Bayesian Analysis of Pentaquark Signals from CLAS Data


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The debate about the existence of the $S = +1 \Theta^+(1540)$ baryon state is still going at this point in time in spite of results from dedicated, high-luminosity measurements. One of these [1], from the CLAS collaboration at the Thomas Jefferson National Accelerator Facility, used the reaction $\gamma d \rightarrow pK^+K^-n$. It showed convincing evidence that production cross sections for such a state are nowhere near the levels implied by an earlier CLAS measurement [2] of the same channel, which had seen a peak in the $pK^-$ missing mass spectrum at 1.542 GeV/$c^2$ with a 5.2$\sigma$ statistical significance. The salient point is that the work of Ref. [1] was a dedicated, high-luminosity repeat of Ref. [2], where the experimental running conditions were as similar as practically possible.

In the whole history of $\Theta^+$ pentaquark searches, there were several independent experiments that claimed to have found evidence, while a similar number claimed to have found nothing. It is impractical to examine the results of all such experiments in a consistent fashion, but the similarity of the two CLAS experiments provides us with an ideal opportunity to investigate apparently contradictory results.

One can examine in detail whether any discrepancy arose from the data quality of the two experiments by making systematic tests on, for example, the effects of different cuts. In the original work for both measurements, however, parallel analyses were carried out to confirm the final spectra, and different internal reviews verified the correctness of the analysis procedures. We therefore assume that the quality of the data in both the experiments was consistent, and that the analyses of both experiments were carried out correctly. We concentrate solely on the endpoints of the analyses, namely, the events passing all cuts, which contribute to missing mass spectra.

To get a feel for the problem, we took the data set from Ref. [1] (hereafter referred to as “$g10$” after the CLAS running period in which the data were obtained) which had been analyzed in exactly the same way as the data from Ref. [2] (hereafter referred to as “$g2a$”). The $g10$ data contained a factor of just under six more events, which could be directly compared. The $g10$ data were then split into five independent subsamples, each containing the same number of counts as the $g2a$ data set, and $pK^-$ missing mass spectra were produced. These missing mass spectra would be where a $\Theta^+$ might be expected to appear. The $g10$ subsample spectra are depicted in Figs. 1(a)–1(e), and the $g2a$ spectrum is depicted in Fig. 1(f).

Peak-like features appear in several of the $g10$ subsamples, but the shapes are by no means consistent. As mentioned previously and in keeping with current convention, the $g2a$ result quoted a “significance” of about 5$\sigma$, which was similar to other experiments claiming evidence of discovery. However, 5$\sigma$ means that the probability that a feature is a fluctuation is of the order of $10^{-6}$. This is a very small number; it does not appear to match the relative ease of generating peak-like features in the subsample spectra. How do we quantify the intuitive feeling that the odds of obtaining the observed $g2a$ peak from fluctuations are not as small as 1 in $10^{6}$?

In this Letter, we attempt to address this problem within a Bayesian analysis framework and to suggest an alternative means of quantifying the evidence for discovery. What is specifically required is a quantitative comparison be-
between two hypotheses: “the spectrum contains a peak,” and “the spectrum does not contain a peak.” One can model the shape of a spectrum as the addition of simple functions, provided that they appear to describe the shape of the spectrum reasonably well, and have plausible physical origins (e.g., Gaussians for resolution effects, etc.). We refer to these as “data models,” to distinguish them from theoretical models. The posterior probability that a data model \( M \) is true given some observed data \( D \) is given by Bayes’ theorem,

\[
P(M | D) = \frac{P(D | M)P(M)}{P(D)},
\]

where \( P(D | M) \) is the probability of the data being observed given the model, and \( P(M) \) represents the prior probability of the model being correct. \( P(D) \) is a normalizing constant, which will cancel out in the ratio that compares the posterior probabilities of two models.

Now the data model will depend on some parameters \( \xi \), and the posterior probability of these taking on specific values is

\[
P(\xi | D, M) = \frac{P(D | \xi, M)P(\xi | M)}{P(D | M)},
\]

where \( P(D | \xi, M) \) is the probability of the data being observed given the model and its parameters, and \( P(\xi | M) \) is the prior probability of the parameters. Fitting parameters to data is a matter of maximizing this posterior. The quantity in the denominator of Eq. (2) is known as the evidence for a model and is obtained by marginalizing (integrating) over the parameters:

\[
P(D | M) = \int d\xi P(D | \xi, M)P(\xi | M).
\]

Since the evidence is an integral over the model parameters, it implicitly implements Occam’s razor. Evidence ratios provide a balance between favoring on the one hand the simpler model, and on the other hand the model that better fits the data.

We construct two very simple data models of the missing mass spectra obtained from experiment: (1) Model \( M_0 \): The spectrum can be described by a 3rd order polynomial in the region of interest. This represents the assumption that there is no new particle. A 3rd order polynomial was employed in the original analysis to model the background shape. This model depends on four parameters. (2) Model \( M_p \): The spectrum can be described by a “narrow” Gaussian peak sitting atop a 3rd order polynomial background in the region of interest. Narrow in this case means that the width is significantly less than the region of interest in the mass spectrum. This model depends on seven parameters.

To compare the different models, a ratio of their probabilities in the light of data can be formed:

\[
R_E = \frac{P(M_p | D)}{P(M_0 | D)} = \frac{P(D | M_p)P(M_p)}{P(D | M_0)P(M_0)},
\]

where Bayes’ theorem has been used to obtain the final expression. This is the ratio of evidences for the models multiplied by the ratio of prior probabilities of the models. If there is no prior preference for either model, the final factor is unity, so the ratio of model probabilities becomes a ratio of evidences. \( R_E \) is known as the “Bayes’ Factor” or “evidence ratio.”

It is computationally convenient and equivalent to examine the logarithms of the evidence ratios:

\[
\ln(R_E) = \ln(P(D | M_p)) - \ln(P(D | M_0)).
\]

Determining what value of \( \ln(R_E) \) to use in deciding between data models is somewhat arbitrary, but Jeffreys established [3] a rough evidence scale versus written descriptors: \( |\ln(R_E)| < 1 \) is weak, \( 1 < |\ln(R_E)| < 2.5 \) is substantial, \( 2.5 < |\ln(R_E)| < 5 \) is strong, and \( |\ln(R_E)| > 5 \) is
So model comparison is quantified by $R_E$, and as constructed means that data favoring a data model with a peak have positive $\ln(R_E)$.

To evaluate evidences, we see from Eq. (3) that an integral over a likelihood $P(D \mid \xi, M)$ and a prior $P(\xi \mid M)$ is required. We calculate the likelihood by evaluating for each bin in a spectrum an “ideal” number of counts, $S_i(\xi)$, for a given set of parameters. The probability of this being correct given the measured counts $n_i$ is calculated using a Poisson distribution. The total likelihood is then a product of these probabilities for each bin:

$$P(D \mid \xi, M) = \prod_i S_i^n \exp(-S_i) / n_i!.$$  \hspace{1cm}(6)

Here, the prior probability is constructed by assuming no initial correlations between parameters, so it is simply a product of priors for each separate parameter. We assume that each prior is a uniform distribution between a lower and upper limit since this represents the least initial bias. The prior parameter ranges were established by performing an initial fit and setting the limits to be $\pm 50\%$ of the values found. This resulted in a large flexibility in the shapes of both background and peak.

To perform the integrations over the many parameters in the models, we utilized the technique of “nested sampling” developed by Skilling [4,5]. Essentially, this is a Monte Carlo integration method developed specifically for Bayesian data analysis. We refer the reader to the original reference for details, and to Ref. [6] for an example application.

We applied the model comparison framework to all the spectra shown in Fig. 1. In addition, we analyzed the spectra shown in Fig. 2, which consisted of: (a) the full g10 spectrum; (b) a “fake” spectrum, constructed by sampling from a combination of signal and background functions in the data model with the peak ($M_{\rho}$), which had the same signal-to-background ratio as the g2a spectrum. This was done to show what the results of this analysis would have been had a resonance been there; (c) and (d) $pK^-$ missing mass spectra from the g2a and g10 data sets, but showing the $\Lambda(1520)$ signal, in order to test how the technique fared for the case of a well-established particle.

The results are quoted in Table I, and displayed graphically in Fig. 3. We omit the results for the $\Lambda(1520)$ from the figure, as they would render the scale unusable. To estimate the uncertainty in the Monte Carlo integrals, we ran at least 20 independent calculations for each spectrum analyzed. The errors listed in the Table represent the standard error of the samples.

With the splitting of the g10 data set, we have shown (Fig. 1) the relative ease with which one can obtain a peak-like feature, given a small number of events. The evidence ratios calculated for the individual subsamples in g10 generally suggest a bias against a peak, which perhaps mirrors an intuitive feeling about how significant such features really are. However, two of the five subsamples (2 and 4) are compatible with the “weak” category, meaning that the results are essentially inconclusive. Whilst the g2a result is more of an outlier, it also falls in the weak category and is inconclusive; the results of the two measurements are therefore compatible with each other.

The $\ln(R_E)$ value for g2a ($-0.408$) indicates weak evidence in favor of the data model without a peak in the spectrum. What this means is that while a data model including a peak gives a better fit by eye to the spectrum, it does not compensate for having had to introduce additional parameters for the peak. This is Occam’s razor in action; simpler models are preferable unless more complex models do much better. One must be careful what to

### TABLE I. Evidence ratios. Calculations are done by nested sampling, hence the need to include standard errors.

<table>
<thead>
<tr>
<th>Data sample</th>
<th>$\ln(R_E)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>g10 sample 1</td>
<td>$-1.56 \pm 0.07$</td>
</tr>
<tr>
<td>g10 sample 2</td>
<td>$-1.09 \pm 0.13$</td>
</tr>
<tr>
<td>g10 sample 3</td>
<td>$-1.64 \pm 0.09$</td>
</tr>
<tr>
<td>g10 sample 4</td>
<td>$-1.11 \pm 0.11$</td>
</tr>
<tr>
<td>g10 sample 5</td>
<td>$-1.82 \pm 0.07$</td>
</tr>
<tr>
<td>g10 full</td>
<td>$-2.87 \pm 0.11$</td>
</tr>
<tr>
<td>g2a</td>
<td>$-0.41 \pm 0.10$</td>
</tr>
<tr>
<td>fake</td>
<td>$5.78 \pm 0.27$</td>
</tr>
<tr>
<td>g2a $\Lambda(1520)$</td>
<td>$96.70 \pm 0.70$</td>
</tr>
<tr>
<td>g10 $\Lambda(1520)$</td>
<td>$549.12 \pm 2.17$</td>
</tr>
</tbody>
</table>
conclude from the $g_{2a}$ spectrum, however, since the evidence ratio does not conclusively rule out a peak; it is simply inconclusive.

We now turn to the question of whether the $g_{10}$ experiment could conclusively discriminate between the two possibilities. The log of the evidence ratio for the full $g_{10}$ spectrum is $-2.9$. This makes it strong evidence against a peak in the spectrum. Another way of looking at this is that with this evidence ratio, the odds against a peak in this spectrum are about 17 to 1. Whilst this cannot completely rule out a discovery, another measurement of this channel is probably not necessary. By comparison, the odds in favor of a peak in the fake spectrum are about 320 to 1, meaning that had a signal really been there in $g_{10}$, the experimental result would have been decisive.

The study of the $\Lambda(1520)$ shows that when a resonance is there, this method picks it out rather readily, with both $g_{2a}$ and $g_{10}$ data sets yielding a decisive result. We take this as a positive test that our method works.

In summary, we have applied a Bayesian model comparison method to analyzing the missing mass spectra produced in pentaquark searches. This has been used to study the relationship between the results of two CLAS measurements, which were taken under almost identical conditions. We have shown that there is no conflict between the results of the two experiments, and that the low number of counts in the first experiment resulted in an ambiguous signal. Furthermore, we have shown that the $g_{10}$ result shows strong evidence against the discovery of a pentaquark in this channel. More generally, this method could be applied to any data set where a search for a new state has been carried out, and can provide a quantitative measure with which to judge whether or not a result represents a discovery.

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