Bayesian curve fitting for lattice gauge theorists

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A new method of extracting the low-lying energy spectrum from Monte Carlo estimates of Euclidean-space correlation functions which incorporates Bayesian inference is described and tested. The procedure fully exploits the information present in the correlation functions at small temporal separations and uses this information in a way consistent with fundamental probabilistic hypotheses. The computed errors on the best-fit energies include both statistical uncertainties and systematic errors associated with the treatment of contamination from higher-lying stationary states. Difficulties in performing the integrals needed to compute these error estimates are briefly discussed.

1. INTRODUCTION

The physical consequences of a quantum field theory can be deduced from the correlation functions of the fields. Usually this is done by matching the correlation functions (or suitable combinations of them) to known behavior which contains the observables of interest. The path integrals which yield the correlation functions are often computed using Monte Carlo estimates. The Monte Carlo estimates of the correlation matrix tend (neglecting boundary conditions) to a handful of damped exponentials whose decay rates yield the lowest-lying energies of the system. To reliably determine these decay rates, the correlators $h^{ij}(t)$ must be calculated for large enough $t$ such that they are well approximated by only small numbers of exponentials. The Monte Carlo estimates of the correlation matrix $h^{ij}_{M}(t)$, where now $D$ denotes Monte Carlo estimates, must then be fit to known asymptotic forms $h^{ij}_{D}(t)$, where $M$ denotes the model fitting functions. The fit is carried out by adjusting the fit parameters, collectively denoted by $u$, in $h^{ij}_{M}(t)$ in order to maximize the likelihood of finding the results $h^{ij}_{D}(t)$ given $h^{ij}_{M}(t)$. This amounts to minimizing

$$
\chi^2 = \sum_{\alpha\beta} \left( M_{\alpha}(u) - D_{\alpha} \right) C_{\alpha\beta}^{-1} \left( M_{\beta}(u) - D_{\beta} \right), \tag{2}
$$

where $M_{\alpha}(u)$ refers to $h^{ij}_{M}(t)$ and $D_{\alpha}$ refers to $h^{ij}_{D}(t)$. The indices $\alpha$ and $\beta$ each include the indices of the correlation matrix as well as the instants in time at which the correlation matrix is sampled. $C_{\alpha\beta}$ is the covariance matrix associated with $D_{\alpha}$. One commonly-used method is to use the fitting function (which neglects boundary conditions)

$$
h^{ij}_{M}(t) = \sum_{p=0}^{n-1} Z^{ip} Z^{jp} \exp(-tE_p), \tag{3}
$$

with $E_{p+1} > E_p$, and fit the $n \times n$ matrix $h^{ij}_{D}(t)$ for $t_{\min} \leq t \leq t_{\max}$. If the $n$ operators have been constructed using a variational approach applied to a much larger set of operators, the off-diagonal matrix elements $h^{ij}_{D}(t)$ are often statistically consistent with zero. In such cases, little is gained by including these correlation functions in the fitting, and hence, one may fit only to the $n$ diagonal correlators using the fitting function

$$
h^{(i)}_{M}(t) = \sum_{p=0}^{n-1} Z^{ip} \exp(-tE_p), \tag{4}
$$

where now the $Z^{ip}$ are usually real and positive.

When using either Eq. (3) or (4) as the fitting function, it is crucial that $t_{\min}$ be chosen such
Figure 1. Effective mass plot showing fits for the lowest four energies of Example 1 using the model fitting function from Eq. (4) and $7 \leq t \leq 14$. The data points in this figure are effective masses as defined by Eq. (5.18) in Ref. [1] such that the lowest effective masses tend to the lowest physical masses as the temporal separation becomes large. Note that the correlation functions themselves are used in the fitting, not the effective mass points which are included only for illustrative purposes.

Table 1

| $t_{\min}$ | $t_{\max}$ | $Q$ | $E_0$ | $E_1$ |
|------------|------------|-----|-------|-------|
| 3          | 10         | 0   | 0.130632(27) | 0.32435(10) |
| 4          | 11         | 0   | 0.130514(24) | 0.32319(10) |
| 5          | 12         | 0   | 0.130449(27) | 0.32268(14) |
| 6          | 13         | 0.23 | 0.130421(33) | 0.32217(21) |
| 7          | 14         | 0.69 | 0.130413(32) | 0.32210(26) |
| 8          | 15         | 0.21 | 0.130404(31) | 0.32206(32) |

that contributions from all higher energy eigenstates are negligible. Generally one is guided by the quality of fit to determine this, but the guidelines for doing so are not very clear. For example, one could start with $t_{\min} = 0$, for which the fit quality is usually very bad, and increase $t_{\min}$ by one temporal lattice spacing at a time, keeping $t_{\max} = t_{\min} + t_{\text{window}}$ for fixed $t_{\text{window}}$, until some acceptable fit quality, such as $Q = 0.2$, is achieved. But the acceptable value of $Q$ and the value of $t_{\text{window}}$ are somewhat subjective, and it is not clear how to incorporate the uncertainties in choosing these values into the final best-fit values for the energy levels. More stringent values of $Q_{\text{acc}}$ and $t_{\text{window}}$ generally lead to larger values of $t_{\min}$, and hence, larger errors in the best-fit energies. The subjectivity of these errors and the loss of information associated with the discarded $t < t_{\min}$ measurements of the correlators are undesirable features of this approach.

For illustrative purposes, consider two examples, one with an excellent signal (Example 1), the other with a questionable signal (Example 2). Both examples are energies of a gauge field in the presence of a static positive and a static negative charge in 2+1-dimensional compact U(1) lattice gauge theory using an improved action on an anisotropic lattice. Typical fit results for the lowest few energies using the standard analysis with the fitting function from Eq. (4) are shown in Figures 1-3. $S_g$ denotes the chosen symmetry channel and $R$ is the number of lattice sites between the charges. Tables 1 and 2 detail the fit results for the lowest-lying energies $E_0$ and $E_1$ in each example. In Example 1, a suitable choice for $t_{\min}$ and $t_{\max}$ is easily guided by the fit quality $Q$ and the final results are reasonably insensitive to minor changes in the fit range. In Example 2, one might agonize over whether $t_{\min} = 4$ or 5 should be used. The standard analysis provides no means of incorporating such an uncertainty into the error estimate.

In order to incorporate information from small time separations, many exponentials must be retained in the model fitting functions. Standard fits using many exponentials quickly encounter severe problems: often, fitting instabilities occur and uncertainties in the estimates of the parameters of interest become very large. The source
of these problems is the fact that unconstrained fits allow physically insensible or impossible values for those parameters which are not adequately constrained by the data. The solution to this problem is to introduce constraints. Bayesian statistics allows the introduction of such constraints in a natural way. The Bayesian approach is now widely used throughout many disciplines, such as economics, medicine, astrophysics, and condensed matter physics.

This talk is a report on progress being made in exploring methods of extracting physical observables from Monte Carlo estimates of correlation functions which incorporate Bayesian inference into the statistical analysis. Such procedures can fully exploit the information present in the correlation functions at small temporal separations and use this information in a way consistent with fundamental probabilistic hypotheses. The computed errors on the best-fit parameters include both statistical uncertainties and systematic errors associated with the treatment of the excited-state contaminations. Example 1 will be used to verify that such new methods work in cases where the signal is excellent. Example 2 will then be used to see if the new methods provide an improved means of extracting the information in cases when the signal is more obscured.

An earlier report on such explorations is given in Ref. [2]. Bayesian methods have been previously discussed in the context of lattice QCD in Ref. [3], and more recently, using maximum entropy methods in Refs. [4–7].

Since these methods employ Bayesian inference, a brief review of the Bayesian approach is given in Sec. [2]. The application of the Bayesian approach to the particular problem of interest...
here, namely, extraction of energies from Monte Carlo determinations of correlation functions, is then detailed in Sec. 2.

2. BAYESIAN REGRESSION

Bayesian regression is different from the classical (frequentist) approach. In the classical approach, the data are the only source of information explicitly taken into account in constructing an estimate or test. In the Bayesian approach, an estimate or test is produced by combining the current data with information from past experience and/or theoretical constraints (prior information). In this section, Bayesian regression is described in general terms. Some introductory textbooks on Bayesian statistics are given in Refs. 8-10. For particular emphasis on Bayesian regression, see, for example, Refs. 11,12, and for a few other discussions of Bayesian methods in physics, see Refs. 13,14.

The Reverend Thomas Bayes was a Presbyterian minister born in 1702 in London, England, and who died April 17, 1761. His theory of probability was published posthumously by Richard Price in 1763 in an article entitled an Essay towards solving a problem in the doctrine of chances in the Philosophical Transactions of the Royal Society. The theorem presented by Bayes was restricted to the binomial distribution, but its generality was most likely recognized by Bayes. Bayes’ theorem was generalized beyond the binomial distribution by Laplace in 1774, most likely without knowledge of Bayes’ work. Standard (frequentist) statistical methods were developed later than Bayesian methods. Although linear regression and goodness of fit first appeared in the late 1800’s, the field blossomed during the 1920’s with the works of Fisher, Neyman, and Pearson, with a flurry of research and applications during World War II. Bayesian methods are much older, but were largely ignored or actively opposed until the 1950’s when they were championed by prominent non-statisticians, such as physicist H. Jeffreys and economist A. Bowley. Their popularity grew rapidly during the 1970’s with the advent of affordable computers, and heated debates over the superiority of either method have raged ever since.

The cornerstone of Bayesian statistics is Bayes’ theorem:

\[
P(M|D \cap I) = \frac{P(D|M \cap I) P(M|I)}{\int dM P(D|M \cap I) P(M|I)},\tag{5}
\]

where \(D\) represents the data, \(M\) denotes the model which one believes should describe the data, and \(I\) represents our prior knowledge or background information about the system. The conditional probability distribution \(P(D|M \cap I)\) is known as the likelihood of the data, \(P(M|I)\) is the prior probability distribution, and \(P(M|D \cap I)\) is called the posterior probability distribution. The term in the denominator of Eq. (5) is independent of \(M\) and so can often simply be absorbed into an overall normalization. Bayes’ theorem essentially states that

\[
\text{posterior} \propto \text{likelihood} \times \text{prior}.
\]

Bayesian regression uses the posterior distribution for all statistical inference. Model parameters are estimated using one’s favorite statistic but evaluated using the posterior distribution. Common measures of central tendency include the mode, mean, and median, and common measures of dispersion include the variance, skewness, and kurtosis. For example, if the model is specified by a set of parameters \(u\), then the mean value and variance of a model parameter \(u_j\) are given by

\[
\langle u_j \rangle = \int du \ u_j \ P(M(u)|D \cap I),\tag{6}
\]

\[
\text{var}(u_j) = \int du \ (u_j - \langle u_j \rangle)^2 \ P(M(u)|D \cap I).\tag{7}
\]

The mode of \(u_j\) is found by maximizing the posterior probability, that is, maximizing the probability of the model \(M\) being correct given data \(D\) and subject to the background information \(I\). The likelihood \(P(D|M \cap I)\) is the probability distribution of the finding the data \(D\) given the particular set of model parameters \(M\) and background information \(I\), and from the Central Limit theorem has the form

\[
P(D|M \cap I) \propto \exp(-\chi^2/2),\tag{8}
\]

where \(\chi^2\) is the familiar form given in Eq. (3).
The new and key feature of Bayesian statistics is the prior \( P(M|I) \). In standard maximum likelihood fits, the Bayesian prior \( P(M|I) \) is taken to be a constant and thus, maximizing the likelihood becomes equivalent to minimizing \( \chi^2 \). When the number of fit parameters is small compared to the number of data points, this is a reasonable thing to do. However, when the number of fit parameters becomes comparable or larger than the number of data points, the above procedure can become unstable and/or yield parameter estimates with overly large uncertainties, as mentioned previously. The role of the Bayesian prior \( P(M|I) \) is to filter out parameter values that are impossible and improbable, given some prior knowledge, but overly large widths could lead to overly large uncertainties in the best-fit parameters.

Prior distributions are specified based on information accumulated from past studies (earlier data), the opinions of subject-area experts, and/or from theoretical constraints. One often considers logical connections between the parameters and symmetries which the prior distribution must obey. To simplify the computational burden, practitioners often restrict the choice of prior to some familiar distributional form. In some cases, the prior can be endowed with very little informative content (such as maximum entropy which amounts to letting a monkey throw balls into bins). Strictly speaking, using the observed data to influence the choice of prior is against the Bayesian philosophy. However, it is legitimate to use a small subset of the data to guide the construction of the prior and to use the rest of the data in the regression itself. It is very important that one should avoid putting in more information than is truly known, since results do and should depend on the prior. The prior is often viewed both as a nuisance and as an opportunity.

3. CHOICE OF PRIOR AND TESTING

By introducing a Bayesian prior, we can now use the model function of Eq. (4) but summing over many more exponentials:

\[
h_{M}(t) = \sum_{p=0}^{N_{\text{exp}}-1} A_{p}^{(i)} \exp(-tE_{p}),
\]  

where \( N_{\text{exp}} \) is the number of exponentials, \( N_{\text{cor}} \) is the number of correlators to be simultaneously fit, and \( i = 0, \ldots, N_{\text{cor}}-1 \). To ensure positivity of the coefficients and to order the energies, define

\[
A_{n}^{(i)} = \left( \frac{h_{n}^{(i)}}{\gamma_{n}} \right), \quad E_{n} = E_{n-1} + \epsilon_{n}^{2}, \quad (10)
\]

so that the actual parameters used in the fitting are \( u_{\alpha} = \{ E_{0}, \epsilon_{n}, b_{n}^{(i)} \} \). In total, there are \((N_{\text{cor}}+1)N_{\text{exp}}\) fitting parameters. We then choose a Gaussian prior

\[
P(M|I) \propto \exp \left( -\sum_{\alpha} \frac{(u_{\alpha} - \eta_{\alpha})^{2}}{2\sigma_{\alpha}^{2}} \right), \quad (11)
\]

In this work, we shall not include a prior for the first \( N_{\text{cor}} \) energies, so that the sum over \( \alpha \) in Eq. (11) includes all parameters except \( E_{0}, \epsilon_{1}, \epsilon_{2}, \ldots, \epsilon_{N_{\text{cor}}-1} \). Hence, there are \( N_{\text{cor}}N_{\text{exp}} \) terms in the summation over \( \alpha \). In using such a prior, we incorporate our expectation that the parameter corresponding to the index \( \alpha \) most likely has a value between \( \eta_{\alpha} - \sigma_{\alpha} \) and \( \eta_{\alpha} + \sigma_{\alpha} \).

To simplify matters, we assume that the energies of the excited-state contamination above the \( N_{\text{cor}} \)-th level are most likely to be equally spaced. Also, due to the variational construction of our operators, we expect that correlator \( j \) is dominated by the \( E_{j} \) exponential. We further simplify matters by taking the same \( \eta_{\alpha} \) and \( \sigma_{\alpha} \) for all the \( b_{j}^{(i)} \). All other coefficients are expected to be small and are given the same expected range of values. To summarize, our prior is specified using

\[
\eta_{\alpha} = \begin{cases} 
\varepsilon, & \alpha = \varepsilon, & j = N_{\text{cor}} \ldots N_{\text{exp}}-1, \\
\Gamma, & \alpha = b_{j}^{(i)}, & j = 0 \ldots N_{\text{cor}}-1, \\
\gamma, & \alpha = b_{j}^{(i)}, & i \neq j,
\end{cases}
\]

and for the Gaussian widths, we similarly use \( \sigma_{\varepsilon}, \sigma_{\Gamma}, \) and \( \sigma_{\gamma} \). Choosing values for \( N_{\text{exp}}, \varepsilon, \Gamma, \gamma, \sigma_{\varepsilon}, \sigma_{\Gamma}, \) and \( \sigma_{\gamma} \) completes the specification of our prior. \( N_{\text{exp}} \) is easily chosen by increasing its value until the energies of interest stabilize. The widths must be chosen sufficiently large so that the fits are not overly constrained beyond our knowledge, but overly large widths could lead to overly large uncertainties in the best-fit parameters. Note that describing the prior is very straightforward, in contrast to describing how one
Figure 4. Effective masses and fits for the lowest four energies of Example 1 using the Bayesian analysis described in Sec. 3. Prior parameter values are $N_{\text{exp}} = 34$, $\varepsilon = 0.2$, $\sigma_\varepsilon = 0.1$, $\Gamma = 0.9$, $\sigma_\Gamma = 0.2$, $\gamma = 0.05$, $\sigma_\gamma = 0.05$.

subjectively chooses fitting ranges $t_{\text{min}}$ and $t_{\text{max}}$ in the standard analysis.

The results of applying this Bayesian curve fitting method in the two examples previously described are shown in Figs. 4 and 5. The fit values shown correspond to the mode of each quantity, that is, the value of the quantity at the maximum of the posterior distribution. The errors have been computed assuming a normal distribution about this maximum for reasons discussed below. These errors automatically combine the statistical errors in the data with the systematic uncertainties in our prior knowledge. The method works remarkably well in both examples, eliminating the need to subjectively determine $t_{\text{min}}$.

The insensitivity of the results to the parameters in the prior probability is demonstrated in Fig. 6. Notice that the error estimates increase significantly only in case (e) which is a drastic change in the prior. This insensitivity indicates that $E_0$ and $E_1$ are largely being determined by the data. Of course, the data contain little information about many of the other parameters, and hence, such parameters depend strongly on the prior. We have found that this new method works well as long as $\varepsilon$ is not set much larger than the expected level spacings in the system and the allowed ranges of the coefficients are not set too large, as in case (e).

Currently, there remains one fly in the ointment, namely, the computation of the error estimates. We have evaluated the multi-dimensional integrals needed to calculate the variances in the fit parameters using both the Metropolis method and HMC and have encountered severely long auto-correlations. The presence of many poorly constrained parameters leads to long, narrow ridges in the probability distribution which hamper the Markov chain Monte Carlo integration. For the Metropolis method, we have commonly observed auto-correlation lengths in the tens of thousands. The problem is somewhat ameliorated with HMC, reducing the auto-correlation lengths by factors of twenty or so, but clearly, this is still not sufficient. Work continues on this topic. An alternative method of computing the
Figure 6. Sensitivity of $E_0$ and $E_1$ to changes in the parameters of the Bayesian prior. For each point, the parameters in the prior are the same as stated in Figs. 4 and 5, except for the following changes:

(a) $\varepsilon = 0.1$, $\sigma_\varepsilon = 0.05$,
(b) $\varepsilon = 0.2$, $\sigma_\varepsilon = 0.10$,
(c) $\varepsilon = 0.3$, $\sigma_\varepsilon = 0.15$,
(d) $\Gamma = 0.8$, $\sigma_\Gamma = 0.3$, $\gamma = 0.1$, $\sigma_\gamma = 0.1$,
(e) $\Gamma = 0.7$, $\sigma_\Gamma = 0.7$, $\gamma = 0.7$, $\sigma_\gamma = 0.7$.

errors (which deviates from a strict Bayesian philosophy) is suggested in Ref. [2].

4. CONCLUSION

Bayesian regression techniques are a viable alternative method for extracting physical observables from stochastically-determined correlation functions. By allowing the regression to take into account prior knowledge of the system from theoretical considerations and/or previous experience, Bayesian methods can make use of information obtained at short temporal separations and can incorporate systematic effects into error estimates of the observables. However, Bayesian methods are not a cure for bad data.

Much of this work arose from a collaborative effort to explore Bayesian methods. Others involved in this effort are Peter Lepage, Bryan Clark, Christine Davies, Kent Hornbostel, Paul Mackenzie, and Howard Trottier. The author wishes to acknowledge especially fruitful discussions with Peter Lepage, and also with Robert Swendsen, Julius Kuti, Jimmy Juge, David Richards, and Mike Peardon. This work was supported by the U.S. National Science Foundation under Award PHY-0099450.

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