What is the statistical significance of the solar neutrino flux problem?

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Abstract

The discrepancy between measured and predicted neutrino fluxes on earth may be less significant than generally believed. Indeed, a technically incorrect method has so far been used to propagate the errors on the standard solar model input parameters through the calculation of the predicted neutrino flux. Input parameters should have been drawn from uniform not Gaussian distributions. The uncertainty on the flux may therefore have been substantially underestimated and the significance of the observed discrepancy correspondingly overestimated. Results from different experiments which cannot be directly compared without reference to a model, may also be less incompatible than currently believed.

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Uncomfortably, an almost trivial remark must be made about the technique employed to compare the measured flux on Earth of solar neutrinos, with the prediction of the standard solar model. For a clear statement of the problem, the reader is referred to John Bahcall’s concise talk at the XXVI International Conference on High Energy Physics [1], with apologies to all other scientists who contributed so much to this subject.

Very briefly, the flux of solar neutrinos on Earth is measured by several experiments using various techniques [2–5]. This flux is independently predicted using the standard solar model [6–8]. The prediction depends on a collection of input parameters derived from prior experiments with substantial uncertainties. These uncertainties are propagated in the neutrino flux prediction, which is found incompatible with the experiments, and further renders incompatible such experiments that cannot be directly compared but are nevertheless related through the model [1].

The complexity of the standard solar model and the subsequent neutrino flux derivation precludes the straightforward analytical computation of the covariance matrix which, together with the errors on the input parameters, would supply the uncertainty associated with a predicted value. Standard solar model authors [6,7] estimate this uncertainty by repeating the calculation of the model with several, typically one thousand [1], sets of randomly selected input parameters, each drawn from a Gaussian distribution defined by the experimental value and error for the input parameter.

This technique is of course perfectly legitimate except that the sets of input parameters should be drawn from uniform, not Gaussian, distributions.

The rest of this note explains why, and what the consequences may be.

The source of this widely spread wrong practice is the confusion between quantities whose nature is random and quantities which are fixed even if unknown.

When an experimentalist measures some physical parameter, the outcome of the measurement is a random variable, but the parameter is a fixed constant of perfectly well defined value even if nothing is known yet about that value. Usually, many independent random sources (of finite variances to be absolutely correct) contribute to the probability distribution of the outcome of the measurement which is Gaussian by virtue of the central limit theorem; this Gaussian probability has a very well defined but unknown expectation value (the actual value of the parameter being measured), and its variance is the sum of the contributing variances.

When the value of the parameter is known, it is legitimate to simulate the possible outcome of a measurement by drawing random numbers from a Gaussian distribution of known expectation value and variance. This is common practice, for example, in High Energy Physics detector simulation, where particle momenta are originally produced by an event generator with strict momentum conservation, and are then smeared, and a detector momentum dependent response guessed, by adding random errors.

Because a physical parameter is not a random number, it is completely meaningless to talk about the probability that such a parameter has some given value [4]. Usual intuitive speculations about the value of an unknown parameter begin by wrongly considering it as a probabilistic quantity. What an experimental result stated as $\hat{P} = \hat{P} \pm \sigma$ teaches us about the actual value of some parameter $P$ which is being measured, is simply this [10]:

“There is no more than 31.73% probability to be wrong when asserting that the interval $[\hat{P} - \sigma, \hat{P} + \sigma]$ covers the actual value of the parameter $P$.”
This is the common definition of a confidence interval. Nothing is gained, no statement is made more conservative, by making the interval twice as large and correspondingly limiting the risk for a wrong statement to $4.55\%$, or any other combination, as long as the error distribution is Gaussian. There is no additional information about where the actual value of $P$ is located; it is even not necessarily covered by the confidence interval. There is therefore no reason to give preference to any value. After having taken a calculated risk to be wrong when taking for granted that the actual value is indeed covered by the confidence interval, there still is no reason to give preference to any value within that interval. Therefore, values to be tried for the parameter should be selected uniformly over the arbitrarily wide confidence interval (without forgetting the risk of systematic wrong choice since the interval may fail to cover the actual value of the parameter), rather than following a Gaussian distribution which very strongly prefers points close to the center of the interval. The common intuition that values close to the center are more likely, simply fails to recognize that the unknown but fixed central value (the parameter) of a Gaussian distribution and one random but known outcome (the measurement) drawn from this distribution are not functionally symmetrical and may therefore not be exchanged.

By taking an infinitely wide confidence interval, the probability to make a wrong statement when saying that this interval covers the actual value of the parameter vanishes. This is the uninteresting situation in which all models fit all data. Remembering that $\sigma$ is the initial experimental error on the input parameter, the width $2k\sigma$ of the interval to be used can however be selected after an arbitrary (reasonable) threshold for wrong statements $\epsilon$ has been set. For example, if the author of this note would always have set $\epsilon$ to the one standard deviation level, $31.73\%$ of the papers carrying his name would contain false results. Before invoking the need for new Physics following a discrepancy between an experiment and a model, one might probably at least want a safer $2\%$ level, still one or two wrong papers in a life time, which corresponds to $k = 2.58$ (2.58 standard deviations).

The extension to more than one parameter is straightforward even if, again, seemingly anti–intuitive. For a point in $n$-dimensional space to be located outside of a given hypercube, it is necessary and sufficient that at least one coordinate be located outside the corresponding confidence interval; more coordinates out of their confidence intervals are not relevant. The probability that at least one confidence interval does not cover the corresponding parameter, which is simply the complement of the probability that all intervals cover their own parameter, is $1 - (1 - \epsilon)^n \approx ne$. The common reduced width of uniform distributions for $n$ parameters is therefore the number of standard deviations at which the Gaussian tail is equal to $\epsilon/n$ where, we remind, $\epsilon$ is the threshold for the acceptance of publishing a wrong result.

Predictions for the solar neutrino flux have been compared with experiment [1] by generating 1,000 solar models with 1,000 sets of input parameters drawn from Gaussian distributions. Although finite width uniform distributions will clearly not let the guessed parameters vary as far as normal distributions, the uniform density in the hypercube is very different from the clustering around the central values that Gaussian distributions provide. Thus, the distribution of predicted fluxes will almost certainly become very substantially wider using uniform distributions. We will however have to wait until the correct method for guessing parameters is applied to existing computer programs [6,7], to see how the significance of the observed discrepancies is affected.
Discussions with colleagues have shown that the very mundane facts about statistics quoted above are not well perceived. A very simple example is therefore presented to illustrate the issue. Consider a simple pendulum, made of a point mass at the free end of a thin string of length $l$. The model to be checked says that the period $T$ is given by the formula $T = 2\pi \sqrt{\frac{l}{g}}$ with $g = \frac{GM}{R^2}$ where $G$ is the gravitational constant, $M$ the mass of a spherical Earth and $R$ its radius. The length $l$ has supposedly been directly measured, and a prediction for $l$ derived from measurements of $\pi, G, M, R, T$. The measured value of $l$ is compared with the prediction, possibly requiring “new Physics” such as massive strings or non spherical earth or highly viscous air etc...

Let us begin with the physical constant $\pi$, which as everybody knows has a very well defined value. Hopefully, nobody will ever ask the question “what is the probability that $\pi = 3.17$ ?”, since $\pi$ is not a random variable. Assume that we get the value of $\pi$ from a very cheap three digits display pocket calculator, showing $\pi = 3.14$. We know nothing about the next digit (a fresh student might perhaps write $\pi = 3.145 \pm 0.005$: this however is not fair, because we do not take 31.73% chances to make a wrong statement when stating that the interval $[3.14, 3.15]$ covers the true value of $\pi$, we take exactly 0% such risk). When estimating the effect on the predicted length of the pendulum of the uncertainty on $\pi$, would it come to anybody’s mind to draw values for $\pi$ from a normal distribution centered around 3.145 and of $\sigma = 0.002$ ($\sigma$ set such that the random number almost always remains in the $[3.14, 3.15]$ interval)? Obviously, one will rather try each of 0, 1, . . . , 9 in turn as the next digit; random numbers from an uniform distribution do the same: the order of the trial is of no importance, nor the exact value if enough trials are made.

Let us now turn to any of the four other parameters in our trivial problem. Is there any reason to handle them differently, just because their 100% confidence intervals are not finite?

This trivial example has been simulated. Figure 1 shows the expected Gaussian distribution of a direct measurement of a string of known length, and simulated model predictions using Gaussian and uniform errors for the input parameters. We have artificially considered $\pi$ as a parameter to enhance the effect of having many parameters. Values and errors for $G, M, R$ have been taken from litterature [10,11], and errors of the same order of magnitude have been set for $\pi$ and the “measured” length and period. The period was set to two seconds, the well known beat of a one meter pendulum. The nominal true length was slightly shifted as shown in the figure so that Gaussian simulated predictions come out incompatible within several standard deviations while uniformly simulated ones “agree” quite often, always at the 1% risk of wrong publication level. This exercise has no other purpose than illustrating the broadening of the distribution of the model prediction when uniform distributions are used. It does not imply anything about the solar neutrino flux. At best, it may hopefully trigger authors of solar models to rerun their programs with uniform errors.

In conclusion, a more reliable estimate of the significance of the discrepancy between experiments and standard solar model neutrino fluxes could easily be obtained by rerunning computer simulations of the standard solar model with input parameters drawn from uniform rather than Gaussian distributions, of widths controlled by the priorily set accepted risk of publishing a wrong result.

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FIGURES

FIG. 1. The expected distributions of direct length measurements (solid line), and of the length predicted from a measurement of the period using Gaussian random input parameters (broken line), or uniform random input parameters (dotted line). A cross shows the nominal length and its one standard deviation error.