# There's no 'sampling bias' for the mutual information

 C. Battistin

<claudia.battistin@gmail.com>

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This note shows that the so-called 'sampling bias' in the estimation of the mutual information for stimulus-response frequencies doesn't exist. If the estimate from the sample forecasts a high mutual information, then the response is indeed very likely to be informative. On the other hand, if we expect the response to be uninformative or if we just want to be conservative, a correct calculation of the estimate leads to a negligible mutual information. A correct frequentist analysis also shows that the frequentist estimator is unbiased. No corrections of any kind are needed.

Note: Dear Reader & Peer, this manuscript is being peer-reviewed by you. Thank you.

#### 1 Bias?

The mutual information between the long-run relative frequency of a signal and that of a response is a measure of how much our uncertainty about the signal is reduced by knowledge of the response. This measure is sometimes used in neuroscience, the response being some characteristic – such as the activity or the firing rate – of a neuron or of a network of neurons.

Recent works in neuroscience (Panzeri et al. 2007) \* add others claim that our estimate about the mutual information for long-run frequencies from a small sample is biased; that is, in frequentist terms: the expected value of its sample estimate is different from its 'true' value. Other works have proposed various corrections to this alleged bias.

In this note we show three main results:

1. The direct estimation of the 'long-run' mutual information from a sample is reliable and doesn't need any kind of correction. On the one hand, if we are equally uncertain about the long-run response frequencies, and the estimate from a small sample suggests a large mutual information, then we should indeed expect the response to be informative. On the other hand, if we initially suspect that the long-run response frequencies should be uninformative, or if we are very

conservative in our inference, then a correct calculation consistently leads to very low estimates of the mutual information.

- 2. For small samples, our estimate crucially depends on the predata probability for the long-run relative frequencies of the response conditional on the stimulus. In making such estimates we must therefore give some thought to assessing this pre-data probability, rather than correcting non-existing biases.
- 3. A correct frequentist analysis proves that the estimator for the long-run mutual information is unbiased.

The main result is first shown in the next section with a straightforward calculation, and explained intuitively in the subsequent section. The calculations use the example given by Panzeri et al. (2007 Fig. 1). The final section briefly discusses the third result and the importance of the pre-data probability in our estimations when the sample is small. Some ways to assess this probability are also discussed.

## 2 Bayes

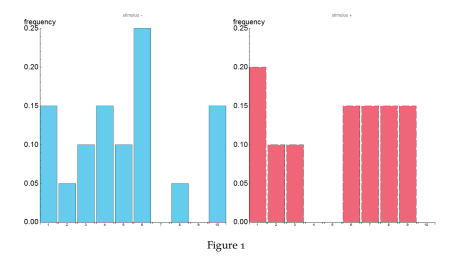
First of all let's state what our inference is about. Given a sample of stimulus-response data we want to assess what's the most probable set of long-run¹ relative frequencies of the response conditional on each stimulus value, and from these assess what's the most probable value of the associated mutual information between stimulus and response. We assume that all stimuli appear with equal relative frequencies.

Let the stimulus s have two possible values  $\{-,+\}$ , and the response r ten possible values  $\{1,\ldots,10\}$ . Let the data D be a set of n stimuli – which yielded n responses  $(r_i^-)$ ,  $i\in\{1,\ldots,n\}$ , and n stimuli + which yielded n responses  $(r_i^+)$ . The ten response state appeared with relative frequencies  $q^- := (q_r^-)$ ,  $r\in\{1,\ldots,10\}$ , for the stimulus –, and with relative frequencies  $q^+ := (q_r^+)$  for the stimulus +.

We can use the concrete data summarized in fig. 1, consisting in n = 20 samples per stimulus.

If the long-run frequencies conditional on stimulus – are  $f^- := (f_r^-)$ , and conditional on +,  $f^+ := (f_r^+)$ , then out of symmetry the probability

 $<sup>^1{}&#</sup>x27;But$  this  $long\ run$  is a misleading guide to current affairs. In the long run we are all dead.' (Keynes 2013 § 3.I, p. 65)



of obtaining the data *D* is

$$p(D \mid f^{-}, f^{+}, K) = \prod_{r} \left[ (f_{r}^{-})^{nq_{r}^{-}} (f_{r}^{+})^{nq_{r}^{+}} \right]. \tag{1}$$

This is also the *likelihood* of the long-run frequencies in view of the sample. Their probability density is proportional to the likelihood, corrected by their initial probabilities  $p(f^-, f^+ | K)$ 

$$p(f^{-}, f^{+} | D, K) \propto p(D | f^{-}, f^{+}, K) p(f^{-}, f^{+} | K) =$$

$$p(f^{-}, f^{+} | K) \prod_{r} \left[ (f_{r}^{-})^{nq_{r}^{-}} (f_{r}^{+})^{nq_{r}^{+}} \right]. \quad (2)$$

We can calculate this probability analytically when possible, or estimate it with Monte Carlo sampling. From such samples we estimate the probability distribution of the long-run mutual information

$$I := \sum_{r} \frac{1}{2} f_{r}^{-} \ln \left( \frac{\frac{1}{2} f_{r}^{-}}{\frac{1}{2} f_{r}^{-} + \frac{1}{2} f_{r}^{+}} \right) + \sum_{r} \frac{1}{2} f_{r}^{+} \ln \left( \frac{\frac{1}{2} f_{r}^{+}}{\frac{1}{2} f_{r}^{-} + \frac{1}{2} f_{r}^{+}} \right). \tag{3}$$

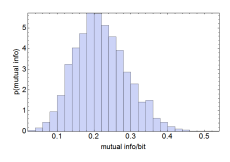
Let's consider two possible probabilities for the long-run conditional frequencies:

### 2.1 Uniform uncertainty about the frequencies

If we initially think that equal ranges  $(\triangle f^-, \triangle f^+)$  of pairs of conditional frequencies are equally possible, then

$$p(f^{-}, f^{+} | K_{u}) = 1. (4)$$

We sample 5 000 pairs of conditional frequencies from the density (2) obtained from our example data and this pre-data probability. The resulting distribution of their associated mutual information is shown in fig. 2. It tells us that the response is likely to be informative; the most



likely values of the long-range mutual information are around 0.2 bit. The next section explains intuitively why this probability distribution is correct.

Figure 2

## 2.2 Conservative uncertainty about the frequencies

If we initially think that the long-run response frequencies conditional on the stimuli should be very similar, or if we simply want to do a conservative estimate, then our pre-data probability must be higher for pairs with similar conditional frequencies; for example \*replace with combination of Dirichlet – same effect

$$p(f^-, f^+ | K_c) \propto \exp\left[\frac{\sum_r (f_r^- - f_r^+)^2}{2\sigma^2}\right].$$
 (5)

This density states that the two conditional frequencies should be roughly equal, but otherwise leaves a uniform uncertainty about the values of each. Smaller values of  $\sigma$  represent more conservative estimates.

A sample of 5000 pairs of conditional frequencies from the density (2) with the conservative initial density (5)  $\stackrel{\bullet}{h}$  specify  $\sigma$  leads to the distribution of mutual information of fig. 3. The estimate now says that

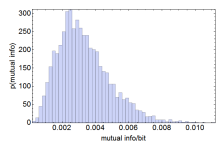


Figure 3

we should expect a negligible mutual information, with a most probable value around 100 times smaller than in the previous case.

#### 3 Because

Let's try to understand why the estimate of § 2.1, fig. 2, is reliable; and to understand what happens in the conservative case of § 2.2.

If we deem all pairs of conditional frequencies equally possible, we can sample  $5\,000$  pairs uniformly. Figure 4 shows the resulting scatter plot for first component of the two conditional frequencies, that is, the frequency of the response value 1.

Let's now visualize the samples of long-run conditional frequencies in two dimensions as follows: for each sample, the horizontal coordinate is the probability that the frequencies assigns to our observed data; and the vertical coordinate is the mutual information associated with the frequencies. We obtain the scatter plot of fig. 5. All points, of three different sizes and colours, are part of the plot. The pair of two uniform conditional frequencies (1/10 probability for each response) is the largest, yellow point. This pair of long-run conditional frequencies assigns probability  $10^{-40}$  to the data and has zero mutual information.

It's clear from this scatter plot that the data strongly suggest a large estimated mutual information, for two reasons:

1. All pairs that assign very low probability to our data, say less than  $10^{-40}$ , are represented by small blue points. Each one is very

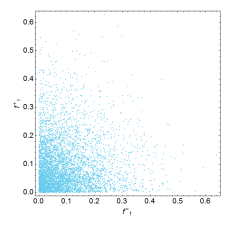


Figure 4

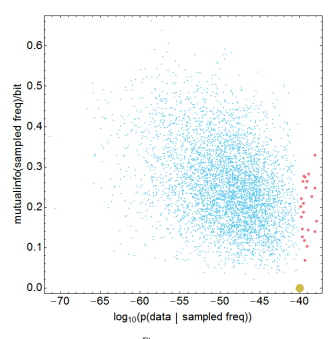


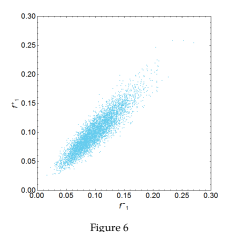
Figure 5

unlikely to be the continuation of our data. But at the same time they constitute the overwhelming majority of possible pairs, and therefore there is a non-negligible probability that the data come from one of them. Most of them have high mutual information.

2. The pairs that assign higher probability to our data,  $10^{-40}$  or more, are represented by the larger red points and the largest yellow point. The majority of these pairs also have high mutual information. In fact, the pair of uniform conditional frequencies is an outlier: it's very unlikely that our data come from it, compared with the other possible pairs of frequencies.

The estimate of fig. 2 is therefore quite correct and reliable.

Now consider the conservative initial density (5), and sample  $5\,000$  pairs of conditional frequencies from it. The scatter plot for the first components of the sampled frequencies is shown in fig. 6. It shows that



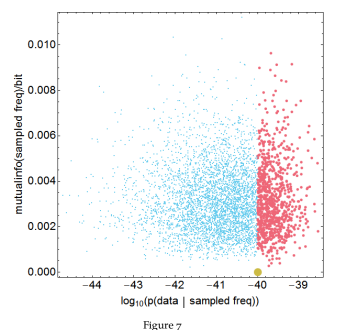
the two long-run frequencies are very similar to each other and close to 1/10, the probability given by the uniform distribution.

Figure 7 shows the pairs of long-run frequencies plotted as in the previous case. There are now many pairs that assign roughly the same probability to the data as the uniform-distribution pair does. The majority of all pairs has a negligible mutual information. This is indeed reflected by the estimate of fig. 3.

#### 4 Brief

We have calculated the probability distribution for the long-run mutual information, given our knowledge of a small sample. The calculation was conceptually straightforward and needed no corrections, even if it may require numerical sampling methods. •• Mention partial exchangeability. We also saw (§ 3) why the distribution thus fund is intuitively correct. The crucial point is the specification of the pre-data joint probability for the long-run conditional frequencies. This specification is especially important if the data are few. It's possible to specify a pre-data probability that express a conservative guess, and the resulting estimate of the mutual information is very close to zero.

This conclusion isn't surprising: the main point is the same as, for example, in the inference of diseases or some phenotypes from genetic peculiarities and vice versa. Imagine that we know the relative frequencies of two gene variants among people who have a particular disease, and the relative frequencies among people who don't have the disease. Now we ask: given that a specific person has one of the gene



variants, what's the probability that this person has the disease? The answer is that in general we don't know until we specify the incidence rate of the disease in the full population. Because even if that gene variant appears more frequently among people with the disease, the number of people having such disease may be so small that the probability still favours the person's being healthy.

The direct calculation via the probability calculus (Jaynes 2003; Sox et al. 2013; Hailperin 1996) makes any discussion about biases superfluous. Note, however, that a serious frequentist statistician would also find that the estimator of the mutual information is *un*biased. The erroneous conclusions reached in papers claiming a sampling bias for mutual information (Panzeri et al. 2007) i add others may be summarized thus: they confuse a non-parametric problem (see e.g. Wasserman 2006, a frequentist book) with a parametric one. The problem is not the estimation of a parameter that identifies a distribution in a family, but the estimation of the distribution itself: the whole distribution – or distributions in our case – is the 'parameter'. The mutual information is a function of this 'parameter'. The procedure of Panzeri et al. (2007 pp. 1065–1066) is therefore wrong.<sup>2</sup> To assess the expectation of the estimator we must sample the 'parameter'; in our case this means drawing *N* samples of the empirical conditional frequencies in n data and then taking their average to estimate the long-run frequency. The latter is a notoriously unbiased estimator. Then we compute the mutual information from this estimate. If we do this with  $N = 5\,000$  and n = 20 for a pair of uniform conditional distributions we obtain a mutual-information estimate of the order of  $10^{-5}$ . As N increases, with n = 20 fixed, the estimate goes to zero. There is no bias.

The topsy-turvy point of view typical of frequentist-like analyses unfortunately leads to this kind of oversights. Our problem is not to infer data from a specific known frequency, but to infer an unknown frequency *among several possible ones* from known data – because our uncertainty about the long-run mutual information is a consequence of our uncertainty about the long-run frequency.

## - Paragraph on shrinkage

<sup>&</sup>lt;sup>2</sup>Note in passing that plotting the distribution of a parameter or of a function thereof, as in Panzeri et al. (2007 Fig. 1, right columns), is anathema from a frequentist point of view: parameters do not have distributions, by divine Fisherian decree.

#### Baraka

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- ('de X' is listed under D, 'van X' under V, and so on, regardless of national conventions.)
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