}{r_v}.$$
 (3)

The distribution of errors of the combined estimator, like the assumed distribution of errors of the individual modalities' estimators, is Gaussian.

First, we determined the distributions of errors of our model given uni-sensory and cross-sensory stimuli. To do that, we fed our network with input in which either only auditory neurons $i_{a,k}$, or only visual neurons $i_{v,k}$ had non-zero activity (according to Eq. 1), or both, as usual. Figure 3 shows histograms of errors (mislocalizations) for uniand cross-sensory localization, as well as Gaussian functions fitted to these errors. It can be seen that the distribution of errors is Gaussian-like, and that combined localization has much greater reliability than either of the individual localizations. Closer analysis reveals that the standard deviations of auditory-only, visual-only, and cross-modal localization are $\sigma_a = 1.966 \times 10^{-2}$, $\sigma_v = 1.473 \times 10^{-2}$, and $\sigma_m = 1.385 \times 10^{-2}$, respectively. The expected crossmodal localization error according to Eq. 3 would be $\sigma_{m,e} =$ 8.978×10^{-3} . We attribute this discrepancy to sampling error,² outliers, and actual learning errors. All in all, both visual inspection of the distribution of errors and this analysis demonstrate that our network effectively integrates the information in its multi-sensory input.

To test whether the behavior of our network is consistent with the MLE model described above, we conducted another experiment. As in the first experiment, we again chose one auditory stimulus \mathfrak{p}_a and a visual stimulus \mathfrak{p}_v in every trial. This time, each trial consisted of three conditions: an auditory, a visual, and a cross-sensory condition. In the auditory condition, we combined the normal auditory population response (Eq. 1) with a flat response of all-zero activity. The visual condition was analogous and in the cross-sensory condition, the population responses were combined as usual. In each trial *n*, the input was presented to the model and the auditory, visual, and cross-sensory localizations $l_{a,n}, l_{v,n}, l_{c,n}$ were recorded.

We then computed the least-squares solution to the equation

$$p_a \begin{pmatrix} l_{a,1} \\ l_{a,2} \\ \vdots \\ l_{a,N} \end{pmatrix} + p_v \begin{pmatrix} l_{v,1} \\ l_{v,2} \\ \vdots \\ l_{v,N} \end{pmatrix} = \begin{pmatrix} l_{m,1} \\ l_{m,2} \\ \vdots \\ l_{m,N} \end{pmatrix},$$

where N = 10.000 is the number of trials. We found $p_a = 3.659 \times 10^{-1}$ and $p_v = 6.400 \times 10^{-1}$ which is close to the optimal values $\hat{p}_a = 3.595 \times 10^{-1}$, $\hat{p}_v = 6.405 \times 10^{-1}$ obtained by inserting σ_a and σ_v into Eq. 2.

Together, these results show that our algorithm is not only statistically well-motivated and shows response characteristics similar to that of a biological information processing system, the SC, as was found in the first experiment; Its behavior on the functional level is also comparable to the optimal cue combination behavior demonstrated in human multi-sensory integration. This is especially interesting for our algorithm as a general machine learning algorithm.

4 Discussion

In this paper, we have shown that the neural learning algorithm introduced previously is able not only to integrate multi-sensory input, but also mimics biology both on the single-neuron and behavioral level. We can therefore interpret our network as a model of the SC, as it develops a representation of sensory input space, integrates percepts from different modalities depending on their reliabilities, uses the statistics of the input to learn this, and incorporates concepts known to be key in the SC, like population coding, winnertake-all, and local interactions.

²Estimation is done by selecting the winner neuron and choosing its preferred value as the estimate. Since there are only finitely many neurons but infinitely many rationals in [0, 1], estimation is bound to make sampling errors.



Figure 3: Histograms of Distances between Visual, Auditory, and Audio-Visual Localization and Stimulus



Figure 4: Research Cycle in Computational/Robotic Neuroscience

We strongly believe that both fields, life sciences and artificial intelligence, will benefit from the approach of modeling observed biology to generate biological research questions, and implementing models in technical systems to validate their fitness and real-world applicability (see Fig. 4). Therefore, the next step will be validating our model's functionality and resemblance of biology in a robotic implementation. Initially, our experiments will mimic classical experiments like the ones due to Stein and Meredith [1993], which originally established the properties of multisensory integration in the cat SC: In our versions of these experiments, a robot will take the place of the feline or human subjects. It will be exposed to very similar multi-sensory stimuli as the subjects in the original experiments. And its behavior and simulated neural processing will be monitored and compared to the original findings.

For these experiments, we will use the virtual reality lab infrastructure recently implemented ([Bauer *et al.*, 2012a], see Fig. 5). Designed and built as a basis for robotic sensory and cognitive experiments, this Virtual Reality for Robots Lab features a 180° projection screen and a matrix of 18 speakers. It allows us to precisely control the conditions of audiovisual localization experiments and still deliver rich and lifelike stimuli to the iCub robot head which is placed at its center [Beira *et al.*, 2006]. With feedback from these experiments, we will extend our model and aim to accommodate attentional effects. This will give our model greater explanatory



Figure 5: 3D Model of the Virtual Reality Robot Environment: A multi-purpose aluminium structure holds four projectors and a 180° projection screen. Around the screen, there is an array of speakers. The robot head is placed at the center of the half-cylinder spanned by the screen.

power and, at the same time, make it more flexible and more widely applicable in robotic and other AI systems. Again, biological experiments like those by Spence *et al.* [2004], which studied the effects of priming on multi-sensory integration, will guide our modeling efforts and serve as models for robotic experiments.

Acknowledgements

This work is funded by the DFG German Research Foundation (grant #1247) – International Research Training Group CINACS (Cross-modal Interactions in Natural and Artificial Cognitive Systems).

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