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NEW SPECTRAL CLASSIFICATION TECHNIQUE FOR X-RAY SOURCES: QUANTILE ANALYSIS

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ABSTRACT

We present a new technique, quantile analysis, to classify the spectral properties of X-ray sources with limited statistics. Quantile analysis is superior to conventional approaches, such as X-ray hardness ratio or X-ray color analysis, for studying relatively faint sources or investigating a certain phase or state of a source in detail, where poor statistics do not allow spectral fitting using a model. Instead of working with predetermined energy bands, we determine the energy values that divide the detected photons into predetermined fractions of the total counts, such as the median (50%), terciles (33% and 67%), and quartiles (25% and 75%). We use these quantiles as an indicator of the X-ray hardness or color of the source. We show that the median is an improved substitute for the conventional X-ray hardness ratio. The median and other quantiles form a phase space, similar to the conventional X-ray color-color diagrams. The quantile-based phase space is more evenly sensitive over various spectral shapes than the conventional color-color diagrams, and it is naturally arranged to properly represent the statistical similarity of various spectral shapes. We demonstrate the new technique in the 0.3–8 keV energy range using the Chandra ACIS-S detector response function and a typical aperture photometry involving background subtraction. The technique can be applied in any energy band, provided the energy distribution of photons can be obtained.

Subject headings: methods: statistical — X-rays: general

Online material: color figures

1. INTRODUCTION

A major obstacle to understanding the nature of the faintest X-ray sources is poor statistics, which prevents the usual practices of spectral analysis, such as fitting with known models. Even for apparently bright sources, as we investigate the sources in detail we frequently need to divide source photons into various phases or states of the sources, and the success of such an analysis is often limited by statistics. The common practice of extracting spectral properties of X-ray sources with poor statistics is to calculate the X-ray hardness or color of the sources (e.g., Schulz et al. 1989; Kim et al. 1992; Netzer et al. 1994; Prestwich et al. 2003). In this conventional method, the full energy range is divided into two or three subbands and the detected source photons are counted separately in each band. The ratio of these counts is defined as X-ray hardness or color, to serve as an indicator of the spectral properties of the source.

In principle, one can constrain a meaningful X-ray hardness or color to a source if at least one source photon is detected in each of at least two bands. However, the equivalent requirement for total counts in the full energy band can be rather demanding, and it is strongly spectral-dependent. Figure 1 shows an example of an X-ray color-color diagram (Kim et al. 2004). In the figure, we divide the full energy range (0.3–8.0 keV) into three sub-energy bands: 0.3–0.9 (S), 0.9–2.5 (M), and 2.5–8.0 keV (H). The ratio of the net source counts in the S–M band is defined as the soft X-ray color (x-axis in the figure), and the ratio of the net counts in the M–H band is defined as the hard X-ray color (y-axis). The energy range in this example is the region where the Chandra ACIS-S detectors are sensitive, but the technique described in this paper is not bounded to any particular energy range, provided the energy distribution of photons can be obtained.

The grid pattern in the figure represents the true location for the sources with the power-law spectra governed by two parameters: power-law index ($\Gamma$) and absorbing column depths ($N_{HI}$) along the line of the sight. The grid pattern is drawn for an ideal detector response, which is constant over the full energy band. The grid pattern appears to be properly spaced according to the changes of the spectral parameters, and such an arrangement suggests that this color-color diagram may be an ideal way to classify sources with poor statistics.

However, the appearance can be deceiving, which is hinted at by the uneven sizes of error bars in the figure. The error bar near each grid node represents the central 68% of simulation results from the spectral shape at the grid node. Each simulation has 1000 source counts in the full energy range with no background counts, and thus the size of the symbol represents the size of typical error bars for 1000 count sources in the diagram.

In the figure, there are two kinds of error bars for some spectral shapes (e.g., $\Gamma = 0$ and $N_{HI} = 10^{22}$ cm$^{-2}$); the thick error bars represent the central 68% distribution of the simulation results that have “proper” color values (defined here as at least one photon in each of all three bands), and the thin error bars show the 68% distribution of all the simulation trials (10,000) for each spectral model, i.e., a distribution of the central 6827 trials. The two error bars should be identical if all the trials produce a proper soft and hard color.

The right panel in Figure 1 shows the minimum counts in the full energy band to have at least 1 count in each of three bands. The figure indicates that the required minimum counts are dramatically different among the models in the grid. For the power-law spectra with $\Gamma = 0$ and $N_{HI} \sim 10^{22}$ cm$^{-2}$, more than 500 counts are required in order to have reasonable color values, while in the case of $\Gamma = 2$ and $N_{HI} \sim 10^{20}$ cm$^{-2}$, a few counts are sufficient. Because of statistical fluctuations, for some models even 1000 counts in the full energy range do not guarantee positive net counts in all three bands, which explains why some trials fail to produce proper colors.

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The spectral model dependence of error bars in this kind of color-color diagram is inevitable, and the dependence is determined by the choice of the sub-energy bands for a full energy range. In this example, the subbands are chosen so that the diagram is mostly sensitive near $\Gamma = 2$ and $N_{\text{H}} < 10^{22}$ cm$^{-2}$. In fact, it is not possible to select subbands to have uniform sensitivity over different spectral shapes.

Figure 1 clearly indicates that the conventional color-color diagram (CCCD) is heavily biased, which is very unfortunate when trying to extract unknown spectral properties from the sources. The heavy requirement on the total counts for certain spectral shapes defeats the purpose of the diagram, since the large total counts may allow nominal spectral analysis such as spectral fitting with known models. One might have to repeat the analysis with different choices of subbands to explore all the interesting possibilities of spectral shapes.

2. QUANTILES

In order to overcome the selection effects originating from the predetermined sub-energy bands, we propose using the energy value to divide photons into predetermined fractions. We choose fractions to take full advantage of the given statistics, such as 50% (the median), 33% and 67% (terciles), and 25% and 75% (quartiles), although any quantile may be used. We use the corresponding energy values—quantiles—as an indicator of the X-ray hardness or color of the source. In particular, we show below that the median, $Q_{50}$, is an improved substitute for the conventional X-ray hardness.

Let $E_{\text{n}}$ be the energy below which the net counts is $x$% of the total counts and we define the quantile $Q_x$ as

$$Q_x = \frac{E_{x\%} - E_0}{E_{\text{up}} - E_0},$$

where $E_0$ and $E_{\text{up}}$ are the lower and upper boundaries of the full energy band, respectively (0.3 and 8.0 keV in this example). The algorithm for the estimation of quantiles and their errors relevant to X-ray astronomy is given in Appendices A, B, and C. Unlike the conventional X-ray hardness or colors, for calculating quantiles there is no spectral dependence of required minimum counts. The required minimum only depends on types of quantiles: 2 counts for the median and terciles, and 3 counts for quartiles.

Figure 2 shows an example of quantile-based color-color diagrams (QCCDs) using the median $m(=Q_{50})$ for the $x$-axis and the ratio of two quartiles $Q_{25}/Q_{75}$ for the $y$-axis (see § 5 for the motivation of the choice of the axes). The grid pattern in Figure 2 is drawn for the same spectral parameters as in Figure 1 with five additional cases for $N_{\text{H}} = 5 \times 10^{22}$ cm$^{-2}$. Note that the approximate axes of $\Gamma$ and $N_{\text{H}}$ are rotated $\sim 90^\circ$ from those in Figure 1. The error bars are drawn for the central 68% of the same 10,000 simulation results of the spectral shape at each grid node, and each simulation run contains 1000 source counts with no background counts in the full energy band. The relatively similar size of the error bars for 1000 count sources in the figure indicates that there is no spectral-dependent selection effect. Note that there is no need for the distinction of thin and thick error bars in this figure because all the trials produce proper quantiles.

The grid pattern of power-law spectra in Figure 2 appears to be less intuitively arranged than that in Figure 1. However, we believe that the proximity of any two spectra in the quantile-based diagram accurately exhibits the similarity of the two spectral shapes (as folded through the detector response, which is constant in this example). For example, in the case of $\Gamma = 0$ in Figure 2, the separation in phase space by various $N_{\text{H}}$ values is much smaller than in the case of $\Gamma \geq 1$. Note that the column depth ($N_{\text{H}}$) in the considered range can change mainly soft X-rays ($\leq 2$ keV), and the spectrum for $\Gamma = 0$ is less dominated by soft X-rays than that for $\Gamma \geq 1$. In other words, the

$^2$ The quantile analysis software (IDL and Perl versions) is available at http://hea-www.harvard.edu/ChaMPlane/quantile.
overall effects of $N_H$ on the spectral shape would be much smaller in the case of $\Gamma = 0$ than in the case of $\Gamma \geq 1$. Therefore, the grid spacing in Figure 2 indeed reveals the appropriate statistical power needed to discern these spectral shapes, despite the degeneracy arising from utilizing only three variables (quantiles) extracted from a spectrum.

3. COLOR-COLOR DIAGRAM EXAMPLE

Now let us consider more realistic examples. We introduce the Chandra ACIS-S response function; we use the preflight calibration data for energy resolution and the energy-dependent effective area from PIMMS, version 2.3.5.

Figure 3 shows the simulation results of the spectra with 100 and 50 source counts with no background counts. The error bars again indicate the interval that encloses the central 68% of the simulation results for each model. The results are shown for both the conventional and new color-color diagrams in the figure. The grid patterns in the figures are for the same power-law models, and they look different from the previous examples because of the Chandra ACIS-S response function.

The analysis of faint sources is often limited by background fluctuations. In contrast to Figure 3, we allow for large background counts in the source region (e.g., a point source in a bright diffuse emission region). The top two panels in Figure 4 show the simulation results of the spectra with 100 source counts and 50 background counts in the source region, and the bottom two panels show the case of 50 source counts and 25 background counts. For background subtraction, we set the background region to be 5 times larger than the source region. The background region contains 250 (top panels) and 125 background (bottom panels) counts. Note that these are relatively high backgrounds for a typical Chandra source and thus illustrate the worst case of a point source superposed on background diffuse emission. The background photons are sampled from a power-law spectrum with $\Gamma = 0$ and $N_H = 0$, which is folded through the Chandra ACIS-S response function.

In the case of the CCCD in Figures 3 and 4, one notes that some of the error bars lie away from their true location (grid node). The severe requirement for the total counts (>100) for some of the spectral models results in only a lower or upper limit for the color in many simulation runs. Cases with proper colors for these models are greatly influenced by statistical fluctuations because of low source counts in one or two sub-bands, and thus the estimated colors fail to produce the true value. It is evident that this conventional diagram is more sensitive toward $\Gamma \sim 2$ and $N_H \leq 10^{22} \text{cm}^{-2}$.

In the case of the new quantile-based diagram, the error bars stay at the correct location and the size of the error bars is relatively uniform across the model phase space, indicating that the phase space is properly arranged. The quantile-based diagram shows more consistent results regardless of background.

4. HARDNESS RATIO EXAMPLE

When we explore the spectral properties of sources with an even smaller number of detected photons, the only available tool is the use of a single hardness ratio, which requires only two sub-energy bands. Here we use net counts $S$ in a 0.3–2.0 keV soft band and net counts $H$ in a 2.0–8.0 keV hard band. Two popular definitions for hardness ratio (HR) exist in the literature: $HR = H/S$ and $HR = (H - S)/(H + S)$. We compare the median with the conventional hardness ratio using the first definition.

The left panel in Figure 5 shows the performance of the conventional hardness ratio as a function of the total counts folded through the ACIS-S response. The plot contains three spectral shapes with $\Gamma = 2$. The shaded regions represent the central 68% of the simulation results for a given total net counts (except for the case of $H$ or $S \leq 0$). The simulation is done for the case without background (top panels) and for the case in which the source region contains an equal number of source and background counts (bottom panels), where we perform a background subtraction identical to the one described in §3. The background follows the same spectrum as in the previous examples.

Similar to the CCCD, the required minimum of the total counts for the conventional hardness ratio depends on spectral shape. The right panels in Figure 5 show the fraction of the simulations that have at least 1 count in both the soft and hard bands (otherwise HR = 0 or $\infty$). The plot indicates that one cannot assign a proper hardness ratio value in a substantial fraction of cases when the total net counts are less than 100. In the case of $N_H = 10^{20} \text{cm}^{-2}$, many simulation runs result in $H = 0$ (too soft), and for $N_H = 5 \times 10^{22} \text{cm}^{-2}$, $S = 0$ (too hard). A set of predetermined bands keeps the hardness ratio meaningful only within a certain range of spectral shapes. In order to compensate for such a limitation, one needs to repeat the analysis with different choices of bands, which in turn will have a different limitation.

Even for the cases with a proper hardness ratio value (Fig. 5, left panel), the hardness ratio distribution of two spectral types ($N_H = 10^{22}$ and $5 \times 10^{21} \text{cm}^{-2}$) overlaps substantially when the total net counts are less than 40 in the case of high background.

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3 See the Chandra Proposers' Observatory Guide at http://cxc.harvard.edu.
4 Approximately 100–250 eV FWHM. See http://cxc.harvard.edu/cal/Acis.
5 Portable, Interactive Multi-Mission Simulator. See http://heasarc.gsfc.nasa.gov/docs/software/tools/pimms.html.
Below 10 counts, it is difficult to relate the distributions to their true value.

Figure 6 shows the performance of the new hardness defined by the median. First, there is no loss of events. Second, three spectra are well separated down to less than 10 counts regardless of background. Even at 2 or 3 net counts, each distribution is distinct and well related to its true value.

5. DISCUSSION

The newly defined hardness and color-color diagrams using the median and other quantiles are far superior to the conventional ones. One can choose specific quantiles and their combinations relevant to one’s needs. We believe the new phase space by the median and quartiles is very useful in general, as summarized in Figure 7.

For a given spectrum, various quantiles are not independent variables, unlike the counts in different energy bands. However, the ratio of two quartiles is mostly independent of the median, which makes them good candidate parameters for color-color diagrams. In essence, the median $Q_{50}$ shows how dense the first (or second) half of the spectrum is, and the quartile ratio $Q_{25}/Q_{75}$ shows a similar measure of the middle half of the spectrum.

For the x-axis in Figure 7, one can simply use $m (\equiv Q_{50})$ on a log scale to explore the wide range of the hardness. However, a simple log scale compresses the phase space for relatively hard
spectra \((m > 0.5; \text{note } 0 \leq m \leq 1)\). Our expression, \(6 \log \left(\frac{m}{(1-m)}\right)\) when \(m \rightarrow 0\), and approximately \(-\log (1-m)\) when \(m \rightarrow 1\).

In this case, the median (x-axis) in the QCCD can be a good measure of line shift, and the quartile ratio (y-axis) can be a good measure of line broadening.

The right panel in Figure 7 shows the overlay of the grid patterns for typical spectral models—power law, thermal Bremsstrahlung, and blackbody emission. For high-count sources, one can use such a diagram to find relevant models for the spectra before detailed analysis.

Finally, we investigate how the detector energy resolution affects the performance of the quantile-based diagram. The left panel in Figure 8 shows the grid patterns of the power-law emission models for detectors with various energy resolutions but with the same detection efficiency of the Chandra ACIS-S detector. Starting with the energy resolution of the Chandra ACIS-S detector in the previous examples (Figs. 3, 4, 5, and 6; Fig. 4.—Realistic example of the color-color diagram with relatively high background: 100 count source with 50 background counts (top) and 50 count source with 25 background counts (bottom). Left: CCCD. Right: QCCD. The background follows a flat spectrum, and both source and background photons are folded through the Chandra ACIS-S response function. The diagrams are generated after the background subtraction (see Appendix C). Similar to the previous results, the new technique performs well throughout the phase space. In particular, note the relatively uniform error bars across the entire grid as well as the relative lack of points outside the grid. [See the electronic edition of the Journal for a color version of this figure.]
Fig. 5.—Conventional hardness ratio calculation. The soft band ($S$) is $0.3–2.0$ keV, and the hard band ($H$) is $2.0–8.0$ keV; the Chandra ACIS-S response function is used. Left: Conventional hardness distribution as a function of total net counts. Right: Fraction of the simulations that have $H > 0$ and $S > 0$ for a given total net count. Top: The case of no background. Bottom: For a given net count, the source region contains an equal number of source and background counts, and we perform a background subtraction identical to the one described in the previous examples. Three spectral shapes ($\Gamma = 2$) are simulated, and the solid horizontal lines in the left panels represent the true hardness ratio of each spectrum. The shaded region represents the central 68% of simulated results (only for $H > 0$ and $S > 0$) at a given total net count. A large fraction of trials do not produce a proper hardness ratio when the total counts are less than 100. Below 10 counts, the hardness distribution hardly bears any relation to its true value. Blending of the central 68% distributions starts at $\sim 40$ counts for $N_H = 10^{20}$ and $5 \times 10^{21}$ cm$^{-2}$ for the case with background. [See the electronic edition of the Journal for a color version of this figure.]
Fig. 6.—Median distribution for three power-law spectra with $\Gamma = 2$. Compare this with Fig. 5. Regardless of background, the three distributions are well separated down to less than 10 counts, bearing a tight relation with their true value. In addition, there is no loss of events in this diagram. [See the electronic edition of the Journal for a color version of this figure.]

Fig. 7.—Overview of various spectral shapes (approximated as concave or straight lines) in the QCCD using $Q_{25}$ and $Q_{75}$ (left) and the grid patterns for power law (PL), thermal Bremsstrahlung (TB), and blackbody (BB) models in a 0.3–8.0 keV range ideal detector or incoming flux space (right). For PL, the grid patterns are for $\Gamma = 4, 3, 2, 1,$ and 0 along the positive x-axis direction; for TB, $kT = 0.2, 0.5, 1, 2, 4,$ and 10.0 keV; and for BB, $kT = 0.1, 0.2, 0.5, 1, 2,$ and 4 keV. For all the models, $N_H = 0.1, 1, 5,$ and $10 \times 10^{21}$ cm$^{-2}$ along the positive y-direction. [See the electronic edition of the Journal for a color version of this figure.]
FWHM $\Delta E \sim 150$ eV at 1.5 keV and $\sim 200$ eV at 4.5 keV), we have successively decreased the energy resolution by multiplying the energy resolution at each energy by a constant factor. Each pattern is labeled by the energy resolution ($\Delta E / E$) at 1.5 keV. As the energy resolution decreases, the grid pattern shrinks. The pattern will shrink down to a point $(x, y) = (0, 1)$ in the diagram for a detector with no energy resolution.

The right panel in Figure 8 shows 68% of the simulation results for a detector with $\Delta E / E = 100\%$ at 1.5 keV. The size of the 68% distribution in this example is more or less similar to that in the case of the regular Chandra ACIS-S detector with $\Delta E / E = 10\%$ (Fig. 3, top right panel). The similarity is due to the fact that, unlike the grid pattern, the dispersion of the simulation results (or the error size of a data model point) for each model is mainly due to statistical fluctuations for low-count sources. In summary, as the energy resolution decreases, the relative distance between various models in the diagram decreases, but the error size of the data remains roughly the same, and thus the spectral sensitivity of the diagram decreases.

Note that the overall size of the grid patterns in the left panel of Figure 8 is more or less similar when $\Delta E / E \leq 20\%$ (at 1.5 keV). We expect that for a detector with energy resolution $\Delta E$, the overall size is dependent on a quantity $\Delta E / (E_{\text{up}} - E_{\text{lo}})$, since quantiles are determined over the full energy range. Therefore, the quantile-based diagram is quite robust against finite energy resolutions of typical X-ray detectors.

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APPENDIX A

QUANTILE ESTIMATION USING ORDER STATISTICS

Many routines have been developed to estimate quantiles and the related statistics (Wilcox 1997). We use a simple interpolation technique based on order statistics to estimate quantiles. For real data, we also need to have a reliable estimate for the errors of quantile values, for which we employ the technique by Maritz & Jarrett (1978). For simplicity, we assume that we measure the energy value of each photon. In the literature, many quantile estimation algorithms often assume that the lowest value (energy) of the given distribution is the (equivalent) lower bound of the distribution and the highest value is the upper bound. In X-ray astronomy, the lower ($E_{\text{lo}}$) and upper bounds ($E_{\text{up}}$) of the energy range are usually set by the instrument or user selection, where these bounds may or may not be the lowest and highest energies of the detected photons. We can explicitly impose this boundary condition by assigning 0% and 100% quantiles to $E_{\text{lo}}$ and $E_{\text{up}}$, respectively.

We sort the detected photons in ascending order of energy, and we assign $E_{x\%}$ to the energy $E_i$ of the $i$th photon as

$$E_i = E_{x\%}, \quad \text{where} \quad x = \frac{2i - 1}{2N},$$

and $N$ is the total number of net counts. Using $E_{0\%} = E_{E_{\text{lo}}}$ and $E_{100\%} = E_{E_{\text{up}}}$, one can interpolate any quantiles from the above relation of $x$ and $E_i$. With the definitions of $E_{0\%} = E_{E_{\text{lo}}}$ and $E_{100\%} = E_{E_{\text{up}}}$, the above interpolation is very robust. One can even calculate quantiles without any detected photons (although not meaningful), the result of which is identical to the case of a flat spectrum. In the
case of only one detected photon at E, the above relation reduces to \( E = E_{50\%} \). Therefore, the distribution of the median from 1 count sources with the same spectra is the source spectrum itself.

The above interpolation for quantile \( E_{\text{50\%}} \) essentially uses only two energy values, \( E_i \) and \( E_{i+1} \), where \((2i-1)/2N < x/100 < (2i+1)/2N\). In order to take advantage of other energy values, one can use more sophisticated techniques, such as that of Harrell & Davis (1982). In many cases, the Harrell-Davis method estimates quantiles with smaller uncertainties than the simple-order statistics technique, but because of smoothing effects in the Harrell-Davis method, there are cases in which the simple-order statistics performs better, such as distributions containing discontinuous breaks (Wilcox 1997). In real data, the finite detector resolution tends to smooth out any discontinuity in the spectra. Our simulation shows that about 10%–15% better performance is achieved by the Harrell-Davis method compared to the simple-order statistics for the cases of 50 source photons and 25 background photons. The results in the paper are generated by the order statistics technique.

APPENDIX B
QUANTILE ERROR ESTIMATION USING THE MARITZ-JARRETT METHOD

Once quantile values are acquired, one can always rely on simulations to estimate their uncertainty (or error) using a suspected spectral model with parameters derived from the QCCD. If no single model stands out as a primary candidate, one can derive a final uncertainty of the quantiles by combining the results of multiple simulations from a number of models. However, error estimation by simulations can be slow, cumbersome, and even redundant, considering that the quantile errors are more sensitive to the number of counts than the choice of spectral shapes (cf. Figs. 3 and 4). A quick and rough error estimate is often sufficient, and so we introduce a simple quantile error estimate technique—the Maritz-Jarrett (1978) method. The results (Figs. 1–6, error bars and shades) in the main text indicate the interval that encloses the central 68% of the simulation results and were not driven by the Maritz-Jarrett method.

The Maritz-Jarrett method uses a technique similar to the Harrell-Davis method of quantile estimation. Both methods rely on L estimators (linear sums of order statistics) using beta functions. We sort the photons in the ascending order of energy from \( i = 1 \) to \( N \). Then, we apply the Maritz-Jarrett method with small modifications7 as follows. For \( E_{\text{x50\%}} \), we set

\[
M = Nx + 0.5, \quad a = M - 1, \quad b = N - M.
\]

We then define \( W_i \) using the incomplete beta function \( B \),

\[
W_i = B(a, b, y_{i+1}) - B(a, b, y_i),
\]

\[
B(a, b, y) = \frac{\Gamma(a + b)}{\Gamma(a)\Gamma(b)} \int_0^y t^{a-1}(1 - t)^{b-1} dt,
\]

where \( y_i = i/N \) and \( \Gamma(n + 1) = n! \) (the gamma function). Using \( W_i \) and \( E_i \), we calculate the L estimators \( C_k \),

\[
C_k = \sum_{i=1}^{N} W_i E_i^k, \quad \text{and} \quad \sigma(E_{\text{x50\%}}) = \sqrt{C_2 - C_1^2}.
\]

The error estimate by the Maritz-Jarrett method requires at least 3 counts for medians, 5 counts for terciles, and 6 counts for quartiles \((a, b \geq 1)\).

Figure 9 shows the accuracy of the error estimates by the Maritz-Jarrett method for \( \Gamma = 2 \) and \( N_{\text{tot}} = 5 \times 10^{31} \text{ cm}^{-2} \). In the case of no background, the Maritz-Jarrett method is accurate when the total net counts are greater than \( \sim 30 \). It tends to overestimate the errors when the total net counts are below \( \sim 30 \). In the case of high background, it tends to underestimate the errors overall, since the adopted background subtraction procedure (see below) does not inherit background statistics for the error estimates. We find that multiplying by an empirical factor \((1 + N_{\text{bkg}}/N_{\text{src}})^{1/2} \) can compensate, approximately, for the underestimation \((N_{\text{src}}: \text{total source counts}; N_{\text{bkg}}: \text{total background counts})\).

For the error of the ratio \( Q_{25}/Q_{75} \), since these two quartiles are not independent variables, the simple quadratic combination from two quartile errors overestimates the error of the ratio (\( \sim 20\%–30\% \) for the examples in Figs. 3 and 4). One can find more sophisticated techniques such as Bayesian statistics to estimate quantile errors in the literature. For example, Babu (1999) and references therein discuss the limitation of bootstrap estimation and show other techniques such as the half-sample method.

APPENDIX C
BACKGROUND SUBTRACTION

For background subtraction, we calculate quantiles at a set of finely stepped fractions separately for photons in the source region and the background region. Then, by a simple linear interpolation, we establish the integrated counts IC(\( \geq E \)) (number of photons with energy greater than \( E \)) as a function of energy for both regions. Now at each energy, one can subtract the integrated counts of the background region from that of the source region with a proper ratio factor of the area of the two regions. Because of statistical fluctuations, the subtracted integrated net counts may not be monotonically increasing from \( E_{\text{lo}} \) to \( E_{\text{up}} \). Therefore, we force a

7 Originally \( M = \lfloor Nx + 0.5 \rfloor \), where \( \lfloor x \rfloor \) is the integer part of \( x \).
monotonic behavior by setting $\text{IC}(>E) \geq \text{IC}(>E')$ if $\text{IC}(>E) < \text{IC}(>E')$ and $E > E'$. Such a requirement can underestimate the quantiles overall, so we repeat the above using $\text{IC}(<E)$ and force a monotonic decrease. Then, quantiles for the net distribution are given by

$$x = \frac{N + \text{IC}(>E) - \text{IC}(<E)}{2N},$$

$$E_{x_{\text{net}}} = E,$$

where $N$ is the total net counts. If there is no statistical fluctuation, $\text{IC}(>E) = N - \text{IC}(<E)$. For the error estimation, we need to know the energy of each source photon, which will be lost from background subtraction. Thus we generate a set of $N$ energy values for source photons matching the above quantile relation and then apply the above error estimation technique using the Maritz-Jarrett method.

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