The extinction law from photometric data: linear regression methods

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ABSTRACT

Context. The properties of dust grains, in particular their size distribution, are expected to differ from the interstellar medium to the high-density regions within molecular clouds. Since the extinction at near-infrared wavelengths is caused by dust, the extinction law in cores should depart from that found in low-density environments if the dust grains have different properties.

Aims. We explore methods to measure the near-infrared extinction law produced by dense material in molecular cloud cores from photometric data.

Methods. Using controlled sets of synthetic and semi-synthetic data, we test several methods for linear regression applied to the specific problem of deriving the extinction law from photometric data. We cover the parameter space appropriate to this type of observations.

Results. We find that many of the common linear-regression methods produce biased results when applied to the extinction law from photometric colors. We propose and validate a new method, LinES, as the most reliable for this effect. We explore the use of this method to detect whether or not the extinction law of a given reddened population has a break at some value of extinction.

Key words. methods: data analysis – ISM: clouds – dust, extinction – stars: formation

1. Introduction

The properties of interesting dust appear to be fairly constant throughout the interstellar medium (ISM) of the Galaxy (Rieke & Lebofsky 1985; Kenyon et al. 1998; Lombardi et al. 2006; Jones & Hyland 1980; Martin & Whittet 1990), reflecting the homogeneous physical conditions that characterize it. In the cold molecular cores, however, under lower temperatures and higher densities, the dust grains are believed to change, namely grow by coalescence and/or develop ice mantles (e.g., Whittet et al. 1988; Ossenkopf 1993; Whittet et al. 2001; Draine 2003; Román-Zúñiga et al. 2007; Steinacker et al. 2010). Measuring these differences using methods other than detailed spectral analysis has proved somewhat challenging, but the advent of larger and more sensitive telescopes has started to reveal extinction laws toward these regions that depart from the ISM typical curves, particularly in the near- and mid-infrared regime$^1$, putting forward the extinction law as a good indicator of grain properties.

Whereas the extinction law in low-density regions and the ISM is well characterized by a power-law ($A_\lambda \propto \lambda^{-\beta}$) of index $\beta \sim 1.8$, several authors have found a pronounced flattening of the extinction law in high density regions. Lutz et al. (1996) and Lutz (1999) first noted a flat extinction law toward the Galactic center in this wavelength range using spectroscopy of hydrogen recombination lines. Nishiyama et al. (2006) confirmed a flat extinction law toward the Galactic center using the colors of red clump stars, later confirmed independently by Fritz et al. (2011). Another example was found by Indebetouw et al. (2005), using a different method based on Spitzer photometry, in the direction of and around the star forming region RCW 49. A number of other studies on the extinction law using Spitzer followed, namely Flaherty et al. (2007), Chapman et al. (2009) and McClure (2009), who found a gray extinction law for star forming regions and molecular clouds; Chapman & Mundy (2009) and Román-Zúñiga et al. (2007), who found a gray extinction law for cloud cores; and Nishiyama et al. (2009), who found a similar law for the Galactic center. Chapman & Mundy (2009), Chapman et al. (2009) and McClure (2009) go a step further by analyzing the dependence of the extinction law with extinction, finding that the extinction law becomes grayer at higher extinction regimes. More recently, Cambrésy et al. (2011) measured an actual change of the extinction law within the same region for a threshold of $A_V = 20$ mag in the Trifid Nebula.

In this paper we address the issue of determining the extinction law from photometric data alone and the biases inherent to some of the fitting methods used frequently in the literature. We begin by defining the mathematical problem and the possible methods to extract a linear fit (Sect. 2), validating each method with synthetic data (Sect. 3). In Sect. 4 we address the issue of detecting a flattening of the extinction law with extinction.

This is the first paper in a series of two. The following paper (Ascenso et al., in prep.) will apply the results reported here to actual observations of cores in the Pipe Nebula.
2. The extinction law from photometric data

2.1. Defining the problem

The problem of deriving the extinction law from photometric data is one of linear regression: the goal is to determine the slope of the line that best fits the reddening-displaced positions of the stars in a color–color diagram. This slope, $\beta$, is the ratio of two color excesses that compose the color–color diagram, e.g., $\beta = E_{\lambda-K}/E_{H-K}$ in a $\lambda - K$ vs. $H - K$ diagram, and is, in this case, related to the extinction law $A_\lambda/A_K$ by:

$$\frac{A_\lambda}{A_K} = \left( \frac{A_H}{A_K} - 1 \right) \frac{E_{\lambda-K}}{E_{H-K}} + 1. \quad (1)$$

Linear regression, however, is not a simple science. The presence of errors in both coordinates, that they vary as a function of the quantities being analyzed (heteroscedasticity), that they may be correlated, and the presence of intrinsic scatter, make the required linear regression analysis far more complex than the typical chi-squared minimization of residuals. Problems of this nature, in particular applied to astronomical analysis, have been debated in the literature for over 50 years (Seares 1944; Trumpler & Weaver 1953), although every specific case seems to require a careful consideration of the methods to use. The particular distribution to which we presently wish to fit a linear function has the following characteristics:

- The $X$ and $Y$ variables are photometric colors, obtained by subtracting the magnitudes of a star in two bands. Both variables are therefore subject to photometric errors that increase with magnitude but not in an entirely predictable way with color.
- Because we observe (parts of) non-homogeneous clouds, the amount of extinction is not the same for all stars, and since most of the area is at low extinction, the distribution of points in a color–color diagram is denser in the bluer end, and scarcer in the redder end.
- The $X$ and $Y$ colors will usually have one band in common (e.g., $J - H$ and $H - K$), which causes the errors to be correlated (anti-correlated in the example).
- Because we observe the colors of a random sample of stars background to the cloud, the data will have intrinsic scatter caused by the range in spectral types of the stars. The intrinsic scatter is unrelated to extinction.

To summarize, the position of each datapoint in the color–color diagram is determined by three factors: the intrinsic scatter from the dispersion in spectral types of field stars, the reddening caused by dust extinction, and the measurement (photometric) error. The first alone would trace the loci occupied by the unreddened colors of the population of stars in the field, mostly giant and main-sequence stars; the effect of extinction is to (dim the objects and) move each point along the line whose slope we want to determine; and the photometric errors scatter the points in an ellipse around the intrinsic, reddened position (not a circle because the errors are correlated). A proper method to measure the slope of this distribution should be robust enough to disentangle these effects and return the single slope of the reddening vector.

2.2. Methods for linear regression

In this paper we test the following methods for linear regression applied to the problem described in the previous section.

2.2.1. Least squares fitting

The ordinary least-squares (OLS) fit is the simplest approach to linear regression. It works by minimizing the vertical distance of all data points to successive lines in the 2-D (slope and intercept) parameter space. Formally, it is equivalent to finding $\alpha_{\text{OLS}}$ and $\beta_{\text{OLS}}$ that minimize the quantity:

$$\chi^2(\alpha_{\text{OLS}}, \beta_{\text{OLS}}) = \sum_{i=1}^{N} (y_i - \alpha_{\text{OLS}} - \beta_{\text{OLS}}x_i)^2$$

where $(x_i, y_i)$ is the $i$th data point, and $\alpha$ and $\beta$ are the intercept and the slope one is trying to find. This method treats all data points the same, even though the position of some points will be more uncertain than that of others. The knowledge of the measurement errors for each data point can be used to attribute different weights to the different data points, thus optimizing the fit, which is equivalent to finding $\alpha_{\text{WLS}}$ and $\beta_{\text{WLS}}$ that minimize the quantity:

$$\chi^2(\alpha_{\text{WLS}}, \beta_{\text{WLS}}) = \sum_{i=1}^{N} \frac{(y_i - \alpha_{\text{WLS}} - \beta_{\text{WLS}}x_i)^2}{\sigma_{yi}^2 + \beta_{\text{WLS}}^2 \sigma_{xi}^2}$$

where $\sigma_{yi}$ and $\sigma_{xi}$ are the measurement errors of the $i$th data point. This method is called the weighed least squares (WLS) method.

2.2.2. Symmetrical methods

Applications of the least squares method have been suggested to be most appropriate when the intrinsic scatter in the data dominates over the measurement errors (see Isobe et al. 1990, for a detailed discussion and formal description). As opposed to the method described above, these treat both variables symmetrically, in such a way that it is no longer meaningful to speak of dependent and independent variables.

Two of these methods are based on calculating the OLS slope of the $Y$ vs. $X$ distribution, $\beta_1 = \text{OLS}(Y|X)$ following the notation of Isobe et al. (1990), and that of the $X$ vs. $Y$ distribution, $\beta_2 = 1/\text{OLS}(X|Y)$. The best fit is the line that bisects the OLS($Y|X$) and the OLS($X|Y$) in the bisector method (Eq. (4)), or the geometric mean of the OLS($Y|X$) and OLS($X|Y$) slopes in the geometric mean method (Eq. (5)). A third method, orthogonal regression (Eq. (6)), minimizes the distances of the points to a model line, but perpendicularly to the line instead of vertically as the regular least-squares method. Isobe et al. (1990) clearly state that these three methods do not produce the same or equivalent solutions. The plus or minus in the equations below refer to the sign of the covariance of the two variables – positive if they are correlated, negative if anti-correlated.

$$\hat{\beta}_{\text{bisorcor}} = \frac{\hat{\beta}_1 \hat{\beta}_2 - 1 + \sqrt{1 + \hat{\beta}_1^2 (1 + \hat{\beta}_2^2)}}{\hat{\beta}_1 + \hat{\beta}_2} \quad (4)$$

$$\hat{\beta}_{\text{geom}} = \pm \sqrt{\hat{\beta}_1 \hat{\beta}_2} \quad (5)$$

$$\hat{\beta}_{\text{orth}} = \frac{\hat{\beta}_2 - \hat{\beta}_1^{-1} \pm \sqrt{4 + (\hat{\beta}_2 - \hat{\beta}_1^{-1})^2}}{2} \quad (6)$$

2.2.3. Binning

Binning data always causes loss of information, but, sometimes, what is lost in information, is gained in simplicity of analysis.
and results. In their studies of extinction by molecular clouds, Lombardi et al. (2006) developed a method to determine the extinction law that consisted in binning the colors in both axes and fitting the bins using the weighted least-squares fitting described in Sect. 2.2.1. The characterization of the data changes entirely, because (1) the data points are no longer measurements but the weighed average of many measurements, (2) the errors are no longer measurement errors but the dispersion of colors within each bin, and (3) the fewer high extinction points are given more weight than before as they are represented by a bin with the same weight as those bins containing more points. The latter point implies, although with the potential of introducing small number statistics issues, that the weight of any extinction range is the same, regardless of where the majority of datapoints lie. In this way, instead of the fit being weighed by the more abundant points at low extinction, as is the case for the fit to all datapoints, it is equally weighed by a much larger range of extinction thus allowing a better constrain of the extinction law. It also implies that part of the intrinsic scatter problem is eliminated as the high extinction population will be dominated by giants whose range in intrinsic colors is much narrower.

For the purpose of these experiments we have binned the data in two ways: (1) along the color in the X-axis in bins of color and (2) iteratively along an assumed reddening vector in bins of $A_V$ following the method described by Lombardi et al. (2006).

2.2.4. BCES method

The standard BCES method (Akritas & Bershady 1996) starts from the simple observation that the slope $\beta$ that minimizes the standard OLS Eq. (1) can be alternatively written as

$$\beta_{\text{OLS}} = \frac{\text{Cov}(x, y)}{\text{Var}(x)},$$

(7)

where $\text{Cov}(x, y)$ is the observed covariance between the data $[x_i]$ and $[y_i]$, and $\text{Var}(x)$ is the variance of $[x_i]$. These two quantities can be evaluated from the usual equations

$$\text{Var}(x) = \frac{1}{N} \sum_{i=0}^{n} (x_i - \bar{x})^2,$$

(8)

$$\text{Cov}(x) = \frac{1}{N} \sum_{i=0}^{n} (x_i - \bar{x})(y_i - \bar{y}),$$

(9)

where the bar indicates the average values. Note that for the specific purposes of the BCES method the variance and covariance of the data should be evaluated using $N$ in the denominator (instead of the more common $N - 1$, used when estimating the variance and the mean from the same dataset).

The equation for $\beta$ above suggests that we can easily take into account the presence of measurement errors on both $x$ and $y$ by simply subtracting their effect from the estimates of the covariance and variance. To this purpose, suppose that each point $(x, y)$ in our dataset is affected by a statistical error, so that the measured values $(\tilde{x}, \tilde{y}) = (x, y) + (e_x, e_y)$. The quantities $(e_x, e_y)$ represent the errors, and are drawn from some distribution; the errors on $x$ and $y$ are not assumed to be uncorrelated here. The presence of the errors changes the covariance and variance in the expression of $\beta$ above as follows

$$\text{Cov}(x, y) = \text{Cov}(x, y) + \text{Cov}(e_x, e_y),$$

(10)

$$\text{Var}(x) = \text{Var}(x) + \text{Var}(e_x),$$

(11)

where we have introduced the variance and covariance of the errors measurements. Since the true, original variance and covariance is relevant for the estimate of $\beta$, the equations above suggest that in presence of measurement errors, we can replace the equation above with

$$\beta_{\text{BCES}} = \frac{\text{Cov}(x, y) - \text{Cov}(e_x, e_y)}{\text{Var}(x) - \text{Var}(e_x)},$$

(12)

Note that, in contrast to $(x, y)$, the variance and covariance of the measurement errors is assumed to be known and cannot be derived from the data alone. For example, in our case, if $(x, y)$ are two colors, say $x = H - K$ and $y = J - H$, we will have

$$\text{Cov}(e_x, e_y) = -\sigma_H^2,$$

(13)

$$\text{Var}(e_x) = \sigma_H^2 + \sigma_K^2,$$

(14)

where $\sigma_H$ and $\sigma_K$ are the mean photometric error in the $H$ and $K$ bands, respectively.

2.2.5. A new method for linear regression: LinES

The standard BCES methods is very simple to implement, but, although, in principle, it should be able to work in presence of an intrinsic scatter, in practice, at least in the original version presented by Akritas & Bershady (1996), it is does not. The authors do mention the possibility of accounting for intrinsic scatter in the data, but only when $x$ is measured without error, which is not applicable to the case of the extinction law. Additionally, in the situations that they consider, the amount of intrinsic scatter is taken to be unknown. In our case, however, we can estimate this quantity directly from a control field with no extinction: the variance and covariance of the colors there is just the sum of the variance and covariance due to the intrinsic scatter in color of stars, and by the photometric errors for the control field. This way we can derive the slope using the following expression:

$$\beta_{\text{LinES}} = \frac{\text{Cov}(x, y) - \text{Cov}(e_x, e_y) - \text{Cov}(e_x, y) + \text{Cov}(e_x, e_y)}{\text{Var}(x) - \text{Var}(e_x) - \text{Var}(y) + \text{Var}(e_x, e_y)}.$$

(15)

We call this estimate the LinES method (Linear regression with Errors and Scatter)$^2$. As shown in Sect. 3.4.7, this method is robust against the presence of correlated errors in both variables, and against the presence of intrinsic scatter.

3. Method validation

We tested the methods described above using controlled sets of synthetic data. The following sections discuss these data, and the effects of varying the data parameters on the results produced by each method.

3.1. Synthetic data

We constructed a set of simulations meant as controlled, realistic datasets comparable to the observed data for the Pipe Nebula cores (Lombardi et al. 2006; Alves et al. 2007). The simulated data consisted of a number of points (stars) characterized by brightnesses in three bands – arbitrarily $J$, $H$ and $K$ – affected by a value of extinction that obeys a predetermined extinction law. We simulated three sets of data to test the different aspects of each method.

$^2$ A code for fitting data with LinES will be available online.
approximately matches that for the observed data of the Pipe cores
the median extinction of the Pipe, and with a width of

\[ \sigma_A = 0.112 \] (Rieke & Lebofsky 1985) and the values of extinction drawn
from Eq. (16) were applied to the J, H and K brightnesses using
the extinction law characterized by:

\[ A_H/A_K = 1.55 \] (17)
\[ A_J/A_K = 1.55(\beta + 1) - \beta. \] (18)

The second equation, derived from Eq. (1) for a (J − H) vs. (H − K)
diagram, defines \( \beta = E(J − H)/E(H − K) \), which means the
reddening vector has a slope of \( \beta \) in a (J − H) vs. (H − K) color–
color diagram. \( A_H/A_K = 1.55 \) is adopted from Indebetouw et al. (2005).

3.1.1. Set 1: Homoscedastic data, no intrinsic scatter
For the first set, the H and K brightnesses were derived from J
assuming that all stars have the typical intrinsic color of giants
(J − H = 0.7 and H − K = 0.15, Bessell & Brett 1988).3 Each star
was then assigned a value of extinction drawn randomly from the
extinction profile (Eqs. (16), (18), Fig. 2).

In addition to reddening, each star was assigned errors in
J, H and K, to simulate the photometric errors inherent to real
data. The errors were first applied in the simplest possible way:
individually for brightness. The magnitudes of the errors were
drawn randomly and independently for J, H and K, from a nor-
mal distribution with \( \mu = 0 \) and \( \sigma = 0.05 \), so that 95% of the
stars will have errors below 0.1 mag, the typical acceptable
errors for photometry. A random value from these distributions
was then added to the magnitudes of each star in all bands inde-
pendently. The synthetic data produced in this way will hence-
forth be referred to as Set 1, or homoscedastic dataset, since the
data do not scale with the variables.

3.1.2. Set 2: Heteroscedastic data, no intrinsic scatter
A second approach was designed to reproduce the fact that the
error in real observations does in fact increase with magnitude.
We modeled this dependence using an error distribution in the
form of a power-law \( S(m) = Cm^\beta \), where \( S \) is the typical error
associated with magnitude \( m \) (Fig. 3). The normalization con-
stant \( C \) was set so that 90% of the 25th magnitude stars have an
error up to 0.3 mag, and the index \( \beta \), that defines how rapidly
the errors increase with magnitude, was set to 4. Both parame-
ters were empirically chosen to produce a curve that resembles a
typical error distribution of the NIR data, but other combinations
around these values would also be good representations of the
general distribution of photometric errors in a sample, and do not
change the results. This function was then used to determine the
width of the Gaussian from which the errors for each star were
drawn according to its magnitude, the error for the bright stars
being drawn from a narrow Gaussian, and that for the faint stars
being drawn from a wider Gaussian (see the insets in Fig. 3). The
synthetic data produced in this way will hereafter be referred to
as Set 2, or heteroscedastic data, since the errors do scale with
magnitude.

Figure 4 shows the construction of the synthetic data in the
color–color diagrams for a given realization of heteroscedastic
data without intrinsic scatter.

3.1.3. Set 3: Heteroscedastic data with intrinsic scatter
The final experiment was meant to represent the fact that the
stars being observed through the cloud do not have a single

\[ \beta = \frac{E(J − H)}{E(H − K)} \] (18)

For all sets, the J brightnesses were drawn randomly from
the distribution of J luminosities from an observed unreddened
control field toward the galactic bulge (Fig. 1).

The synthetic extinction profile was defined as a lognormal
distribution (Eq. (16)) centered at \( \mu = \log(2.5) \), the logarithm of
the median extinction of the Pipe, and with a width of \( \sigma = 0.46 \),
chosen so that the number of stars at high extinctions approxi-
mately matches that for the observed data of the Pipe cores
(Fig. 2)

PDF = \frac{1}{\sqrt{2\pi} \sigma} e^{-\frac{(\log A_K - \mu)^2}{2\sigma^2}}. \tag{16}

Because the probability distribution function (PDF) for the ex-
tinction is a lognormal, there will be many stars at low extinction
and progressively fewer stars at high extinction.

\( A_K \) was then calculated from the relation \( A_K/A_V = 0.112 \)
(Rieke & Lebofsky 1985) and the values of extinction drawn
3 The exact value of these colors is not relevant for the tests.

Fig. 1. Observed J-band luminosity function of a control field, used as
a model distribution for the unreddened J luminosities of the synthetic
data.

Fig. 2. Log-normal model distribution of extinction \( A_V \) for a generic
cloud. The dotted line corresponds to \( A_V = 2.5 \) mag, the median extinc-
tion of the Pipe Nebula.
For each set intended to pose as science data, we generated a corresponding set intended to pose as data from a control field. For Sets 1 and 2 the synthetic control fields had equivalent homoscedastic or heteroscedastic errors, respectively, drawn randomly but independently from the same distributions as the errors for the synthetic “science” fields. For Set 3, apart from the heteroscedastic errors, the control field simulation also included an amount of intrinsic scatter equivalent to that of the synthetic “science” field. Neither of the synthetic control fields included extinction, as they are meant to represent the population background to the cloud that is causing the reddening on the “science” field stars.

3.2. “Real” data

There is one aspect of real observations that we cannot test with the synthetic datasets described above: whereas at low extinction both main-sequence and giants can be observed, at high extinction the main-sequence stars are more efficiently dimmed below the detection limit, leaving a population dominated by the intrinsically brighter giants at redder colors. In our definitions for the synthetic data, this means that \( f \) should change with extinction, being larger at low extinctions and progressively smaller at high extinctions. As already hinted in Sect. 3.1.3, in set 3 of our synthetic data we do simulate intrinsic scatter, but the brightness of each star does not scale with its spectral type, which translates into having a constant \( f \) throughout the entire extinction range. To test the effect of a varying amount of intrinsic scatter with extinction on the methods we applied them to actual observations of control fields (courtesy of C. Róman-Zúñiga), which we reddened in the same way as we did the synthetic data. We refer to these datasets as “real”, keeping the quotes to make clear that the actual observed data were then artificially modified.

The first dataset contains data taken with the SOFI instrument at ESO’s New Technology Telescope, in the direction of the galactic disk \((10°38′12″, −59°12′02″, J2000.0)\), on the night of March 31st, 2006. This dataset, which we refer to as “disk dataset”, contains 548 stars with PSF photometry in the \( J, H \) and \( K_s \) filters. This is not the ideal dataset for two reasons: first, it contains few stars, and second, since this is a field in the galactic disk, it already has some extinction.

The second dataset contains data also taken with the SOFI instrument at ESO’s New Technology Telescope on the night of June 22nd, 2002, but in the direction of the galactic bulge \((17°08′10″, −28°03′03″, J2000.0)\). This dataset, which we refer to as “bulge dataset”, contains 1071 stars with PSF photometry in the \( J, H \) and \( K_s \) filters.

5000 “science” subsets were drawn randomly from each of these datasets, and extinction was applied to each star from the extinction distribution (Eq. (1), Fig. 2). Similarly, 5000 “control field” subsets were drawn randomly from each dataset, and used as they were. 450 and 800 stars were drawn from the disk and the bulge datasets, respectively, in order to keep the possibility of the “control” and “science” subsets being made of different stars.

For the effects of these experiments, the two sets differ in three aspects: the bulge dataset contains more stars, does not have extinction, and is made up mostly of giant stars, whereas the disk dataset contains fewer stars, has already some extinction, and is more likely to have a higher fraction of main-sequence objects. The \((J − H)\) vs. \((H − K)\) color–color diagrams of the two datasets are shown in Figs. 5 and 6, and illustrates these differences. In both figures, the left panels are the original datasets, and the middle and right panels show one realization of the extracted “control field” and “science” subsets used for the tests, respectively.

The only caveat regarding these datasets is that, when reddening the stars to pose as science subsets, their magnitudes change, so their associated errors should change accordingly. However, since these are real observations and given the statistical nature of the errors, that adjustment is not possible. In practice, this means that there will be stars at high extinction with an underestimated associated error, but for the effect of our tests this is not critical, since there continues to be no clear dependence of the error with extinction, as is the case in real data subject to extinction.

Fig. 3. Width of the error distribution as a function of magnitude used to generate the “photometric errors”. The insets show the actual error distribution for stars of magnitude 15 (left) and 25 (right).
**Fig. 4.** Example of the generation of synthetic heteroscedastic data without scatter. From *left to right*: all synthetic stars have the same intrinsic color; they are reddened according to the log-normal distribution of $A_V$; photometric error that depends on magnitude is added; and finally a magnitude cut is implemented, reducing the size of the sample and the coverage in $A_V$.

**Fig. 5.** ($J - H$) vs. ($H - K$) color–color diagrams for the disk dataset. *Left*: all stars in the dataset. *Middle*: 450 stars chosen randomly from the dataset, representing the “control field”. *Right*: 450 stars chosen randomly from the dataset, and reddened to represent the “science field”.

**Fig. 6.** ($J - H$) vs. ($H - K$) color–color diagrams for the bulge dataset. *Left*: all stars in the dataset. *Middle*: 800 stars chosen randomly from the dataset, representing the “control field”. *Right*: 800 stars chosen randomly from the dataset, and reddened to represent the “science field”.

A139, page 6 of 15
Table 1. Effective number of stars as a function of magnitude cut.

<table>
<thead>
<tr>
<th>( m_c )</th>
<th>( N_S )</th>
<th>( N_{S_E}^{\text{eff}} )</th>
<th>( r_{\text{eff}} )</th>
<th>( (H-K)_{\text{max}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>5000</td>
<td>4929</td>
<td>0.99</td>
<td>2.32</td>
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<tr>
<td>21</td>
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<td>4550</td>
<td>0.91</td>
<td>1.65</td>
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<tr>
<td>19</td>
<td>5000</td>
<td>2930</td>
<td>0.59</td>
<td>1.35</td>
</tr>
<tr>
<td>17</td>
<td>5000</td>
<td>680</td>
<td>0.14</td>
<td>1.04</td>
</tr>
</tbody>
</table>

3.3. Parameters

To test the robustness of the methods, the synthetic datasets were generated using a range of parameters, namely input slopes of the reddening vector, amount of intrinsic scatter, size of the sample, and other specific parameters only relevant to some of the methods.

Input slope. Each set was generated with seven values of input slope \( \beta \) in the range \([-1, 0, 3.0]\) to cover the range expected for an extinction law in the near- and mid-infrared. The methods were tested under ideal conditions of number of stars and \( A_V \) coverage to test only the ability of the methods to deal with different values of \( \beta \). The input slope was varied to guarantee that our conclusions are not only valid for one specific value of \( \beta \). While varying the remaining parameters the input slope was fixed at 1.8.

Magnitude limit. The magnitude limit was parameterized by \( m_c \). It corresponds to setting a brightness limit in real data, and it was applied identically in \( J, H \) and \( K \) such that all the synthetic stars fainter than \( m_c \) in any band after applying the extinction, are discarded from the fit (see rightmost panel in Fig. 4). A magnitude limit is naturally set in real data (detection limit), but is also something one might consider doing artificially to eliminate those stars with the largest photometric errors.

Decreasing \( m_c \) is equivalent to reducing the size of the sample, both in number and in range of extinction (the stars at high extinction will likely be fainter), while simultaneously reducing the range of errors in Sets 2 and 3 (fainter stars will have larger errors). By construction, a value of \( m_c = 25 \) corresponds to allowing the largest errors to be 0.3 mag (see Sect. 3.1.2).

Table 1 shows the average\(^4\) number of stars from Set 3 that survive each magnitude cut (effective number of stars, \( N_{S_E}^{\text{eff}} \)) from an initial sample of \( N_S = 5000 \) synthetic stars. The ratio \( r_{\text{eff}} = N_{S_E}^{\text{eff}} / N_S \) reflects the functional form of the model distributions and, as such, would be the same for any other input number of stars \( N_S \) for each magnitude cut. The table also shows the average \( H - K \) color of the most heavily reddened datapoint attained for each magnitude cut, illustrating the loss in \( A_V \) coverage with magnitude limitation. The same magnitude cuts were applied to the control field, but because the control field does not have extinction, the effect of the cut in the effective number of stars is much more subtle. The ratio of effective to initial number of stars for the control field is 1.00 for \( m_c = 21 \) mag, and drops to 0.90 and 0.25 for \( m_c \) of 19 mag and 17 mag, respectively.

Number of stars. Generating fewer stars in the first place also changes the size of the sample. The (subtle) difference with respect to implementing magnitude cuts is that the sample with fewer stars and no magnitude cut will most likely have a broader range of \( A_V \) than would a richer sample with magnitude cut, as some of the fewer stars that are generated and kept may still be faint and heavily extincted from the random drawing process. In real observations, generating fewer stars without imposing magnitude cuts would be comparable to observing a sparse region of the sky where the faint and highly reddened stars can be detected. Imposing a magnitude limit, on the other hand, would correspond to having shallow observations regardless of the richness of the observed field; in this case, the fainter and more reddened stars would not be detected.

We tested the methods against varying number of stars in the range \([100, 5000]\), both in the science and the control field.

Amount of intrinsic scatter. The amount of intrinsic scatter for Set 3 was parametrized by \( f \), the fraction of stars in the main-sequence locus with respect to those in the giant locus, taking values of 0.01, 0.15 and 0.50. Given the shape of the loci, a larger scatter is obtained for larger values of \( f \), although the giant locus alone still produces some intrinsic scatter. Larger values of \( f \) would correspond to a large fraction of the stars behind the cloud being main-sequence in real data, as opposed to having mainly giants. When varying the remaining parameters, \( f \) is fixed at 0.15 for Set 3 (\( f \) is not a parameter in Sets 1 and 2).

Number of control-field stars. The synthetic control field is only used for the LinES method, as the other methods rely solely on the science data. Unless explicitly stated, the control field was generated with the same number of stars and was subject to the same magnitude cuts as the synthetic “science” datasets. This means that, after applying a given magnitude cut, there will effectively be more stars in the control field sample than in the science sample, since a fraction of the stars in the science dataset will have been dimmed below the magnitude cut by the effect of extinction. This mimics real observations in that the control field should be obtained from a region of comparable background stellar density (same number of stars generated in the simulations) and using the same instrumental setup (same magnitude cut in the simulations) as the science field. To test the effect of a less than ideal control field, we tested the LinES method against varying numbers of stars also in the control-field datasets.

3.4. Results from synthetic data

Each method was applied to 5000 realizations of the synthetic data, producing an average value \( \hat{\beta} \) and a dispersion \( \sigma_\beta \) around the average for each parameter within sets 1, 2, and 3. We define bias \( b \) as the difference between \( \hat{\beta} \) and the input value of the slope (\( b = \hat{\beta} - \beta \)), and consider a method unbiased if it produces an estimate within the 1-\( \sigma \) dispersion of the input value (i.e., \( b \leq \sigma_\beta \)).

We note that the quoted absolute values of the biases are formally only valid for the conditions of these simulations, namely for the magnitude of the error and the functional form of the error distribution with magnitude.

The results are summarized below. Figures 7 through 13 show the bias as a function of varied parameters.

3.4.1. Ordinary least-squares

The OLS method has long been known to be biased when there are errors in both coordinates. This was observed also in our simulations (Fig. 7), with the method failing to recover the right
Ordinary Least Squares (OLS)

Bias (b) vs. Magnitude cut (mc [mag])

Set 1
Set 2
Set 3

Input slope (β)

Number of stars (Nstars)

Intrinsic scatter (f)

Weighed Least Squares (WLS)

Bias (b) vs. Magnitude cut (mc [mag])

Set 1
Set 2
Set 3

Input slope (β)

Number of stars (Nstars)

Intrinsic scatter (f)

Fig. 7. From left to right, bias in the slope as a function of the magnitude cut, input slope, number of stars in the science field, and intrinsic scatter for the ordinary least squares (OLS) method. The black, red and blue lines represent the results for the 5000 realizations of Sets 1, 2 and 3, respectively. The error bars correspond to the 1-σ dispersion in the results.

Weighed Least Squares (WLS)

Bias (b) vs. Magnitude cut (mc [mag])

Set 1
Set 2
Set 3

Input slope (β)

Number of stars (Nstars)

Intrinsic scatter (f)

Fig. 8. From left to right, bias in the slope as a function of the magnitude cut, input slope, number of stars in the science field, and intrinsic scatter for the weighed least squares (WLS) method. The black, red and blue lines represent the results for the 5000 realizations of Sets 1, 2 and 3, respectively. The error bars correspond to the 1-σ dispersion in the results.

value of the slope in the majority of the tests performed. It did provide an unbiased result for the following combinations of input slopes and datasets: β = −0.5 for Set 1, β = −1.0 and β = −0.5 for Set 2, and β = 1.0 for Set 3, making this method unsuitable if one is trying to find precisely β.

This method is also not robust against variations in mc, the bias and the dispersion both increasing for brighter mc in the heteroscedastic sample, except for the brightest magnitude cut considered, where the bias is suddenly reduced. It also reacts, although to a lesser extent, to changes in the amount of intrinsic scatter in the heteroscedastic sample (Set 3), the bias increasing with f.

This method is extremely robust against variations in the number of stars within the same magnitude cut, although the dispersion in the slope increases for fewer stars as would be expected from poor statistics, and the absolute value is biased.

Overall, this method is not a reliable estimator of the extinction law.

3.4.2. Weighed least-squares

The WLS method performs well in homoscedastic or heteroscedastic data without intrinsic scatter (Fig. 8), suggesting that, under these conditions, the method can deal properly with the presence of errors in both coordinates, and with them being correlated. For these samples the dispersion in the slope is remarkably small, making it a very accurate estimator. In the presence of intrinsic scatter, however, the method systematically fails to recover the input slope whatever its value in the range [−1.0, 3.0], although it does come close around β = 2.5. The bias as a function of input slope plot for this method and dataset (Fig. 8, blue line) suggests that there could be another unbiased value of β between −0.5 and 0.5, but tests suggest that there is instead a discontinuity around β = 0.

Although biased for Set 3, this method is robust against variations in the number of stars to within 1.5% in the considered range, but the dispersion increases steadily for fewer stars.

Biased in the presence of even a small intrinsic scatter, this method is not a reliable estimator of the extinction law.

3.4.3. Symmetrical methods

The three symmetrical methods returned biased results for all tests (Fig. 9), except for input slopes of −1.0 and 1.0 in datasets 1 and 2 (without intrinsic scatter). The bisector and geometric mean methods produce very similar results. The orthogonal regression method presents the largest biases of the three.

Besides being biased, neither method is robust against variations in mc or f in the presence of intrinsic scatter, the bias increasing for bright magnitude cuts and more intrinsic scatter. The methods are highly robust to variations in the number of stars, the dispersion increasing steadily for fewer stars.

These methods are not reliable estimators of the extinction law.
3.4.4. Binning in \((H-K)\)

This method is biased for most datasets and parameters tested; the exceptions are for Set 3 with the brightest magnitude cut \((m_c = 17 \text{ mag})\) or for \(\beta = 0.5\) and \(1.0\), and for \(\beta = -0.5\) and no intrinsic scatter (Fig. 10).

The slope is mostly underestimated for all datasets, with the bias increasing for brighter \(m_c\) for Set 1, and keeping relatively stable against varying \(m_c\) for Sets 2 and 3. For Set 3 the bias slightly increases with \(f\), as does the dispersion. This method reacts to the number of stars for a given magnitude cut, the bias increasing toward fewer stars. Overall, this is not a reliable estimator of the extinction law.
3.4.5. Binning in $A_v$

This method is formally unbiased for all tested values of $\beta$ when applied to the three datasets (Fig. 11). However, whereas the bias is always close to zero for the datasets without intrinsic scatter, it becomes slightly large for most values of $\beta$ when intrinsic scatter is introduced.

The method is robust against variations in the amount of intrinsic scatter within the considered range, and it is stable against variations in magnitude cut, except for the brightest value of $m_c$ for Set 3, where the smaller number of stars and range in $A_v$ result in very few bins for the fit. The dispersion increases toward brighter magnitude cuts and amount of intrinsic scatter. Because it is based on binning, this method reacts significantly to the number of stars within the same magnitude cut, the bias and the dispersion increasing for fewer stars.

Small variations occur when the size of the bin is varied, with the slope being increasingly overestimated for smaller bins, but the method continues to be unbiased within the dispersion.

This method is a reliable estimator of the extinction law.

3.4.6. BCES method

This method is highly reliable and unbiased for homoscedastic data and for heteroscedastic data without intrinsic scatter (Sets 1 and 2). However, in the presence of intrinsic scatter (Set 3), it becomes biased for all input slope values except for $\beta = 2.1$ (Fig. 12) for $f = 0.15$. Since the other parameters were tested using an input slope of 1.8 (very close to 2.1), the bias seems small in the $m_c$, $N_{\text{stars}}$, and $f$ plots, but we can nevertheless analyze the sensitivity of the method to these parameters. For Set 3, the bias is mildly sensitive to $m_c$, robust against variations in the number of stars, and only minimally sensitive to variations in the amount of intrinsic scatter $f$.

Since real data will be similar, in essence, to Set 3, we do not consider this method a reliable estimator in the specific case of the extinction law.

3.4.7. LinES

This method is the most unbiased and robust of all presented here for homoscedastic or heteroscedastic data, with or without intrinsic scatter, for all tested values of the input slope (Fig. 13).

The method is robust against variations in $m_c$ and $f$, number of stars in the science field, and number of stars in the control field, although the dispersion follows the same tendency as before: increases for brighter magnitude cuts, slightly increases with $f$, and decreases with number of science and/or control field stars. The bias is larger when there are simultaneously very few...
science and control field stars, and small $A_V$ coverage, but is nevertheless better than any of the other methods for the same
displacement. The performance is significantly smaller than the next
closest unbiased method, the binning in $A_V$, in all cases, granting it more precision.

Since LinES relies on the characterization of the data
through the properties of a control field, we tested the stability
of the method against variations in the number of stars in the
control field. Given a reasonable number of stars in the science
field, the method is robust against variations in the number
of control field stars. However, if the science field itself does not have enough stars or $A_V$ coverage, the bias increases further
for few control field stars. Inevitably, the dispersion increases
toward fewer control field stars. This method has proven to be
robust as long as the control field is a good representation of the
underlying population on the science field, even if containing a
smaller number of stars.

The excellent performance of this method while varying all
relevant parameters validates the LinES method for our case
study. In general, it will provide accurate results for observations of cores with either rich or poor background populations,
regardless of their spread in spectral types, even for relatively
shallow observations, as long as there is a reasonable spread in
extinction and the control field is a good representation of the
reddened, background population.

Limitations. The simulations show that LinES is not reliable for
distributions that do not cover a wide enough range of extinction, if the errors are too large. For reasonable errors, like those
described for the simulations, LinES starts overestimating the
slope by more than 10% for ranges in $x$-color (i.e., the color
plotted on the $x$-axis) smaller than 0.25 to 0.45 mag for slopes
between 0.6 and 3.0, respectively, and underestimating the slope
by more than 10% for the same ranges in $x$-color for slopes be-
tween 0.3 and 0.5. This method should therefore not be applied
to data that span less than these values in $x$-color.

Error estimation. We used the bootstrap method (e.g., Wall &
Jenkins 2003) to estimate the uncertainty in the slope derived by
LinES. This method consists in randomly dividing each sample
in two equal-number subsets, and measuring the slope of the extin-
tion law in each subset. This produces two values of $\beta$ from
which we derive the standard deviation $\sigma_{\beta}$. This was repeated
$N = 1000$ times and the uncertainty in the slope was defined as:

$$\sigma_{\beta} = \frac{1}{\sqrt{2N}} \sum_{i=1}^{N} \sigma_{\beta_i}. \quad (19)$$

We applied this method on the synthetic data and compared the
results with the dispersion $\sigma_{\beta}$ from the simulations. We found
that the bootstrap uncertainty is typically 80% of $\sigma_{\beta}$, the disper-
sion from the synthetic data, and the value we believe is a better estimate of the actual dispersion expected for real data. Given
the consistency of the results against the variation of the differ-
ent parameters, we take the uncertainty on the LinES result of
our case study to be 1.25 times the uncertainty derived by the
bootstrap method described above.

3.5. Results from “real” data

Figures 14 and 15 show the bias of each method vs. varying input
slope and varying magnitude cut, respectively, for the “real”
datasets described in Sect. 3.2. Except for the cases described
below, these results follow those from the synthetic data, main-
taining the same behavior and changing only the magnitude of
the bias. The methods that deserve some further consideration
are the binning in $A_V$ method, the BCES, and LinES.

The BCES (bottom-middle panel in the figures) applied to
these datasets shows the same type of behavior as with the syn-
thetic data against varying magnitude cuts (Fig. 15). The bias
is slightly larger than before because there are fewer stars and
dispersion. For varying input slopes, however, the bias is con-
ciderably larger, showing that the method is severely biased
for some ranges of $\beta$ when applied to less than ideal data.

The binning in $A_V$ method, which we concluded was reason-
ably unbiased when applied to the synthetic data, shows a severe
bias also for some values of the input slope $\beta$, in particular for the
disk dataset (bottom-left panel in the figures). However, this hap-
pens because this dataset has very few stars, and a very limited
range in $A_V$ (see Fig. 5), which translates into too few $A_V$ bins to
constrain the extinction vector slope adequately. This presents a
limitation of the method, away from which it can be used with
relatively good precision.

In the “real” datasets, LinES continues to perform beauti-
fully. Except for the first point in Fig. 15 (bottom-right panel),
where the magnitude cut is such that limits the $A_V$ to a very
narrow range, the bias is consistent with zero for all other magnitude cuts and for all values of the input slope we tested.

4. Detecting a break in the extinction law

Grain growth at high densities has been proposed by a number of studies (see references in Sect. 1). If it does occur, then in dense cores the extinction law will become grayer toward higher extinctions, which should translate into a variation of the slope of the reddening vector with extinction, either smooth or abrupt depending on the nature of the transition between grain sizes. This break was detected and measured in the Trifid Nebula for an extinction of $A_V = 20$ mag by Cambrésy et al. (2011). We used the synthetic and “real” datasets to test whether we could detect such a break in the extinction law using LinES. The data was generated using the exact same method as described above for Set 3 with $f = 0.15$ and for the “real” dataset (see Sects. 3.1.3, 3.2), but the reddening vector was made flatter at some value of $A_V$, or conversely, at some value of $(H-K)$ color, since the color scales linearly with $A_V$. This produced color–color diagrams similar to that of Fig. 16, where the break is more or less obvious depending on the difference between the two slopes.

For each realization, we divided the sample into low-extinction ($(H-K) < (H-K)_{\text{limit}}$) and high-extinction ($(H-K) > (H-K)_{\text{limit}}$), and determined the best fits to the reddening vector in the two groups using the LinES method, obtaining two slopes $\hat{\beta}_{\text{low}}$ and $\hat{\beta}_{\text{high}}$. This was done for increasing values of $(H-K)_{\text{limit}}$ in steps of 0.2, and starting at $(H-K)_{\text{limit}} = 0.4$.

Figures 17 and 18 show the behavior of $\hat{\beta}_{\text{low}}$ (black) and $\hat{\beta}_{\text{high}}$ (red) as a function of $(H-K)_{\text{limit}}$ for the synthetic dataset 3 and the “real” bulge dataset, respectively, when applied to 5000 realizations. The solid lines show the median curves, and the shaded regions represent the 1–σ scatter from the 5000 realizations. The upper panels are for a single-slope reddening vector with slopes 1.0 (left) and 1.5 (right). The bottom panels show the same distributions but for broken extinction laws, with slopes of 1.5 at low-extinction and 1.0 and high-extinction, and breaks located around $A_K = 0.4$ mag (left) and $A_K = 1.5$ mag (right). For completeness, we find the same results using the disk dataset, albeit with a larger dispersion.

In the absence of a break, the two curves are indistinguishable except for the lowest and highest values of $(H-K)_{\text{limit}}$; this is because the subsets used for the fits in these two extremes contain too narrow ranges in $(H-K)$ to constrain the LinES method. When a break does exist, however, the two curves separate distinguishably, even if the break is at low extinction. This then provides a simple method to test whether the same extinction law applies to the full $A_V$ range of a given dataset, or if it would rather best be described as a two-segment law. Unfortunately this method does not allow for the determination of the actual value of the break, but the figures show that the slope of the extinction vector at high extinction can be determined with reasonable accuracy. In particular, the procedure of measuring the slopes in the low-$A_V$ and high-$A_V$ regimes provides a much better handle...
on the extinction law at high extinction (red line in the figures) than measuring the slope of the entire dataset as a whole (blue line in the figures).

5. Summary

We tested several methods of linear regression associated with the problem of measuring the extinction law from photometric data. We found that many of the commonly used methods provide biased results caused by the presence of errors in both coordinates (which are colors), by the fact that they are correlated, and by the presence of scatter intrinsic to the underlying distribution.

We adapted the BCES method of Akritas & Bershady (1996) to allow a compensation for intrinsic scatter, using a control field to characterize the background, unreddened population. We called this method LinES (Linear regression with Errors and Scatter). Using synthetic data, we showed this method provides unbiased and correct results, and that it is robust against the variation of all relevant parameters (at least) within reasonable limits, such as size of sample, range of extinction, and amount of intrinsic scatter.

We found that dividing any subset in sliding values of $A_V$ and measuring the slopes of each subset can robustly differentiate between an extinction law characterized by a single slope and one with a break.

These results can be applied to observations of background stars seen through dense cores of molecular clouds, or through regions that span a reasonable range of dust density. The characterization of the extinction law through deep, photometric data is a very useful tool to probe the properties of the dust grains.
Fig. 17. $\hat{\beta}_{\text{low}}$ and $\hat{\beta}_{\text{high}}$ as a function of $(H-K)_{\text{limit}}$ for 5000 realizations of the synthetic dataset 3. The solid lines show the median curves, the shaded regions represent the 1-$\sigma$ scatter from the 5000 realizations, and the dashed lines mark the input values $\beta_{\text{low}}$ and $\beta_{\text{high}}$ of the slopes. Upper panels: $\hat{\beta}_{\text{low}}$ and $\hat{\beta}_{\text{high}}$ for a single-slope reddening vector with slopes 1.0 (left) and 1.5 (right). Bottom panels: same distributions but for broken extinction laws, with slopes of 1.5 at low-extinction and 1.0 and high-extinction, and breaks located at $A_K = 0.4$ mag (left) and $A_K = 1.5$ mag (right). The blue line indicates the average value of the slope if LinES is applied to the whole range of $A_V$.

Fig. 18. $\hat{\beta}_{\text{low}}$ and $\hat{\beta}_{\text{high}}$ as a function of $(H-K)_{\text{limit}}$ for 5000 realizations of the bulge dataset. The solid lines show the median curves, the shaded regions represent the 1-$\sigma$ scatter from the 5000 realizations, and the dashed lines mark the input values $\beta_{\text{low}}$ and $\beta_{\text{high}}$ of the slopes. Upper panels: $\hat{\beta}_{\text{low}}$ and $\hat{\beta}_{\text{high}}$ for a single-slope reddening vector with slopes 1.0 (left) and 1.5 (right). Bottom panels: same distributions but for broken extinction laws, with slopes of 1.5 at low-extinction and 1.0 and high-extinction, and breaks located at $A_K = 0.4$ mag (left) and $A_K = 1.5$ mag (right). The blue line indicates the average value of the slope if LinES is applied to the whole range of $A_V$. 

A139, page 14 of 15
in these regions, and a “cheap” one when compared with, for example, spectral analysis of many individual sources.

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