A Likelihood Method for Measuring the Ultrahigh Energy Cosmic Ray Composition

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Abstract

Air fluorescence detectors traditionally determine the dominant chemical composition of the ultrahigh energy cosmic ray flux by comparing the averaged slant depth of the shower maximum, $X_{\text{max}}$, as a function of energy to the slant depths expected for various hypothesized primaries. In this paper, we present a method to make a direct measurement of the expected mean number of protons and iron by comparing the shapes of the expected $X_{\text{max}}$ distributions to the distribution for data. The advantages of this method includes the use of information of the full distribution and its ability to calculate a flux for various cosmic ray compositions. The same method can be expanded to marginalize uncertainties due to choice of spectra, hadronic models and atmospheric parameters. We demonstrate the technique with independent simulated data samples from a parent sample of protons and iron. We accurately predict the number of protons and iron in the parent sample and show that the uncertainties are meaningful.

Key words: cosmic rays — acceleration of particles — composition

1 Introduction

Measurements of the energy spectrum, composition and arrival direction distributions elicit clues to the origin of cosmic rays. Since a direct measurement and identification of the cosmic ray primaries is not possible at these energies, we depend on indirect methods to understand the chemical composition of the cosmic ray flux. The methods vary with the detector type. In ground array-type cosmic ray detectors, composition analyses typically focus on the energy-dependence of the muon to charged particle ratio, which is thought to signify the change in primary particle composition as a function of energy [1]. Air fluorescence experiments, on the other hand, image the shower development in the atmosphere and obtain information about the primary composition from $X_{\text{max}}$, the slant depth at which the particle count in the cosmic ray air shower is a maximum.

Showers induced by protons tend to reach the maximum of the shower development at larger slant depths than showers induced by heavier elements. Since shower-to-shower fluctuations are large, a determination of the primary type for individual showers is not possible, but averaged quantities like the so-called elongation rate have been successfully used to understand the composition of the cosmic ray flux on a statistical basis. The elongation rate is defined as the slope of the mean height of shower maximum measured in units of slant depth,

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$< X_{\text{max}} \rangle$, versus the logarithm of the primary cosmic ray energy observed. The measured elongation rate is compared to that of simulated proton and iron primaries to assess whether the cosmic ray composition is proton-like or heavier [2,3,4].

Due to the relative insensitivity of ultrahigh energy cosmic ray detectors to small shifts in composition between various low-Z or high-Z primaries, their observations have historically been compared to what is expected for “protons” and “iron,” with the two elements as stand-ins for all light and heavy elements. For $< X_{\text{max}} \rangle$ (see Section 4), we expect $\sim 100 \text{ g/cm}^2$ difference between protons and iron. However, as shown in Fig. 1, large shower-to-shower fluctuations and the detector resolution cause a significant overlap in their distributions ($\sim 40 - 70 \text{ g/cm}^2$).

Here, we present a method to answer the question: in a fluorescence data sample where one has measured $X_{\text{max}}$ for each event, what is the best estimate for the expected mean $^2$ number of protons and iron from that experiment? This question is not directly answered through other methods. Once the answer to this question is obtained, the mean number can be converted to a flux for protons and iron. This method can be expanded to evaluate the contribution of any chemical element. Further, this method uses more information than averaged quantities like the elongation rate; it makes use of the shape of the $X_{\text{max}}$ distribution rather than simply its mean. Lastly, the technique allows the introduction of systematic uncertainties and prior knowledge or lack thereof concerning the composition spectra, hadronic generators, parameters concerning the extensive air shower and atmospheric uncertainties.

In brief, the technique for extracting the cosmic ray composition essentially requires us to find the best normalizations for the “hypothesized” $X_{\text{max}}$ distributions for protons and iron. By “best” we mean the sum of the expected mean events for the hypothesized distributions that best match the expected mean for the $X_{\text{max}}$ distribution for data. The chance that the combined hypothesized distributions match the parent distribution for data is proportional to the likelihood, $l(D|p, f)$. The likelihood $l(D|p, f)$ is maximized for the best estimate for the expected mean number of protons ($p$) and iron ($f$) where $D$ denotes the data sample. The variables $p$ and $f$ sum to the expected mean number of data events, $p + f = d$. Note the expected means for protons and

\[ \text{Here we will refer to the mean although we are taking a Bayesian approach. The number of protons, iron, etc. predicted by the likelihood can better understood as the best estimate of the “true” number of protons and iron given the universe’s cosmic ray flux and the HiRes aperature rather than the mean number expected for an infinite number of HiRes experiments. (Here the universe’s cosmic ray flux is assumed to be homogeneous and isotropic as there is currently no evidence to the contrary. However, there is nothing to prevent us binning $X_{\text{max}}$ as a function of sky coordinates.) Regardless, for brevity, we use frequentist language.} \]
Fig. 1. Simulated cosmic ray events show shower-to-shower fluctuations which cause a large overlap between proton (black) and iron (red) events when plotted as a function of \(X_{\text{max}}\) and \(\log(E/\text{eV})\). The QGSJet hadronic model was used under the CORSIKA extensive air shower simulator to model the shower profiles. These events were then passed through a simulation of the HiRes stereo detector response.

iron add to the expected mean number of events and not to the total events in the given data sample.

We develop the method using simulated stereoscopic data for the High Resolution Fly’s Eye (HiRes) experiment in Utah. The method is tested with many independent data sets that originate from a simulated parent data set with \(p_{\text{true}}\) proton events generated with an \(E^{-\alpha_p}\) spectrum and \(f_{\text{true}}\) iron events generated with an \(E^{-\alpha_f}\) spectrum. Each bin of this parent distribution is fluctuated according to Poisson statistics to build a set of independent simulated data distributions with which we test the method of predicting \(p\) and \(f\). A
general description of this technique is given in [5]. To include typical instrumental uncertainties in our study, we will apply the method to a particular cosmic ray experiment, the HiRes stereo experiment. All simulated showers used in this paper have been subjected to the full HiRes stereo simulation and reconstruction.

The paper is organized as follows: Section 2 gives an overview of the HiRes detector, which we use as the detector in the simulated data. Section 3 gives a description of the likelihood method used to find the most likely mixture of protons and iron in the simulated data. In Section 4, the likelihood method is first modified to account for our ignorance of the proton and iron spectra, and then demonstrated with simulated data samples. Finally, Section 5 states conclusions and results.

2 The HiRes Detector

HiRes is an air fluorescence experiment with two sites (HiRes 1 & 2) at the US Army Dugway Proving Ground in the Utah desert (112° W longitude, 40° N latitude, vertical atmospheric depth 860 g/cm²). The two sites are separated by a distance of 12.6 km.

Each of the two HiRes “eyes” comprises several telescope units monitoring different parts of the night sky. With 22 (42) telescopes with 256 photomultiplier tubes each at the first (second) site, the full detector covers about 360° (336°) in azimuth and 3° − 16.5° (3° − 30°) in elevation above horizon. Each telescope consists of a mirror with an area of about 5 m² for light collection and a cluster of photomultiplier tubes in the focal plane.

A cosmic ray primary interacting in the upper atmosphere induces an extensive air shower which the detectors observe as it develops in the lower atmosphere. The photomultiplier tubes triggered by the shower define an arc on the sky, and, together with the position of the detector, the arc determines the so-called shower-detector plane. When an air shower is observed in stereo, the shower trajectory is in principle simply the intersection of the two planes. This method can be further improved by also taking advantage of the timing information of the tubes, and in our analysis the shower geometry is determined by a global $\chi^2$ minimization using both the timing and pointing information of all tubes.

The next step in the reconstruction is to calculate the shower development profile. However, light arriving at the detector is collected by discrete PMTs, each of which covers about 1° × 1° of the sky. The signal from a longitudinal segment of the EAS is thus necessarily split among many PMTs. For profile fitting, the signal must be re-combined into bins that correspond to the lon-
gitudinal segments of the EAS in HiRes 1 and HiRes 2. The re-binned signal, corrected for atmospheric extinction and with Čerenkov light subtracted is fit to a Gaisser-Hillas functional form (Eq. 1) using a $\chi^2$ that fits the function to the profiles measured in the two detectors. This form has been shown to be in good agreement with EAS simulations [6,7,8] and with HiRes data [9].

$$N(X) = N_{\text{max}} \left( \frac{X - X_0}{X_{\text{max}} - X_0} \right)^{(X_{\text{max}} - X_0)/\lambda} \exp \left( \frac{(X_{\text{max}} - X)}{\lambda} \right).$$ (1)

From measurements of laser tracks and stars in the field of view of the cameras we estimate that the systematic error in the arrival direction determination is not larger than $0.2^\circ$, mainly caused by uncertainties in the survey of mirror pointing directions.

Various aspects of the HiRes detector and the reconstruction procedures are described in [10,11,12].

3 The Likelihood

To calculate the best estimate of $p$ and $f$, the mean number of protons and iron expected (at a HiRes experiment), we maximize $l(D|p, f)$, the likelihood of the data sample $D$ given $p$ and $f$. To this end, Bayes’ theorem allows us to convert the probability of a particular data sample $D$ given the composition mixture, $P(D|p, f)$, to the probability of the composition mixture given the data, $P(p, f|D)$:

$$P(p, f|D) = \frac{P(D|p, f)q(p, f)}{\sum_{p,f} P(D|p, f)q(p, f)}.$$ (2)

In this expression, $q(p, f)$ is the prior probability of $p$ and $f$. Initially, the prior probability will contain information about the expected distributions for protons and iron and their respective energy spectra. However, in principle, it can contain any number of (un)knowns concerning the hadronic generator, uncertainties in atmospheric parameters, etc. We effectively maximize $P(p, f|D)$ by maximizing $l(D|p, f) \equiv P(D|p, f)q(p, f)$.

To calculate $l(D|p, f)$, we divide the $X_{\text{max}}$ distributions into $N$ bins. We let $P_i$ and $F_i$ be the number of events in the $i^{th}$ bin of the simulated proton and iron samples, respectively, and $D_i$ be the number of data events in the $i^{th}$ bin. (The capitalized quantities are ones which we know at the outset of the calculation.)
While we typically generate samples of simulated events that are much larger than the data sample, the sets $P_i$ and $F_i$ nevertheless represent single instances or fluctuations around the true, unknown means $\tilde{p}_i$ and $\tilde{f}_i$ for the simulated samples. Similarly, the number of data events $D_i$ in the $i^{th}$ bin fluctuates around the true mean count $\tilde{d}_i$. Assigning a Poisson probability to the counts in each bin $D_i$, $P_i$, and $F_i$, we can write the likelihood function as:

$$l(D|\{\tilde{d}_i\}, \{\tilde{p}_i\}, \{\tilde{f}_i\}) = \prod_{i=1}^{N} \left( \frac{(\tilde{d}_i)^{D_i}e^{-\tilde{d}_i}}{D_i!} \right) \left( \frac{(\tilde{p}_i)^{P_i}e^{-\tilde{p}_i}}{P_i!} \right) \left( \frac{(\tilde{f}_i)^{F_i}e^{-\tilde{f}_i}}{F_i!} \right)$$ \hspace{1cm} (3)

The actual values for $\tilde{p}_i$ and $\tilde{f}_i$ will not interest us; these are nuisance parameters which we will eventually eliminate. However, the mean number of data counts expected in the $i^{th}$ bin is expressed as a weighted sum of these parameters:

$$\tilde{d}_i = \epsilon_p \tilde{p}_i + \epsilon_f \tilde{f}_i.$$ \hspace{1cm} (4)

The purpose of the weights $\epsilon_p$ and $\epsilon_f$ is to 1) scale down the (presumably larger) simulated sample sizes to the data sample size, and 2) set the relative mixture of protons and iron expected in the data. Hence, the quantities which we want to estimate are $\epsilon_p$ and $\epsilon_f$. Inserting Eq. (4) into Eq. (3) and marginalizing (integrating out) the nuisance parameters $\tilde{p}_i$ and $\tilde{f}_i$ we define the global likelihood function $l(D|\epsilon_p, \epsilon_f)$ as:

$$l(D|\epsilon_p, \epsilon_f) = \prod_{i=1}^{N \text{ bins}} \int d\tilde{p}_i \int d\tilde{f}_i \left( \frac{(\epsilon_p \tilde{p}_i + \epsilon_f \tilde{f}_i)^{D_i}e^{-(\epsilon_p \tilde{p}_i + \epsilon_f \tilde{f}_i)}}{D_i!} \right) \times \left( \frac{(\tilde{p}_i)^{P_i}e^{-\tilde{p}_i}}{P_i!} \right) \left( \frac{(\tilde{f}_i)^{F_i}e^{-\tilde{f}_i}}{F_i!} \right).$$ \hspace{1cm} (5)

In fact, the integration can be performed exactly (see [5] for details), and reduces to:

$$l(D|\epsilon_p, \epsilon_f) = \prod_{i=1}^{N} \sum_{n=0}^{D_i} \binom{P_i + D_i - n}{D_i - n} \frac{(\epsilon_p)^{D_i-n}}{(1 + \epsilon_p)^{P_i+D_i-n+1}} \times \binom{F_i + n}{n} \frac{(\epsilon_f)^n}{(1 + \epsilon_f)^{F_i+n+1}}.$$ \hspace{1cm} (6)
In practice, then, the function we maximize is not \( l(D|p, f) \) directly; rather, we maximize \( l(D|\epsilon_p, \epsilon_f) \) (or, more precisely, we minimize \( -\log l(D|\epsilon_p, \epsilon_f) \)) with respect to both \( \epsilon_p \) and \( \epsilon_f \).

With our best estimates for \( \epsilon_p \) and \( \epsilon_f \), we can now estimate the mean number of proton and iron events \( p \) and \( f \) expected for the experiment. The estimates suggested in [5] are

\[
p = \epsilon_p (\sum_{i=1}^{N} P_i + N)
\]

which of course reduce to the more obvious estimates \( p = \epsilon_p \sum P_i \) and \( f = \epsilon_f \sum F_i \) in the usual situation when the number of simulated events is much greater than the number of bins in \( X_{\text{max}} \).

### 4 Demonstrating the Method

To demonstrate the method, we create many simulated data samples from a single parent distribution made with \( p_{\text{true}} \) proton and \( f_{\text{true}} \) iron simulated events with energies in excess of \( 10^{18.5} \text{ eV} \). First, we generate a library of showers of various energies and primary particles with the CORSIKA simulator using the QGSJET hadronic model. For a given composition, these showers are selected randomly according to their energy spectrum and then simulated in the detector with a random geometry. We expect the energy spectra to have an energy dependence between \( E^{-2} \) and \( E^{-3} \). In our first application of the method, we will pick \( \alpha_p = 2 \) and \( \alpha_f = 3 \), a choice motivated by previous composition analyses [3]. This will just serve as an example; we will drop this assumption in the next section, where we show how to eliminate the spectral dependence. The simulated data samples are subject to the same quality cuts that we would apply to real data (see e.g. [13]). We require a minimum track length of \( 3^\circ \) in each detector, an estimated angular uncertainty in both azimuth and zenith angle of less than \( 2^\circ \), and a zenith angle less than \( 70^\circ \). We additionally require an estimated energy uncertainty of less than 20\% and \( \chi^2/\text{dof} < 5 \) for both the energy and the geometry fit, and that \( X_{\text{max}} \) appear in the field of view one of the detectors. Lastly, the reconstructed primary particle energy must be greater than \( 10^{19} \text{ eV} \).

After applying the quality cuts to the parent sample, we fluctuate each bin in the \( X_{\text{max}} \) distribution in the parent data sample many times according to Poisson statistics to obtain many independent data samples. For the moment, we assume that we know that \( \alpha_p = 2 \) and \( \alpha_f = 3 \). If the method works, we
Fig. 2. Iron and proton $X_{\text{max}}$ distributions generated with $E^{-2}$ and $E^{-3}$ spectra with energies above $10^{19}$ eV.

expect the means for $p_{\text{max}} - p_{\text{true}}$ and $f_{\text{max}} - f_{\text{true}}$ to be 0, and the calculated uncertainties for $p$ and $f$ should be meaningful; that is, the quoted values for $p$ and $f$ are within 1 $\sigma$ of $p_{\text{true}}$ and $f_{\text{true}}$ in 68% of the data samples. Figure 2 shows the proton and iron distributions used to calculate the likelihood. We use the same events for generating the parent distribution and for the hypothesized distributions in the likelihood. We generate 2571, 773, 665 and 1121 events for $E^{-2}$ protons, $E^{-3}$ protons, $E^{-2}$ iron and $E^{-3}$ iron. A similar but independent set of distributions were combined to build the ‘fake’ data sample.

We first test the method with a mixture of 50% protons and 50% iron. Figure 3 shows $-\log l(D|p, f_{\text{max}})$ and $-\log l(D|p_{\text{max}}, f)$, for one simulated data set with 86 protons and 117 iron events fluctuated from a parent sample of 100 proton and 100 iron events; $p_{\text{max}}$ ($f_{\text{max}}$) is where $l(D|p, f)$ is maximized with respect to $p$ ($f$). From the parabolic shape of these distributions one might suspect that the 1 $\sigma$ uncertainties for $p$ and $f$ can be calculated through the
Fig. 3. $-\log l(D|p, f_{max})$ (top) and $-\log l(D|p_{max}, f)$ (bottom) evaluated for one simulated data sample with 86 proton and 117 iron events fluctuated from 100 proton and 100 iron events. Although regions about the minima are parabolic, the distributions have substantial tails.

The equations

$$-\log l(p_{max} \pm \sigma_p, f_{max}) = -\log l(p_{max}) + \frac{1}{2}$$  \hspace{1cm} (9)

and

$$-\log l(p_{max}, f_{max} \pm \sigma_f) = -\log l(f_{max}) + \frac{1}{2}$$  \hspace{1cm} (10)

where $\sigma_p$ and $\sigma_f$ are the uncertainties for $p$ and $f$, respectively. However, we find that $l(D|p_{max}, f)$ and $l(D|p, f_{max})$ have substantial tails and so their uncertainties have to be calculated numerically.

Figure 4 shows the corresponding distributions for protons and iron normalized to the number of predicted events, the total predicted mean and the simulated...
Fig. 4. The distributions for protons and iron normalized to the number of predicted events, the total predicted mean, and a simulated data set of 86 proton and 117 iron events created from a parent distribution of 100 proton and 100 iron events. The $\chi^2$ between the parent data set and the predicted sum is 6.7 with 8 degrees of freedom.

data set. We calculate the $\chi^2$ for this fit to be 6.7 with 8 degrees of freedom while keeping in mind that the total events predicted by the sum of $p$ and $f$ are not meant to predict the total number of events, but rather the expected mean number of events.

To artificially separate the systematic and statistical uncertainties, one can, for example, remove any prior knowledge concerning the uncertainties in our models for protons and iron (i.e. remove $q(\{p_i\}, \{f_i\}, \epsilon_p, \epsilon_f)$ from
\[ l(D|\{p_i\}, \{f\}, \epsilon_p p, \epsilon_f f) \]

and define statistical uncertainty as

\[ l_{\text{stat}}(D|p, f) \equiv \prod_{i=1}^{N} \int dp_i \int df_i \frac{\tilde{d}_i^{D_i} e^{-\tilde{d}_i}}{D_i!} \]  \hspace{1cm} (11)

Then, by calculating the total uncertainty (\(\sigma_{\text{tot}}\)) from \(l(D|p, f)\) and \(\sigma_{\text{stat}}\) from \(l_{\text{stat}}(D|p, f)\), the systematic uncertainty (\(\sigma_{\text{sys}}\)) can be deduced from

\[ \sigma_{\text{sys}} = \sqrt{\sigma_{\text{tot}}^2 - \sigma_{\text{stat}}^2}. \]  \hspace{1cm} (12)

Notice we have assumed that the \(l_{\text{stat}}(D|p, f)\) and \(l(D|p, f)\) both have maxima at the same \(p_{\text{max}}\) and \(f_{\text{max}}\). This is not the case in general. However, in the instance where the maxima do not occur at the same \(p_{\text{max}}\) and \(f_{\text{max}}\), the difference arises from choosing what is in fact an arbitrary definition of the statistical uncertainty, \(l_{\text{stat}}(D|p, f)\).

Figure 5 shows the distributions of \(p_{\text{max}} - p_{\text{true}}\) and \(f_{\text{max}} - f_{\text{true}}\) histogrammed in blue. The means of these distributions do not deviate more than 1.3% for protons and iron - an insignificant deviation considering the mean uncertainties in Fig. 6. Figure 6 shows the distribution of \(|p_{\text{max}} - p_{\text{true}}|/\sigma_p\) and \(|f_{\text{max}} - f_{\text{true}}|/\sigma_f\) where \(\sigma_p\) and \(\sigma_f\) are the uncertainties for \(p\) and \(f\), respectively. As expected, 68% of the predictions for \(p\) and \(f\) deviate less than 1\(\sigma\) from the \(p_{\text{true}}\) and \(f_{\text{true}}\). Figures 7 and 8 show the relative error for protons and iron, respectively. The most probable error for \(p\) and \(f\) is \(\sim 13\%\).

We have tested the method with other mixtures of iron and protons and find that it gives reasonable results for all two-component mixtures. For instance, in Fig. 9 we find that the method still gives reasonable uncertainties in the case where there is an 80:20 proton:iron mixture in a 200 event parent distribution. The mode of the fractional error is \(\sim 8\%\) (\(\sim 28\%\)) for protons (iron).

### 4.1 Eliminating Spectral Dependence in the Likelihood

In practice, the energy spectra for protons and iron are unknown. To include our lack of prior knowledge of the energy spectrum, a slightly more sophisticated likelihood is used.

The method allows us to float \(\alpha_p\) and \(\alpha_f\). The likelihood can be maximized as a function of \(\alpha_p\) and \(\alpha_f\) along with \(p\) and \(f\), but, due to limited statistics in the hypothesized simulated samples, we choose to marginalize these parameters. In principle, one would integrate over an infinite number of spectra. However, the CPU time for such a calculation is unrealistic, so here these spectra will
Fig. 5. Histograms of $p_p^{\text{true}}$ (top) and $f_f^{\text{true}}$ (bottom) for $p_p^{\text{true}} = 100$ and $f_f^{\text{true}} = 100$. The green (blue) histograms were calculated with Eq. (15) (Eq. (5)). The black histogram makes use of Eq. (15) that is applied to a parent data sample made of 250 events with 100 proton, 100 iron and 50 gammas. The red histogram includes an additional term in Eq. (15) to account for gammas. The histograms have overflow events (e.g., most of the events in the black histogram lie outside the abscissa limits).

be marginalized using $E^{-2}$ and $E^{-3}$ spectra. Eq. (2) then becomes

$$P(p, f|D) = \frac{\sum_{\alpha_p=-2, -3} \sum_{\alpha_f=-2, -3} P(D|p, f, \alpha_p, \alpha_f) q(p, f, \alpha_p, \alpha_f)}{\sum_{p, f, \alpha_p, \alpha_f} P(D|p, f, \alpha_p, \alpha_f) q(p, f, \alpha_p, \alpha_f)}$$

(13)

Other priors such as those concerning the interaction model can be introduced and marginalized in a similar manner. Therefore, maximizing the probability $P(p, f|D)$ is to maximize the likelihood

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Fig. 6. Deviation from $p_{\text{true}}$ and $f_{\text{true}}$ expressed relative to the proton and iron uncertainties, $\sigma_p$ and $\sigma_f$, respectively. The green (blue) histograms were calculated with Eq. (15) (Eq. (5)). The black histogram makes use of Eq. (15) that is applied to a parent data sample made of 250 events with 100 proton, 100 iron and 50 gammas. The red histogram includes an additional term in Eq. (15) to account for gammas. The histograms have overflow events (e.g., many of the events in the black histogram lie outside the abscissa limits).

\[
l(D|p, f, \alpha_p, \alpha_f) \equiv P(D|p, f, \alpha_p, \alpha_f)q(p, f, \alpha_p, \alpha_f) \\
= l(D|\{p_i\}, \{f_i\}, \epsilon_p, \epsilon_f),
\]

or

\[
l(D|\epsilon_p, \epsilon_f) = \sum_{\alpha_p=-2,-3} \sum_{\alpha_f=-2,-3} \prod_{i=1}^{N \text{ bins}} \int dp_i \int df_i \left( \frac{dD_i e^{-\tilde{d}_i}}{D_i!} \right) \left( \frac{\tilde{p}_i e^{-\tilde{p}_i}}{P_i!} \right) \left( \frac{\tilde{f}_i e^{-\tilde{f}_i}}{F_i!} \right).
\]

(14)

(15)
Fig. 7. Fractional uncertainty for the mean number of protons ($p$). The green (blue) histograms were calculated with Eq. (15) (Eq. (5)). The black histogram makes use of Eq. (15) that is applied to a parent data sample made of 250 events with 100 proton, 100 iron and 50 gammas. The red histogram includes an additional term in Eq. (15) to account for gammas. The histograms have overflow events.

We can see that the introduction of priors does not change the distributions in Figs. 5 and 6 substantially as seen by comparing the blue and green histograms. Its effect can be best seen by the upward shift in the mean relative uncertainty (between the blue and green histograms) in Fig. 7 and 8.

We conclude that the method gives an accurate prediction of the mean number of protons and iron. For instance, with 100 proton and 100 iron events in a data sample with primary energies over $10^{19}$ eV, the uncertainty in $p$ and $f$
Fig. 8. Fractional uncertainty for the mean number of iron ($f$). The green (blue) histograms were calculated with Eq. (15) (Eq. (5)). The black histogram makes use of Eq. (15) that is applied to a parent data sample made of 250 events with 100 proton, 100 iron and 50 gammas. The red histogram includes an additional term in Eq. (15) to account for gammas. The histograms have overflow events.

is $\sim 12\%$ with a better chance of obtaining a larger uncertainty. This increase is expected as we have introduced some “ignorance” in the choice of spectra. Further, the likelihood is able to accurately predict the 1σ uncertainties.

In the instance where we wish to sum over spectra, we would ideally integrate over every possible spectrum for protons and iron. Computationally, this is obviously impossible. Instead, we compromise by summing over “extremes” of what we would expect for protons and iron. In this first approximation, we calculate $L(D|\epsilon_p, \epsilon_f, \alpha_p = 2, \alpha_f = 2)$, $L(D|\epsilon_p, \epsilon_f, \alpha_p = 2, \alpha_f = 3)$, $L(D|\epsilon_p, \epsilon_f, \alpha_p = 3, \alpha_f = 2)$ and $L(D|\epsilon_p, \epsilon_f, \alpha_p = 3, \alpha_f = 3)$. That is, we cal-
calculate the likelihood with every combination of spectra for the proton and iron distributions. Then, we sum these 4 likelihood and maximize the sum with respect to $\epsilon_p$ and $\epsilon_f$. Had we summed over generators (say QGSJET and SIBYLL), we would sum over 8 likelihoods where each likelihood is calculated with a different combination of spectra and hadronic generators. The calculation of uncertainties are made in the usual way with the summed likelihood.

5 Discussion

We now investigate the case where the sample made of proton and iron is “contaminated” with other components. We start the discussion with gamma primaries. AGASA has estimated the upper limit in the $\gamma$-ray flux to be 28% for events above $10^{19}$ eV [16]. We therefore create a 250 events sample with 100 protons, 100 iron, 50 gammas. The hypothesized distributions for gammas are based on 1352 and 490 gammas with $E^{-2}$ and $E^{-3}$, respectively. We use the gammas with an $E^{-3}$ spectrum to generate the parent distribution. The result of the contamination is a large bias in $p$ and $f$ found in the black histogram in Fig. 5 and wrong uncertainties as seen in Figs. 6, 7 and 8. However, if the likelihood is modified to

$$l(D|\epsilon_p, \epsilon_f) = \sum_{\alpha p = -2, -3} \sum_{\alpha f = -2, -3} \prod_{i=1}^{N \text{ bins}} \int d\tilde{p}_i \int d\tilde{f}_i \left( \frac{\tilde{d}^p_i e^{-\tilde{d}_i}}{D_i!} \right) \left( \frac{\tilde{p}_i}{P_i!} \right) \left( \frac{\tilde{f}_i}{F_i!} \right) \left( \frac{\tilde{g}_i}{G_i!} \right)$$

Fig. 9. Fractional uncertainty for $p$ (black) and $f$ (red) calculated for simulated data sets with a parent sample comprised of 160 proton and 40 iron events.
where $\epsilon_g$, $G_i$, etc. carry the same definitions for gammas as $\epsilon_p$, $P_i$, etc. carry for protons, we see that the uncertainties once again make sense and the bias becomes negligible as shown by the red histograms in Figs. 5 and 6. In Figs. 7 and 8 we see an increase in the relative uncertainties for iron while the proton uncertainties do not change significantly. An increase in the uncertainties would not be surprising as there is an overlap in the $X_{\text{max}}$ distributions between protons, iron and gammas. The lack of significant change is a result of the right combination of protons with an $E^{-2}$ spectrum and iron with an $E^{-2}$ spectrum having a much larger likelihood value than the alternatives in the summation of $\alpha_p$ and $\alpha_f$. This demonstration shows that one must account for gamma-like primaries if there is a reasonable expectation that they may be in the data.

Accounting for all possible contributions from every imaginable primary is clearly impossible. A number of questions then arise. For instance, is it sufficient to assume a proton and iron mixture when analyzing a real data distribution? Is it necessary to assume, for instance, a proton, helium, carbon and iron mixture? Would such complex mixtures be more or less useful than simply quoting the result assuming a proton and iron mixture?

These questions can be approached in two ways. If we had good reason to believe ultrahigh energy cosmic rays were composed exclusively of (say) protons, carbon and iron, then we would consider it useful to measure the relative contributions from these three elements regardless of their uncertainties.

On the other hand, if we consider protons, carbon and iron as stand-ins for “light,” “medium” sized and “heavy” primaries, then we can ask how sensitive the likelihood method is to these elements. That is, if there were something resembling carbon mixed with protons and iron, could it be detected to any accuracy?

In the latter vein, we build a “fake” data sample with 100 proton, 50 carbon and 100 iron primaries. We then apply the full likelihood in Eq. (16), replacing the gamma term with carbon, and evaluate the distribution of uncertainties (Fig. 10). Although we find the uncertainties are meaningful, the large uncertainties for carbon indicate that the resolution is not sufficient to make statements about the fraction of medium-sized primaries in this type of data sample by inserting the carbon hypothesis. From this perspective, it is less important to account for carbon primaries than gamma primaries.

We also study the case where two hypothesized elements are indistinguishable. We create a fake data set of 250 events with 150 proton and 100 iron events. A proton distribution, a second identical proton distribution and an iron distribution act as hypotheses. That is, the likelihood is calculated using Eq. (16) substituting a second proton distribution for the gamma distribution.
Fig. 10. Distribution of the relative errors for proton, carbon and iron hypotheses evaluated using Eq. (16) over simulated data sets fluctuated from a parent sample of 100 proton, 50 carbon and 100 iron events.

The algorithm that maximizes the likelihood, unable to distinguish how many of the 150 proton are attributable to the first proton distribution as opposed to the second, settles on an arbitrary number of proton events between 0 and \(\sim 150\) for the first proton hypothesis \((p_1)\). What remains of \(\sim 150\) events is attributed to the second proton hypothesis \((p_2)\). The uncertainties are then calculated for \(p_1\) and \(p_2\) by fixing the number of events for \(p_1\) \((p_2)\) fluctuating \(p_2 \ (p_1) \pm 1\sigma\) according to Eq. (9). As there are large correlations between \(p_1\) and \(p_2\) that are not considered when calculating the uncertainties in this way, the method underestimates the uncertainties (1 \(\sigma\) corresponds to 15% confidence region). Therefore, before an element is inserted as a hypothesis, the correlations between the various distributions need to be understood or one needs to verify that the distributions are sufficiently uncorrelated such that
the results have meaningful uncertainties.

Also, one may consider other hadronic models. In principle, the hadronic models can be marginalized by simply “summing” over them. $X_{\text{max}}$ distributions for protons, iron and gammas with $E^{-2}$ and $E^{-3}$ spectra would be generated using QGSJET and SIBYLL hadronic models. The complexity of the likelihood and the amount of Monte Carlo increases sharply with the number of priors. For the sake of time and simplicity the full calculation is not made here. However, to show that it is important to consider different hadronic models, we generate a simulated data set with 100 proton and 100 iron events with the SIBYLL hadronic model. We then fluctuate this distribution many times in the same fashion described above each time calculating the maximum likelihood using proton and iron events generated with QGSJET. Figure 11 shows the $\sim 60\%$ shift in the mean number of protons and iron resulting from calculating the likelihood using QGSJET. Regardless, this shift shows that the composition fractions cannot be properly calculated without considering at least the QGSJET and SIBYLL hadronic models.

We have discussed widening the measurement to include such knowledge and given an example of how one might introduce prior knowledge by introducing and then marginalizing the unknown spectral dependence of iron and protons. Other parameters concerning the extensive air shower and atmospheric uncertainties can be introduced and marginalized provided sufficient data and CPU time. In so doing, uncertainties are rigorously compounded in the calculation.

Further, we have investigated the consequence of contaminating the simulated data with gammas and carbon. Proton, iron and gamma hypotheses are sufficient if one is only interested in stand-ins for “heavy,” “light,” and “lighter” elements. However, it is important to consider all elements that one reasonably expects to be in the data.

This method of measuring the composition is different from the method using elongation rate. It uses the full distribution of $X_{\text{max}}$ to determine the composition as opposed to the elongation rate which uses the mean. Also, the measured elongation is independent of any model assumptions although its interpretation requires one to compare the measurement to what is expected for various compositions. The method is currently being applied to HiRes stereo events and results will be presented in a separate paper where the composition will be evaluated for events in intervals of $10^{18.5}$ eV, $10^{19.0}$ eV, $10^{19.5}$ eV, etc.

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Fig. 11. Histograms of $p - p_{\text{true}}$ (top) and $f - f_{\text{true}}$ (bottom) for $p_{\text{true}} = 100$ and $f_{\text{true}} = 100$ where the simulated data is generated with SIBYLL and the likelihood calculation uses proton and iron events generated with QGSJET.

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